CANADIAN THESES ON MICROFICHE

THESES CANADIENNES SUR MICROFICHE

National Library of Canada Collections Development Branch

Canadian Theses on Microfiche Service

Ottawa, Canada KIA 0N4 Bibliothèque nationale du Canada Direction du développement des collections

I.S.B.N

Service des thèses canadiennes sur microfiche

NOTICE

The quality of this microfiche is heavily dependent upon the quality of the original thesis submitted for microfilming. Every effort has been made to ensure the highest quality of reproduction possible

If pages are missing contact the university which arouted the degree.

Some pages may have indistinct print especially if the original pages were typed with a poor typewrite of the original pages were typed with a poor typewrite.

Previously popylighted materials (in 1997 States) rechtlehed tests als Jans of filmed

Reproduction in full or in part of this film is governed by the Canadian Copyright Act, R.S.C. 1970, c. C.30. Please read the authoritation for one which a constraint the states in the states of the second states in the stat

THIS DISSERTATION HAS BEEN MICROFILMEN EXACTL AS RECEIVED

AVIS

La qualité de cette microfiche dépend grandement de la qualité de la thèse soumise au microfilmage. Nous avons tout fait pour assurer une qualité supérieure de reproduction.

S'il manque des pages, veuillez communiquer avec l'université qui a conféré le grade.

La qualité d'impression de certaines pages peut laisser à désirer, surtout si les pages originales ont été dactylographiées à l'aide d'un ruban usé ou si l'université nous a fait parvenir une photocopie de mauvaise qualité

Les documents qui font déjà l'objet d'un droit d'auteur (articles de revue, examens publiés, etc.) ne sont pas microfilmés.

La reproduction, même partielle, de ce microfilm est soumise à la Loi canadienne sur le droit d'auteur, SRC 1970, c. C-30. Veuillez prendre connaissance des formules d'autorisation qui accompagnent cette thèse.

> LA THÈSE A ÉTÉ MICROFILMÉE TELLE QUE NOUS L'AVONS REÇUE



		×		0-315-19360	»-3
	National Library of Canada	Bibliothèque national du Canada	e	· ·	· ·
$\langle D \rangle$	Canadian Theses Division	Division des thèses ca	anadiennes		
	Ottawa, Canada K1A 0N4				\$
	, ,	672	.6 8	n,	
	PERMISSION TO MIC	ROFILM - AUTO	RISATION DE MI	CROFILMER	
				•	· ·
• Pleas	e print or type — Écrire en lettre	es moulées ou dactylograp	hier		. ,
Full Nar	ne of Author Nom complet de	a l'auteur			
•	MANDHARA PIYA.	ILRI JAYATILAN	CA LENARATNE	~	
Date of	Birth — Date de naissance		Country of Birth Lie	u de naissance	
÷.	& JUNE 1953	7	SRI, LANKA	9	
Perman	ent Address - Résidence fixe				· · · · ·
	AMPAGRAZIA	SIGNATURE	₽1 /		
	No 60, MI	92 WATTE ROA	D		
	Dei	ALWATTE ROA HIWELA, COLOI	MBD, SKI LAN	KA.	
Title of	Thesis — Titre de la thèse				
	THE INTERACT	TON BETWERV	FrogENOUS N	JORP DECNALT	Nº 4
	AND TRANSMU				
	SAPHENDUIL				-
•					
5 . .		•			
Univers	ity Université				
	('/ / 3 /,	REFIA, EDMON	721		
Degree	for which thesis was presented	- Grade pour lequel cettr	thèse fut présentée		
•	Ph.D. MEDICAL	SCIENCE" (M11	MYINE)		
Year th	is dearer underred - Accés it	intention of a sector	Tame of Supervisor	Nom du directe de th	,), , , ,
	1984		DR (T IN	じじりをかりう	
	-				-
	sion is hereby granted to the N A to microfilm this thesis and t		OUE NATIONALE D	par la présento, accordé U CANADA de microfilm des exemplaires du film	ler i atta thành at ita
thesis r	thor reservés other publication nor extensive extracts from it m nor durant interaction with	in the reintert or other	ni de longs extraits	les autres droits de pub s de polleici un doivent le com Lautorication Acc	t ê're imprimés 👘
			,		:
Dete				0 1	17
			Minot	ANT PARAN	,

.....

••

•

HE DI 14 TO

₽

22

3.4

THE UNIVERSITY OF ALBERTA

THE INTERACTION BETWEEN EXOGENOUS NORADRENALINE AND TRANSMURAL NERVE STIMULATION IN THE CANINE SAPHENOUS VEIN

.

by

MANOHARA P.J. SENARATNE

A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH

ΤN

HUNTCH SCIENCES (HEDICINE)

EDMONTON, ALBERTA

FALL 1984



UNIVERSITY COLLEGE LONDON DEPARTMENT OF ANATOMY AND EMBRYOLOGY GOWER STREET LONDON WCIE 6BT

Head of Department PROFESSOR G. BURNSTOCK, D.Sc., F.A.A.

TELEPHONE 01-387 7050

GB/AE

4

29 February 1984

۵

Mr. M.P.J. Senaratne, Room 8-104, Clinical Sciences Building. University of Alberta, Department of Medicine, Edmonton, Alberta, Canada T6G 2G3.

Dear Mr. Senaratne,

Thank you for your letter of 5 February. I am happy to give you permission to include the illustration (Br. Med. Bull. 35: 257, 1979 Plate VI Fig. Ca and Cb) in your thesis. Good buck!

With best wishes.

Yours Fincerely. Long Parter 1

8 Pin tool

SUPER STRUCES COL

DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

, National Institutes of Health National Heart, Lung, and Blood Institute Bethesda, Maryland 20205

÷,

ंग

1

Ċ,

÷.,

February 28, 1984

Mr. M.P.J. Senaratne Room 8-104 Clinical Sciences Building University of Alberta, Dept. of Medicine Edmonton, Alberta T6G 2G3 CANADA

Dear Sir:

`I am more than happy to give you my permission to include the following illustration(s) in your thesis:

ANN. REV. BIOCHEMISTRY 49:1980 Fig. 1, page 925 Fig. 2, page 928

Sincerely yours,

Robert S. Adelstein, M.D. Chief, Laboratory of Molecular Cardiology National Heart, Lung, and Blood Institute National Institutes of Health

RSA:kre

M.P.J. Senaratne Room 8-104 Clinical Sciences Building University of Alberta, Dept of Medicine Edmonton, Alberta, Canada T6G 2G3

50/Feb/1984

The Editor Annual Review of Biochemistry Dr. R.S. Adelstein

Dear Sir,

I am a graduate student attached to the Department of Medicine of the University of Alberta (Edmonton, Alberta, Canada) and studying for a PhD in Medical Sciences. My research area of interest is vascular smooth muscle pharmacology. I am writing up the thesis (title: Interaction between exogenous noradrenaline and transmural nerve stimulation in the canine saphenous vein) at the present time and would be grateful if you could grant me written permission to include the following illustration(s) in my thesis.

Апп Rev Biochem 49: 1980 Fig 1 Pg 925 Fig 2 Pg 92£

Thanking you,

Manchera fereralis

M.P.J. Senaratge

Permission is granted provided you use the following acknowledgement on the first rage of the reprinted material: "Reproduced, with permission, from the Annual Review of <u>Biochconistry</u>, Vol.49. C 1980 by Annual Reviews Inc."

M15/84 Permissions, Department Annual Reviews Inc

M.P.J. Senarathe Room 8-104 Clinical Sciences Building University of Alberta Edmonton, Alberta, Canada T6G 2G3 20 (March 1984

DR PM Vashoutte Dept of Physiology & Biofohysics / POBoxin CH 4009 Basel Mayo Clinic and Founde Hons / Switzer land

Dear Sir,

I am a graduate student attached to the Department of Medicine of the University of Alberta (Edmonton, Alberta, Canada) and studying for a PhD in Medical Sciences. My research area of interest is vascular smooth muscle pharmacology. I am writing up the thesis (title: Interaction between exogenous noradrenaline and transmural nerve stimulation in the canine saphenous vein) at the present time and would be grateful if you could grant me written permission to include the following illustration(s) in my thesis.

Fig. 1 Page 117 Blood Versels 16:113-125,1979

Thanking you,

Manchara Scharalus

M.P.J. Senaratne

Paul M. Vanhante

M&P.J. Senaratne Room 8-104 Clinical Sciences Building University of Alberta Edmonton, Alberta, Canada T66 2G3 15/ March / 1984

Dear Sir,

1

S Karger AG,

O Box; CH

Witzerland

-4009

I am a graduate student attached to the Department of Medicine of the University of Alberta (Edmonton, Alberta, Canada) and studying for a PhD in Medical Sciences. My research area of interest is vascular smooth muscle pharmacology. I am writing up the thesis (title: Interaction between exogenous noradrenaline and transmural nerve stimulation in the canine saphenous vein) at the present time and would be grateful if you could grant me written permission to include the following illustration(s) in my thesis.

> Fig. 1 Page 117 Block I FILFIS IL 112 125, G. K.

Thanking you.

Manchara Ferratio

M.P.J. Scharathe

Permission dianted under the condition that the exact original source and the under the source of the source of the mentioned.

Basel April 4. 1984

S. KARGER AG, BASEL

M.P.J. Senaratné Room 8-104 Clinical Sciences Building University of Alberta Edmonton, Alberta, Canada 166 263 5/Feb/19P4

Dr. DJ Hartsborne Mutale Biology Groub University of Avizona

The Editor American Physiological Society

Dear Sir.

I am a graduate student attached to the Department of Medicine of the University of Alberta (Edmonton, Alberta, Canada) and studying for a PhD in Medical Sciences. My research area of interest is vascular smooth muscle pharmacology. I am writing up the thesis (title: Interaction between exogenous noradrenaline and transmural nerve stimulation in the canine saphenous vein) at the present time and would be grateful if you could grant me written permission to include the following illustration(s) in my thesis.

Fig. 2 Pg 100 Chapt 4 Handboot of Hyprology, Carchiovarca las system IT Varcalar Imooth Muscle American Physicological Porce fy 1900

Thanking you.

Manchava figaratio

M.F.J. Senarative

Transition is printed to rear i love illustration, ". 11.7/e t Alter those

M.P.J. Senaratne Room 8-104 Clinical Sciences Building University of Alberta Edmonton, Alberta, Canada T6G 2G3 5 Feb 1984

Dr. J. A. G. Rhodin Dept. of Anatomy, University of Michigan American Physiological Ann Arbor, Michigan Cociety

Dear Sir,

I am a graduate student attached to the Department of Medicine of the University of Alberta (Edmonton, Alberta, Canada) and studying for a PhD in Medical Sciences. My research area of interest is vascular smooth muscle pharmacology. I am writing up the thesis (title: Interaction between exogenous noradrenaline and transmural nerve stimulation in the canine saphenous vein) at the present time and would be grateful if you could grant me written permission to include the following illustration(s) in my thesis.

Fig 1, 11, 12 Chapter 1 Handbook of Physiology Condicingention Lystem I, Varcalow Smooth muscle ing you. Am l'hysiclogical Scelety 1400 Thanking you.

Maschara Sever Ali Senarathe Permission granted. Johannes A.G. Rhodin, M.D., Ph.D. Professor & Chairman

Professor & Chairman Department of Anatomy University of South Florida College of Medicine 12901N 30th Street

40 B 20/1984

THE BRITISH COUNCIL

10 Spring Gardens London SW1A 2BN telephone 01:930 8466 ext 2697/2667 telex 3866522 8952201

Mr M P J Senaratne Room 8-104 Clinical Sciences Building University of Alberta Department of Medicine Edmonton ALBERTA Canada T6G 2G3

43

our ref GEN/109/9

your ref

please quote our reference

23 March 1984

Dear Mr Se

21

Thank you for your letter of 5 February in which you seek permission to reproduce an illustration that appeared in the British Medical Bulletin of 1979.

Professor Burnstock as the author of that article in fact also holds the copyright, notwithstanding the printed notice in the Bulletin of that date which is not valid. If you receive his permission, therefore, that is all that will be necessary.

Yours sincerely

O

Dr J D Crowlesmith Senior Medical Adviser

cc: Professor G Burnstock PhD DSc FAA Department of Anatomy & Embryology University College, London

i,

M.P.J. Senaratné Room 8-104 Clinical Sciences Building University of Alberta Edmonton; Alberta, Canada 166 263 5 / Feb/ 1984 The Editor Dr Hartsborne DJ Dr JAG Rhodin American Physiological Society Betherda, Maryland, 20 F14 9650, Rockville Ake AMERICAN PHYSIOLOGICAL SOCIETY . FEB 1 3 1984 Dear Sir, I am a graduate student attached to the Department of Medicine of

the University of Alberta (Edmonton, Alberta, Canada) and studying for a PhD in Medical Sciences. My research area of interest is vascular smooth muscle pharmacology. I am writing up the thesis (title: Interaction between exogenous noradrenaline and transmural nerve stimulation in the canine saphenous vein) at the present time and would be grateful if you could grant me written permission to include the following illustration(s) in my thesis.

le and would be yrac. to include the following illustration () Fig. 2 Pg 100 Chapter 4 CF (G. 11,12) Pg Handbook of Physiology Fig 1 1 Pg Cardiovascular Mysfem II Vascular 1940 14 Chapter 1 Ø Smooth muscle

Thanking you,

Manchera Scharala

M.P.J. Senaratne

THE AMERICAN PHYSIOLOGICAL SOCIETY 9650 Rockville Pike - Bethosda, MD 20014

Permission is granted for use of the material specified above, contingent upon consent of the author and provided the publication is credited as the source.

Stephen R. Geiger Publ Mgr & Ever Editor

Date

THE UNIVERSITY

RELEASE FORM

NAME OF AUTHOR: Manohara P.J. Senaratne

n

TITLE OF THESIS: The Interaction Between Exogenous Noradrenaline and Transmural Nerve Stimulation in the Canine Saphenous Vein

DEGREE FOR WHICH THESIS WAS PRESENTED: Doctor of Philosophy

YEAR THIS DEGREE GRANTED:

Fall 1984

Permission is hereby granted to THE UNIVERSITY OF ALBERTA LIBRARY to reproduce single copies of this thesis and to lend or sell such copies for private, scholarly or scientific research

The author reserves other publication rights, and neither the thesis nor extensive extracts from it may be printed or otherwise reproduced without the author's written permission.

and Fraena (Signed)

PERMANENT ADDRESS:

No. 60/1, Malwatte Road Dehiwela SRI Lanka (Coston)

Pated: June 04, 1984.

THE UNIVERSITY OF ALBERTA

FACULTY OF GRADUATE STUDIES AND RESEARCH

The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research, for acceptance, a thesis entitled THE INTERACTION BETWEEN EXOGENOUS NORADRENALINE "AND TRANSMURAL NERVE STIMULATION IN THE CANINE SAPHENOUS VEIN submitted by Manoharn Separation in partial fulfillment of the requirements for the doct of Dect of Milesophy in Medical Sciences.

Supervisor Dr. C.T. Kappagoda Research Professor of Medicin Department of Medicin University of Alberta Edmonton, Canada

Dr. M. Wolowy's Professor Department of Pherica University of Al

1

5. External Examiner

Dr. J.T. Shephord Professor Department of Physiclog: an Elon Mayo Graduate hard of Main Dr.

Dr. D.A. Cook

6 · · · ·

Chairman Department of Pharmacology University of Allows Wimpton Canada

Dr. T. Wet an Professor 1 Does 1 0 1

This thesis is dedicated to my wife, Savitri, and to my To parents who went through a lot of hardship in helping me

whieve this goal.

4

•

÷

.

ABSTRACT

The present investigation was undertaken to study the interaction between contractions produced by exogenous noradrenaline (NA) and contractions produced by transmural nerve stimulation (TNS) in isolated canine saphenous vein rings/strips Four protocols were carried out. Protocol 1.1: Response to a single dose of exogenous NA was determined as a control. After washing, TNS was applied to produce a contraction between 10-90 per cent of the control value and the same duse of NA applied against this background and the response determined. Finally, after washing, control response with NA cas repeated. The observed contraction to NA against a background of INS was significantly less than the expected contraction (78 observations in 12 veins, p 4 0.001), Imhus the response to exogenous NA was inhibited by background Further, the inhibition appeared to be greater with increasing TNS. magnitude of background TNS, Protocol 1.2. The reciprocal of Froto of 1.1 was done here. The response to TNS was determined against a background contraction by exogenous UA. The observed contraction to TNS was significantly greater than the expected correction (7) observed ins in 12 motion p < 0.0011. Thus backer with ingeneric 11 was found to enhance the response in TVS. The control of the fold information of the UV contraction observed a to two it was not deeped in the magazine of guanethild a or dility of the history is after to a set the rest of restance Wheel at a second to be a second second state of the second second second second second second second second se 100 and the second second second second second adled against curre !! Lorde . 11 I I I I the states of an first " "heoxarfu: the third the the tenters

phenomenon observed in Protocol 1.1 was no longer evident with any of the above agonists. Protocol 1.5: The inhibition of exogenous NA contraction was not significantly altered by propraholol $(10^{-5} \text{ mol}/1)$, indomethacin (10^{-5} mol/1), cimetidine (10^{-5} mol/1) and aminophylline (10⁻⁵ mol/1). Protocol Two: Protocol 1.1 and Protocol 1.2 carried out in a superfused saphenous vein strip (following labelling of NA stores with ³H-noradrenaline) demonstrated inhibition of the TNS induced release of ³H-noradrenaline by exogenous NA. Protocol Three: The response to TNS in a vein ring pre-contracted with prostaglandin F_{2lpha} (10) wol/1) following sympathetic (guanethidine 10^{-4} mo1/1.phenoxybenzamine 2 x 10^{-5} mol/1) and muscarinic blockade (atropine 5 x 10 6 mol/1) was examined here. TNS applied as intermittent trains of stimuli (9V, 1.0 ms. 1-32 Hz for 30 seconds) produced a frequency dependent relaxation (marimum relaxation, mean 3.28+0.18 SEM g, n=17). This relaxation was not significantly altered by tetrodotoxin (10^{-6}) mol/1), cimetidine, aminophyline, indomethacin, catalase or ascorbic acid. The relexation was not present in rings following cold storage of the caphonous veins at 4 C for ? days. The relaxation was also abolished by matain (2 m 10.4 met 1) and zero-K" "Krebs bicarbonate" buffer solution. Tratecal Jours Sub-threshold (for contraction) con entrations of MA was found to potentiate the response to trains of the (ov 3 H-, '.) me (or 5 econde) to a concentration depredent manner Transform potent offor: many 246.2433.27 SEM. of control). The present study indicator (1) by fground TNS indibits the contractions produced by or services NA. (2) "I contacts a non-adrence gto, the cholinergic . See in which would be not a start by a petrolotoxin registant nervo or $= (-1)^{n} + (-1)^{n$ (nen neus

- - , ,

ł

7. Ŧ

account for the inhibition of NA contraction mentioned above. (3) Background exogenous NA (including sub-threshold concentrations) potentiate the responses to TNS. (4) Thus, there is a significant interaction between the effects of exogenous NA and TNS in the canine saphenous vein.

 $\sum_{i=1}^{n}$

ACKNOWLEDGEMENTS

I would like to thank the following:

Dr. C.T. Kappagoda, my supervisor, for his invaluable advice, encouragement and guidance during both the research and the writing of this thesis;

Dr. D.A. Cook and Dr. A.S. Clanachan for their advice during the research;

Mr. Alvin Todd for his technical assistance and for his companionship during the long hours spent in the laboratory;

Dr. P.M. Vanhoutte and Robert R. Lorenz for providing me with an opportunity to learn the technique of superfusion, at the Mayo Clinic, Rochester, Minnesota. The time spent by Bob Lorenz in teaching me the technique and in helping me set up the same in Edmonton is particularly appreciated;

Paula Priest, Priscilla Chin, Moira McCubbin and Marilyn Coulomb for their belp during the study:

The Alberta Heritage Foundation for Medical Research for providing financial support by means of a Fellowship:

Non-her Lenz and Lolitho Jayakody for their help during

1. 1. 1. 1

. • •

TABLE OF CONTENTS $\hat{\sigma}_{\mathcal{A}}$

- marine a

•

.

	TABLE OF CONTENTS	
	*4	
		PAGE
I	INTRODUCTION	1 -
[]	LITERATURE REVIEW	8
	MORPHOLOGY OF THE VESSEL WALL	10
	Morphology of the canine lateral saphenous vein,	18
	ULTRASTRUCTURE OF SMOOTH MUSCLE CELLS	18
,	Ultrastructure of the canine lateral saphenous vein	23
<i>9</i> 2	STRUCTURE AND CHEMISTRY OF THE CONTRACTILE PROTEINS	24
	Thick filaments	25
	Thin filaments	28.
	Intermediate filaments	300
	REGULATION OF CONTRACTION IN SMOOTH MUSCLE	31
'	Myosin-linked regulation	34
	Actin-linked regulation	35 ,
	Dual-linked regulation	39
	Phosphorylation theory	39
	Leiotonin theory	44
	AUTONOMIC INNERVATION OF VASCULAR SMOOTH MUSCLE	47
	Autonomic innervation of the canine lateral saphenous vein	52
	saphenous vern,	12
	FLECTROPHYSIOLOGY OF SMOOTH MUSCLE	53
	Resting membrane potential	54
	Na ⁺ /K ⁺ membrane pump	56
	Action potentials	50
	Electro-mechanical and Pharmacormechanical coupling.	64
	CATCHUM REGULATION IN SMOOTH MUSCLE	68
	Activator calcium in smooth muscle	69
	Intracellular calcium stores	78
	Calcium sequestration during relaxation	80
	Role of cyclic nucleotides in the regulation of	R /1

		PAGE
	ADRENERGIC NEUROEFFECTOR INTERACTION	89
	Transmitter synthesis and storage	.90
	Transmitter release	95 ·
	Transmitter disposition and termination of	·
	its effect	105
	Diffusion	105
	Uptake ₁ (neuronal uptake)	107
	Uptake ₂ (extraneuronal uptake)	115
	Catabolism	117
	Binding of catecholamines to connective tissue	122
	Relative importance of the disposition pathways.	123
	Transmitter disposition in the canine saphenous vein	123
	Pre-synaptic modulation of transmitter release	129
		127
LII	METHODS	143
	GENERAL METHODS	143
	Preparation of venous strips and rings	143
	SPECIFIC METHODS	147
	Interaction Experiments: Protocol One	149
	Protocol 1.1	149
	Protocol 1.2	154
	Protocol 1.3	159
	Protocol 1=4	160
A	Protocol 1.5	161
	Superfusion Experiments:Protocol Two	163
	Protocol 2.1	166
	Protocol 2.2	173
	Column chromatographic analysis	176
	Relaxation Experiments: Protocol Three	193
	Protocol 3.1.	193
*a1	Protocol 3.2.	e 195
•••ф	Protocol 3.3.	196
	Protocol 3.4.	

	PAGE
Protocol Four	
STATISTICAL ANALYSIS	199
RESULTS	
Protocol One	200
Protocol 1.1	
Protocol 1.2	205
Protocol 1.3	
Protocol 1.4	213
Protocol 1.5	218
Protocol Two	223
Protocol 2.1	224
Protocol 2.2	229
Protocol Three	234
Protocol 3.1	
Protocol 3.2	
Protocol 3.3	251
Protocol 3.4	253
Protocol Four	255
V DISCUSSION	
Calculation of expected contraction i	n Protocol One 262
Influence of sympathetic nerve activi sensitivity to exogenous noradrenal	
Influence of exogenous noradrenaline sensitivity to transmural nerve sti	-
Influence of electrical current on th	
to exogenous noradrenaline	
α_1/α_2 receptor interaction	
Investigation of the possible mediato inhibition observed in Frotocol 1.1	
Role of pre-synaptic α_2 inhibition in in Protocol 1.1	
Relaxation to TNS	····· ?яя
Mechanism responsible for the relaxat	ory response
i se	

/		
		·
•	Site of origin of the post ated inhibitory	PAGE
	neurotransmitter	291
	Possible mediators of the relaxatory response to TNS	293
	Influence of mode of TNS on the relextion	294
	Role of TNS induced relaxation in Protocol 1.1	295
	Considerations for the future	299
VI	BIBLIOGRAPHY	302
VII	APPENDIX I: Drugs and chemicals	331
	APPENDIX II: Solutions used for column	
	chromatographic analysis	335

APPENDIX III: multions used during chemical sympathectomy with 6-hydroxydopamine.... APPENDIX IV.....

336

337

٠,

LIST OF TABLES

مربر

.

1.

١

"

....

١

'ABLE		PAGE
. •	The form used for recording of data during a column chromatography experiment	191
! •	The form used for recording of results during a column chromatography experiment	192
3.	Summary of results of column chromatographic analysis: samples from Protocol 2.1	228
۴	Summary of results of column chromotographic analysis: samples from Frotocol 2-2	233

.

.

LIST OF FIGURES

.

·

FIGUR	Ε	PAGE
1.	Postulated locations for α-adrenoceptor sub-types in adrenergically innervated smooth muscle	6
2.	Schematic diagram of the structural characteristics of blood vessels in mammals	9
3.	Diagrammatic representation of the wall of an elastic artery	11
4.	Diagrammatic representation of the wall of a muscular artery	14
5.	Diagrammatic representation of the myosin molecule	26
6 .	Diagrammatic representation of the structure of the thin and thick filaments	37
7.	Schematic diagram of the phosphorylation theofy	40
8.	Summary of the lejotonin theory	45
9.	Model of the vascular smooth muscle neuro-muscular junction	49
10,	Steps in the synthesis of noradrenaline and adrenaline in the sympathetic nervous system	91
11	Apparatus used for the pharmacological experiments in the present study	2 145
יו.	Summary of the steps in Protocol 1.1	151
רו.	A diagrammatic representation of the method of calculation of the expected contraction in Protocol 1.1	153
14.	Summary of the stops in Frotocol 1.2	155
15.	A diagrammatic representation of the method of calculation of the expected contraction in Protocol 1.2	157
16.	Steps of Protocol 1.4	162
1	A diagrammatic representation of the apparatus used for the superfusion experiments	165
19	Summery of Froteent 2.1.	168

20. Re- ana 21. A difor 21. A difor 22. An 23. Sum 24. The con 25. An 26. Sum 27. The con 28. A r 29. Sum 30. A r 31. Sum 32. Sum 33. Sum	mary of Protocol 2.2 pipette dispensers used for the column chromatographic lysis iagrammatic representation of the glass columns used the column chromatographic analysis example from Protocol 1.1 mary of the results from Protocol 1.1 individual data points for the observed/expected tractions in Protocol 1.2 mary of the results from Protocol 1.2 individual data points for the chserved/expected tractions in Protocol 1.2	175 180 181 202 203 204 203 204 203 204 203 204 203 204 203
ana 21. A di for 22. An 23. Sum 24. The con 25. An 26. Sum 26. Sum 27. The 28. A r 29. Sum 30. A r 31. Sum ago	<pre>lysis iagrammatic representation of the glass columns used the column chromatographic analysis example from Protocol 1.1 mary of the results from Protocol 1.1 individual data points for the observed/expected tractions in Protocol 1.1 example from Protocol 1.2 mary of the results from Protocol 1.2 individual data points for the observed/expected tractions in Protocol 1.2 example from Protocol 1.2 individual data points for the observed/expected tractions in Protocol 1.2</pre>	181 202 203 204 204 207 208 209 201
for 22. An 23. Sum 24. The 24. The 25. An 25. An 25. An 26. Sum 27. The 28. A r 29. Sum 30. A r 31. Sum ago 17. Sum	the column chromatographic analysis example from Protocol 1.1 mary of the results from Protocol 1.1 individual data points for the observed/expected tractions in Protocol 1.1 example from Protocol 1.2 mary of the results from Protocol 1.2 individual data points for the observed/expected tractions in Protocol 1.2	202 203 204 204 207 208 209 209 209
23. Sum 24. The con 25. An 26. Sum 26. Sum 27. The 28. A r 29. Sum 30. A r 31. Sum ago	mary of the results from Protocol 1.1 individual data points for the observed/expected tractions in Protocol 1.1 example from Protocol 1.2 mary of the results from Protocol 1.2 individual data points for the observed/expected tractions in Protocol 1.2	202 203 204 207 207 208 209 209 201
24. The con 25. An 26. Sum 27. The con 28. A r 29. Sum 30. A r 31. Sum 32. Sum 31. Sum 32. Sum	individual data points for the observed/expected tractions in Protocol 1.1	204 201 202 203 211
con 25. An 26. Sum 27. The 28. A r 29. Sum 30. A r 31. Sum ago 17. Sum	tractions in Protocol 1.1 example from Protocol 1.2 maty of the results from Protocol 1.2 individual data points for the observed/expected tractions in Protocol 1.2 epresentative example from Protocol 1.3	201 202 202 211
26. Sum 27. The 28. A r 29. Sum 30. A r 31. Sum ago 17. Sum	mary of the results from Protocol 1.2 individual data points for the chaerved/expected tractions in Protocol 1:2	211 211
27. The con 28. A r 29. Sum 30. A r 31. Sum ago 17. Sum	e individual data points for the cheerved/expected tractions in Protocol 1:2	۱۱ د در ۶
28. A r 29. Sum 30. A r 31. Sum ago 17. Sum	epresentative example from Fritecol 1.3	211
29. Sum 30. A r 31. Sum ago 17. Sum		
30. Ar 31. Sum ago 12. Sum		יוי
31. Sum ago 12. Sum	mary of results from Protocol 1 3.	
ago 17. Sum	epresentative example from Protocy 1 1 4.	· 1 -
	mary of the results from Protocol 1.4 (background mists:tyramine, histamine and methoxamine)	211
	mary of the results from Protocol J.4 (heckground mist phenylephrine)	, ?1 ⁻
33. Sum	many of the results from Protocol 1.5 (propramatal)	• • •
Sun and	mary of the reality from from $p = 1.1.5$ (interaction).	221
۲ <u>۲.</u> ۲۰۱۱	mary of results from Protocol 1. " (astrophylitics)	· · · · ·
16 541	mmary of regulta from firsto of 2.1	· > 4.
37 Sur	more of results from tr to all 1.2	1 11
The		

Congramping Constants Congram

FIÇU	RE	PAGE
41.	Scanning electron micrograph of the intimal surface of saphenous vein rings	, 241
42.	Effect of mechanical de-endothelialisation on the relaxatory response to transmural nerve stimulation	242
43.	Response of saphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$, to transmural nerve stimulation applied "cumulatively" following sympathetic and	~ / /
	muscarinic blockade	244
44.	Effect of tetrodotoxin on the relaxatory response to transmural nervé stimulation	245
4ና.	Effect of cimetidine and indomethacin on the relaxatory response to transmural more stimulation	247
46.	Effect of aminophylline on the relaxatory response to transmural nerve stimulation	248
	Effect of ouabain and zero K ⁺ "Krebs" buffer solution on the relaxatory response to transmutal nerve	
	stimulation	250
48-	Effect of ascorbic acid and catalase on the relaxatory response to transmural nerve stimulation	252
49.	Effect of cold storage of the isolated veins on the relaxatory response to transmural nerve stimulation	254
	Effect of chemical sympathetic denervation with 6-hydroxydopamine on the relaxatory response to transmutal	·
-	norve stimulation	256
51	A representative crample from Frotocol Four	257
25	Cummary of results from Protocol Four	259
	Effect of a radrenalian on tension, total radioacti '' ' super' tate and efflux of "He prodrenaliane and state of the first electric to the lation of coving	
		284

INTRODUCTION

.

The sympathetic nervous system everts its off on the body via the catecholamines noradronaline, advertise unit dopamine (1). Noradrenaline and adrenaline, the predominant neuroliguemitters is the peripheral sympathetic nerves, are released it two majo altest the post-ganglionic sympaticatic nerve endings and the adrenat glands which contain specialised rost gaugiton's neuronal coll bodies that secrete these to bermones. New your ne is the probations ""ectclamine released of the follow offer while e advocative than no Arene 11 ip die secreted of the latter give in vaccily and th "UR(Acres 1. noradronalies and rightalies a set thefe a traine of and that attimulation of high reaction loopt to be month made all membrane. The attant contraction on factate market market as been investigated to the the second title the method of the states in a noredree lies to a state on a the times by the adjoint from • - C mitalle I c sei and a advantine to the transmission oppose and the second second the forther than the forther than the 1999 - N. S. S. S. S. S. S. S. It tops is The second second

trenent Is 11- 1-11 1 i 1 Ing is 1.7 11.1 11.00 and the second second neurotecore · 6 · 10 history and the second second second 10 present in 11 The second se 1 1 11.2 5 a 1 45 **.** . ''' t ··· • • • • · · · · · · · · · · ,

4

1

5

۰.

believed to exert contractile effects in vascular smooth muscle through a single population of α -receptors(3). Evidence emerged in the early 1970's that the a-receptors located at pre-synaptic sites might be pharmacologically different those located at post-synaptic from sites(4.5). Drugs such as clonidine and α -methylnoradrenaline were found to be more effective at the pre-synaptic α -receptor, in causing inhibition of release of noradrenaline from the sympathetic nerve endings than on the post-synaptic arrementor sites on the smooth muscle mombrane mediating muscle contraction. On the other hand, drugs such as methomamine and phenviephrine were far more effective as agonists at the post synaptic membrane, than at the pre-stuaptic membrane. Alpha autogenicia also showed a similar differential sensitivity at the preand post synaptic receptor sites. In order to compare the potency of different Hrogg at the two receptor sites, the effectiveness of both $\dot{\alpha}$ agoniate and resurgenters it the procland post examptic remeceptors was extremed as a catio. This pre/post ratio ared from 0.01 to 100 with different druce fullys for a wid spectrum of differential sensitivity at he two receptores those finition led in the classification of exhaptly inceptors is and the process of receptors as ap-

۲)

Powe is difficulties a non with this erminology; as alphareception low about on the physical characteristics of u_2 denotes a second of the processor inhibition the release of acception pullars. For the average receptors inhibition the release of acception, were found it correspond to retrace and their the location of Wage reception is in a contractive prior to relation to sympathetic neces indicate the forget in all the relation to sympathetic neces indicate the forget in all the relation to sympathetic neces indicate the forget in all the relation to sympathetic nerve

2

system also posed a problem in this pre-synaptic/post synaptic classification of the a-receptor sub-types. Thus it was proposed that the sub-types of the alpha receptors be classified as α_1 and α_2 on a functional basis rather than on an "anatomical" basis(6). Nevertheless, the arreceptors on the vascular smooth muscle cell membrane were considered to be of a single (α_1) type. This concept was questioned by the findings of Bentley. Drew and Whiting (7) in an in vivo study using pithed rate and anaesthetieed cate which demonstrated that two types of a receptors mediated the present responses of the advenergic agonists used: one type was selectively stimulated by phenylephrine and blocked by provisin $(\alpha_1$ thereptor): the other was stimulated by noradrenaline and blocked by phentolamine, but registant to prazosin (", receptor). Further in vivo studies by Drew and Whiting(8), Docherty, McDonald and McGrath(9) and Timmamann, Kwa and Von Zwieten(10,11) provided more definition evidence for the existence of both α_1 and α_2 receptors in vascular smooth muscle. Inter, Do Mey and Vanhoutte demonstrated evidence for the existence of both α_1 and α_2 post sympthy advanceptors in vitro in control subscripts and verts(12).

.

Presible locations for these two types of post symptic a receptors of the endered straigestic in the engression of the discourt studies of the and Weild in the minor pin i torrial sub-momental attraction of the component of a 300 000 µm each of an entertain to the component of a 300 000 µm each of an entertain to the torrial of the transmission of the interthe torrial of the transmission of the interthe torrial of the transmission of the interthe torrial of the transmission of the interent torrial of the transmission of the interation of the transmission of the transmis

and the first second second

. 3

common response observed. At other sites of application, a depolarization similar to an excitatory junction potential was observed. This led to a generalised constriction of the arteriolar segment if an action potential was initiated by this depolarization. The former type of response was abolished by phentolamine but the latter type of response could be detected in the presence of phentolamine. Transmural nerve stimulation applied to this preparation produced excitatory junction potentials: increasing the stimulus strength, increased the magnitude of the junction potential leading to the triggering of an action potential and a generalised constriction. The junction potentials were not reduced by phentolamine, tolazoline or prazosin. Thus, the depolarization and generalised constriction produced at occasional sites by iontophoretically applied noradrenaline seemed to mimic the effects of transmural norve stimulation. A mapping procedure using flu rescence microscopy to visualise the nerves was employed to find the anatomical relationship of the adrenergic nerves to the different sites producing the two types, of responses to iontonhoistically applied noradronaline. This demonstrated that the Sites at which depolarization was eldered by iontrophoratically applied pereduceeline may reprinted to regions close to the summathetic nerves supplying the sus less the changes in membrane potential were detected which the print of application of the noradrenaline was greater than 10 or from a floor-acent nerve. Although a few depolarization responses vero dole ed at regime 5-10 pm distant from a northe, the metority of associative irors acre within 5 1" of a fluoregent proc. Thus the come of the opportunal alpha months located at the averathetic and the state of a second s

4

nerve endings (i.e. outside the synaptic cleft) emerged (Fig. 1). However, Hirst and Neild concluded that the intrajunctional receptor may not be an α -type receptor because of its resistance to α -antagonists.

Based on the above findings it could be postulated that endogenous noradrenaline (released at the sympathetic nerve endings into the junctional cleft) would exert a greater portion of its effects through intrajunctional a-receptors and exogenous noradrenaline would exert a greater portion of its effects through extrajunctional a-receptors. In isolated vascular smooth muscle experiments in tissue baths, endogenous and exogenous noradrenaline would be equivalent to transmural nerve stimulation (assuming a predominantly sympathetic innervation) and noradrenaline added into the bath from outside respectively. Although the individual effects of transmural nerve stimulation (TNS) and exogenous noradrenaline in isolated blood vessels have been extensively investigated, not much information is available about the interaction between the two "effects. This interaction has been studied with respect to the pre-synaptic inhibition produced by exogenous noradrenaline on the release of endogenous notadrenaline via TNS(15) However, little attention, has been paid to the post-synaptic expect of this interaction(16). This is probable due to the fact that the differences in the mediation of the responses of the two water of establishing in the protect number of the and apparent only buring the last free concerns i'l the Heconom of both of and as receptors in amosth muscle.

The present furestigation cas undertaken to study the interaction between a openanc notadronalific and ransmunal merge at interaction in "anoular amoth messle; different concentrations of more con-

. 5.



Figure 1. A diagrammatic representation of the postulated locations for readrenoceptor sub-types in adrenergically innervated smooth muscle. Frequenctional a receptors located on the axonal membrane are believed to be α_2 in type. Post junctional arreceptors are of two types (1) α_1^{-1} receptors believed to be located at the synaptic cleft i.e., intrajunctionally. (2) α_2 receptors: believed to be located away from the arconditional for the extrajunctionally. whole range of the dose-response and stimulus-response curves. This permitted any progression of the interaction to be studied. Canine lateral saphenous veins were selected for the present investigation for the following reasons:

- 1. the canine saphenous vein has been shown to possess a good sympathetic innervation with nerves penetrating almost up to the tunica intima: refer to literature review for details(17,18).
- the presence of post-synaptic α₁ and α₂ receptors have been demonstrated in the isolated canine saphenous vein(12).
 the isolated saphenous vein responds well to both exogenous
 - noradrenaline and TNS.

LITERATURE REVIEW

The vascular system can be divided into the arteries, the veins and the microvasculature(19,20)(Fig. 2). The arteries are generally subdivided into two categories, the elastic arteries and the muscular arteries. Elastic arteries contain many elastic laminae in their walls and generally have large luminal diameters (e.g. aorta, carotid Potential energy, stored during cardiac contraction in the arteries). elastic tissue of the aorta and its branches is reconverted into kinetic energy for the circulation during the diastolic phase. This elastic recoil of the vessels sustain the pressure head better and renders blood flow to the periphery steadier than it would otherwise be. Muscular arteries which are formed by branching of elastic arteries have less muscle cells in their walls. tissue and smooth elastic more arterioles, the pre-capillary sphincters, the Collectively, the capillaries and the post-capillary venules are referred to as the The arterioles which are 0.1-0.8 mm in diameter in microvascular bed. the dog(19), have the smallest lumen: wall ratio (approximately 0.4) among all blood vessels and thus contribute most to the resistance to These pre-capillary resistance blood flow in the vascular system. vessels (arteriales) usually exhibit an efficient local myogenic control of their own vascular radius, and on this myogenic tone is superimposed an extrinsic neural control effected by autonomic nerves. The precapillary sphincters, themselves part of the pre-capillary resistance vessels, are particulary important in determining the size of the. capillary exchange which is perfused at any moment in the tissue: an-----increase in the potency of the sphincters causing an increase in the number of capillaries open. The capillaries function as exchange



Figure 2. Schematic diagram summarising the major structural characteristics of principal segments of blood vessels in mammals [Reproduced with permission; from Rhodin JAG, Handbook of

Physiology(20)].
vessels where exchange of nutrients and metabolic waste products between the blood and the tissues take place. The veins are generally divided into large and medium-sized veins. They tend to have the greatest lumen: wall ratio among blood vessels and these offer little or no resistance to the flow of blood. They also possess valves to prevent back-flow of blood within the venous system. The veins or capacity vessels are important sites of change in the capacity of the vascular system. Changes in luminal configuration (from elliptical to circular cross-sectional profiles) and changes in myogenic tone of the veins induced by sympathetic constrictor nerves are of great importance in adjusting the capacity of the vascular system, particularly in postural changes.

MORPHOLOGY OF THE VESSEL WALL(20)

The walls of blood vessels (with the exception of the capillaries) consists of three layers: tunica intima, tunica media, tunica adventitia (Fig. 2, Fig 3). The tunica intima, the innermost layer of the wascular wall is composed of the following structures:

- a basement membrane 80 nm in thickness
- 3. the sub-endothelial layer composed of collagen fibres, elastic fibrils and smooth muscle cells; this sub-endothelial layer is usually present only in the large elastic arteries.

The endothelial cells are flat and elongated with their long axes parallel to that of the blood vessels. They are approximately 10-20 μ m in length and 5 μ m in width at their widest point. They have a thickness of about 0.2-0.5 μ m with a slight bulge at the region of the bucleus. Their morphology can be best studied by pressure-perfusion


Figure 3. A diagrammatic representation of the wall of an elastic artery with well organised elastic laminae in the medial layer (turica media) [Reproduced with permission: from Rhodin JAG, Hardbook of Physiology(20)] with the fixative at a pressure equal to the normal intravascular pressure in the vessel studied. Collapse and wrinkling of the endothelial layer takes places unless pressure-perfusion is used during fixation. The endothelial cells are bound to each other by tight junctions(zona occludentes) and communicating junctions(gap junctions; maculae communicantes). The tight junctions are areas where the opposing endothelial cell membranes have fused along ridge-like protrusions of the individual cells. In section, these appear as punctate fusions of the two cell membranes. The gap junctions are patches (maculae) of opposing cell membranes consisting of a polygonal lattice of cell membrane sub-units. They function as sites for cell-tocell transfer of ions and metabolites (ionic, electrotonic and metabolic couplings)." Both these types of junctions are more common in arterial than venous endothelial cells.

•**___**__.

The sub-endothelial layer, as stated before, is present only in the large elastic arteries. In the human aorta it undergoes a series of changes during life. At birth it is a thin structure with a narrow layer of connective tissue fibres. In voing adults it increases in thickness and becomes fibrous and cellular by middle age. In senile subjects the layer is thick, fibrous and hyalinised. In some species (e.g. pig, man) smooth muscle cells are present in the sub-endothelial layer of the aorta. In small arteries and arterioles (where the subendothelial layer is lacking), endothelial cells, and smooth muscle cells in the media are closely connected through myo-endothelial junctions. These are cytoplasmic processes from the endothelial cells or the smooth muscle cells and they penetrate the basal lamina and if present, the internal elastic lamina. There are few of these processes

in small arteries but they become increasingly numerous as the arterioles approach their terminal ramifications and the pre-capillary sphincters.

"The tunica media is the middle layer of the vaseular wall. Smooth muscle cells are the principle constituent of the media but it also contains a varied number, of elastic laminae, collagen fibrils and elastic fibrils(Fig. 3, Fig. 4). It is usually bounded by the internal and external elastic laminae and is thicker in the arteries than in the veine. The tunica media of the human aorta contains about 40.60 fenestrated elastic laminae. These elastic laminae which are shout 3 µm in thickness are concentrically arranged and spaced equidistantly. They are interconnected by a network of elastic fibrils and this highly structured and elaborate elastic framework gives the media its great resilience and strength in the elastic arteries. The smooth muscle cells are found within this framework. The elastic laminae diminish in number and in organisation, the smaller the size of the artery with arterioles generally lacking elastic laminae. Most of the smooth muscle cells in the media are ordented obliquely funning diagonally at small angles between the elastic lominae.

The smooth muscle cells in the matta tend to form a artical although the orientation may vary with the distonling forces(21). Welinsky and Glagov(21) in a study of the rabbit ports fixed helps distortion pressure, showed that the smooth muscle cells are oriented chliquely or even perpendicularly with respect to the elastic laminae. However, if the preparations were fixed at pressures equal to be greater than the diastolic pressure, the smooth rack while other production is greater than the



14





Figure A diagnam the reprisentation of the will strugged at a strugged at the will strugged at the strugged at

of the helix can vary depending on the contractile status of the vascular wall. It is conceivable that in a muscle spiral with extensive mobility, the pitch of the helix may change considerably and the direction of the muscle cells may alter from longitudinal to diagnomal to perpendicular. It to the shared deriveding on the functional state of the block percent.

This apparent below as sugement $\widehat{\mathbf{v}}$ the smooth models calle in the media in the basis in the belically out it is first introdue at by Furchy it a ' Bhadra on the first'. The helf-al cutting would wield atrips that prove a considered lie which were extended more on loss parallel to the consistence (is which were extended more on loss the smooth multiple in the set of the dimension of the smooth multiple in the set of the dimension of the smooth multiple in the set of the dimension of the smooth multiple in the set of the dimension of the smooth multiple in the set of the dimension of the dimension of the dimension of the dimension of the set of t

A STAR NO

...

man to the area call of the test ۰. şt nodia(Fig. 11 and the second 2 Pro 1 1 1 . · · · · a transformer and the T to a second the property of the property of the second sec 4.1 S I have a set of the • • • • • • • • • • . • Max I C . all for the second . . 1. 1. A. 1. A. 1. · · · $(t_{i,j}) \in [0, 1]$

•

some medium and large sized veins, smooth muscle cells are present in abundance although these cells are often located outside the zone that is classically regarded as the media (i.e. in the sub-endothelial tissues and in the tunica adventitia). However, as the external and internal elastic laminae are not present in veins, the distinction between the intimal, medial and adventitial layers is not as clear as in arteries. The smooth muscle cells in veins are arranged longitudinally and circularly as well as helically in different types of veins. Thus, the rat portal vein contains two distinct layers of smooth muscle cells, an inner, narrow sub endothelial layer with circularly oriented cells, and an outer, wider layer with longitudinally arranged smooth muscle colle(25). In the boying mesenteric vein the contractile response of a longitudinally out strip to transmiral nerve stimulation was about 20 times that of a vircularly cut strip indicating the predominance of longitultually arranged smooth miscle cells in these veine(26). The salls of motion contraits threadout collagen fibred separating individual th man's cells as we'l at lavers of smooth muscle.

The Ma the stille to the enternost layer of the vaccular wall. The a had a barren of the in a contractionable depending on the type and the still of the big there is a new transfer the coupling blood vessels theory of a attraction the theory to a net s of dense fibre Agents of the second of the second second send of the blood vessel wall and " attend - court and trips the 'ssue, together the to sporth muscle cells in ref and to an the sina garagement. alvortitia except in large when which outsin longitudinally and smcath 10 States and the second nisegeog in nerves. 1 2 22 1.5 1.1 100 11 1 1 1 1 in the - 1 g

.

for sensory organs, muscles and visceral organs 2) vasomotor and sensory nerves for the vascular wall itself. The adventitia functions to anchor the blood vessel to its surrounding tissues by way of loose connective stasue. In the elastic arteries, the adventitia tends to be thin especially in the larger vessels and contains a loose network of elastic The tunica adventitia tends to be wider in the muscular fibrils. arteries often occupying half of the vessel wall. The adventitial layer is thin and inconspicuous in arterioles and venules merging with the surrounding connective "issue. This layer is best developed in mediumsized and large veins comprising up to 75 per cent of the vascular Here the adventitia is made up of collager bundles, elastic wall. fibres and smooth muscle cells. The collagen bundles are arranged in a belical fashion forming an intimate relationship with the smooth muscle cells in the adventitia. This relationship and the firm attachment of the adventitia to the purrounding round the times allow the adventitial layer to play a more direct of the second of its surface to ates an compare with the stories.

···· . ·

Top Tortes comprises a force action of an endered abort (750 pm tobor ofth tota and arrian tona which in commence in relation to the longt' the orithment takes The fotal copilized -whong outfor and the control through the weto's farball of the bo 15: 600 square motore. The at the sure have a grane we streated to which all systemic conflicted is not which is loost a are an thely "no cent fille that and That . 1 10 try and the same plant press of 1) 1.1.1.1.4 - 1 g 111. . · · · 57 To 5 T and tr . . .

المحادثة والمحاور والمحاور والمحاور

associated with these vessels: the pericyte is neither a smooth muscle cell nor a fibroblast. Structurally, it is surrounded by an external lamina, and it contains a small number of micro-filaments in its cytoplasm. It could represent a potential smooth muscle cell or function as a phagocytic cell.

Morphology of the canine lateral saphenous vein

Most of the studies on the histology of this vein has been done by Osswald & Cuimaraes(17). This is a medium sized vein 3-5 mm in diameter containing an abundance of smooth muscle fibres. The tunica intima in this vein is made up of a single layer of endothelial cells and the basement membrane, with no definite sub endothelial layer. The tunica media having a thin, discontinuous elastic lamina as its innermost laver is relatively thick containing 8-12 layers of smooth muscle These smooth muscle cells are mostly circularly oriented cells. although there are longitudinal as well as oblique fibres. This predominantly circular arrangement of the smooth muscle cells accounts for the very much greater tension developed by ring preparations of the yos al compared with holdenly out strips. The smooth muscle fibres of the soft wall were found to make contact with each other at their ends and also laterally of though spaces of variable width containing collagen and tenuous elastic fibrils were often found between the muscle cells in the above studies. The tunica adventitie in the arthonous very is well developed with numerous elastic fibron.

ULTRA STRUCTURE OF SMOOTH MUSCLE CELLS (28, 29, 30, 31, 32)

Smooth muscle cells have an elongated but irregular shape with multiple cellular protructions. They are a 40 100 μ m in length and 2-5 μ to the character of the amplies rempared with cardiac muscle cells which.

are 10-20 μ m in diameter but approximately of the same length. The smooth muscle cells have a single nucleus situated in the middle at the widest portion of the cell body. Each cell is surrounded by its plasma مده ريار membrane (80 A in thickness) and there is no protoplasmic continuity between cells. The plasma membrane and the basement membrane, when the latter is present, constitute the sarcolemma. The presence of a basement membrane is regarded being one of the as distinct ultrastructural diagnostic features of smooth muscle, and it has been demonstrated in a variety of blood vessels. In fight junctions between smooth muscle cells, the fused plasma membranes of adjacent cells are devoid of basement membranes(31). The small intercellular space between smooth muscle cells contains blood vessels, nerve fibres, extracellular matrix and reticular fibres. Connective tissue cells are rarely found in the interstices of the sheets of smooth muscle. These muscles do not possess a distinctive tissue "capsule" or perimysium characteristic of skeletal muscle fibre(33).

and a second second

۰,

The plagma membrane of smooth muscle cells contains multiple small flask shaped invaginations termed caveolae or surface vesicles. These are 50.80 nm in diameter and are also present in endothelial cells and fibroblasts. The entracellular space continues into these coveolae as demonstrated by the fact that extracellular markers such to colloidel Lanthanum (La¹¹) and tan to acid enter these weathlos(12,34). These use not randomly acattered over the cell surface but tend to have a longitudinal intentation in empoth muscle(32). These caveolae increase the surface membrane area for cell volume by 25.70 per cent. Nevertheless, they are far less extensive compared to the transverse (1) tubular system in shalets, and cardiac muscle. Receive observations (1) 19

. . ..

the mouse coronary artery indicated that these surface vesicles may occur singly or in more complex forms that resemble chains of beads fused end to end. These beaded tubules may be composed of as many as 17 the end and an end of the mathematical and the record vesicles and they often extend deep into the cells(35). Smooth muscle lacks T-tubules and some investigators believe that caveolae are analagous to the T-tubules although no conclusive evidence is available . at present. The surface vesicles may not be fixed structures and their distribution may reflect the functional status of smooth muscle(36). Occasionally in electron micrographs, some of the surface vesicles appear to be intracellular and have therefore been described as However, the penetration of extracellular markers into pinocytotic. these vesicles after fixation and the examination of tilt pairs of micrographs indicate their true extracellular nature(32). This artifactual intracellular appearance is probably due to their narrow necks which may not "enter" into the plane of section. Nevertheless, the possibility that they may become pinocytotic under some conditions cannot be definitely ruled out. Both mitochondria and sarcoplasmic reticulum have been observed to be closely associated with the surface vesicles, with the intervening distance being as small as 3-6 nm in some cases(34). However, the association of the sarcoplasmic reticulum with these surface vesicles is far less common compared with the association. between the sarcoplasmic reticulum and T-tubules in skeletal muscle.

The sarconlasmic reticulum in smooth muscle, as in striated muscle can be divided into the rough endoplasmic reticulum, with its associated ribosomes and the smooth endoplasmic reticulum. These are continuous with one another and also with the nuclear membrane, but not with the lumer of the golgi apparatus. Extracellular markers such as colloidal

lanthanum, horse radish peroxidase and ferritin do not enter the sarcoplasmic reticulum confirming their true intracellular nature(34). The sarcoplasmic reticulum of smooth muscle is not as well organised as However, the tubules of the reticulum are in striated muscle. distributed throughout the cell and they approach the sarcolemma forming The two membranes may be as close as 10-12 nm at surface couplings. these surface couplings and this intervening space has been observed to traversed by electron-dense bridging structures which have a be periodicity of 20-25nm (34,37). The sarcoplasmic reticulum also forms close couplings with the surface vesicles (29). It is thought that the twitch contractions of vascular smooth muscle, triggered by action potentials are mediated via the release of calcium from sarcoplasmic 5.0 Rettor reticulum at these couplings(38), but no conclusive evidence is available present. The volume of the sarcoplasmic reticulum shows a at Significant variation between different types of smooth muscle(39). In the rabbit portal-anterior mesenteric vein and the taenia coli it constitutes only 2 per cent of the cytoplasmic volume. On the other hand, in the rabbit main pulmonary artery and aorta it amounts to 5.0-7.5 per cent of the total cell volume(33,38). There is a direct correlation between the volume of the sarcoplasmic reticulum and the 1 n calcium-free ability of smooth muscles to contract solutions(32,33). The sarcoplasmic reticulum is regarded as a major calcium source and sink (i.e. a sequestration-site) in smooth muscle. Strontium has been used to demonstrate the ability of the reticulum to accumulate divalent cations as it is more electron dense than calcium due to its higher atomic number(37). The electron-dense deposits can be identified as strontium by electron probe analysis.

21

, ¥

The mitochondria in smooth muscle were previously considered as a possible source and sequestration site for calcium in smooth muscle. They too are often closely associated with the surface vesicles, a 4-5 nm distance separating the two membranes. Recent studies, however, fail to show, any "evidence for a role for mitochondria in the physiological (as opposed to pathological) regulation of cytoplasmic calcium. In the presence of Mg^{2+} , isolated vascular smooth muscle mitochondria(39,40) have a rather low affinity for Ca^{2+} (apparent Km = 17 µmql), that is inconsistent with the requirements of a physiological relaxing system. However, in damaged muscle fibres, massive mitochondrial calcification in the form of granules has been readily demonstrated(41). In these cells intracellular Na⁺ and Ca²⁺ were high and the intracellular K⁺ low, suggesting that mitochondrial calcium loading may have been due to the abnormally high cytoplasmic Ca²⁺ caused by cell damage. Smooth muscle cells contain an elongated 'ellipsoidal nucleus

containing one or two nucleoli and a double nuclear envelope in the relaxed cells. However, the nucleus becomes highly compressed and convoluted in contracted cells(36).

A variety of cell junctions are present between smooth muscle These structures, are believed to play a role in ionic and cells. metabolic cell-to-cell communication and are also present in other excitable cells such as cardiac and nerve cells and in non-excitable " cell[®] such 88 endothelial cells, liver parenchymal cells. and fibroblasts(32). Gap junctions or nexuses having a 2-4 nm gap between the outer leaflets of the opposed cell membranes can occur between two parallel cell membranes, between opposed cell protrusions and between invaginations of one cell into its neighbour. The entire width of the

junction (including gap and membranes) is 15-19 nm. Extra-cellular markers such as horseradish peroxidase and colloidal lanthanum penetrate the gaps in these junctions. The second type of junction that is believed to be present in smooth muscle cells is the tight junction or zona occludens which is not penetrated by these extracellular markers. Freeze fracture studies of the tight junctions show a strikingly different appearance from gap junctions. They show a meshwork of ridges with furrows on the complementary face as compared with the 8.5 nm drameter particles and corresponding pits seen in freeze fractured gap junctions(32). Other types of junctions such as intermediate contacts (attachment plaques), septate junctions and simple appositions are also found in smooth muscle. In electrically excitable cells, gap junctions provide a low-resistance pathway for the spread of depolarization throughout the tissue. In smooth muscle, although all cells appear to be electrically coupled to one another, the extent of this coupling varies widely from tissue to tissue(42,43). The structure responsible for this coupling is also not established but the gap junctions are favoured by many workers as the evidence for their involvement in ionic coupling in other tissues is very strong(42,43). The gap junctions are also believed to participate in metabolic coupling allowing transfer of small molecules such as amino acids, sugars and nucleotides from cell to cell.

Ultrastructure of the canine lateral saphenous vein

Electron microscopic study of the tunica media of this vein(18) demonstrated that the numerous smooth muscle cells present contained many short projections and indeptations on their surfaces. The projections from ediacent or the sector opposed (comber cell

junctions. At these junctions, the intervening space was approximately 20 nm in width and the opposing sarcolemmal membranes often demonstrated an electron density. The cells contained many oval shaped surface vesicles 80 nm in width and 90 nm in length. The interstitial space between the smooth muscle cells contained a variable amount of collagen fibrils, an amorphous filamentous background rich in 10 nm thick microfibrils and rare elongated fibrocytes.

1. •

STRUCTURE AND CHEMISTRY OF THE CONTRACTILE PROTEINS

As in skeletal and cardiac muscle a sliding-filament mechanism is believed to operate in smooth muscle contract fon. The elements of the contractile apparatus include three types of filaments (thick, thin and intermediate), dense bodies and attachment plaques. When isolated smooth muscle is observed to contract, small closely spaced blebs form on the cell surface interspersed with undistorted regions(44). It is believed that the latter are sites 🗣 attachments of the contractile elements to the sarcolemma and have been named attachment plaques. Dense bodies, too, are similar structures but dispersed throughout the sarcoplasm and believed to be intracellular attachment sites for the contractile elements analogous to the Zolines in cardiac muscle. αactivin similar to that found in skeletal muscle has been isolated from some types of smooth muscle. In skeletal muscle amactinin is believed to form part of the 7-line structure. In smooth muscle a-actinin has been localised in the dense bodies (using an antibody technique).

Smooth miscle cells contain 3 types of filaments in their contractile apparatus: the thick filaments (13-20 nm in diameter), the thin filaments (5.8 nm in diameter) and the intermediate filaments (approximately 10 nm in diameter). The force generating apparatus in

a service a

and the second second

smooth muscle is organised less intricately than in cardiac muscle, in that highly structured parallel arrays of interdigitating filaments characteristic of cardiac muscle are not seen(45). Rather, one finds throughout the cytoplasm, large numbers of filaments arranged in directions roughly parallel to the longitudinal axis of the cell. When the muscle is relaxed, the filaments tend to lie parallel to the longaxis of the smooth muscle cells. However, when the muscle cells contract, the filaments tend to assume a progressively oblique direction to the long axis. At maximal contraction the angle between the long axis of the cells and the filaments may be as much as 25-40°(46).

ويتجرب والمراجع والمتعا والمعاجم والرارين والمراجع

Thick filaments

The thick filaments are fibrous(α -helix) in structure and contain protrusions with globular heads along their length (Fig. 5). They are comprised of upwards of 200 myosin molecules (relative molecular mass:470,000). The myosin molecule can be enzymatically cleaved into two parts(29).

- 1. Light meromyosin (LMM, relative molecular mass: 150,000) this is a fibrous protein which forms the "tail" of the myosin molecule (150 \pm 20 mm in length) and lines up in a clustic with other similar molecules to form a thick filement.
- Heavy moromyosin (1994) this appears as protructions from the thick filaments. These protructions form the cross buildes between the thick and thin filaments during contraction. The heavy meromyosin can be further sub (ractionated into a fibrous S-2 segment (relative molecular mass: 60,000) and a globular " I segment contrativing the two heads of the myosin molecule (? y 120,000). The " J maginal is comprised of two identical outer.

25 🖂

\$



total length of myosin molecule ~1,400 Å

Figure 5. Diagrammatic representation of the myosin molecule composed of the light meromyosin (LMM); heavy meromyosin subfragment-1 (HMM S-1) and heavy meromyosin sub fragment-2 (HMM S-2). The arbelical portion of the molecule is dericted as a rope-like structure. At the origin of the cross-bridge, i.e., one end of the HMM S-2 molecule, it is not known whether the arbelical structure is retained (dashed line). The position of the four light chains (shown as the smaller finder in the globular head) is completely arbitrary. Conformation of the globular head differs from the rest of the molecule in that it is not predominantly an α helix, and this difference is remented diagrammatically. [Reproduced with permission, from Hartshorn: Di and Gart to A, Handbook of the top (GD)]. each containing an attachment site for actin, an enzymatic site that hydrolyses adenosine triphosphate (ATP) to liberate energy used up for the contraction, and two "myosin light chains" that are involved in the function of these segments. Although striated muscle also contains myosin light chains, there are distinct differences in structure and function from those of smooth muscle. The light chain composition is to a degree characteristic of a particular myosin. Smooth muscle myosin contains two light chains of relative molecular mass 20,000 and two light chains of relative molecular mass 17,000 (one of each in the two heads); myosin from cardiac muscle her two classes of light chains of relative molecular mass 27,0 then two classes of light chains of relative molecular mass 27,0 then two classes of muscle type in its light chain composition.

and the second second

The heavy meromyosin and the light meromyosin appear the form a flexible "hinge between them. The fibrous heavy meromyesis functions to transmit the force generated by the operational site of the mosin molecule, to the light more youin. Tweein mole also are accombled into fliaments as a tail to tall biopelar "gment at the contre, and extende to a bead-to tail fights on of the edde of the central of strigted must be (19). Thus the sign consist of the prophy extremely in the best of the stricture mutual elle is crossing to the the the start (it note entruscient). In smooth maging content here the here are be upgational domination of a star of the or the star 1 1 1 1 1 1 that form to the form of the solution of the advantation of the ' 1 t testa Identifial, . .

smooth muscle, myosin molecules are packed so that their heads are oriented in only one direction: i.e. each half or face of the filament has the same polarity (unipolar arrangement)(28,47).

Thin filaments

The thin filaments contain the contractile proteins actin and tropomygain in smooth muscle. Troponin present in striated muscle as a protein which regulates contractile activity, is absent in smooth mus le. Actin, the major component of the thin filament, is present in all muscale cells and in almost all sukaryotic cells. It is therefore not surprising that there appears to be little lifference in the chemical composition of actin from different sources. The thin filament is composed of a double-strauded helical filament (Filamentous/Fibrous notto. Flastin) with an avial repeat of about 36-38 nm(32). The Fractin is a polymer of the (globular) Gractin molecules (5.5 nm in diameter). The is the motocule is a single chain protein with a relative molecular mass of 2,000 the SI subfragment of the heavy merchayosin (i.e. the) glob lar berde of the mostin molecule) binds to the thin filaments to form cross ridges. Tay functional roles suggested for the actin in the contractile concess as : (1) the transmission of force delivered by the nor nont of the same bridge of the monath molecule. I contraction of the result of the ATE are to strage frie energy (a section of the unter Control Ite Mertine ne to Smooth Prested

terminal like as front is indicated in procession of types of anomal parts. Although all states that there is an antireport of the main of the different of the theory those are accorded on the terminal terminant is the

· · · · · · · · · · · ·

12.1

۰.

to Practin(48). Tropomyosin forms an alpha-helix lying in the two grooves of the double-strand of F-actin and is composed of tronomyosin polecules (relative melecular mass: approximately 70,000 The molar stoichiometry of actin to tropomyosin (approximately 7:1 is cimilar in all types of marks. Because of the absenta of tropomy is for rion of the molar is another because of the absenta of tropomy is for rion

•

In reneral, amoth more is contained in market the objectal manufactor. The myorin content is an other solar is in the object 20 mg per griell verification on parel is fith 12 mg per griel of solar for skeletal more of the investor opered is distered muchly. It is the content is higher do related is the class of the object of the figure higher do related is the class of the object of the figure higher do related is the class of the object of the figure higher do related is the class of the object of the figure higher do related is the class of the object of the figure higher do related is the class of the figure of the figure higher do related in the figure of the figure of the figure higher do related in the figure of the figure of the figure higher do related in the figure of the figure of the figure higher do related in the figure of the figure of the figure double of hy month me is in the figure of the figure of the figure is related by month in the figure of the figure of the figure is related by the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by the figure of the figure of the figure of the figure is related by

at the second second

the group are private to be defined on the second of the 1. the company of the Section of the sector p } . and provide the first second second 1 en la seconda de la seconda the state of the second s · · · المركزة بي المراجع الم . . 1 . . + **1** . . .

'h (···

н

.

smooth muscle could result from such an arrangement of cells whose individual ability to develop force is more in keeping with their myosin content(48). The oblique arrangement of the filaments during contraction in smooth muscle may also offer a mechanical advantage of suggested by Rosenbluth(50).

A longer contact time of the encode bridge with actin during the cross-bridge cycla

Intermediate filaments

The intermediate filamente are the third type of filamentous struct is found in smooth mugcle, and have a diameter of approximately 10 op These filments are generally ensociated with dense bodies, often uniounding them. In transverse section they appear hollow. "Inital fitments have been chaerve! in a variety of non-muscle cells, o a gliat filamenta, filamenta of fibroblasts and endothelial cells. The intermediate filements are not attached directly to either thick or thin filaments and do not enpear to play a direct role in the contractile process. They are discorred throughout the sarcoplasm and are believed to play a sole in the gell architecture as a cytoskeleton is the hidgen among dince hold a. The intermediate filaments are nade part restain need acaletta (relativo molocular maeas 53,000) their men are furieger to a oral growth macle fibres and growd out (1) bound of (2) (1) (1) the followed are boliored the second second to anallable of . .

actinin has been localised in the dense bodies, using an antibody technique. The dense bodies form an anchor for the thin filaments in smooth muscle and are believed to serve as an equivalent to the 7-lines of striated muscle.

 $\langle \rangle$

REGULATION OF CONTRACTION IN SMOOTH MUSCLE

In 1954, A.F. Huxley & Niedergerke(51) and H.E. Huxley & Hanson(52) proposed independently that the shortening of striated muscle was 'the result of a relative sliding between two sets of filaments. This method shortening known as the sliding filament mechanism, firmly of established in stricted muscle, is also believed to operate in smooth muscle contraction. The two sets of filaments have been identified as the thick and thin filaments. Noither filament type altern in longth during shortening of the muscle, the change in length of the muscle being achieved by varying amounts of overlap between the think and thin filamenta. The thick filaments are composed of moster melecules arranged in a manner such that the ensumatically active portion of the molecule motrudes from the body of the filament. Their motividing portions, known as the crock-hride s, hind to the actin of the thin filments during contraction to form after of concise for ly ont. This interaction between the actin and mostly to responsible to the tentor and termentify trigted type? preases An in distributions to security and a distribution of standing in month much to the state state state of the (Mp IT) The Mp In It to be being a dertage marches are appendent to the our in the education of the spherice offer Alline of the factorial to the case transing to the east on the CI of the set to action A term of a plan been granted as a second • • • · · · ·

correlation with the enzyme activity(29). Vascular smooth muscle has a shortening velocity which is very much less than in skeletal muscle suggesting a marked difference in the actomyosin-ATPase activities in the two muscle types. The velocity of contraction is believed to be a function of the cycling rate of the cross-bridge. A cross-bridge cycle consists of the attachment of the bridge to the actin molecule, the sliding of the two filaments, detachment of the cross-bridge and the re-attachment of the bridge at a different site on the thin filament. On the other hand, the force generating ability of smooth muscle is equal to or greater than that in stricted muscle as explained under Structure and Chemistry of the Contractile Proteins. The force of contraction (as opposed to velocity of contraction) is believed to be a function of the number of simultaneously active cross-bridges between the thick and thin filaments, and the efficiency of the transduction of the force generated by a single cross-bridge to overall force generation by the muscle.

Apart from the lower magnitude of the Mg^{2+} -ATPake activity in smooth muscle there are other in vitro features of this enzyme which are different in smooth muscle as compared with striated muscle(53). The smooth muscle inzyme requires a higher concentration of Mg^{2+} to generate marinum enzyme requires a higher concentration of Mg^{2+} to generate northous enzyme requires a higher concentration of Mg^{2+} to generate the uncentration of the easen for this is not vet clear. Another the uncentration of myosin. At low concentrations of the enzyme the rate of primet formation follows a linear dependence on the concentration of the enzyme in skeletal muscle. In smooth muscle this decendence is not linear. This effect may be related to the solubility of the note and the transmission of set measing the proteins are detendence is not linear. This effect may be related to the solubility of the note and the concentration of set measing the proteins are detendence is not linear. This effect may be related to the solubility of the note and the concentration of set measing the proteins are protein forms a precipitate that has a higher ATPase activity. Finally the Mg²⁺-ATPase activity of smooth muscle is less affected by an increase in ionic strength in the surrounding medium (e.g. addition of KC1) than that of skeletal muscle.

As in striated muscle, an increase in intracellular (sarcoplasmic) free ionic calcium (Ca²⁺) leads to contraction of the muscle cells and a decrease to relaxation. The increase in intracellular Ca²⁺ is regarded as the final common step in initiating contraction. Furthermore, the ionic concentration of Ca²⁺ required for activation are comparable in the two muscle types with about half-maximal activity at a Ca²⁺ concentration of about 10^{-6} mol/1.

Recently, free calcium indicators have been utilised to study the relationship between intracellular ionic Ca²⁺ concentration and the contractile state in smooth muscle cells(54,55). The bioluminescent photoprotein aequorin was used in these studies. Aequorin, first isolated in 1961 from the jellyfish aequorea forskalea (aequorea aequorea) is a calcium-activated photoprotein that emits light when it binds Ca²⁺ ions(56). Morgan and Morgan(55) using a microinjection technique to load vascular smooth muscle cells from Amphiuma tridestvlum with acquorin demonstrated that contractions caused by exogenous drugs as well as electrical stimulation were associated with onlanced light responses which preced d the testion responses. The light reprise to electrical estimulation should a supported perfect which appeared to return to the local lovel by the time to tension reach is peal. However, enhanced leading of the colle with security implying the colle Experiperion to the compound) demonstrated a conflex, guarafied the second second the second second

The activator Ca^{2+} for contraction in smooth muscle comes from either intracellular or extracellular sources and these sources seem to differ between smooth muscle cells from different vessels and between different agonists used to initiate contraction. At present there is considerable controversy regarding the sources and sinks (i.e. sequestration sites) of Ca^{2+} in smooth muscle(30).

Although it is well established that increases in intracelfular free Ca^{2+} leads to contraction of smooth muscle cells ultimately by activation of the actomyosin-ATPase, how Ca^{2+} effects this activation remains controversial. The molecular mechanisms by which Ca^{2+} regulates actin-myosin interaction have been examined in a variety of muscle types, e.g. vertebrate and invertebrate striated and smooth muscles, as well as in various non muscle motile systems e.g. platelets. Based on these studies a number of different regulatory mechanisms have been identified. These mechanisms can be divided into 3 groups:

1. myosin (or thick filament) linked regulation

2. actin (or thin filament) linked regulation

3 dual (or actin and myosin linked), regulation

Myosin-linked regulation

The myosin linked regulatory system can be divided into two subtypes(29).

Regulation by myosin, with Ca^{2+} acting as a disinhibitor, e.g. molluscan emoth muscle(29). The regulatory system in this Qmuscle lies in the myosin light chains (relative molecular mass: 20,000) which inhibit the activation by actin of the actomyosin-ATFash. This inhibition is removed in the presence of Ca^{2+} long. However, if the myosin light chains are detached from the

ethyleneglycol-bis(βtreatment with myosin molecules by aminoethyl ether) -N',N',N',N'-tetraacetic acid (EGTA, 10 mmo1/1) the muscle cells can be activated by actin in the absence of Ca^{2+} When the light chain from molluscan muscle or smooth ions. muscle is added back to this system the Ca^{2+} requirement for activation is restored. Myosin light chains from mammalian skeletal or cardiac muscle cannot substitute for the former. Phosphorylation of the myosin light chains of molluscan muscle does not regulate the actomyosin-ATPase activity in this system. Regulation by phosphorylation and dephosphorylation of the light chains of myosin, with Ca^{2+} acting as an activator of the phosphorylation reaction. The phosphorylation of the myosin light chain allows the activation by actin of actomyosin-ATPase leading to contraction. This theory known as the phosphorylation theory is believed to operate in vertebrate smooth muscle by the majority of investigators. The theory is discussed in detail below.

Actin-linked regulation

Β.

Ð

Three sub-types of actin-linked regulation have been identified(57) Regulation by troponin is believed to be the primary regulatory evstem in vertebrate striated muscle. A mixture of pure actin and pure myosin exhibits a near maximal activity of actomyosin-ATPase (thus maximal contractile activity). Thue, Ca²⁺ is not necessary for the activation of the pure system. However, in intact resting muscle and in the extract of crude (native) getemyosin the Mg²⁺-ATPase activity is inhibited by a complex cooperative influence of troponin and tropomyosin.

inhibitory action of troponin and tropomyosin is prevented by Ca^{2+} which therefore acts as a disinhibitor rather than a direct activator. Ebashi and his collaborators discovered this inhibitory effect of the troponin-tropomyosin molecules, which are bound to actin in the thin filament both <u>in situ</u> and when extracted in the native system(58). Thus, <u>in situ</u> and in the native system the formation of cross-bridges between actin and myosin are prevented by the presence of troponin and tropomyosin.

Striated muscle tropomyosin has a relative molecular mass of 66,000 and is composed of two helical sub-units that coil about each other. Tropomyosin molecules are polymerised end-to-end and form a strand that lies in the grooves of the actin filament (Fig. 6). The troponin in striated muscle exists as a globular complex of three proteins: troponin I which acts to inhibit the actomyosin-ATPase activity, troponin T which serves to bind the globular complex of troponin to tropomyosin, and troponin C which serves as a reversible binding site for Ca²⁺ ions.

In the absence of Ca^{2+} , troponin I binds tightly to actin and tropomyosin, maintaining tropomyosin in a position on the actin molecule that prevents the actin-myosin interaction(59). Binding of Ca^{2+} to troponin C results in a reduction in the binding affinity of troponin I to actin and tropomyosin leading to dissociation of troponin I from the latter. This in turn results in a shift in the position of tropomyosin in the actin groove. This new position no longer prevents the actin-myosin interaction leading to the formation of cross-bridges. During relavation, Ca^{2+} is removed from the troponin C which ultimately 36



Figure 6. Diagrammatic representation of the structure of the thin and thick filaments and their interaction in muscle. The diagram shows components of three known regulatory systems for actomyosin-ATPase activation. (1) The thin filament regulatory system in which three troponin molecules (i.e. troponin-T, troponin-I and troponin-C) plus tropomyosin act cooperatively, in response to an increase in free ionised calcium to permit the activation of the actomyosin-ATPase of the globular portion of the myosin molecule by actin. (2) Actin activation of actomyosin-ATPase by a direct binding of calcium to myosin (3) The phosphorylation of the 20,000 dalton myosin light chain, thereby disinhibiting the actomyosin-ATPase. Not shown in this diagram is a second thin filament regulatory system that involves leiotonin molecules, which are bound to actin. Note: smooth muscle does not contain troponin [Reproduced with permission from the Annual review of Biochemistry, 1980: 49:925. Adelstoir PS (Annual Reviews Inc.)]

results in a shift of the tropomyosin to the "old" position in the actin groove and thus an inhibition of the actomyosin-ATPase activity.

Β.

с,

- Regulation by leiotonin, a protein that is believed to be located in, the filments. This theory, known as the leiotonin an alternative to the phosphorylation theory theory, is (discussed above) in smooth muscle regulation. Leiotonic, in the presence of Ca²⁺, activates the actomyosin-ATPase activity leading to contraction. Thus, leiotonin can be compared to troponin in striated muscle. However, unlike in striated muscle (which demonstrates near maximal Mg²⁺-ATPase activity in a mixture of pure actin and pure myosin - without troponin - in the absence of Ca²⁺) pure actin and pure myosin in smooth muscle does not have any ATP-ase activity in the presence of Ca²⁺ unless leiotonin and tropomyosin are present. This leiotonin theory is discussed in detail below.
- Recent studies suggest the presence of a unique thin-filament linked regulatory system in certain smooth muscle types (e.g. pig aorta) that is characterised by phosphorylation of a basic protein (relative molecular **mass**: 21,000) in thin filaments(50). This phosphorylation of the thin filament protein results in an approximately four-fold increase in Ca²⁺ binding by the thin filaments. Further, the eight-fold decrease in the Ca²⁺ concentration which is required to switch on the thin filament activation of skeletal muscle actomyosin Mg^{2±}ATPase, was also produced by this phosphorylation

.Dual-linked regulation

Dual (actin and myosin-linked) regulation has been shown to occur in some invertebrate muscles(61). For instance, the skeletal muscle of Limulus (the horseshoe crab) has been shown to possess a thick filamentlinked system (myosin phosphorylation) and a thin filament-linked system (troponin)(62).

In summary, the two most important theories are the phosphorylation theory and the leiotonin theory. These are discussed in detail below.

Phosphorylation theory

This, the more popular of the two theories, states that the key event in the activation of the Mg²⁺ ATPase activity is the phosphorylation of the myosin light chains. The basic concents behind this theory are(28,29,63) (Fig. 7):

- I. In the relaxed muscle the sarcoplasmic Ca²⁴ is about 10⁻⁷ mol/1: myosin is in the non-phosphorylated state and cross-bridge interactions between actin and myosin are detached.
- Contraction is initiated by an increase in earconlasmic free Ca^{2+} ions. This increased Ca^{2+} concentration activates the enzyme myosin light chain kinase (MLCK) which ploschorylates the two 20,000 relative molecular meas light chains f myosin marine of 2 moles of phosphete can be in the chain per million of the two 1 mole for each light chain)

The light chain proceeding latton all is the activitien by actin of Mg^{24} -ATPace activity of month. This leads to cross bridge formation and cross bridge cycling between active ad myosiate 't is resumed that ha long as the $Co^{(1)}$ concentration training the training to the sector the sector training to the secto



Figure 7. Schematic diagram of the phosphorylation theory of regulation of actomyosin-ATFase in smooth muscle and its modulation by cyclic AMP [Reproduced with version from the Annual review of Blochemistry 1980; [Theorem 1980]

ŧ

. 🕰 .

4. Relaxation follows a return of the sarcoplasmic Ca²⁺ to below, the activation threshold with resultant inactivation of the myosin light chain kinase.

et et al.

5. Phosphorylated myosin is dephosphorylated by one of more phosphatases astisfying the requirements for the reversibility of the system.

The myosin light chain binace is composed of two sub units(65). The larger subminit has a relative molecular main ranging from 80,000 to 125,000 depending on the anir o of the envyme. The emalion, regulative sub-unit has been identified as calculate. The Ga21 i condency of my all light chain klusse to determined by the calmodally. Colmodulty is a single chain of 148 amino acida with a calatiza molecular mass of 16,700, and has four (a" binding stree(65). The main use appears to be folded into four roughly matching longing, eich of which has a -2' binding ofter. On binding Ca2+ the calmodulity materials tries on a new more compact as we and so become activited. This is the statue he larger of up's of wosin the shows three with ear second it or the are entryme. By the later is the linger sector of the sector she and the france property of a state to the state of the second to the second and monotones of the standard to be plan in the little of the standard the the phone is the phone phone is the second seco and the second to have been able to be shown in ortho light about the north phophorals a set ATT dependent metric bases 1 4 the men provide the second physical states and the second states of the second

dispersion is a single of a state of

and the second sec

. . .

appenzyme. The phosphorylation of Site A leads to diminished binding affinity for malmodulin, which is reflected by a diminished myosin light chain binase activity(63). Smooth muscle relaxant activity of cyclic-AMF is in port the to the above event - Cyclic-AFF is of course, also believed to act in locating the free touth Co²¹ concentration in the sarcoplasm.

Depherphonelation f phosphoryl bed myosin light chains is performed by a number of energies usual myosin light chain phosphatases (HEFF). The aptidue of the energies are not effected by the concentration of e^{2} , thus they are active during both the related and contentiation of e^{2} , thus they are active during both the related and contentiation of e^{2} , thus they are active during both the related and contentiation of e^{2} , thus they are active during both the related and contentiation of e^{2} , thus they are active during both the related and contentiation of the light chains are is on the myosin light chain belong autistic groups out the activity of the phosphatases. When the associated Ca^{2} concentration folly during to pay ion, the light chain belong outstick groups out the activity of the phosphatases. When the associated Ca^{2} concentration folly during the page is with ack in factor of lar entry is inhibit do ability is indicated in the set of m^{2} is smooth ruscle the is the in subject on the phosphatase of m^{2} is smooth ruscle with mass verying $\frac{1}{2}$ is indicated in the phosphatase of m^{2} is smooth ruscle with mass verying $\frac{1}{2}$ is indicated in the set of m^{2} is in the first of m^{2} is the phosphatase.

(hendral, electronic of the contract of the state of the transference of the physical density of the contract of the state of the physical density of the physical

calmodulin dependent myosin light chain kinase from turkey gizzards to a limited digestion with α -chymotrypsin. This generated a kinase fragment (relative molecular mass:80,000) that was fully active in the absence of free Ca²⁺ ione. Phosphorylation of the invocint in the absence of Ca²⁺ allowed diasociation of phosphorylation from other potential (a²⁺ dependent regulatory metharisms Ca²⁺ independent regulatory metharisms Ca²⁺ independent regulatory metharisms from other potential (a²⁺ dependent regulatory from the sector of the sector of

tithough it we in finde to classe in support of the contraction the come of the contract finds of the first of the contraction first of the contraction of the contra

the level of myreth chomphe station line with alw or parallel in decree a actomy of a True period of

In some graveth ments is presented althouse pheaphe short a procedure the constant of the large of the characteristic tests and the second of the second of the second of the second procedure the second of the second of the second of the second procedure the second of the second of the second of the second procedure the second of the second of the second of the second second of the second of

and the second second

Theorem 1997 of the second sec

. .

cost of energy maintenance in smooth muscle. However, no conclusive evidence for the shown is available at present.

Leiotonin Theory

This theory was put forward by Ebashi and co-workers in 1975 with the discovery of a 80,000 relative molecular mass protein component of native actomyosin from the chicken gizzard(67). This protein, named leigtonin, was essential for activation of actomyosin in the presence of Ca²⁺ tong. It differed from troponin of striated muscle in that its affinity for actin was greater than that for tropomyosin. Nevertheless, tropomination was also required for the full activation of actomyosin-Lefetonin also differed from troponin in that pure ATPage agt with actin and pure much from chicken gizzard did not have any ATPase activity. even in the presence of Ca2+ ions unless the regulatory proteins letoto in and propomyosin were also present. As the leiotonin is located in the thin filaments this theory differs from the ab ashered than theory which is a mosin (thick) filament-linked The vertice Wosin light chain thesphorylation is not involved · Intribute theory of summarized by Floght(68) The essential r of this "heavy at ("ty ");

relations of the second perific for emoth models (sheletal transmus) is the the regulatory system.

"rinto in is one of two submanite: leiotonin A, the commonly (strift molecular mass: 80,000) and leiotonin control (strift molecular mass: 10,000).



LEIDTONIN C







,

.







· "7
3. Leiotonin C is a Ca²⁺ binding protein, which, although clearly different from calmodulin, can be substituted for by calmodulin in the activation of the actomyosin-ATPase by leiotonin A. However, leiotonin C cannot substitute for calmodulin in activating enzymes.

The effective leiotonin/actin molar ratio is less than 1:50.
Leiotonin has no affinity for tropomyosin, but only for actin.
During contraction or superprecipitation (an <u>in vitro</u> analogue of contraction) of smooth muscle; Ca²⁺ binds to leiotonin C. This allows the activation by leiotonin A of the actomyosin-ATPase activity resulting in the actin-myosin interaction.

In addition to providing evidence for the leiotonin theory Ebashi and co-workers have presented the following evidence which argues against the phosphorylation theory(69). (1) Leiotonin, while fully activating the actomyosin-ATPase activity of a mixture of actin, myosin and tronomyosin in the presence of (a^{21}) , phosphorylates myosin only workly. On the other hand, a mixture of myosin light chain kinase and calmodulin phosphorylates the myosin completely, but only weakly activates the ATPase. Thus, the phosphorylation of myosin appeard to be discontained from the actomyosin-ATPase activity in these experiments. (2) When a chosenhatase was added to native actomyosin superprecipitation occurred in the presence of (a^{21}) ions without any phosphorylation of the myosin light chains. (3) The pH dependency of this system also provided evidence against the phosphorylation theory. At pH 6.7 the ATPase was (1) active with only minimal phosphorylation.

The medianism of action of the leforonin is not clear at present. An lefot of the functional of a corr low lefotening actin ratio, it is

believed that its role is unlikely to be structural. If this is proven it would be another difference from the striated muscle regulatory protein, troponin, which plays a structural role in the activation actomyosin-ATPase. This low (1:50) leiotonin: actin ratio has been used as an argument against the leiotonin theory as the effective troponin: actin ratio is 1:7. However, two other proteins, β -actin and gelsolin have been found to be effective at a molar ratio (to actin) of less than 1:50(68). These two proteins do not require tropomyosin for effective function whereas tropomyosin is required for activation of ATPase by leiotonin. The exact role of tropomyosin in the leiotonin theory is not well understood.

In summary, there seems to be good evidence for and against both phosphorylation and leiotonin theories. It is possible that both regulatory mechanisms may be functional in smooth muscle either in different muscles or in a single muscle as a dual regulatory system.

AUTONOMIC INNERVATION OF VASCULAR SMOOTH MUSCLE

Autonomic nerves have been identified in the walls of most blood vessels(70). Must of the norve fibred are sympthetic unreducingic in origin although sympthetic cholinergic fibres are found in some vessels such as the skeletal muscle vescular bed. A parasymptihetic cupply is present in other blood vessels such as the vasculature of the erectile ticsues and is corobial circulation(11). Here recently, the existence of primersic veryes in non vascular emothermic is here been down downented (72). On the other hand, some bloc' schede such as the umbiliget artery of the lumon deep time vessels such as the umbiliget of the intervention of the bound deep time vessels such as the umbiliget artery of the bound deep time vessels and include the lumon deep time vessels of the other hand is not account to be a scheder of the bound deep time vessels on the time of the include of the bound deep time vessels and the scheder of the include of the bound deep time vessels and the scheder of the bound deep time vessels and the time of the bound of the bound deep time vessels of time of the time of the time the scheder of the bound deep time vessels and the scheder of the time of the time of the time of the time of the scheder of the time of t

indicating the presence of an adrenergic nerve supply(70). As a great deal more is known about the adrenergic nerve supply compared to the other two types the ensuing discussion relates to the former unless specified otherwise.

The adrenergic nerves to blood vessels are non-myelinated, postganglionic fibres. These fibres usually form two plexuses in the adventitia: a primary plexus in the middle or outer third of the adventitia and a terminal plexus typically restricted to the adventitiomedial junction(73). This adrenergic terminal ground plexus tends to be similar to that found in non-vascular smooth muscle, being irregular, multiaxonal and of varying density. Non-myelinated axons, 0.25-0.5 μ m in diameter with a surrounding schwann cell sheath form the plexon which surrounds the tunica media like a sheath. The axons contain varicosites 1.5-2.0 µm in diameter at intervals of 3-10 µm along their lengths. These are the storage and release sites for noradrenaline which is contained in a large number of storage vesicles within the varicosites. A nerve impulse propagating along an axon successively depolarizes a series of variocosities resulting in the release of transmitter at each of these sites. The released neurotransmitter brings about changes in vascular smooth muscle tone. The nerves themselves terminate at the adventitio-medial junction or in the outer third of the media in most vessels. A model for the innervation of smooth muscle has been proposed by Burnstock on the basis of electrophysiological, histochemical and electron-microscopical studien(74). The equential features of this model are as follows (Fig. 1. 1.





Figure 9. Model of the vascular smooth muscle neuromuscular junctions. Vascular smooth muscle is controlled by both nerves (....) and circulating catecholamines (arrows) and contains three types of cells (1) directly innervated cells (2) directly coupled cells (3) indirectly coupled cells [Reproduced with permission from Burnstock G, British Medical Bullerin, (711)

- 1. The effector unit is a muscle bundle rather than a single smooth muscle cell.
- 2. Unlike those of the skeletal muscle neuromuscular system, autonomic nerves in smooth muscle run long distances, containing varicosities which have high levels of neurotransmitter: the transmitter is released <u>en passage</u> during the conduction of an impulse.
- 3. Individual muscle cells are connected by low-resistance pathways or gap junctions which allow electrotonic spread of activity within the effector bundle.
- 4. In most smooth muscle bundles, some (but not all) muscle cells are directly innervated, i.e. in close (20-120 nm) apposition with the the varicosites, and are directly affected by the transmitter released from them ("directly innervated cells"). The adjoining cells have been named "coupled cells" as they are electrotonically coupled to the former by low resistance pathways so that excitatory junction potentials can be recorded in these cells. These junction potentials are very slow (nearly 1.0s), resulting in a whole area of the effector bundle depolarizing almost simultaneously, triggering an action potential. The action potential propagates through the effector bundle to activate a third group of cells named the "indirectly coupled cells" which are neither directly innervated nor directly coupled and yet respond on stimulation of the nerves supplying the organ. The density of innervation tends to vary widely in different parts of the vascular system. In general, arteries receive a richer innervation than veins with large arteries having a relatively poor

.

nerve supply compared to small arteries and arterioles. Further, there is considerable species variation in the innervation of the same type of Rat aorta is sparsely innervated, whereas the rabbit blood vessel. aorta contains a terminal plexus at the adventitio-medial junction. The large elastic arteries usually do not contain nerves penetrating the tunica media. Muscular arteries such as the rabbit ear artery tend to be more heavily innervated than elastic arteries. However, the innervation is usually confined to the adventitio-medial junction in In some muscular arteries, particularly in most muscular arteries. certain species such as sheep and man, the nerve fibres penetrate about one-third of the thickness of the media. Although veins in general have an innervation which is less dense than arteries, some medium sized veins such as the small saphenous vein of the rabbit have an extensive Medial innervation is commonly found in submedial innervation. cutaneous limb veins. A summary of the innervation characteristics in a variety of blood vessels is provided by Bevan et al. ("3) who also make the following points regarding neuronal density in macular entry muscle.

1. Nerve density does not very systematically with versal liew ter or wall thickness.

> show a widely different nattern of The waaael may same which the cract location at the innervation depending on morphology is investigated, e.g., the first few millimetres of the rabbit saphenous artery is not innervated. It then gives off a small muscular branch after which the innervation becomes deuse and medial in nature. Passing Hatalls, the innervation takes on Les the A consola advontitio melial nottorn 11 ... nn

excised for <u>in vitro</u> studies, they have to be removed from the * same anatomical site.

- 3. Nerve density, provided other parameters of transmission are similar, can be related to the maximum level of neurogenic tone of which a vessel is capable. However, there are exceptions to this rule.
- 4. Nerve density does not remain constant with age.
- Neuronal density of a particular vascular segment is not the same in different species.

Autonomic innervation of the canine lateral saphenous vein

An investigation on the innervation of this vein using fluorescence microscopy (with formaldehyde) and autoradiography (with tritiated noradrenaline) was reported by Osswald and co-workers(17,18,75). This study demonstrated the presence of adrenergic nerve fibres spread across the whole media. The nerve fibres were unmyelinated axons enwrapped by Schwann cells. Sometimes the axoplasm was seen to contain microtubules anly, but in most cases it contained numerous dense-core vesicles (made up of large granular and small granular vesicles) agranular vesicles and mitochondria. The large granular vesicles constituted approximately 43 per cent of all dense core vesicles and were 80-200 nm in diameter whereas the small granular residles were 40-60 nm in diameter. This considerable proportion of large granular vesicles (43 per cent) found in this vein is different from most advenergic nerves, which contain. predominantly small dense core vesicles. The diameter of the large dense-core vesicles observed is also larger (80-200 nm) than in most The agranular vesicles present had a diameter adronorale vervee. -"1 200 mm. renortion from The areas of the axons containing dense-core

vesicles were often found to be partially lacking Schwann cell sheaths and were located in close apposition to smooth muscle cells. The neuromuscular distance at these points usually varied from 100-300 nm with a much shorter distance (20 nm) sometimes being observed rarely. There was no thickening of the opposing plasma membranes at these neuromuscular junctions, however, the basement membranes were continuous at these points. This narrow cleft width observed in the sachenous vein accounts for the predominant role played by uptake₁ in the transmitter disposition in this vein. It may also account for the relatively short delay (less than a second) between the commencement of electrical stimulation and the beginning of the contractile response, observed in this preparation.

The sympathetic innervation to the name of the dog hind-limb originates from the lumbar spinal cord('6,77). The nerves leave the cord via the upper lumbar roots and run in the main sympathetic ganglia. The nerves join the sciatic corve via the rami to the sixth and seventh lumbar nerves and the second sacral nerves. Thus, surgical sympathetic denervation of the veins in the hind limb is a bioved by dividing the sympathetic chain at the second or third lumbar certe ral body and dissecting the chain free from the surrounding the the fifth lumbar vertebrail (78).

ELECTROPHYSIOLOGY OF SMOOTH MUSCLE

Vascular smooth mumbe calls, like all other living cells printain a potential difference a rose their cell membranes, with the inside Being negative compared to the outside. This resting membrane potential (Em) varies from -40 to -75 my in different smooth model alls. This potential difference is dependent on the faction is the up of

distribution of ions across the sarcolemmal (cell) membrane with different permeabilities to the respective ions. (2) The electrogenic transport of ions across the cell membrane with the aid of the sodium/potassium membrane pump (Na^+/K^+ pump).

Resting membrane potential

The unequal distribution of ions across the sarcolemma results in the generation of a diffusion potential for each ion depending in its concentration difference across the sarcolemma and the membrane permeability to that particular ion. If one assumes an unrestricted movement of a particular ion, the diffusion potential (at which there is no neith movement is the ion across the sarcolemma) is given by the Nermal movement

 $E_{I} = \frac{RT}{2F} \frac{n}{|I|i}$

Eı	•	equilibrium diffusion prevital for ion I
R		gas constant (8.316 1) per legres)
r	•	absolute temperature
		valency of the 💏
च		Faraday constant (96,500 coulombe/mel of 'an'
9 71		untural logarithm (2.313)
[T]e	•	concentration of ion in extracellular fluid
["]#	•	concentration of ion in intracellular fluid
	At a	remperature of 30°° the Nerver equation for the V ¹ ion can b

E TARTER TO:

54

be

Thus the membrane potential would depend on the resultant effect of all the diffusion potentials for the different ions in the muscle cell and the extracellular fluid. However, the potential difference each ion contributes to the Em depends on the permeability of the schoolemma to that portioniar ion. In the resting state. This is is below from account is the Nernst equation which asgum a unrease to the school from the ions. The mesting membrane potential is the product of the ions, the mesting membrane potential is the product of the ions (by Na', K⁺, Cl') across the still membrane constraint is fell mation of foldman(72).

 $E_{m} = \frac{P_{K}(K^{+})o + F_{Na}(Na^{+}) + F_{(1)}('1'')}{(r^{+})}$

to the type where near the

If wents one were relaction to all is (the ment of potential cost to appr the cost open to be applitude of the potential as a second second of a file of the second by points equal of a constraint of the c where a fit of drange of the forth change to fit to be a proceeded and theory of all of a the spectrum gets a star with a second ور د د و د $\frac{1}{2} \left[r + r \right] = \frac{1}{2} \left[r + r \right]$ $\left\{ r = 1, \alpha \right\} = \left\{ r = 1, \beta = 1, \beta$ the line of the two for a fill, 81 the second se mombanes to prove set and the provest set and the set 1 1 $(1, 2, \dots, 2^{n-1}) = (1, 2^{n-1}, \dots, 2^{n-1})$ • • * 1 e · . 1

muscle compared with skeletal muscle. Secondly, as $[K^+]_0$ is increased in vascular smooth muscle, K^+ conductance (\mathbf{g}_K) increases (\mathbf{g}_K) : the conductance if an four through a membrane is the reciprocal of its resistance - It is defined as the nett current flow per unit voltage)(80°. Such an increase in \mathbf{g}_V tends to hyperpolarize the membrane of any given elication in $[V^+]_0$. In other words, the amount of inpolarization produced by a given elevation in $[K^+]_0$ would be less be more of the increase in \mathbf{g}_K . This fractor houds to be more prominent in arterial smooth muscle cell which have $1 \leq \mathbf{g}_K$ and \mathbf{P}_K since \mathbf{F}_m (the membrane cotential) is further \mathbf{F}_K the fractor could be less to be more prominent in arterial smooth muscle cell which have $1 \leq \mathbf{g}_K$ and \mathbf{P}_K since \mathbf{F}_m (the membrane cotential) is further \mathbf{F}_K the fractor could be reserved.

is copiate the inclutant equilibrium diffusion potential for a group of the construction uning the Coldman equation. The realized or and i amouth scale often tends to be less results the second new potential $\left(\begin{array}{c} \mathbb{R} \\ \mathbb{R} \end{array} \right)$ recorded hv atelested at the tendies. The lifference by year the two values is istication of the station of the alectrogeness of the pump. A major performance of the two approach is the time to the former actions and \sim and \sim remarkability of ε_m where ε_m is the station of ε_m time spectre are lift in the state as wataly. The No /K rempt for success to attend the second to an use of in exchange for entrarel tar " in level difficult of the laws of electrochemical and fout and the second difference of all all going for measuration and Prove in the set concerned in Na 200 and to maintain the "for talk "gets cont of the exchange · · · · · · for the product of the state of the my

56

· · · • • • • •

electrically neutral. However, in most cases the exchange is not one for one, more Na⁺ ions being extruded than K⁺ ions brought in, i.e., the pump is electrogenic. The coupling ratio for the pump in red blood cells is 3Na⁺:2K⁺. Estimates in other tissues though variable, generally approximate the 3:2 ratio. The pump is energy dependent and is associated with the membrane bound enzyme Na⁺/K⁺-ATPace which hydrolyses the Mg-ATP. The presence of Mg-ATP inside the cell is essential for the activity of the pump(79) with one molecule being hydrolvged for every three Na⁺ ions pumped out. The level of cell ATF can be lowered by removal of substrate from the bathing medium (, and glucose being replaced by N2 and serie (tot). Since prooth muscle cells are capable of anaerobic respirates a combination of indecenate and dinitrophenol is required to reduce cell ATP to unmeasurable levels. Under these circumstances cells "r'ily lose " and rain Na'(83). The activity of the Nat/K pump is a function of the intracellular Nat concentration and the optracellular " concentration, the nump being inhibited if the P' concentration of the bathing medium to made zero-It is also indified on low teams atures bacause of its energy topordo and in the glycogides (a qualiste digitalis) the bit the supported the standard matter to take the state of the same set of

There is considerable debute a to the magnitude of the constraint of the Norl/K^d nump to the contribution potential in amouth muscle. Two approaches have been used to estimate this contribution (82). The first multiplication is less the intermination is the difference the constraint of the constr

involves the determination of the change in membrane potential during the inhibition of the pump. Both methods are not without error(82) making the quantification of the contribution of the pump to the resting membrane potential difficult. The contribution is probably much less than was formerly believed amounting to about 4mV(84).

In smooth muscle, where tension is a function of the membrane potential, the Na^+/K^+ pump may function as a mediator of the vasoconstriction and vasodilation associated with certain "drugs"(82). Hyperkalaemia and hypokalaemia (moderate) are associated with vasodilatation and vasoconstruction respectively. This is probably mediated through modulation of the activity of the Na^+/K^+ pump in the muscle cells. Thus, high extracellular K⁺ leads to stimulation of the pump with resultant hyperpolarization and a decrease in tension. These responses are antagonised by ouabain. Shepherd and co-workers showed that the Na /K ATEnse inhibitor acetylstrophanthidin potentiated the contractions of the lateral saphenous veins produced by noradrenaline, 5-hydronitivitaming of hydrholine and harfum(85). These effects of cardian given and trobably due to changes in resting membrane ' te (1al(82))

The inwardly Hrecked Na⁺ gradient created by the Na⁺/K⁺ pump is used in transporting other ions or molecules against their concentration gradient. The Ma⁺/Ca²⁺ exchange, Na⁺/amino acid exchange and Na⁺/Mg²⁺ exchange are some of the transport processes in smooth muscle, which are helf of the function using this Na⁺ gradient(83). The Na⁺/K⁺ pump may also plate a table in the post-junctional supersensitivity in smooth muncle. Thus a decrease in Na⁺/K⁺-ATPase activity with a partial

reactivity in some cases of post-junctional supersensitivity(82).

Action potentials

Action potentials of different configurations have been recorded in a wide variety of blood vessels(81) and they occur in bursts or at regular intervals. The action potentials may resemble those in skeletal muscle with a sharp rise and fall or they may have a prolonged time course with a plateau phase similar to that observed in cardiac muscle, e.g. turtle aorta. The spike potentials in smooth muscle may have an overshoot of up to 20 mV. However, the maximum rate of rise of the spike is very much slower (5-20 V/s) compared to striated muscle or nerves (1000 V/s). The maximum rate of repolarization is often similar to that of depolarization or it may be slightly faster(84).

Some smooth muscles such as in the guinea-pig urinary bladder and in the porto-mesenteric veins of many species, exhibit spontaneous action potentials. Others demonstrate action potentials only in response to stimulation with drugs. Tetraethylammonium ion (TEA) which reduces K^+ conductance (gk) induces spontaneous action potentials in some smooth muscle with no inherent activity. In other smooth muscles which do not respond to transmural nerve stimulation with action potentials, it induces spikes in response to stimulation(80).

The ionic basis of the action potentials in smooth muscle has been the subject of some debate(81). As vascular smooth muscle action potentials have a variety of configurations with different time courses, it is possible that the ionic mechanisms may be different in different musles. In the majority of preparations reduction of the extracellular Na⁺ conducts for does not have an appreciable effect on the action reduction in

the rate of depolarization and the amplitude of the overshoot in nerves and skeletal muscle. This argues against Na⁺ ions playing a major role in the action potentials in smooth muscle. The lack of effect of the fast Na⁺ channel inhibitor tetrodotoxin on the action potential supports the above concept. The smooth muscle spikes are blocked by the bivalent ions cobalt (Co^{2+}) , nickel (Ni^{2+}) and Manganese (Mn^{2+}) . They are also abolished by the removal of Ca²⁺ from the extracellular fluid and by calcium anatagonists. Both strontium (Sr^{2+}) and barium (Ba^{2+}) ions are able to replace Ca²⁺ as carriers of the inward current(84). The above findings point to Ca^{2+} ions being the major ion involved in action potentials of vascular smooth muscle. However, in the quies fint sheep carotid artery, Keatinge was able to induce electrical activity in Ca²⁺ and Mg²⁺ free solutions(86). These action potentials were abolished when extracellular Na⁺ was replaced by Tris or choline, and they also ceased when Ca^{2+} or Mg^{2+} were added. It was concluded that Na^+ was the principal ion carrying the depolarization current in the action potential in this preparation. However, tetrodotoxin, the fast Na⁺ channel blocker, did not abolish these action potentials.

In addition to action potentials, some smooth muscle demonstrate a regularly occurring, rather slow changes in membranes potential (lasting up to several seconds) which are referred to as slow waves(84). These slow waves vary in frequency and time course from one tissue to another and may not mecessarily exceed the threshold to trigger action potentials. Slow waves are present in intestinal smooth muscle of many species and their amplitude is generally about 20 mV, though it may be up to 40 mV rarely. They may occur without any accompanying changes in tension. The slow waves increase muscle tension either if the waves

crosses the threshold and initiates (one or more) action potentials or if they cross the contraction threshold in gradedly responsive muscle (see below). Unlike action potentials, slow wave activity is little affected by changes in the membrane potential and they are also not propagated in the tissue(87). The depolarization phase of the slow waves are believed to be mediated via a Na⁺ influx which is resistant to tetrodotoxin. There is good evidence that these slow waves are a function of spontaneous changes in electrogenic pumping(82). Ouabain, K^+ -free solutions and cooling inhibits slow waves, supporting the role of the Na⁺/K⁺ pump in the generation of these waves.

Effectrophysiologically smooth muscle can be divided into two broad

(1) spike generating the muscle

(2) non-spike generating smooth muscle.

The spike-generating smooth muscle can be sub-divided into the single-unit type and the multi-unit type on the basis of their ability to generate and propagate electrical activity(81). In both single-unit and multi-unit spike generating smooth muscle, the contractions are brought about by action potentials. However, in the single-unit variety, the potentials action are triggered by slow-wave depolarizations or pacemaker potentials resembling the phase depolarization of cardiac muscle, but having a much slower time course. The portal-mesenteric veins of many species provide a good example of this type of vascular smooth muscle. Smooth muscles of the intestine and urinary tract are other examples of single-unit spike generating smooth muscle. The action potentials may occur in bursts or at regular intervals in this type of smooth muscle and they travel along

the tissue at a conduction velocity of 10-80 mm/s via low resitance pathways between the muscle cells. However, the spread is usually decremental. Therefore large preparations show multiple pacemaker sites and variable patterns of contraction(81). These action potentials originate within the muscle itself as they are unaffected by tetrodotoxin. The contractions in this type of muscle is triggered by the action potentials, with the basal tone in the muscle being proportional to the frequency of the spikes in the resting state. The frequency of the action potentials tends to be decreased by cooling. On the other hand, the frequency is increased in some tissues by stretching This may provide a possible basis for myogenic of the muscle. autoregulation of blood flow. Neurotransmitters released at nerve endings within the muscle and exogenous drugs car also modulate the frequency of the spikes with either an increase or decrease depending on the agent involved. These drugs may exert their action by changing the membrane potential: depolarization causes an increase and decrease in frequency or the abolition of hyperpolarization a spontaneous activity(84). This is probably due to the fact that raising lowering the membrane potential would or increase or decrease (respectively) the time required for the spontaneous depolarization wave to reach threshold potential, thus altering the frequency of the Stimulation of portal-mesenteric veins by noradrenaline and spikes. adrenaline has been shown to result in an increase in the frequency of action potentials(81,88). With high concentrations of these agents, the marked depolarization produced, obliterated the action potentials. However, the strong contractile response remained intact. On the other hand, the action of isoproterenol in portal veins of different species

the above(81). clear The drug caused cut **a**8 was nòt **a**8 hyperpolarization with cessation of spontaneous activity in the rabbit However, in the rat portal vein, and guinea pig portal veins. isoproterenol resulted in the action potential pattern changing to one of short, frequent bursts, each containing fewer spikes associated with depolarization of the cell membrane. It is possible that isoproternol may have other actions which alter the relaxatory mechanisms in smooth muscle more directly via the β -receptors and cyclic AMP.

In the multi-unit variety of spike generating smooth muscle action potentials are initiated by transmitter induced excitatory junction potentials. The transmitter released at nerve endings produce miniature excitatory junction potentials at the post-synaptic membrane. These miniature junction potentials summate to produce an all-or-none action potential when the membrane potential reaches threshold. Mouse vas deferens typifies this type of smooth muscle. There are no good examples of vascular smooth muscles with similar electrical properties and predominance of neural control(81).

Non-spike generating vascular smooth muscle contract in the absence of action potentials. This muscle is characterised by graded depolarizations and contractures instead of spikes and twitches. The contraction is accompanied by a membrane depolarization without a true action potential with the contraction being maintained as long as the membrane remains depolarized. A true cause and effect relationship between the electrical and mechanical events is more difficult to prove in this type of smooth muscle: it is possible that the two responses just occur as simultaneous parallel phenomena. Rabbit pulmonary artery provides an example of this type of vascular smooth muscle. In smooth

muscle, which shows a graded response, electrotonic spread of the depolarization produced by neurotransmitters probably does not play a significant role in the cell-to-cell spread of activity. The absence of action potentials is probably due to the fact that the mechanisms necessary for the fast, regenerative changes in ion conductance that mediate spikes are, in fact, not normally operating in the membranes of these cells(81). However, some smooth muscle of this type can be 'stimulated' to produce action potentials by the tetraethylammonium ion.

In both spike-generating and non spike-generating muscles, an inward Ca^{2+} current occurs during the depolarization phase via Ca^{2+} There appears to be a number of voltage sensitive Ca²⁺ channels. channels in smooth muscle: rapidly activating and inactivating channels involved in the upstroke of the action potentials, and slowly activating maintained Ca²⁺ but channels functional during prolonged depolarization. Drugs which produce depolarization by a mechanism other than an inward Ca²⁺ current may also secondarily activate any voltagesensitive Ca²⁺ channels if the membrane potential is moved into the correct range. This would generate tension in the smooth muscle by increasing the Ca²⁺ influx. The exact, mechanisms by which drugs to influence the membrane potential have not been established in most cases. In principle they probably act primarily through an effect on passive ion permeabilities and in some instances electrogenic ion transport(80).

Electro-mechanical and Pharmaco-mechanical Coupling

In both spike generating and non spike-generating muscle described above, the contraction of the muscle cells is coupled with an electrical event at the cell membrane: an action potential in the former and a

graded depolarization in the latter. This type of coupling is referred to as electro-mechanical coupling and is similar to the phenomenon found in skeletal and cardiac muscle. A quantitative relationship between the membrane potential and the tension can be demonstrated in skeletal and cardiac muscle after blockade of the action potential. An S-shaped relationship has been found defining the mechanical threshold and the characteristics of the electromechanical coupling(81). These curves are Attempts at similar characterisation in shifted by drugs and ions. smooth muscle has been less successful because of the difficulty in blocking the action potential without losing the contractility(89). However, experiments carried out at 10°C were more successful in demonstrating a S-shaped relationship between tension and the \star xtracellular K⁺ concentration in the rat portal vein(90). The fact that normally spike generating vascular smooth muscle maintain tonic contractures in depolarizing high K⁺ solutions demonstrates that the action potentials are not essential for evoking contraction in these tissues.

Although changes in membrane potential can initiate changes in tension, in smooth muscle certain hormones and drugs can cause contraction without any change in the membrane potential. This phenomenon known as pharmaco-mechanical coupling was first described by Somlyo and Somlyo in 1968(91). Pharmaco-mechanical coupling has been defined as a process or processes through which drugs can cause contraction or relaxation of smooth muscle without a necessary change in the resting membrane potential or in action otential frequency. This definition does not imply that pharmacomechanical coupling cannot also act concomitantly with a change in the membrane potential. It is quite

probable electro-mechanical and pharmaco-mechanical coupling that processes contribute to activation (or inhibition) simultaneously. Quantitative investigations are needed to ascertain the relative magnitudes of the two processes. For instance, if the contraction produced by an agonist which also produces a simultaneous depolarization is greater than that elicited by an equivalent depolarization by high K^+ solution (or an electrical current), it is probable that the excitatory actions, of the agonist is due to both electromechanical and pharmacomechanical coupling (or to pharmaco-mechanical coupling alone). Farley and Miles in recent experiments carried out in the dog trachealis muscle, demonstrated that although acetylcholine in a concentration of 10^{-7} mol/1 produced some membrane depolarization with a contraction approximately 15 per cent of the maximum, an equivalent depolarization by high K^+ did not generate tension(92). Further, a recent study on the guinea-pig main coronary artery demonstrated that the contractions caused acetylcholine were by accompanied by a membrane hyperpolarization(93). The evidence for pharmaco-mechanical coupling has been summarised as follows by Johansson and Somlyo(81):

1. the same blocking and potentiating agents are effective in polarized as well as depolarized smooth muscle.

2. depolarization by drugs may be less, but the maximum contraction greater than the respective effects of high K⁺ solutions.

3. the differences in the maximum contractile effects of different drugs are maintained after depolarization.

4. drug-induced contractions may be sustained in smooth muscles that respond to depolarization by high K^+ with a transient phasic

contraction.

5. relaxing agents can relax polarized smooth muscle without evidence of hyperpolarization or inhibition of spike electrogenesis.

These authors comment that 'whereas theoretically the possibility of excitatory pharmaco-mechanical coupling without depolarization in polarized smooth muscle is possible, a rigorous demonstration of, this phenomenon requires experiments in which membrane potential and contractile responses of single cells are monitored simultaneously'. The mechanism by which pharmacomechanical coupling occurs is not clear at present but the suggested possibilities are(81):

1. mediation by the influx of extracellular Ca^{2+} . As by definition the membrane potential should not change during pharmaco-mechanical coupling, for the above mechanism to account for this process there has to be a simultaneous movement of another ion to balance the membrane potential change that would be produced by the influx of Ca^{2+} .

- 2. mediation via release of Ca^{2+} from the sarcoplasmic reticulum in response to a change in permeability of the surface (plasma) membrane at the couplings between the reticulum and the surface membrane.
- 3. _____mediation via a direct effect of the drug on intracellular Ca²⁺ storage sites. It is unlikely that large polypeptides such as angiotensin could act via this mechanism but more permeant molecules such as the prostaglandins could exert intracellular effects.

These mechanisms are not mutually exclusive, thus two mechanisms could occur together. In summary, although all available evidence suggests that pharmaco-mechanical coupling probably occurs in smooth muscle, no absolute proof is available at present.

CALCIUM REGULATION IN SMOOTH MUSCLE(29, 30, 81, 83, 94)

As tension development in muscle is a function of the concentration of free intracellular Ca²⁺, the regulation of the latter is of prime importance in all muscle cells. Compared with striated muscle, very little is known about the sub-cellular regulation of Ca^{2+} in smooth The specific role of the various Ca^{2+} binding sites muscle. (sarcoplasmic reticulum, mitochondria, sarcolemma) is not clear at present. Further, the immediate source of activator Ca^{2+} is not known, nor is the mechanism of relaxation known. Smooth muscle cells are activated when the ionised intracellular Ca^{2+} rises above 10^{-7} mol/1 with maximal activity between 10^{-5} and 10^{-4} mol/1(83). The total tissue content of calcium in smooth muscles may, however, be 100 fold of the amount required for maximum contractile activity. The total tissue calcium extracellular free, extracellular bound, 18 made up of intracellular free and intracellular bound fractions. Brading and Widdicombe estimated these calcium fractions in the taenia coli of the guinea pig(95). The total tissue calcium was found to be 2.73 ± 0.13 mmol/kg fresh weight of tissue (Ca^{2+} in the bathing medium during the experiment = 2.5 mmo1/1). $\frac{14}{100}$ the free Cq²⁺ present in the [¹⁴C] sucrose. space (used to estimate the tracellular space) was substracted, the Ca remaining in the tissue was 1.8 ± 0.13 mmol/kg fresh weight of tissue. The extracellular bound Ca was estimated by treating the tissue with Lanthanum (La³⁺ 5 mmol/1) for 60 min (this may be an overestimate as some cellular Ca is lost during the La³⁺ treatment). If this amount is subtracted the remaining Ca, which can be considered as the total intracellular Ca, was found to be $0.43 \pm 0.09 \text{ mmol/kg}$ fresh weight of tissue. This is still an order of magnitude greater than that necessary

to activate maximally the contractile apparatus in smooth muscle cells. The bound intracellular Ca is located in the organelles such as the mitochondria, the sarcoplasmic reticulum, the nucleus and in the In the resting muscle, the concentration of free Ca²⁺ in sarcolemma. the myoplasm is less than 10^{-7} mol/1 although the concentration of free Ca^{2+} in the extracellular fluid is greater than 10^{-3} mol/l. Thus, for vascular smooth muscle to be relaxed, its plasma membrane must support a 10,000 fold concentration gradient of the Ca²⁺ ion. Recent studies in . myocardial tissue by Langer have formed the basis for a model whereby the glycocals (a coating external to the unit membrane) participates in the transmembrane ionic exchange(96). The glycocalyx is composed of two layers: an inner. less dense. 20 nm thick surface coat and an outer, slightly more dense, 30 nm thick external lamfna. A high concentration of negatively charged sites are present in the givencalvx. Stalic acid residues which form the terminal groups in the objestaccharide portions of the glycoproteins and glycolipids composing the glycocalyr form a majority of these negatively charged sites. Treatment of myocardial tissue with neuraminidase, which selectively removes sialic acid residues, increases the uptake and vashout of radioactive Ca2+ fivefold. Further, following this enzymatic treatment, In3" ior , which are normally restricted to extracellular space enters the cell feeding to displacement of more than 80 per cent of the Verchangeable calcium, Langer proposed that the glycocalyx may be necessary for the prevention of uncontrolled calcium entry from the extracellular space.

Activator calcium in smooth muscle

The Ca²⁺ responsible for activation in smooth muscles comes from intracellular and/or extracellular sources. The source of activator 69

-

Ca²⁺ seems to differ between smooth muscles from different vessels and between different stimulants used to initiate contraction. Vascular smooth muscle, which shows spontaneous contractions accompanied by action potentials (spike generating smooth muscle, see above), depend on extracellular Ca²⁺ for this spontaneous activity. The action potentials porto-megenteric veins are blocked by the removal of the extracellular Ca²⁺ ions(81). Agonists, which cause depolarization of the cell membrane, lead to entry of Ca²⁺ from the extracellular space through Ca²⁺ channels resulting in muscle contraction(97). к+ depolarization has been used for the investigation of depolarization induced contraction in smooth muscle. Contractile responses produced by high concentrations of K' are considerably inhibited in a Ca^{2+} -free medium lending further support to the role of extracellular Ca^{2+} in this contraction. The contraction in the presence of K⁺ consists of two components: an initial, large, transitory phasic component, and a second slower increase in tension which subsequently diminishes to a lower plateau level of tension over 90-120 minutes(98). The latter is referred to as the tonic component. These phasic and tonic components of the contractile responses have been identified in many agonist induced contractions in smooth muscle. The phasic component has been found to be dependent on intracellular Ca^{2+} ions and the tonic component on extra-cellular Ca²⁺(99). However, both components in the case of the contractile response to K^+ appear to be dependent on extracellular Ca^{2+} ions(98). Thus both components of the response were highly sensitive to the Ca^{2+} channel antagonist verapamil and also inhibited by La^{3+} which competes with Ca²⁺ ions primarily at extracellular sites on the These findings do not imply that extracellular Ca^{2+} is the membrane.

only source of activator ions for the contractile apparatus in K^+ induced contraction. It is quite possible that the inflow of Ca^{2+} ions from the extracellular medium may trigger a further release of Ca^{2+} from intracellular Ca^{2+} stores (calcium-induced calcium release). Hurwitz et al(98) suggested that the two components of the K^+ induced response in intractinal emotion muscle are probably mediated via two separate Ca^{2+} channels in the membrane. The preferential inhibition of the phasic component by a lower concentration of La^{3+} (both components being inhibited by a higher concentration of La^{3+}) suggested the above conclusion. It should be clear that the terms phasic component and indic component are purely descriptive terms used to identify components is gonist induced entractions in muscle. The source of Ca^{2+} conclusion the two components is nearly depending on agonist as well is indiced for the two components may many depending on agonist as well is investigated.

"I connect of a helerogeneous group of the channels in the roth marche off methropic har developfill ar the recent wears a lability for a collectivity of that inhibit of " i ist out that the second second

(1) In the notation of the probability of the probability of the operation of the probability of the sector of the terminate (probability of the sector of the

1 - 1 1 - 1 1 - 1

.71

change in the membrane potential, e.g. low concentrations of noradrenaline. The channels are blocked by sodium nitroprusside and amrinone but are relatively resistant to the organic Ca²⁺ channel antagonists cited above. When potential operated channels are activated in the presence of depolarization produced by an agonist, one cannot exclude a similar pour activation of receptor operated channels independent of the depolarization.

.

3. Ca²⁺ channels responsible for resting Ca²⁺ curve this the to be postulated to the resting Ca²⁺ entry is unaffered by either organic Ca²⁺ channel ortagonists of nitrormaside.

In addition to these these (2^{24} entr) should superficially located high and low affinity (2^{24} entr) should are present in the agreelemme(100). Uptake of $(2^{27})^{-2}$ at the high affinity site is preferentially blocked by structure $(2^{27})^{-2}$ allowing a plan qualitative separation of the two allow $(2^{27})^{-2}$ blocked by structure $(2^{27})^{-2}$ blocked by affinity site and an qualitative separation of the two allow $(2^{27})^{-2}$ blocked by affinity of a site of a site of the two allows $(2^{27})^{-2}$ blocked by affinity of a site of a site of a site of the two allows $(2^{27})^{-2}$ blocked by affinity of a site of a site of a site of the two allows $(2^{27})^{-2}$ blocked by a structure $(2^{27})^{-2}$ blocked by a structure $(2^{27})^{-2}$ block and be affinity of a site of the two allows $(2^{27})^{-2}$ blocked by a structure $(2^{27})^{-2}$ block between the site of a site of the two allows $(2^{27})^{-2}$ blocked by affinity of a site of the two allows $(2^{27})^{-2}$ blocked by a structure block between the site of a site of a site of a site of the two affinity of a site of the two affinity of a site of a sit

and the second sec

a service of a structure of the service of the serv

bath was approximately equal to the sum of Ca²⁺ influxes when either of the activators was present alone.

Several techniques have been developed in the last decade to measure the shift of Ca²⁺ between the extracellular and intracellular compartments. Van Breemen and co-workers pioneered a method(102,103) in which tissues could be exposed to a variety of agonists in the presence of radioactive $Ca^{2+}(45Ca)$ and subsequently placed in wash out solutions containing La³⁺ ione. The concentration of La³⁺ in these solutions is kept high enough to replace all superficially bound 45Ca and to prevent any further uptake or efflux of 45 Ca. The assumption that La³⁺ would replace the extracellular and superficially bound 45Ca and prevent further ⁴⁵Ca uptake is justified. However, the assumption that La³⁺ would block ⁴⁵Ca efflux proved to be incorrect(100). Thus modifications such as the use of a higher La^{3+} concentration and a lower temperature during the wash-out period were introduced. The steps of the La³⁺ method as used by Godfraind et al(104) (for the measurement of the is the of California ortic mighter alle stimulated by Kt. and the effect tagentite in this uptake) are ligied below:

atrice of our a weighing 6.11 mp were to obtain with physiclogical and columns contribute togethe comparison (experimental civile) for an operation with possible control atrice kept in that physicligical of a collific the control for proceeding to the buffer column to the civil fill for and control of the original atrice only.

the extent of ends over the basis to provide the physical edit colution connected x^{-3} and prime 1 for 5 where card the for grapher 2 strateging with x^{-1} is the transmitted over the source 1 and 1 preparations were washed for 5 minutes in 500 ml of tris buffer solution (pH 6.8) containing lanthanum chloride (50 mmol/1
 the strips were then weighed and ⁴⁵Ca extracted using perchloric

acid and hydrogen peroxide.

аŞ

5. the radioactivity extracted from the tissue was counted in a liquid scintilation counter.

A modification of this technique can be used to estimate the efflux of 45 Ca produced by agonists. Steps of the procedure as used by Godfraind et al(104) (to measure the 45 Ca efflux produced by prostaglandin $F_{2\alpha}$ with or without antagonists) are listed below.

- *1. aortic strips were incubated with physiological salt solution containing ⁴⁵Ca (3 Ci ml⁻¹) for 120 minutes with the autagonist heing present during the last 30 minutes (control strips without the antagonist).
 - 7. tissues were rinsed for minutes in a large volume of nonradioactive physiological and colution with a without the antagenist.
 - itssues were transferred to a physical site solution containing rightsdata $F_{2\alpha}$ (in 3 minutes (with other states),

the tipenes were thereafter in the to the the second of th

A different method is thinking a Gratchar' plot has been doubled to identify and observation disconservations of the speak (100). This is type of plot was brighted by if all by Gratche (1005) to result the binding of e all sole does to compare the and here one used extensions to product the speak of the state of $\frac{12}{3}$.

ų,

of the 45 Ca taken up by the cells would be present in bound form and (2) that the extracellular Ca^{2+} concentration is equivalent to the free Ca^{2+} level in the solution. Thus the bound ⁴⁵Ca and free ⁴⁵Ca are measured after incubation of the muscle tissue with different extracellular 45Ca concentrations. A plot of bound ⁴⁵Ca/free ⁴⁵Ca vs bound ⁴⁵Ca constitutes the Scatchard plot. This usually results in a curyed plot which has at least two linear components in smooth muscle(100): a high affinity binding site for Ca²⁺ apparent at low (0403, mmol/i) extracellular ⁴⁵Ca concentrations and a low affinity binding site for Ca²⁺ ions (apparent only at high (1.5-5.0 mmol/1) extracellular ⁴⁵Ca concentrations. Sr²⁺ in an appropriate concentration can be used to block the high affinity Ca²⁺ ions uptake site while deaving the low affinity uptake sites relatively unaltered. 'With prespect to these results it is also of interest that Sr²⁺ ions can be used as a substitute for Ca²⁺ in supporting a K⁺ induced tension response more than in maintaining a noradrenaline induced tension response in some smooth muscles(100,106).

Although extracelluar Ca^{2+} is an important source of Ca^{2+} for contraction, some vascular smooth muscles respond to agonists with contraction in the absence of extracellular Ca^{2+} ions(29). In the rabbit contraction produced by high concentration of K⁺ declined rapidly in Ca^{2+} -free medium, whereas those to noradrenaline and histamine persisted with less reduction(107). Extracellular La^{3+} too has a losser inhibitory effect on contraction produced by noradrenaline than on contractions produced by K' depolarization lending further support to the reline of intracellular Ca^{2+} during noradrenaline induced in antime(29). Thus an intracellular pool probably provides the Ca^{2+}

However, the contractions induced by these agonists in the absence of extracellular Ca^{2+} are reduced to one or at the most, two contractions suggesting that the intracellular Ca^{2+} pool is a limited one because it must be refilled from the outside(102). This intracellular pool takes up Ca^{2+} from the extracellular medium by a process sensitive to lanthanum blockade but not to Ca^{2+} antagonists such as methoxyverapamil in the rabbit aorta(108).

Deth and Casteels(109) in some elegant experiments in the rabbit aorta, investigated the effects of different agonists on the intracellular pools of Ca^{2+} involved in contractions as described above. As the method used in these experiments is another technique used in the measurement of Ca^{2+} fluxes in smooth muscle the steps of the procedure as used by Deth and Casteels(109) are listed below:

•

1.

tissue strips weighing approximately 15 mg were incubated for 3 hours in 10 ml of physiological salt solution containing ^{45}Ca , 1.5 mmol/1 (5 x 10⁶ cpm/ml).

- 2. the tissues were then rinsed for 5 seconds in a large volume of physiological salt solution to remove the adherent loading solution.
- 3. the tissues were transferred at 5- or 10 minute intervals through a series of previously vached a intilation with a Ca²⁺ concentration of physiological salt solution with a Ca²⁺ concentration of 1.5 mmol/1. Oxygen was bubbled through the solution to maintain oxygenation of the tissue. Deth and Van Breemen used Ca-EGTA in this efflux medium to minimize backflux of ⁴⁵Ca luting the efflux(110).

. 5

notadrenaline or caffeine or 2,4-dinitrophenol were introduced into the physiological salt solution in the scintilation vials at various times after the commencement of the efflux measurements. the tissue strips were dried and weighed at the end of experiments and the ⁴⁵Ca remaining in the tissue extracted by digestion.

5.

The efflux of ⁴⁵Ca measured in this fashion follows an exponential pattern. Addition of noradrenaline or caffeine resulted in an immediate increase of the efflux which reached a maximum within 10 minutes. In contrast the response to 2, 4-dintrophenol (DNP) appeared more slowly reaching a maximum only after 20 minutes. A similar difference in the time course of action was noted on tension development in rabbit aortic rings in separate studies carried out simultaneously. Further, a plot of the amount of ⁴⁵Ca efflux (estimated from the area under the efflux curve above the control) and the concentration of noradrenaline showed a good correlation with a plot of tension generated against the concentration of noradregaline The intracellular Ca²⁺ pools utilised by noradrenaline and caffeine appeared to be a single pool as under extracellular Ca²-free conditions, a prior transient caffeine exposure eliminated the ⁴⁵Ca efflux produced during a subsequent exposure to Estimates of the ⁴⁵Ca pool sizes and the rates of noradrenaline. exchange for the three agonists suggested that the noradrenaline and caffeine probably shared a single pool while DNP appeared to ⁴⁵Ca from a separate pool. K^+ depolarization utilised during the ⁴⁵Ca loading procedure to increase cellular uptake of ⁴⁵Ca also showed a differential effect on the three agonist induced effects. The ⁴⁵Ca efflux produced by DNF was increased three to four-fold by this manoueuvre while. rather surprisingly, the offlur produced by caffeine and noradrenaline a start at the share study lies a constructed that at Japat two

different pools of intracellular Ca are involved in the contractions produced by the three agonists in the rabbit aortic muscle. Other studies too suggest at least two intracellular pools: a low affinity (rapid release) pool responsive to noradrenaline, histamine, acetylcholine, caffeine, and a high affinity (slow release) pool somewhat sensitive to these agents but more specifically sensitive to DNP(30). A separate study on the rabbit aorta by Deth and Van Breemen(110) demonstrated that although noradrenaline produced an increase in 45 Ca efflux this dould not be reproduced with a second exposure to the drug. Thus the 45 Ca released from the intracellular, store by noradrenaline is unlikely to be taken back into the same store.

Intracellular calcium stores

8

þ.

Although the role of intracellular Ca2+ pools in providing calcium for contractions produced by some agonists is accepted, there is considerable debate as to the exact source(s) of this intracellular Ca²⁺. The possible sites mentioned are the sarcoplasmic reticulum. The mitochondria and the sarcolemum. The surropleanic retionium is developed to a varying extent in different and the concle colle. constituting approximately 2.0 1.5 per cent of the coll stime. The higher values found in the large sinetic as either include a st ufflerer proportions of rough andoplasmic intertion these for storage provide may not be as much as that of the smooth interter turn. Here is if the sarcoplasmic reticulum constituted only 2 par cost for be well volume, assuming that like the reticulum of all interior model of containing 25 mmol/l f (a) a complete relates of the latter will be more that sufficient to produce manifrom actions to of a constitution to 11(1). The volume of series local netting and the area of the end of convolute option the start print of the constant

extracellular Ca^{2+} ions(32,33). As these measurements did not distinguish between rough and smooth endoplasmic reticulum in the different smooth muscles and as the permeability of the sarcolemma is known to differ between different smooth muscle it is possible that this correlation is spurious(81). Surface couplings in smooth muscle are specialised regions similar to those in cardiac muscle where the sarcoplasmic reticulum is separated from the surface membrane by only a 10-12 nm space traversed by periodic electron opaque processes. These have been suggested as sites where depolarization spreading along the sarcoplasmic reticulum is a possible candidate as a Ca^{2+} source, no definite structural evidence for the actual release of Ca^{2+} from the reticulum is available at present(111).

Isolated mitochondria from vascular smooth muscle are capable of accumulating Ca^{2+} and other divalent cations such as Ba^{2+} and $sr^{2+}(81)$. This has been demonstrated by electron probe analysis and electron microscopy. The apparent Km for the Ca^{2+} uptake by mitochondria use 17 µmol(39). This high Km of isolated mitochondria for Ca^{2+} suggests that mitochondria are unlikely to regulate cytosolic free Ca^{2+} concentration during the normal contraction relaxation cycle. At a intracellular free Ca^{2+} concentration of 1 µmol/1 (which is helieved to produce about 50.8° per cent of the maximum contraction in smooth muscle) the offact of the mitochondrial transport system would be visual.

An othe nation pos thildty is that the inner surface of the plasma membrane is the principal storage site from which Ca²⁺ ions are released during contraction. Suzuki, Gugi and co-workers(112,113) there is a function of the microgropy) that is calls treated with potassium oxalate, the intracellular Ca^{2+} was localised to the peripheral cytoplasm adjacent to the plasma membrane in the relaxed state: following drug induced contraction in the absence of external Ca^{2+} , the Ca^{2+} stain was found diffusely throughout the cytoplasm. This suggested that the Ca^{2+} bound to the inner surface of the sarcolemma was released into the sarcoplasm during the contraction. The role played by the surface vesicles or caveolae in possible Ca^{2+} regulation by the sarcolemma is not clear at the present time.

Calcium sequestration during relaxation

0

From the foregoing discussion it is apparent that there is considerable controversy about the source of Ca²⁺ for activation in smooth muscle. However, there is an even bigger debate about how 'he free Ca²⁺ ious are removed from the sarcoplasm during relaxation in smooth muscle Relaxation in smooth muscle as in striated muscle is brought about by a lowering of the arcoplasmic free Ca2+ leading in turn to a decreased actomyosis Albase activity and a detachment of the cross ridges. This lowering of the carcoplasmic Calicon be brought about 's account processing afmiliar to these cheared in cardiac muscle First, at the parcolomna the finand Ca current that ecoure during depolarization is colleptering(30). This is brought shout by an increase in remembrility to P¹. Which follows the rise in Ca²¹ and Na nermephility. In most muscle with a finn norantials, and occurs atmittance ity in amonth miscle shich show araled depolarizations outs . The itse in intravalliter Calimon that is the the trigger for the the bear's presentifit which results to an outward 'restiffeting' R' current. To resulting it the ration reduces we mean permeability to the contraction to the same after a two to the contract

accumulated in the sarcoplasm during the inward current has to be removed from the vicinity of the contractile proteins. This Ca²⁺ is either extruded out of the cells at the sarcolemma or taken up into intracellular storage sites.

Transport of Ca^{2+} ions out of the cells is an energy consuming process as it involves a transfer against a large electrochemical gradient (There are two principal mechanisms through which Ca^{2+} is believed transported out of the cells: (1) the Na⁺/Ca²⁺ exhange system and (2) the Mg-ATPase dependent extrusion system. The features of the Na⁺/Ca⁺ exchange system can be summarised as follows(115):

- 1. Na⁺ ions move down its electrochemical gradient from the extracellular fluid to inside the cell.
- 2. coupled to this inward movement of Na⁺ ions, Ca²⁺ ions are extruded from the cell.
- 3. the energy liberated by the inward movement of three Na⁺ ions is required to move one Ca^{2+} ion out of the cell.
- ATP may modify the kinetics but is not a primary source of energy for the transport system: the energy being provided by the electrochemical gradient for Na⁺ ions.
- 5. for the pump to function the electrochemical gradient of Na⁺ has to be maintained via the Na^+/K^+ membrane pump. ATP is of course. utilised for this.
- the transport system can also move the ions in the reverse direction (i.e., Na⁴ outwards and Ca^{2+} inwards) across the membrane when the gradients are suitably altered, because of the competition between Na⁴ and Ca^{2+} at sites on both sides of the membrane.

The evidence for the existence of the Na²/Ca² pump in smooth
dependent ⁴⁵Ca fluxes have been shown(114). In vascular smooth muscle lowering of external Na⁺, for instance by replacement with Li⁺ leads to an increase in tension. Further, introduction of ouabain, the Ma_{+}^{+}/K^{+} -ATPase inhibitor, often leads to contraction in vascular smooth muscle associated with an elevation of intracellular Na⁺ concentration. `Both these manoeuvres lead to the loss of the Na⁺ gradient which would inhibit the Na⁺/Ca²⁺ pump as proposed. Re-establishing the Na⁺ gradient by re-introduction of the Na⁺ or removing ouabain in the above experiments leads to a relaxation. Although the evidence is strong that Na^+ plays some role in the control of intracellular Ca^{2+} , it is never theless quite clear that an inwardly directed Nat gradient is not essential for the tissues to be able to regulate intracellular $Ca^{2+}(83)$. Brading(116) demonstrated that smooth muscle can not only remain in the relaxed state but also contract and relax again to stimuli in the absence of a Na' gradient, although the response may be qualitatively different from the control state. Other workers too have found similar results(114) Because of these inconst tencies, AIF dependent extrusion of Ca2+ jour has been suggested as an iternation to the Ma /Ca2 transport soutem. This Ca2 pump to believe to be fuller to that describe! Is not blood calle by Cabot we still) - Again, direct evidence for the strenge of such a (?' pump to sacular smooth must is lacking("). It is possible that both systems described it is are present in amount meaning. [1] has been suggested that the Call All son against again to constant on the sareopleamle rett when membrane the structure fully in Ca²⁺ from the garciplann in 11 is taken take the restantion of other is "gapping and and easy the state is a entry attributes ftutt · ., '

82 -

 l_{ℓ}

Sarcoplasmic reticulum, mitochondria, nucleus and the inner, surface of the sarcolemma are the possible intracellular storage sites for Ca²⁺ ions. The majority of the studies designed to investigate the possible Ca²⁺, storage sites have used differential centrifugation of smooth muscle homogenates to isolate subcellular fractions that have been proposed to be potential sinks and sources of activator calcium. Mitochondria are isolated by sedimentation between 9500 and 15,000 g and a microsomal fraction obtained at 40,000-15,000 g(29). The 'latter fraction, however, contains a mixture of sarcoplasmic⁵ reticulum and sarcolemmal fragments. A fundamental problem in identifying specific sites of calcium accumulation in smooth muscle has been the fact that it has been difficult to isolate either pure or enriched fractions of sarcoplasmic reticulum and sarcolemma. The greatest difficulty has been reliable separation the reticulum fractions from sarcolemmal of fractions. This subject has been recently reviewed by Allen and Bukoski(94). Mitochondria have been isolated in sufficiently pure form during the last few years allowing the characterisation of the Ca²⁺ accumulation by these organelles. These studies suggest the presence of an every dependent mitochondrial Ca2' transport system(81). As ment' not before, this system has a few affinity but a high capacity; It is unlikely that mitochondria rlay a major role in the thing sequestration of Ca fors during relexation in smooth muscle. However, insufficient data is available at present to definitely exclude a greather role for mile to this to a sub-type of mosth muscle such as the vertate(RT).

difficult to draw a firm conclusion as to the membrane fragment responsible for the transport system(94). The requirements for the uptake of Ca²⁺ by microsomal fractions in emooth muscle are ifmilar to those for stillated muscle with both ATF and Mg2+ jone being required This uptake 'spends on the temperature. I fig higher at 37 ' time at off optimum pH values reported have been 7.6 and 7.8(2)). Although sercepteenic reticulum is a likely alle for the Call seque tration during relaxet mit to has we have I thing is nearly had . It has been rearristed that he been are use on function of the printpal intra illular volcome elte for in hurlin contracti e alth the for being university some forel by the man opta of the performance (10). Current antidence asygnate that the alegand then little ellutar et ter during a strager an induced by some upor fairs to a 21 the modifier to module not search and hank to the a state only the state tue, contractions can be indeed, I be a conter to the free we draw; in $\frac{k^{2}}{2}$ the efflort constraints in all each to be the first of an exponsive is a contraction dependent of the contraction of the second second entering to the terminate of the spectrum of the second to the second state of the second sec encounter the theory of the theory and the second Contraction of the second s 1 $\mathbf{t} = \mathbf{t} + \mathbf{t} +$

. . .

.

12

's for 14 mile 1 + for in again i i i i sector for file 1 m
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state hold the sector state
's hold the sector state hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state
's hold the sector state hold the sector state hold the sector state
's hold the sector state hold the sector state hold the sector state
's hold the sector state hold the sector

produced by the action of the enzyme adenylate cyclase on ATP. Beta adrenergic stimulation is known to cause an activation of the adenylate cyclase system with the production of cyclic-AMP. The degradation of cyclic nucleotides including cyclic-AMP is carried out by cyclic nuleotide phosphodieternsen. These enzymes or family of isoenzymes, catalynes the hydrolysis of the 3' bond of the 3':5' cyclic phosphate to give the 5' monophosphate. Cyclic-AMP exerts most of its effect through the arrivation of protein kineses located in the cells(119). The protoin linages are composed of two sub-units: the catalytic sub-unit and the solutions and unit. The lattor inhibits the action of the former on long as it remains bound to it. Cyclic-AMP activates the enzyme by combining with the gogulatory sub-unit, thereby causing its dissociation from the estabeth sub-upit of the enzyme. The activated prototo blane can throupon phosphorylate a variety of substrate proteins ' transfer of a gamma thosphate group from ATP, e.g. glycogen synthese, phosphorylase kinase, etc. One such substrate protein phosph vlat d by the cyclic AMP dependent protein kinase, is the enzyme myoria light chain lines. Thorobarylatics of the light chain kinase in his affinite for calmodel's leading to decreased phosphorylation of my offer light followed thus a terrage to operation frefer above. other than the disting of the termination to be one and the Fill of poly strike synthe AM - Induces colonation machantam 1 ... core th · 12010(120) the other excledion maps of him been president of many of a film by the or plaget anticulum tarlog relevant of to musth much inclution for mosth much microsomer with with Although the scheme to a structure the energy-dependent G_{3}^{2+} -1 $v = t^{-1}$ SA Fra , · TINAN "CTUE MAL "act frie 1 -

plasma membrane as mentioned before. It is possible that cyclic-AMP activates the Ga^{2+} -ATPase in either membrane which, in turn, could increase Ga^{2+} uptake into the reticulum or the extrusion of Ga^{2+} across the plasma membrane or both. It has been suggested that cyclic-AMP is a mediator of the relaxation produced by certain drugs. The evidence for the cyclic-AMP by pathesis () be summarised as follows ()0.151).

.

- accompanied by an increase in cyclic AME lovely.
- A quantitative and a temporal correlation between the elevition of evolic AMP and the relaxation has been shown with forty nal no(114).
 A blocking agents induce both the second sec
- Pelanatory response to Prefres rate ethnilation from entlated by agents that the the second of the Press of the second of the se

Inhibit we do the phodesterage theory same relaxation of amonth much with an inclusion to each A . In the element of the action is inclusion to each A . In the element of a strict is inclusion the second of the theory of the second of the track of the second of t

th min

 As mentioned above, two possible mechanisms of action for cyclic AMP induced relaxation has been demonstrated in smooth muscle.
 Other drugs which produce relaxation in smooth muscle also increase cyclic-AMP levels, eg. prostacyclin, adenosine(123). 87

Although the above evidence appears substantial, some doubts remain the results have not been consistent from different centres(125). instance, during a K⁺ depolarization, propranolol, while blocking teorrenalize induced rise in cyclic AMP, leaves the relaxation the produced by the isoprenaline intact(126). In demonstrating that cyclic nuce villes have a function in a cellular response, it is necessary to show not only that evolve nucleotides are altered by the same agents that evoke the threfological response, but also that these alterations correlated quantitatively and temporally. are Unfortunately, the majority of the studies in support of a role for cyclic-AMP in smooth munchs relevation do not fulfill this requirement. Cyclic-AMP levels have been often measured at only the time point of contraction or relaxation, at only one concentration of the agouist or under incubation conditions when the time was not under a mechanical tension(118). Interpretation of the love gations our do demonstrate the direct relayant effects of yolds AME and difestyryl cyclis AME have is and difficult as positions, one raing the ability of the maleotides to scool the allo of action, and quicktions concerning the effects of bud size' prove to or imprision have been coldom dealt with, in these station thus, controle using advicetor, butyrate, other nucleoridee and in alble contaminants larg not been tested adequately (1'8). This receiving the particularity furnitant as high concentrations (100-1000 for the the methodist 1 of or evelte AMP and Hbutyry]

The have the the method to theme at stick the forther

*

complication arises because cyclic nucleotide derivatives are also capable of inhibiting phosphodiesterase activity. Thus a derivative of cyclic AMP might cause elevation of endogenous cyclic-GMP and vice With regard to the studies using phosphodiesterase inhibitors, · versa. again, no clear demonstration of the relationship between the relaxation and the time-course and magnitude of the changes in cyclic-AMP levels has been carried out. Cyclic~AMP is helieved to increase the uptake of Ca^{2+} by microsomal fractions of smooth muscle via a protein kinase. However. different investigators have resulted studies by in inconsistent findings: some showing phoshorylation and uptake of Ca2+; others showing phosphorylation with no C_{λ}^{2+} binding or vice versa; and rarely no effect on phosphorylation or uptake of Ca²⁺(118).

role of cyclic-GMP in smooth With regard the to muscle contraction/relaxation, less information is available at the present Early studies reported an elevation of cyclic GMP time. levole accompanying contractions in smooth muscle produced by a variety of agonists including prostaglandin $F_{2\alpha}$, acetylcholine and alpha advenergic agents(118). Further, it was shown that Color was becomeriv for the abile effort. On the best of three studies cylife Off val considered ng a monitation of ensely music contractions. Hereiner enheaguent studies 'omentaried a class diene tota holynon the contraction produced and the exclipion levels. In addition, pitroplycerine, at represente and accession of regen containing compounds have been shown to pushing lings increases in cyclic-GMP (but not of cyclic AMP) with an acconstruction relayanteers (128). These agains do not generally require tal for their efforts on synlin GMT levels. Thus during the list fow veers solt (MT has bee vestors ed a prele er of amouth me ela Pt7 / ··· · · · and the second second prove for

absence and presence of M&B 22948, a potent inhibitor of the hydrolysis of cyclic-GMP(123). Acetylcholine, on its own caused a contraction of the arterial strip accompanied by an increase in cyclic- GMP. In the presence of M&B 22948 the dose response curve to acetylcholine was shifted to the right accompanied by a further increase in cyclic-GMP levels. Thus, a higher level of cyclic-GMP was associated with a lesser contractile response. This study casts serious doubts on any role for cyclic GMP in the contractile effects produced by acetylcholine; it suggests a possible role on a relaxatory effect for cyclic-GMP. Further support for the latter is provided by the fact that methylene blue, while attenuating the increase of cyclic-GMP produced by acetylcholine in the bowine coronary artery, increased the contractile response produced by the drug(123). Although an interesting association between relaxation and cycic-GMP in smooth muscle has been demonstrated, no definite proof for a cause and effect relationship is available at the present time.

In summary, it could be said that it appears likely that cyclic-AMP is involved in the relaxation in smooth muscle produced by at least the beta adrenergic agonists. However, it has not been conclusively shown that this cyclic nuceotide is an absolute necessity in the mediation of the relaxation, although the association is strong. With regards to cyclic GMF, much more work has to be done before any conclusion can be arrived at. Further, it is not likely that all smooth muscle relaxing drugs would act through a cyclic nucleotide methanism.

ADRENERGIC NEUROEFFECTOR INTERACTION

Several excellent reviews on the subject have been published during recent years (15,73,129). Almost all blood ressels are innervated with roat gaugitude computation parger although the density of innervation

varies widely. As in other adrenergically innervated tissues the ability to synthesise, store, and release noradrenaline resides in the adrenergic nerve terminals found in the walls of the blood vessels. Transmitter synthesis and storage (Fig. 10)

Noradrenaline and adrenaline are synthesised from the precursor amino acids, phenylalanine and tyrosine(130). The sequence of events confirmed the enzymes involved identified and been and has Tyrosine, which is generally available in the body characterised. fluids is taken up by the nerve terminals and converted to 3,4dihydroxyphenylalanine or DOPA by the enzyme tyrosine hydroxylase. This enzyme, present in the neuroplasm, regulates the biosynthes is of the catecholamines, as the hydroxylation of tyrosine is slower than the Several physiological mechanisms are present to subsequent steps. modulate the activity of this enzyme and thereby regulate noradrenaline In the next step in the synthesic pathway DOPA is synthesis. decarboxylated to form dopamine in the neuroplasm. The dopamine is then taken up by the storage vesicles contained within the adrenergic nerve endings, where it is converted to horadrenaline by the enzyme dopamine As inpamino f hydrovulaso is not 8 hvdroxylann. present in the neuroplasm, dopanting must be taken up into the vertels before uoradramaline could be formed. In the adrenal modulla, most of the noradrenaline leaves the granules and is methylated in the evtoriasm to adrenaline and then rementers a different group of intracellular storage residue, where it is stored until released. In peripheral sympathetic nerve sudjugs the enzyme phenvlethanolamine Nomethyltransferase(PNMT) is absent and the principal transmitter is notadrenalize. However, shout the part of the strength the effective strength the strength of an integral of



Figure 10: Summary of the steps in the synthesis of noradrenaline and adrenaline in the sympathetic nervous system. Steps 1-3 take place in the neuroplasm and step 4 in the adrenergic storage vesicles subsequent. to transport of dopamine into the storage vesicles from the neuroplasm. consist of adrenaline; this probably reflects adrenaline from the circulation that is taken up into the herve endings (when plasma concentration of adrenaline is raised) and released subsequently together with noradrenaline.

It should be noted that some of the enzymes involved in the pathway of catecholamine synthesis are quite non-specific, capable of acting upon many other endogenous substances as well as certain drugs (130). instance 5-hydroxytryptamine (5-HT, serotonin), tyramine and For histamine can be produced by L-aromatic acid decarboxylase (or DOPA decarboxylase) from their corresponding amino acids. Tyramine, in turn can be oxidised to octopamine, the phenol analogue of noradrevaline: while octopamine is present only in small amounts in mammals, it is probably the major adrenergic neurotransmitter in certain invertebrate species. The antihypertensive drug a-methyldopa is also metabolised by this pathway forming α -methyldopamine and subsequently the 'false transmitter" α -methylnoradrenaline. The enzymes involved in the pathway are probably synthesised in the perikaryonal cell bodies of the adrenergic neurones and then transported along the axons to their terminals. A slow transport system (1 to 3 mm per day) and a fast (1 to 10 mm per hour's system have been documented. Ivrostos b troxylase fo helieved to with the the close transport avelow and depending $-\beta$ hydroxylage the factor system. Ifferet chalar system pres of factors are move tlay a role in this (ast transport overem (130,100)

Several mechaniams regulate the synthesis of noradrepaltic via modulation of the twosing believelase activity either on the short term or on a long-term basis(73). First the end product noradionaltic midulates the activity of the curves by feed back fabricities. This came

mechanism may account for increased activity of the enzyme with accelerated synthesis of noradrealine during nerve stimulation. The total geuronal content of noradrenaline need not necessarily be decreased for the enzyme to be activated: thus a specific compartment of noradrenaline in the cytosol probably participates in this regulating In addition, the influx of Ca^{2+} that takes place during mechanism. nerve stimulation may also induce the enzyme. This may help to accelerate the synthesis of noradrenaline during continued sympathetic nerve stimulation. Trans-synaptic mechanisms may also inhibit (via presynaptic α_2 receptors) or activate (via pre-synaptic β -receptors) the Other compounds such as angiotensin, prostaglandin and enzyme. acetylcholine too may regulate the tyrosine hydroxylase activity in a similar manner(130). Acute (short-term) regulation probably involves allosteric effectors that alter the affinity of the enzyme for cofactors, substrate or end-product inhibition. Long-term regulation occurs by other mechanisms such as alteration in the synthesis of the enzyme and covalent modification of the structure of the enzyme. The effect of these long-term modulations usually takes a few days to develop, but they can persist for several weeks; the delay probably reflects the time required for transport of newly synthesised enzyme " m cell body to the nerve terminal.

Moradrenaline control of the walls of blood vessels ranges from $0.00 \ \mu g/g$ wet weight in the rat sorts to nearly $10 \ \mu g/g$ wet weight in the durinous pig uterine artery(73). Although such measurements may provide a rough indication of the density of adrenergic innervation they can be misleading, especially when expressed as a function of tissue weight. Noradrenaline present in adrenergic verve endings is stored in

granular vesicles (also called storage vesicles or dense-core vesicles) with the vesicular membrane providing protection from the intraneuronal enzyme monoamine oxidase. The noradrenaline in the vesicles is complexed with ATP and proteins; of which chromagranin A is the most important. ATP and noradrenaline are present in a molecular ratio of 1:4(130). Dopamine- β -hydroxylase is also present in the vesicles. The vesicles, at least in their initial state, are believed to be synthesised in the cell body and transsorted down the axon.

· · ' •

Two types of storage vesicles can be identified by electronmicroscopy in advenergic nerve endings; small dense-core vesicles, 30-60 nm in diameter and the large dense-core vesicles, 60-120 nm in diameter(132) The large dense-core vesicle can constitute 5.50 per cent and the small dense core vesicle 50.95 per cent of the vesicle population in an advancegic nerve terminal, although the fatter predominate in most cases. In general, the larger enimals seem to possess higher numbers of large dense-core vesicles in their adrenergic nerve endings. It should be understood that a much smaller percentage of large dense core vesicles may provide a large core volume (i.e. storage capability for noradinalitie) at the normal and ingly. Illing, in a nerve ending containing to is per oral large vesicles and RS 20 per cent am 11 west las, the core volume for the two types of "galilas oppear to he equal. In man, the large mest les may comprise in to 80.90 per cent of the total storage expanits in certain advanced variabilities(132). Small vesicles are bollowed to contain up to a 1000 molecules of noradrenalize proventile of ange verticity may con ato up to 16 000 molecules to represent a stale or about a con of the dischard had by compared the second of the second se

. . . 94

density peak of noradrenaline is enriched in small dense-core vesicles and the high density peak is thought to correspond to the large densecore vesicles, the correlation between electron-microscopy and the biochemical methods is by no means certain(73).

Transmitter release

The stored noradrenaline may exit from the nerve endings via the neuroplasm or directly by exocytosis(15). In the resting state, a small amount of the stored noradrenaline diffuses into the neuroplasm continuously through the vesicular membrane. Most of it is deaminated by the enzyme monamine oxidase (MAO) to 3,4 dihyhydroxyphenylglycol (DOPEG), but a small fraction escapes into the synaptic cleft as intact noradrenaline together with DOPEG and the other metabolites formed within the nerve ending. The minute amounts of intact noradrenaline leaking out produce miniature excitatory junction potentials which can be recorded. However, the amounts released are usually not great enough to produce any active tension in most arteries. Drugs such as tyramine, amphetamine, ephedrine and guanethidine are taken up into the adrenergic storage vesicle and lead to the displacement of the noradrenaline into the neuroplasm. They probably evert their effect by having a greater affinity for the storage proteins than noradrenaline(15). As with the spontaneously leading transmitter, part of the displaced noradrenaline is deaminated to DOPEG but a sizeable fraction is released as intact noradrenaline to produce effects on the smooth muscle. Pharmacological displacement of the transmitter, unlike the exocytotic process, is not dependent on an indrease in neuroplasmic free Ca²⁺ concentration. As the indirectly acting sympathomimetic drugs have to be taken up into the makes terminal by untake, (acc below) before they could exert their

effects, their action is blocked by inhibitors of uptake₁. On the other hand, inhibitors of MAO, by preventing the deamination of the displaced noradrenaline, potentiate the effects of indirect sympathomimetic agents.

S. 1

During increased sympathetic activity, propagated action potentials spreading along post-ganglionic sympathetic nerves result in the release of noradrenaline in exocytosis at the adrenergic verve corminate. Ar the nerve terminal the inward current of the action potential is articl by Nat and Can'towa. Increase in the neuroplasmin Can't regults in the exocytoels of the storng vestiles with release of poradrenaling and other vestcular contents (such as the enzyme decambers hydroxylase) into the expertic clift. In isolated emooth muste, transmural entry stimulation filmer release of prodremaline by ecceptoria, by excitation of the intramutal nerves contained in the isolatel much with induction of active potentials(?). High " estations on on feolated emonth muscle cont infor advanced out on the leading to depolarization of the perce endings with release of the stored noradrevaline by excepteals: as with an action potential an invard Cal current as untinted it. The deforance 'on product' by the 2 tong subscripting the relation of the model frequence to should be used that R' and execution indices of external sections in factors and the munche(1)6. The active point of this weat a lease of noradi alter t blo bet by controdote in the fait in Connet inhighter(113) However the release produced by high " i must shared by tetrodotoxin as the P causon a literit depolation of the how topole institut withe The setting of post of the later is the the set of the set of the set

The second se

•

96

- 3 ''

channel blockers verapamil, prenylamine and D600 although at concentrations higher than the usual therapeutic ranges(137). This results in a decrease in transmitter release.

During exocytotic release of the transmitter, only a small (up to 5 per cent) percentage of vesicles within a varicosity release their contents Further, any single varicosity does not respond to every stimulus by releasing transmitters; thus the release is intermittent and holieved to occur once every 7-8 action potentials(132). It is not known whether the intermittency of release is because the varicosity is nor allows invaded by the action potential, or because it is incapable of (or refractory to) transmitter release for a finite period of time following release. Negodrenaling is released from both large dense-core and small dense-core vesicles during exocytosis. The release of transmitter by executoris from the "wo types of vesicles has been. observed in electron micrographs taken following superfusion fixation of fine smooth movel' strips during transmural nerve stimulation(132). Howaver, there is a debate at the present time as to whether each recipion releases all its contents (total content release) or only part of the intents with exceptions (retrial content release)(73). Vesicle to could growld be a hencegery occurrence with the latter theory alth uph 't would take place clop aft total content release and thus dues not constitute of dence against the former. There is evidence that weater recording comments at norve terminals including adrenergic aynapses du ing transmitter releaso(138). Extracellular markers such as tory redish corovidase have been demonstrated inside vesicles following ""ter less indu i is transmural serve stimulation.

in the second of the second deference (139)

÷ 97

demonstrated that the adrenergic varicosities contained approximately 93 per cent small vestcles and 7 per cent large vestcles; 73 per cent of the former and 9" per cent of the latter contained electron-dense cores. The core sizes varied from 50-80 per cert of the vesicle diameter in the small dense more vesicler and were more uniform and typically measured 80 per cont in the lass denses re vestcles. In muscle segments fixed impodiately following transmurph nerve stimulation a 60 per cent reduction in the shall vestiles with no change to the large vestels important diversed. Moreover, in the suill readels that remained, cut is per ce contained dense where and in was of these the core its was multiplic commendation the provident the ported of the decrease to the call of the production was accompanied adjust the room of the second state of the interstates. The amail matche made of and the second ender the protocol the returned to control a pltic " we all the the we from had along a construction of the second second program in the second seco of Imil at feat to Imil is the state of the st demonstrates and a second of the second of and prove the second المراجع والمنافع ومعتورة المراجع والمراجع والمنافع والمنافع والمنافع والمنافع والمنافع والمنافع والمنافع والمنافع r en li and the second and standard the second end of the second and the second of the transformer of the second the state of the second st the second se the second se • • • • a state of the second second

1 1 1

•• •

in the mean perimeter of the varicosity following the transmural nerve stimulation. (3) However, vesicle recycling (or the formation of entirely new vesicles) at the cell membrane of the varicosity, appears to fake place during transmitter release, as evidenced by the presence of horseradish verioxidase in the vesicles following the nerve stimulation. (4) Formation of hew vesicles at the synaptic membrane appears to continue following the stimulation period: the mean perimeter of the mathosities and the vesicular number returning to control values during a fix hour next stimulation period. (5) Replanishment of the mathos in the vesicles is not dependent on to uptake of itemative eleand during nerve stimulation.

Altimate only ampli dense core restales were observed to be released to the above study others believe that large-dense core sectores are also functed in transmitter release. Nelson and Malin (C.133) lamonatisted that the indirectly acting sympathonimetic agent to amina closed aqual depletics of coradionaline from both high and the density frontians instated from success density gradients. Further, some bollows that the stall dense core vestele are formed at the shapile moments schools at the enceytotic release from the large tones core set of the 150). In allow offer hyperboats to that direct The second the large lines of modelies within the caricosity gives the to be enclose on the there is the forest in the elifence for the first having to the prost time(132). The former herethering the operated by the fact that very little depending A hydroxy free and ther eclusive materna (which could have been released at the fri that are percent in the small dense core -1 is a final state of the second \mathbf{y}^* . Here, $\mathbf{x}_{\mathbf{y}}^*$ is a second seco

correlation, or lack of it, between the released catcholamine levels and the dopamine- β -hydroxylase levels during sympathetic stimulation. As both populations of vesicles contain the transmitter and only one type contains the enzyme, the ratio of the released transmitter to the released enzyme would depend on the relative proportions of the two populations of vesicles: this differs from tissue to tissue as well as with the functional status even in a single tissue as explained above. The hypothesis could also explain how antigenic activity right be retained between the large and amall dense-core residles. At the present time, the exact origin of small denon core vestcles is still uncertain Large deuse-core seicles, on the other hand, are known to be formed in the second cell body and transported down the area to the nerve terminel. During this passage the vestimilar content of nor-drengline increases while the depamine forbydrows have activity and the AIF content of the vestele remain unchanged(141).

Another controversial area in neurotransmis for in smooth muscle is the d'fferent sub-cellular compartments involved in transmitter release: these whicelluly compartments may be different types of vesteles (e.g. large and amolt dance core), a sub-portulation, of a single type of vestcle (e.g. veetcles which have gone through a cycle of exos and endneytosta bring pofilled with nonintransmitter, and preferentially released), is different poole of the transmitter within a vestele. for (-iborted related it putty south intent to the was demonstrated to a to a challing of an incontinue "apty of al(147) domo at atal the come, to advecce a constant of the last terms of eple of The three and the second and the second second

1.15

nerve terminal to ¹⁴C-noradrenaline). Stimulation of the splenic nerves at a high frequency (30 Hz) resulted in release of ¹⁴C-noradrenaline having a greater specific activity than that found in the spleen indicating that the 'newly synthesised noradrenaline is selectively When spleens from cats which had been treated with $^{3}H^{-}$ released. noradrenaline (inspead of ¹⁴C-tyrosine) were perfused, the specific activity of ³H-noradrenaline initially was similar to that in the spleen. With continuous stimulation the specific activity fell to about one-third that found in spleen. If the perfusing solution contained α methyl tyrogine (an inhibitor of noradrenaline synthesis), the initial release was not altered. At later times however, only about one-third as much noradrenaline was released and the specific activity of the released ³ H-noradrenaline was similar to that in the spleen. These results suggest that, during continuous rapid stimulation, it is the new synthesis of the transmitter which plays the key role in maintaining a stable output of noradrenaline. It should be noted that, experiments comparing the specific activity of released compounds with that in the organ face a number of difficulties which may lead to erroneous findings(139). For instance; the homogenisation procedure (utilised in measuring the specific activity in the tissue) used may lead to the release of the transmitter. Mechanical shearing that perce endings are exposed to during there disruption and homogenization will (in the presson a of California the release of trapemitter from a large and a of sectore

A study by Hugher and Roth(143) in the rabbit portal wein and yaka loferanti whold 'a lifere t result from the study of Kopin et al using a <u>combination</u> of 3 H-noradrenaline and 14 C-tyrosine. The output per pulse of newly synthesised 14 C-noradrenaline remained constant as the train length of TNS was increased while the output per pulse of 3 Hnoradrenaline increased under the same conditions. This phenomenon was independent of the stimulus frequency. Thus, in this study, the newly synthesised noradrenaline did not appear to be preferentially released during stimulation. Although this study, too, is not without criticism, it is apparent that different sub-cellular compartments of noradrenaline exists in adrenergic nerves. Further, these compartments are probably mobilised in a differential faction during stimulation of the sympathetic nerves.

A model summarising the characteristics of catecholamine synthesis, catecholamine pools and permeability was put forward by Klein and Lagercrantz(132). According to the model the large densercore vesicle is believed to contain two noradrenaline pools; a readily saturable fast-release pool for newly synthesised noradrenaline, which exclusively accumulates in this pool and a slower release, ATP-facilitated uptake pool. The fast release pool may be localized at the vestoriar membrane together with the enzyme dopamine β hydroxylese which is localised at the inner surface of the vestoular membrane. Popamine is taken up into this pool by a high affinity mechanism which is relatively unaffected by noradrounting and doos a functive ATE. On the other hast the slow release post takes up poradionaltus and dependice by a Mg²¹ AIr facilitated uptake contom where noradrenaline competes very effectively with departmen. The restoral membrane too contains a corrier meliated uptable for attaching them ("pt 'e_). This uptake ave' maped be large t and z is a territory on the second state of the second system is a second system.

similar, high affinities (K_m approximately 1.0-1.5 µmol) for noradrenaline. The small dense-core vesicles too have an Mg²⁺-ATP facilitated uptake system but the Km value is much higher (approximately However, the Km of the intraneuronal metabolising enzyme 22 µmol). monoamine oxidase for noradrenaline is approximately 100 µmol. Thus, both vesicle types especially the large vesicles compete very favourably noradrenaline in the neuroplasms compared to potential for free inactivation by monoamine oxidase. Further, a considerably higher level of free noradrenaline in the neuroplasm is also probably needed to inhibit the induction of tyrosine hydroxylase (end-product feed-back inhibition, see above), the rate-limiting enzyme in the synthesis of catecholamines (Km for tyrosine >10 μ mol). Thus the events during the synthesis and storage of noradrenaline in the adrenergic nerve endings can be summarised as follows. With the activation of noradrenaline synthesis by dopamine-β-hydroxylase in the large dense-core vesicles, there ensues a rapid, ATP-enhanced overflow of newly synthesised noradrenaline from the fast release pool into the neuroplasm. The noradrenaline in the neuroplasm is taken up by the ATP-facilitated uptake system into the dense-core vesicles, which may have been depleted and are ready to be re-used or which may be locally formed from specialised endoplasmic reticular elements in the nerve terminals. With the relative saturation of the above uptake system with noradrenaline, neuroplasmic levels of noradrenaline would rise and "turn-off" the two sine hydroxylase activity.

Although the above mentioned model is both a plausible and an attractive one more investigations are needed before it can be accepted in total.

The noradrenaline released from the varicosities enters into the synaptic cleft between varicosities and the smooth muscle. The synaptic cleft width can range from 20 nm in the guinea pig vas deferens to 1900 nm in the rabbit pulmonary artery. In general, in the circulation, the smaller the blood vessel, the narrower the cleft, although there are exceptions(144) with the cleft in the canine saphenous vein being about 100-300 nm. Although neuroeffector separation is quite small (15-25 nm) in most non-vascular synapses, there are exceptions; thus the closest neuro-muscular distance in the longitudinal muscle layer of the. intestine is about 100 nm(55). The neurogenic response of smooth muscle can be related to the width of the synaptic cleft and the density and distribution of the nerve varicosities. Blood vessels containing a dense innervation have a greater maximum response (in relation to the maximum response to exogenous noradrenaline) during maximum nerve activity and a steeper slope in the frequency response relation. For instance, in the rabbit proximal saphenous artery and in the rat portal vein with dense adrenergic innervations, the maximum response to TNS was 85-90 per cent of the maximum contraction produced by exogenous noradrenaline. In contrast, in the rabbit pulmonary artery which has it nerves confined to the adventitio-medial junction with wide clefts, the maximum response to TNS was approximately 50 per cent of that to exogenous noradrenaline(145). On the other hand, a negative correlation is found between the innervation density and the ED_{50} to exogenous noradrena]ine. Bevan determined the concentrations of noradrenaline inside and outside the cleft during the transmural nerve stimulation in terms of steady-state exogenous noradrenaline concentration. In the rabbit pulmonary artery with a wide cleft the two estimates were similar ter a ser a se المراجع والمعرو

with 1.5 x 10^{-7} and 5.5 x 10^{-7} mol/l inside and outside the cleft respectively. With a narrow cleft as in the rat portal vein the intrasynaptic concentrations were estimated as high as $1.5 \times 10^{-5} \text{mol}/1$ while the outside was estimated at 6.0×10^{-9} mol/l during stimulation of the adrenergic nerves(146). Thus, a high transmitter gradient may exist between the inside and the outside of the neural cleft when the This is probably due to a mechanical synaptic distance is small. barrier to transmitter egress with a narrow eleft ... These studies, also. suggest that the noradrenaline must be released only from the area of the varicosity facing the 'smooth muscle, rather than from any part of the surface of the varicosity as otherwise such a big gradient is unlikely to be produced. It is this area only which is usually devoted of a surrounding Schwann cell sheath. It has been observed that the neuronal uptake of tritiated noradrenaline (per unit noradrenaline content) diminishes by as much at 75-80 per cent with diminishing synaptic cleft width(144).

Transmitter disposition and termination of its effect

Diffusion, neuronal uptake, extraneuronal uptake, tissue binding and catabolism constitute the important routes for disposition of released noradrenaline at adrenergic nerve endings. The relative importance of the different pathways varies depending on the density and pattern of innervation, and the width of the synaptic cleft.

In blood vessels with a wide synaptic cleft the immediate movement of the released transmitter into the perisynaptic area is governed by diffusion. However, in a narrow cleft, the movement is restricted within the cleft, and diffusion outward can only occur, from the junction of the synaptic slit with the 'general' extracellular space.

Once the transmitter reaches the perisynaptic area it diffuses in all directions, although initially there is some tangential and longitudinal movement of the transmitter within the nerve plexus due to the scattered sites of release. The transmitter reaches either the tunica adventitia or the tunica media by diffusion, and its distribution between these two layers depends on their relative thickness and on their resistance to transmitter diffusion. The escape of tritiated noradrenaline from the adventitial and intimal sides of the blood vessel during TNS has been investigated by Bevan and Su(146) in the rabbit thoracic aorta and rabbit ear artery. An isolated segment of vessel was perfused (to estimate the overflow of transmitter from the intimal surface) as well as superfused (to estimate the overflow from the adventitial surface) with buffer solution, during transmural nerve stimulation - after labelling the noradrenaline stores with ³H-noradrenaline. In the steady state, in the presence of phenoxybenzamine, the ratio of intimal to adventitial outflow was 0.1 for the pulmonary artery and 0.16 for the ear artery. Presumably in vivo, just as in vitro, there are large sinks at the adventitial and intimal surfaces, the adventitial overflow of noradrenaline being carried away via the adventitial capillary plexue. intimal overflow entering directly into the circulation. and This transmitter entering the circulation constitutes the most important fraction of circulating noradrenaline (most circulating the adrenaline originates from the adrenal glands). In isolated blood vessels studied under controlled conditions, there is а linear correlation between the changes in overflow of endogenous noradrenaline and changes in the effector response during activation of the adrenergic nerve endings(15;147). Diffusion of transmitter out of the blood vessel

wall has not been adequately investigated in most studies of noradrenaline disposition. This is partly because of the difficulties encountered in obtaining proper estimates for diffusion compared with the other disposition mechanisms. It should be noted, however, that the rate of diffusion may exceed that of all other mechanisms.

Uptake₁ (neuronal uptake)

tissues.

1.

Neuronal uptake or uptake₁ is the active transport of noradrenaline and adrenaline into the axoplasm of the adrenergic nerve endings. This is believed to be one of the main pathways responsible for termination of the action of released noradrenaline at the neural cleft. The characteristics of this membrane carrier system has been reviewed by Iverson(148,149) and are summarised below.

- Uptakel process appears to have identical properties in the noradrenaline containing neurones of the peripheral and central.
 - nervous systems and results from the activity of a membrane carrier system requiring metabolic energy (Note: The dopamine containing meurones in the mammalian central nervous system have a membrane carrier system analagous to uptake₁. However, this uptake system has a very high affinity for dopamine and a lower affinity for noradrenaline).
- 2. The uptake process is saturable and has a very high affinity for noradrenaline, the transport constant or apparent "Km" being between 0.2 μmol and 1 μmol in most rat tissues.
- 3. It is stereochemically selective in rat tissues having five times the affinity for the naturally occuring (-)noradrenaline compared with the (+)enantiomer. However, such stereoselectivity seems to be lacking in guinea pig and rabbit

The necessary structural requirements for uptake₁ are: (a) the absence of bulky N-substituent groups (isoprenaline is not a substrate) (b) absence of methoxyl groups on the aromatic ring (normetanephrine and methoxamine are not substrates for uptake₁) (c) presence of at least one phenolic hydroxyl group (amphetamine, phenylethylamine, phenylethanolamine and norephedrine are not substrates).

4 -

· · ·

Depending on the above, various substances are taken up by the adrenergic nerve endings with varying degrees of affinity, e.g. metaraminol, armethyl noradrenaline, tyramine, actopamine. Adrenaline is also taken up by uptake, and has an affinity of about half of that of noradrenaline.

- As with most other membrane transport systems for organic compounds, uptake₁ is temperature sensitive (approximately doubling of the rate of uptake for an increase in temperature of 10°C) and can be inhibited by metabolic poiseus such as dinitrophenol and cyanide or by anoxia.
- 6. It is dependent on extracellular Na⁺ with a marked reduction in uptake when extremel Na⁺ ions are removed. It also requires the presence of a low concentration of \tilde{F}^{T} long (approximately 5 mmol/1) but is inhibited by high concentrations of F^{+} (50 mmol/1). Uptake₁ is also inhibited by inhibitors of membrane Na⁺/K⁺ ATPase such as ouabain.

(a) affinity for untake, sites is decreased by the prospect of

bulky substituent groups on the terminal nitrogen of phenylethylamine side chain, by the presence of methoyl substituents on the ring, and by the presence of a hydroxyl group on the β -carbon of the side chain. For the latter compounds, affinity for uptake₁ sites is greatest for the isomer (corresponding to (-)noradrenaline (b) affinity is increased by the presence of phenolic hydroxyl

groups, particularly in para- and meta- positions, and also by methylation of the α -carbon of the side chain. In the latter case, affinity is highest for the isomer correspondidng to (+) amphetamine.

It should be noted that the structure activity relationships for inhibition of uptake, by sympathomimetic amines are not identical with the requirements mentioned above, for substrates for the uptake process. For instance, amphetamine or β phenylethylamine, which lack phenolic hydroxyl groups, do not appear to be substrates for uptake,, but are nevertheless ³H-noradrenaline uptake. competitive inhibitors of This suggests that such compounds, like competitive enzyme inhibitors, are able to bind with high affinity to uptake, sites in the axonal membrane of adrenergic nerves, but lack the structural features needed for the inward transport further stages which follow the binding. Alternatively, many sympathomimetic amines may compete with noradrenaline for binding to the uptake, sites and this may be followed by transport of the competing amine into the axoplasm. Thus, indirectly acting sympathomimetic amines not only release

noradrenaline from adrenergic nerve-ending, but they also potentiate the actions of the released catecholamine by inhibiting its recuptake, e.g. tyramine.

8.

٦

Many other compounds inhibit uptake, apart from the close structural analogues given above. Tricyclic antidepressants such as imipramine, amitryptaline and desmethylimipramine (desigramine) are potent inhibitors with the latter ompound being one of the most powerful inhibitors anailable at present (50 per cent inhibition of ³H noradrenaline uptake to the rat heart at a concentration of 10^{-8} mol/1). Other inhibitors include the local anaesthetic drug cocains, advenergic receptor blocking drugs phenoxyber amine and oblovpromazing, the monnamine, oxidase inhibitory franvicypromine and phonelsine and the adrenergic neurone blocking druge bretvilium 3111 guanerhidine. Notabe, in not inhibited by reserving.

There appears to be a temporal dissociation between the activation of the nerve termin 1 and derivation between the with the latter complete only when the nerveral combinance polarized(15,150). In the matrix of the defer of the opticie of ³H noredrinating was shown to be foldable to be astmited to the effector pervection for many dependent methods.

The transmitter taken up by the ortike, process is alther stored in the vesicles or deaminated by the neuropliants enzyme when the exidase. The noradionaline is the reproplasm is there early in the storage patches by an energy depend of up its press which is price expensive is allowed or the price is and the press which is price

dependent on the presence of ATP and Mg^{2+} and is accompanied by a splitting of the terminal phosphate groups of ATP. This vesicular uptake system is potently inhibited by reserpine, tetrabenazine and prenylamine which are relatively ineffective as uptake₁ inhibitors. Reserpine is porticularly potent being effective in vitro at a concentration of 10⁻⁹ mol/1. The vesicular uptake system too, appears to be temperature dependent, energy requiring and stereochemically collisive to () hore/tenaline. The ATP-dependent uptake systems of both the large and small dense care vesicles have lower transport constants 'Km'. 1.5 µmol and 10 µmol respectively) for noradrenaline than the Km of menoamine (vidase for noradrenaline (100 µmol)). Thus, most of the noradrenaline "shere up into the nerve terminal by uptake₁ presumably goes back into the vesicies.

Following transport of the noradrenaline into the vesicles it forms a tetrace echolamine-ATE complex (molar ratio 4:1) with ATP. This in turn may be bound to the soluble protein chromographin(157), This resigning therage function appears to be distinct from the vestoriar the both functions are inhibited by uptake Chants pvi recorption (192). The exiting of recorption on the regionalis uptake we have assume to be competities with estecholamines, so the processes of blab out bolaming concentrations in the viginity of the reservine require can compute with the drug. On the other hand, once reservine has become doubth the aborage every the latter remains fulthing even to the pressure of corrections. These, the initial action (on realizing unitake) of recently is a coaffic by the lifer action in storage to be particular the second of the sought of the spect

111

 \mathbf{S}

the arrival in the adrenergia nerve terminal, of new vesicles (from the neuronal cell body) not affected by the drug(152). However, it is not clear as to how the arrival of a relatively few new vesicles could, by themselves, allow the almost complete restoration of the uptake The storage function in the vesicles exhibits a greater function. structural specificity as compared with the vesicular uptake of amine which is relatively a n-specific. Structural requirements for storage in advenningic vehicles are physically limines with a f-hydroxyl group and at least one phenolify group. The moneyle also much be of the correct optical configuration on responding to that of a moral renaline (152). It approve that the issue function, as opplied to the prate for it to of the woof I are then price, shown for the core member and the absolute latinfnint the start for an incompany case there peradreesline. Cores of the second structures of the second 1.011 to the respect of a · · · · · · · · · · ·

ب يو المريد

•

and the second sec T+ 1. 71 , · no finn ait and the second · · · · · 101-11- 11and the second second 1 endings and the second 11.00 the second se net te sta sa 1 11 1 april 1 and 7 1 follow for . 1 . 1 #1 forigt is an 1 : 1 1.

.

deaminated metabolites especially 3,4-dihydroxyphenylglycol(DOPEG) is reduced; DOPEG is formed by the action of monoamine oxidase (MAO) on the noradrenaline, subsequent to uptake $1 \cdot 3$ a moderate augmentation of the contractile response to low-frequency stimulation occurs - there is usually no augmentation with higher frequencies; this minimal effect is probably due to the fact that uptake, is inhibited during sympathetic nerve stimulation in a frequency dependent manner(151). Thus, an uptake, inhibitor may not change the transmitter concentration at the avaantic cleft during nerve stimulation to any appreciable degree. 4) the contractile response, however, tends to become prolonged and the relaxation delayed: this occurs as uptake, plays an important role in removing the transmitter from the receptor site once the stimulation is reiminated. Plockade of this would lead to a prolonged effect while not significantly affecting the magnitude of the concentration of the nurshroughing (and thus the machitude of the contraction) at the cleft.

Uptake, individual also potentiate the responses to exogenous catecholemines. The mightude of this potentiation too depends on the density of the vation and the vidth of the sumptic cliff and also on the particular catechole is and within to a groater entent than those of straights dutch has been effective for the uptake process. Interview of the conductive are fiftill to be against the also is proposed to the state of the sumptic cliff and also on the catechole is a groater entent than those of straights dutch has been effective for the uptake process. Interview of the effect of a reduce of the straight on the effect of an affect is a reatest of a straight of the time of the straight of

, **Þ**.

. • ,

.

113- -

in tissues such as the cat nictitating membrane with a dense adrenergic innervation, and not at all in tissues lacking sympathetic innervation. Also in tissues in which the neuromuscular interval is small the potentiation by uptake inhibitors will be greater(153); the uptake process in general would significantly alter the cleft concentration of noradrenaline only in the case of a narrow cleft in spite of the fact that the rate of diffusion of exogenous noradrenaline into a narrow cleft would actually be less than that into a wide cleft (see above).

. The end of 114 is a second secon

In addition to the above, the route of administration of the catecholamine may play an important role in the magnitude of the potentiation produced by uptake, inhibition in isolated blood vessels. De La Lande et al. (154) in a study on the isolated, perfused, central ear artery of the rabbit, demonstrated that while cocaine greatly potentiated the effects of extraluminally administered noradrenaline, it had little effect on the consitivity to intraluminally administered peradropaline Surgical interruption of the sympathetic supply to the artery 14-24 days prior to the overiment simulated the effecte of construction above. Countre Uself lad up atout load offer following the autitool denore that there bear estiona are graphily avilated . The for that the estimator fully control nors is alter has to diffuse the eight the elemental of a the main produce to a evert its offer on the smooth murches of the device of the glo when pleases supplying the media of the reason is been done to the trace ad outside across for the second frequencies of the conditional the removed and the second second

and the set of a

media thus limiting the response to the drug. Cocaine and denervation, by blocking the uptake process, leads to a supersensitivity to extraluminally administered noradrenaline only. However, studies using cocaine are complicated by the fact that the drug has a post-junctional sensitising action in addition to the uptake, blockade. Thus, it potentiates the responses to the α -agonist methoxamine (which is not a substrate for uptake,) in the rabbit sorta(153). Osswald in a similar. study in the canine lateral saphenous vein(17), demonstrated that cocaine potentiated the effects of both intraluminally and extraluminally applied noradrenaline to the same extent. This is probably due to the fact that in the canine saphenous vein adrenergic nervos are distributed throughout the tunica media.

• • • .

""" it ites (Fatraneuronal uptake)

This is a transport system for catecholamines, present in smooth muscle. cardiac muscle, endotheldum and certain glandular tissues(148,155). This system has a much lower affinity for noradrepuline and adrenaline than uptake, (Km for (-)noradrenaline 0.27 μ mol and 252 μ mol for uptake₁ and uptake₂ respectively in rat heart tissue). However, uptake, has a very much higher capacity than uptake, although it is enturable. An the accumulated catecholamine is not firmly retained but repidly metabolised by MAO and catechol-omethyltransferase (CONT), uptake, is not usually detectable at low concentrations of catecholamines. Uptake, does not demonstrate a stereochemical specificity for (+) or ()noradrenaline or adrenaline and has a higher affinity for the latter. It has quite a different substrate specificity from uptakel: isoprenaline, an amine not taken up In uptains, is a better substrate than even adrenaline for uptake2. The

structure-activity relationship for the inhibition of uptake₂ by sympathomimetic amines are almost the converse of those found for uptake₁(148); thus inhibition of uptake₂ was enhanced by N-substitution and by 0-methylation, normetanephrine and metanephrine being potentinhibitors. On the other hand (-)metaraminol, a potent inhibitor of uptake₁ had no inhibitory effect on uptake₂. Phenoxybenzamine too is a potent inhibitor of uptake₂ although its usefulness as a tool to block uptake₂ is limited due to its other actions such as blockade of uptake₁ and the blockade of α -receptors. Steroid compounds have also been discovered as uptake₂ inhibitors, β -coestradiol and corticosterone being quite potent. Uptake₂ is less dependent on Na⁺ ions compared to uptake₁. The effects of anoxia, cooling and ATP deprivation are also not as clear as with uptake₁(155).

Unlike uptake₁, the role played by uptake₂ in the termination of the actions of catecholamines released at sympathetic nerve endings is likely to be minor. However, under conditions in which uptake₁ is blocked (eg. by administration of tricvelic anti-depressants) uptake₂ may assume a bigger role. On the other hand, because of the widespread distribution of uptake₂ sites in vascular smooth muscle, and the preference for adrenaline as substrate, it is likely that uptake₂ plays an important role in the rapid remeval and inactivation of circulating ratecholarines(148).

Some inclated smooth muncle such as the rabbit sorts and the rabbit car arts v show a potentiation of the effects of catecholamines with blockade of uptake₂(153), although the potentiation is much less compared with that observed following blockade of uptake₁. In the perfused rabbit car artery, responses to both intraluminal and

a ser para ana ang ang at 🕯 👘 🖓

_____116

extraluminal noradrenaline are potentiated and the supersensitivity is not decreased by denervation or cocaine. Another important feature of the uptake, process is that in contrast to the adrenergic nerve ending, gextraneuronal uptake has "little or no capacity for storage of unchanged amines (Note: capacity for uptake is however, high). The extent to which uptake₂ influences the concentration of catecholamines at the receptors is ultimately dependent on metabolism by COMT. Thus if COMT is inhibited, blockade of extraneuronal uptake does not lead to supersensitivity. These findings suggest that uptake2 and metabolism by COMT, are arranged in series so that blockade of one process eliminates. the effects of blocking the second. Thus, for supersensitivity to occur the agonist has to fulfill two criteria: (1) it must be a substrate for uptake₂. (2) it must be a substrate for the enzyme COMT(153). For instance, phenylephrine, although , substrate of uptake2, is not metabolised by COMT as it is not a catechol compound. Thus, the response of the nictitating membrane to phenylephrine is not potentiated by hydrocortisone, an uptake, inhibitor.

Inhibition of uptake, may have one or more of the following effects on the response of blood vessels to sympathetic stimulation(15): 1) no, or modest increase in the overflow of ³H-noradrenaline. 2) reduced appearance ٥f extraneuronal metabolites of noradrenaline (eg. normetanephrines), 3) augmentation of the contractile response, 4) moderate prolongation contractile response of the and delayed relaxation.

Catsholism(156,157)

Chemical inactivation of noradrenaline in adrenergic neuroeffector systems is controlled by two enzymes, monoamine oxidase(MAO) and
catechol-o-methyltransferase(COMT). The enzyme MAO which catalyses the deamination of a wide variety of amines is widely distributed in the body with the liver, kidney, intestines, stomach and aorta constituting rich sources (156). It is either absent, or present in very low amounts in skeletal muscle, plasma and erythocytes, although it occurs in -platelets. It has also been demonstrated in a variety of blood vessels(15). Despite this wide distribution; monoamine oxidases isolated from different organs in species show considerable differences in specificity. The enzyme is located almost exclusively in the outer membrane of the mitochondria, inside cells. The minimum molecular weight of the enzyme has been estimated at 100,000 although it can vary up to one million depending on the preparation(156). The enzyme exists in two main forms, MAO-A and MAO-B. Adrenergic nerves contain mainly type A and the vascular smooth muscle cells, probably type B. The total MAO content of the rabbit ear artery is reduced only 10 per cent by chronic sympathetic denervation indicating the larger size of the extraneuronal stores of the enzyme. However, this relatively small amount of neuronally localised MAO, metabolises the transmitter released at the nerve ending and is thus functionally important in the adrenergic neuro-effector interaction(73). Extraneuronal MAO is effective only against high concentrations of noradrenaline, and its inhibition has minimal influence on the responses to endogenous or exogenous noradrenal(ne(73, 158).

The enzyme COMT, like MAO, is also widely distributed in the body including the brain. with high concentration in the liver and kidnev(130). However, unlike MAO, it is located in the cytoplasm and bas no selective association with adrenergic nerve endings. Next of the

COMT activity of the blood vessels is extraneuronally located, presumably in the vascular smooth muscle cells. Catabolism of catecholamines in the body results from a combination of actions from the two enzymes at neuronal and extraneuronal sites. Oxidative deamination of noradrenaline by MAO results in the formation of 3,4dihydroxyphenyl glycoaldehyde (DOPAE). The aldehyde metabolites of

catecholamines can be demonstrated in vitro, but are rarely detected in This is because these aldehydes are immediately tissues or urine. metabolised to more stable products. The oxidation of DOPAL by aldehyde dehydrogenase leads to the formation of 3,4-dihydroxymandelic acid (DOMA). The reduction of DOPAL by aldehyde reductase (alcohol dehydrogenase) results in the formation of 3,4-dihydroxyphenylglycol (DOPEC). On the other hand, action of COMT on noradrenaline results in methylation 'at the ortho-position with the formation of normetanephrine The NMN may be acted upon by MAO to form an unstable aldehyde (NMN). which in turn can be oxidised or reduced by aldehyde dehydrogenase and aldehyde reductase respectively. This results in the formation of 3methoxy-4-hydroxymandelic acid (incorrectly called vanillylmandelic acid-VMA) and 3-methoxy-4-hydroxyphenylglycol (MOPEG) respectively. 3methoxy-4-hydroxymandelic acid ("VMA") constitutes the major metabolite of catecholamines excreted in the urine. The corresponding product of the metabolic degradation of dopamine, which cantains no hydroxyl group in the Gide chain, is homovanillic acid (HVA).

A significant portion of noradrenaline spontaneously leaking out of the storage vesicles in the resting state and that displaced by indirectly acting sympathomimetic agents are acted upon by MAO, with the ultimate production of DOPEG as the major metabolite under those

· circumstances. Some of the noradrenaline that is taken up (by uptake 1) following exocytotic release too is subjected to oxidative deamination by MAO with the production of DOPEG and DOMA as the end products. These two compounds may possibly be o-methylated (by COMT) in extraneuronal tissues with the formation of MOPEG and VMA. Some of the noradrenaline released during sympathetic nerve stimulation is taken up by uptake, into smooth muscle cells as explained before. This noradrenaline is degraded by COMT which is predominantly located extraneuronally with the formation of NMN. As extraneuronal uptake plays a major role in the disposition of circulating catecholamines, relatively more NMN and the other two o-methylated prducts (MOPEG and VMA) are formed under these circumstances. The relative proportions of the different metabolites formed seem to differ not only with the mode of adrenergic stimulation (exocytotic release, displacement release, stimulation by circulating catecholamines) but also with tissue and species variation(157,159).

Inhibitors of monoamine oxidase may have one or more of the following effects on the response to sympathetic nerve stimulation(15): 1) decreased appearance of deaminated and o-methylated deaminated metabolites, and moderate augmentation of the overflow of intact 2) moderate (or no) augmentation of the contractile transmitter. response. 3) moderate (or no) prolongation of the contractile response and delayed relaxation. In addition they may augment and prolong the response to Indirect sympathomimetic amines and to. exogenous noradrenaline. Analysis of the effects of these inhibitors are made difficult' by the fact that these drugs by themselves can cause pharmacological displacement of the stored transmitter and facilitate the release evoked by adrenergic stimulation. Supersensitivity to

exogenous noradrenaline produced by MAO inhibitors usually occurs only in innervated muscle. Moreover, in the perfused rabbit ear artery preparation, nialamide (a MAO-inhibitor) potentiates the response to extraluminally administered noradrenaline but not to intraluminally applied noradrenaline. These findings suggest that the supersensitivity produced by MAO inhibitors is probably, due to inhibition of the neuronally localised MAO, rather than the extraneuronally localised. enzyme(153). Thus, the supersensitivity following inhibition of MAO is more pronounced in those tissues in which uptake₁ plays a major role in the disposition of the transmitter.

1

Inhibition of COMT by drugs such as pyrogallol, tropolone and U-0521 may have the following effects on the response to sympathetic nerve stimulation in smooth muscle(15): 1) decreased appearance of omethylated and o-methylated deaminated metabolites. 2) augmented outflow of intact transmitter and deaminated metabolites. 3) moderate (or no) augmentation of the contractile response. 4) moderate (or no) prolongation of the contractile response. They may also augment and prolong the contractile response of smooth muscle to exogenous catecholamines. This supersensitivity is probably due to the build-up of noradrenaline in the muscle cells with resultant slowing down of the extraneuronal uptake process, as the two processes seem to be coupled in series as explained before. Conversely, if extraneuronal uptake is blocked by steroids, then inhibition of COMT does not produce supersensitivity. In the isolated perfused rabbit ear artery, inhibition of COMT potentiates the responses to both intraluminally and extraluminally applied noradrenaline to the same extent. In addition, the effects of COMT inhibitors are not affected by prior treatment with

uptake₁ blockers. These findings indicate that the supersensitivity is probably due to the inhibition of (extraneuronal COMT rather than theintraneuronal enzyme (c.f. with MAO inhibitors)(153). Supersensitivityfollowing COMT inhibition is related to the sensitivity of the tissue tothe catecholamine rather than to the functional state of the uptake₁mechanism. This occurs because the o-methylating system is easilysaturated. Thus, supersensitivity occurs.only in tissues in which thesensitivity to catecholamines is high.

Binding of catecholamines to connective tissue

A part of the noradrenaline released at nerve endings is believed to bind to connective tissue. The role of catecholamine binding to collagen and elastin was investigated by Powis(160). Collagen showed no specificity towards binding of either (-) or (+)adrenaline and noradrenaline while elastin bound the (-)isomers to a greater extent. Both elastin and collagen demonstrated two sites of binding: a high affinity, limited capacity site and a low affinity, high capacity Tetracyclines inhibited this binding, with oxytetracycline in a site. concentration of 10⁻⁴ mol/1 inhibiting the binding of noradrenaline to collagen by 68.4 per cent. Oxytetracyline (10 4 mol/1) was shown to potentiate the amplitude of the response of the rabbit car artery to noradrenaline and to very atimulation ten and six fold respectively. As blood vessels contain a relatively high proportion of clustin and collagen (over 60 per cent of the dry wright in some ressels) connective t fague binding of catecholamines cuild theoretically Avort considerable effect on the magnitude of the responses to these compounds. In the rabbit ear artery the potentiation produced by blockade of uptoke, and antake, were much loss than the potentiation

produced by oxytetracycline. However, some studies did not show any potentiation of the responses to neural stimulation and exogenous catecholamines by tetracyclines(161). Thus the importance of the connective tissue binding of catecholamines (as with the other mechanisms of disposition) may vary depending on the tissue as well as with the relative role played by each of the other mechanisms in a particular tissue.

Relative importance of the disposition pathways

•

the existence of several disposition pathways Although of noradrenaline in vascular tissue is well established, the relative role played by each in a particular blood vessel remains unresolved in most preparations(73). The relative roles are particularly affected by 1) the density of the adrenergic innervation, 2) the anatomical arrangement of the nerve endings within the blood vessel, 3) the width of the junctional cleft, 4) the content of collagen and elastin in a particular blood vessel and 5) whether neurogenic noradrenaline or circulating noradrenaline is being investigated(15). If the cleft is narrow, uptake, usually plays a prominent role; if it is wide, diffusion and extraneuronal uptake assume a greater role. The investigations designed to answer this question in a particular blood vessel are complicated by the fact that interactions between these mechanisms are frequent. Therefore, controlling one mechanism by itself may alter the magnitude of the role played by another mechanism, thus confounding the results.

Transmitter disposition in the canine saphenous vein

The relative role of the disposition pathways in the canine lateral caphenous very were investigated by Guimaraes, Osswald and co-

 $(10^{-5} mol/1)$ shifted the dose-response curve to exogenous noradrenaline and the stimulus-response curve to TNS, to the left by a factor of 7.1 and 4.5 respectively(162). The difference in the extent of potentiation between exogenous noradrenaline and TNS, which was significant, may have been due to the local anaesthetic effects of cocaine leading to a decrease in the release of endogenous noradrenaline during TNS. Thus, cocaine would have two opposing effects during TNS: blockade of uptaket leading to potentiation of the contractile response and inhibition of release of the transmitter leading to a decrease in the response. This discrepancy between the extent of potentiation of TNS as compared with exogenous noradrenaline could also be explained by the fact that uptake, is already blocked to some extent during TNS (uptake, is believed to be not operative when the nerve terminal is depolarized). This would result in a lesser potentiation by the cocaine. The potentiation of exogenous poradrenaline responses in canine saphenous veins was also produced by surgical denervation 8.10 days prior to the experiment. Cocaine had no further offect in these veins following the surgical denorvation.

The authorn also massured the time required for half relaxation following TPS by the off immersion technique(7),1621 forsing in reared this time 2.2 times. The uptake, Inhibitors T-0521 and moder to be up significant effect on the time of relaxation forgetting that fifth and uptake, do not represent innortant pathwave for the in times of the released transmitter. The Michigh form the in times of the increase to the flue needed for helf the uptake, of the interval increase to the flue needed for helf the uptake, of the interval increase to the flue needed for helf the uptake, of the interval increase to the flue needed for helf the uptake, of the interval increase to the flue needed for helf the uptake of the interval increase to the flue needed for helf the uptake of the her for the interval increase to the flue needed for helf the uptake of the her for the interval interval.

increases in the relaxation times with prior cocaine treatment. Thus both COMT metabolism and uptake₂ are able to compensate for the loss of uptake₁. This may, in fact, be due to the increase in the sensitivity to endogenous noradrenaline produced by cocaine resulting in an effective concentration range for the noradrenaline, at which COMT and uptake₂ are no longer saturated as explained above. In summary, it appears that neuronal uptake represents the most important pathway of inectivation for endogenous noradrenaline in the saphenous vein.

1.17

1 m

Osewald or al(17) also investigated the relative roles played by the different disposition pathways, during the relaxation after a contraction induced by exogenous noradrenaline by the oil-immersion rechnique. Throniazid prestreatment increased the time to 50 per cent relaxation 1.6 fold while consine and imipramine sugmented the time only 2.6 fold Tropolene (a COMT inhibition) caused only 1.38 fold increase. When the three drugs (MAO inhibitor, uptake, inhibitor and COMT inhibitor) were used in combination (changing the order of addition), iprovinzid added after tropolone and cocaine produced the higgost prolongation of the relaxation time. The authors concluded that avidative 'sputnation to MAO was the most important pathway for Happetition i expansion correctionaline in the canine appendue vern-Honover, when this continue in estigated by studying the extent of potentiation of the cost action to regenerate retrenaline as the tool of measurement, he had a funtakey and the time the proceeds n contraction for the control anchonomy rota

Cutmanned to define the property of the transfer of the matched the of $\eta_{\rm H}$ and metabolism of $\eta_{\rm H}$ and $\eta_{\rm H}$ is an indicated the property weight of $\eta_{\rm H}$ and $\eta_{\rm H}$ is an indicated the property weight. Of the

per cent was metabolised. Deaminated e-methylated metabolites (MOPEG and VMA) represented 59 per cent and a methylated metabolites (metanephrine) respresented 30 per cent of the total metabolised. The deaminated metabolites (DOPEG and DOMA) represented the balance 10 per cent. Cocains reduced the removal of ¹H adrenaline by the tissue 4 80 per cent of the control value. This isduction is mostly of the unmetabolised (stored) fraction which would treatmable by located in the adrenergie usive terminals and this blocked in constant. There were us major charges in the proportions of different i shall the incert by blocker isometrices using on the other hand, for used the unmetabolised Fither, the per cost different is shall to the using the blocker isometrices using on the other hand, for used the unmetabolised Fither, the per cost different is shall to the using the blocker isometrices using on the other hand, for used the unmetabolised Fither, the per cost different is shall to the using the metabolised for a methylated metabolises of and cells is the using the metabolised is methylated metabolises of and cells is the using of the domainstead of methylated metabolises of and cells is the using the shall be a methylated for the to 6 n the

1

In the gass of "R constrained for taken up by the theory, 10 per set was metabolized with 6 per contining as intact a directly the deem(rated metabolites (NOTED NOM/) represented 20 per an an mathy tod, d and no it notices or (MOPRC and VMC is a cost of the f(x,t) = f(x,t) + f(x,t) + f(y,t) + fis a first of the second of the control strength the strength the second s and the second of the second of the star for a charge to a short to prove the contract of the alificant dates and the state of the first second the end of the provide the state of the second second and the second sec Compress of the second second 1 1 1 1 1 metabolic 1 . .

1

'Hton (Party

abolished with a reduction also in the o-methylated, deaminated fraction. In summary, it could be stated that deamination predominated in the metabolism of ³H-noradrenaline and o-methylation in the metabolism of ³H-adrenaline in the saphenous vein. The metabolism of ³H-adrenaline was markedly reduced by extraneuronal uptake blockade. The study also demonstrated that a significant amount of DOPEG formation occurred extraneuronally too when higher concentrations (2.3 μ mol/1) of the tritiated ompounds were used during the incubation period. Thus, in contrast to many other tissues in which extraneuronal uptake plays a miner role in the disposition of caterbolamines, in the dog saphenous wein it plays a significant role in inactivation of exogenous atorbolamines especially adrenaline.

In a more secont study Muldoon et al(159) analysed the metabolic fractions during INS, following labelling of the tissue stores with ³Hnor transline. Under basal conditions intact noradrenaline accounted for only a small fraction of the total spontaneous efflux of tritlated mata 191, The dearlingtod metabolites (DOPEG and DOMA) represented the main fraction with the amount of DOFEG being 4-8 times that of DOMA. the given MOFFO ine the largest component among the o-methylated (MOFTE VMA, NIM) and NIM the emallest. During TNS the met about t man . I wondrevalue in the superfugate increased, and it amount of formed the large i fraction of frequencies of 2Hz and higher. Among the Int chered the most i riking increase followed by MOPEG. metalille Auth I I will what it increases the increasing frequency while VMA During TNS following blockade of with end i Increased above 1 sine () foot is ell moraboline fractions increased 1 is a star in the last offlin approxighty. Or

the other hand, the MAO-inhibitor pargyline produced a decrease of the total radioactivity in the basal state with a decrease in all deaminated compounds except DOMA. However, the efflux of intact noradrenaline and NMN showed a significant increase. During TNS at 2 Hz after pargyline pre-treatment, there was an increased amount of total radioactivity in the superfusate accompanied by increases in intact ³H-noradrenaline and NMN compared with the control state. The amount of DOFEG was less than in the controls. In this study the amount of DOMA in the superfusate was consistently less than the amount of DOFEG. This suggests that, in the saphenous vein, after deamination of noradrenaline, the formed aldehyde is reduced by aldehyde reductase (to the given DOPEC) rather than oxidized by aldehyde debydrogenase to the acid DOMA. ۸n alternative explanation is that DOMA formation takes place mainly extraneuronally. This would explain why conside did not reverse the efflux pathern of DOMA. In the case of the commethylated metabolites, the glycol MOPEG was always found in excess of VMA in the superfusate. Thus, here too the aldehyde reductase appears to clay the more dominant role. The ormethylated, deaminated metaboliton could be croduced by either the nerve terminals or in the extraneuronal tissue. As consine augmonth the evoked thisass of emethylstel doomfus of a me er di Ina TNS (this during the of DOFECT the main strains) + tota for these actional to a contract of the end of the end of the second s

le 11e.

It should be opted that to all oupertments utilizing (featers) and choose of the second state of a second state of the second

different techniques are used to investigate the relative role of the disposition pathways. The studies by Guimaraes and Osswald(17) where the effector responses were investigated, suggested a predominant role deamination for uptake, and by MAO in the disposition of noradrenaline. The other studies(159,163) where metabolite fractions were measured, do not indicate a major role for the above pathways, but stress the role of extraneuronal uptake. This is probably due to the fact that the criterion measured is different in the two sets of studies (effector response and metabolite fractions) and thus do not mean the same in physiological terms.

129

Pre-synaptic modulation of transmitter release

Three groups of receptors are generally involved in chemical neurotransmission(164). 1) The receptors on the some and dendrites of the innervating neurone which determine the frequency of impulses carried down to the axon terminals, 2) The post-synaptic receptors which recognize the transmitter and mediates the response of the innervated cells, 3) While these two receptors would be adequate for the system to produce an effector response, there is evidence that a third group is located on the norme endings modulating the release of the transmitter and in some many its synthesis. These receptors are known as prejunctional or pre-synaptic receptors. Biochemical experiments that retrogrectively can be explained by pre-synaptic receptors date back to the 1950's(165.16() Since these initial reports a large number of publications have confirmed that a adrenoceptor blocking agents, like phenom heugamine, increase the overflow of norepinephrine elicited by nerve elimitation. These findings were initially attributed to, is live a after of is a for the transmitter at the generators, the

V

blocking of neuronal uptake and the blocking of extraneuronal uptake by these a-adrenoceptor blocking agents. The first of these possibilities was later found to be untrue and the other two possibilities, although true, not sufficient by themselves to account for the magnitude and the pattern of increase in overflow of noradrenaline and its metabolites observed with α -receptor blocking agents. The evidence against these hypotheses has been reviewed by Langer(5) and Westfall(129), 1) Cocaine and desmethylimipramine (desipramine) which are potent inhibitors of uptake, produce only a small increase in efflux of ³H-noradrenaline with nerve stimulation in contrast to phenoxybenzamine which produced a marked increase in efflux. 2) α -adrenoceptor blocking agents produce a further increase in efflux when added, following blockade of uptake, by a maximal dose of cocaine or desipramine. 3) α -adrenergic blockers increase the efflux of noradrenaline in doses that do not block neuronal uptake. 4) The α -adrenoceptor antagonist phentolamine which does not block extraneuronal uptake also brings about an increased efflux of noradrenaline during nerve stimulation. Thus extraneuronal uptake cannot account for the increased release of the transmitter. 5) Inhibitors of COMT and inhibitors of untake, such as normetanephrine produce a small or no increase in the stimulation-induced efflux of noradrenaline. 6) a Phaloalkylamine GD 131, similar to phenoxyhenzamine, that inhibits both uptake, and uptake, without blocking a recentors produces only a slight increase in the stimulation induced efflux of noradronaline(129). 7) Accompanying the increased efflux of noradrenaline produced by a advenorgic antagonists, an increased overflow of the intravesionlar enzyme dopamine-B-hydroxylase is also beausel. Since this ensume is a rather large melocule that is

not taken by uptake₁ or uptake₂, or subjected to inactivation in the tissue after its exocytotic release, an increase in overflow of the enzyme does indeed represent an actual increase in release at the adrenergic nerve terminal.

These findings led to the hypothesis of pre-synaptic regulation of transmitter release at the adrenergic nerve ending put forth independently by four laboratories in 1971(167,168,169,170). The hypothesis proposed the existence of α -adrenergic receptors on the pre-synaptic membranes at the adrenergic nerve endings; these, when stimulated by the released noradrenaline, would lead to a decrease in further release of the transmitter, i.e. a negative-feedback control of release.

Evidence available for the existence of pre-synaptic α -receptors as proposed could be summarised as follows(15,129). 1) A number of α adrenergic antagonists including phenoxybenzamine, phentolamine. dihydroergocryptine have been shown to increase the overflow of noradrenaline during nerve stimulation in concentrations that have a " minimal or no effect on neuronal and extraneuronal uptake. In some cases, but not all, the increase in overflow of noradrenaline is accompanied by an increase in the effector response in the preparation. 2) α -antagonists lead to a facilitation of the release of noradrenaline from tissues (such as the heart) which are believed to contain relatively few post-synaptic a-receptors. This suggests that the receptors involved in the regulation of the transmitter release are located pre-synaptically (and argues against a trans-synaptic regulation where the post-synaptic effects of the transmitter leads to the formation of mediator substance which subsequently acts pre-

synaptically to regulate the transmitter release). 3) Phenoxybenzamine shown to increase the stimulation induced efflux of been has noradrenaline from axonal sprouts of cultured ganglia where there is no evidence of post-synaptic innervation. Thus, the site of action has to Similar experiments with synaptosomes (resealed torn-off be neuronal. axon terminals) too suggests a direct-action. However, it should be noted that synaptosomes often have post-synaptic membranes attached, so a trans-synaptic action cannot be definitely excluded(171). 4) High K^+ (external) solutions lead to release of neurotransmitters by a direct depolarizing action on nerve terminals: tetrodotoxin, the fast Na⁺ channel inhibitor blocks the traffic of action potentials. Hence, when a drug (e.g. α -agonist or α -antagonist) modifies transmitter release induced by high K⁺ in the presence of tetrodotoxin, it cannot be acting by way of action potentials in interneurones or by an action potential propagated down to the nerve terminal from the nerve cell body. It must act either on the terminals under study or on neighbouring cells. This model has been used to exclude interneuronal pathways in the α adrenergic inhibition of noradrenaline release(171,172). 5) A wide range of a-agonists including noradrenaline, clonidine and oxymetazoline have stimulation-induced efflux of demonstrated to decrease the been noradrenaline in a variety of tissues unrelated to a post-synaptic effect(129). This effect can be blocked by the simultaneous use of α adrenoceptor antagonists.

÷ .

Further investigations on the α -receptor mediated inhibition of transmitter release resulted in the discovery that pre- and postsynaptic α receptors differed from each other in their pharmacological characteristics(129). When the two types of receptors were compared in

the same preparation, a marked difference in potency was noted between different α -agonists on the two receptor types. For instance, in strips of rabbit pulmonary artery(173), methoxamine and phenylephrine either did not change or enhanced, but never reduced, the contractile response tò transmural nerve stimulation. In contrast oxymetazoline, amethylnoradrenaline and tramazoline at low concentrations selectively inhibited the response to transmural nerve stimulation. The rank order of potency for the reduction of stimulation induced efflux of ³Hnoradrenaline by 20 per cent (EC₂₀ pre) was adrenaline>oxymetazoline >tramazoline>a-methylnoradrenaline>naphazoline>phenylephrine >methoxamine. On the post-synaptic side the rank order of potency for producing 20 per cent of maximum contraction (EC₂₀ post) was adrenaline>oxymetazoline>naphazoline>phenylephrine>tramazoline>a-methylnoradrenaline>methoxamine. The ratio EC20 pre/EC20 post was calculated for each agonist as an index of its relative pre- and post-synaptic According to the ratio the agonists were arbitrarily potency. classified into three groups: group 1 (ratio about, 30) - preferentially post-synaptic agonists e.g. methoxamine, phenylephrine; group 2 (ratio around 1)-similar pre-and post-synaptic potencies e.g. adrenaline, naphazoline; group 3 (ratio below 0.2)-preferentially pre-synaptic agonists e.g. oxymetazoline, α -methylnoradrenaline and tramazoline. Although this method is not without error it illustrated the differences between the pre- and post-synaptic receptors with regards to potency of different agonists. Similarly, the effectiveness of different α antagonists at the two receptor sites in a single tissue was found to be different. These differences in pharmacologic profiles suggested that there is a fundamental difference in the structural requirements for

binding at the two receptors. This resulted in the classification of the post synaptic receptor as α_1 and the pre-synaptic receptor as α_2 as originally suggested by Langer(5) and later adapted universally following the review by Berthelsen and Pettinger(6). The studies involving the determination of relative pre- and post-synaptic potencies of agonists demonstrated a fairly consistent pattern of the relative order of potency for the pre-synaptic effects of different agonist between different tissue preparations. However, the relative postsynaptic potencies of the different drugs showed discrepant findings in different vessels. In particular, some apparently pre-synaptically selective drugs were found to be quite effective on the post-synaptic This was not well understood at the time the above side as well. experiments were carried out. However, with the scovery of highly selective α_1 and α_2 antagonists it became clear that α_2 -receptors were present on the post-synaptic membrane as well. As the relative numbers of α_1 and α_2 receptors on the smooth muscle cells (i.e. postsynaptically) would differ from tissue to tissue the above mentioned discrepancies would no longer be entirely unexpected.

The mechanism by which pre-synaptic α_2 receptor stimulation leads to a decrease in the release of noradrenaline is believed to be mediated by Ca²⁺ ions(129). Exocytotic release of catecholamines is triggered by an increase in neuroplasmic Ca²⁺ ions. Pre-synaptic α_2 receptor stimulation probably leads to a decrease in the availability of the Ca²⁺ ions involved in this neurosecretory coupling. The available evidence can be summarised as follows.

1) The release of noradrenaline from adrenergic nerve endings produced by both transmural nerve stimulation and high V solution is dependent on an influx of Ca^{2+} ions with exocytosis of the storage vesicles. In contrast, the indirectly acting sympathomimetic agents release noradrenaline from nerve endings by displacement without producing an increase in neuroplasmic Ca^{2+} ions. Forenz et al(174) in a study in the isolated canine saphenous vein demonstrated that while noradrenaline inhibited the release of ³H-noradrenaline produced by both K⁺ and Table it had no effect on the release produced by tyramine, an indirect sympathomimetic agent. Starke and Montel(175) found a similar differential effect in the isolated, perfused rabbit heart with both α -agonists as well as α -antagonists. 135

The magnitude of the pre-junctional inhibitory effect produced by α -agonists has been found to be a function of the extracellular Ca²⁺ concentration(129). Thus, lowering the extracellular Ca²⁺ from that normally present in the superfusion fluid (2.5) to 1.0 mmol/l resulted in a dramatic potentiation of the inhibitory effect of clonidine on stimulation induced noradrenaline efflux(176). Conversely, raising the Ca^{2+} concentration to 5.0 mmol/l prevented the inhibitory effects of cronidine. A similar Ca^{2+} dependency has been observed in the pre-junctional inhibition of adrenergic neurotransmission produced by adenosine in the canine saphenous vein(177).

The pre-junctional inhibition of noradrenaline release by activation of α -receptors has a negative correlation with the frequency of stimulation of adrenergic nerves(129,178,179). This too is consistent with a mediation via Ca²⁺ ions.

2.

Presumably, at higher frequencies of stimulation, more Ca^{2+} is available in the neuroplasm overriding the inhibition produced by the activation of pre-synaptic α -receptors.

Although the evidence in favour of Ca^{2+} as the mediator during presynaptic α_2 inhibition is convincing, how the α_2 -receptor activation limits the availability of Ca^{2+} is not known at present. Theoretically, there could be a decreased influx or an increased efflux of Ca^{2+} or some alteration in Ca^{2+} utilisation (e.g. by affecting its affinity or binding to sites important for exocytosis) at the adrenergic nerve terminal.

Pre-junctional inhibition may partly account for the observation that uptake₁ blockers such as cocaine or designamine produce only a mild to moderate increase in ³H-noradrenaline overflow during TNS although uptake₁ is believed to play an important role in the disposition of neurally released transmitter. This may be due to the higher cleft concentration of noradrenaline produced by the uptake₁ blockede inhibiting its own release during TNS. Thus, the actual release of noradrenaline may in fact be less during blockade of neuronal uptake. The other factors that might contribute to this apparently anomelous finding is the fact that uptake₁ is inhibitied during TNS even in the absence of the uptake₁ blockers (thus not such further effect sould be expected when the blockers ar incrediential only the onse in alto its local anaesthetic effects

Although the evidence for pre-supplie resceptor mediated inhibition of noradronaline appears strong on the above evidence, its existence has been questioned by Kalener(180) resulting is it elu discussions via journal articles with the proponents of the theory(191). The evidence against the theory is commarised below

Angus and Korner(182) investigated the effects of α -antagonists on the chronotropic response of guinea pig atria to a single pulse or a train four pulses (one pulse following each of four consecutive atrial electrograms) applied during the refractory Under control conditions a tachycardia was observed. period. Although phenoxybenzamine produced a marked potentiation of this response, two other α -receptor antagonists, phentolamine and yohimbine were without effect on the response to both single and four pulse stimulation. The same result was obtained when the four pulses were applied within one refractory period. As the concentrations of phentolamine and yohimbine used were adequate for effective blockade of α -receptors, and as yohimbine is known to be a relatively selective α_2 -antagonist, these findings are unexplainable on the basis of the pre-synaptic a-receptor The uptake blockers designamine and β -oestradiol (used theory. a _____iar in combination) produced potentiation to phenoxyhenzamine in this prepararation. This suggests that the potentiation produced by phenoxybenzamine may have resulted from uptake blockade rather than pre-synaptic a receptor blockade in this preparation. Thus, no evidence was available for presynaptic invibition in the guinearrig atrium during the experimental conditions of the shove study. Rand and coworkers(181) in a separate study investigated the effects of four pulses at frequencies of 0.125, 0.25, 0.5, 1 and 2Hz in the guinea-pig strium measuring the overflow of "H-noradrenaline. Flenthland the efflux and the cardiac responses at and 1 17 but wet a 0.12" or 2Hr. The conclusion

1.

· N

reached was that the pre-synaptic inhibitory mechanism had a latency in excess of 1.5 seconds (as the inhibitory feedback was not present at 2 Hz) and had a persistence of between 4 and 8 seconds (as the inhibitory effect is present at 0.25 Hz but not at 0.125 Hz(181). It was further postulated that a mechanism with such temporal characteristics would probably involve a second-messenger formation and destruction, with the prejunctional receptor probably coupled to an anzyme generating the second messenger(181). Powever, it should be noted that there is no evidence available at the present time for an of the above conclusions reached by Pand, McCullogh and Story.

Kalsper(183) investigated the effect of phonoxyhouzamine (3.3 v 10"5 mol 1) on the efflux of ³H noradrenaltie and the mechanical response of the guinea-pig was deference to a single pulse and a train of four pulses at "Pa - Thenovybrossodine 'nervasai the efflux of "H poradrepultion to components a note pulle t 328 per cent to introl alice. This mechanics response inc 100 increased time for a A station regult we stration to its four pulse to the Plack to of both uptake, and uptake, attant stanificantly if the aschulant reasons to attract a sing) pilor the nate that the offects of the combines are can and is exident of an offer this uptake blockade. There is not a also increased the basal of flux of 311-paratraph to the able The artest patent that the sime of the target ne a stille 1.15 1 . 111 e te severe e la pagette de la companya de la

inhibit further release by the same pulse appears unlikely as the inward Ca²⁺ current (which triggers the exocytotic release) is essentially terminated with the end of the action potential: not commence for another the release ltself may few mtlltseconds. Rand and co-workers(181) explained these ohser ations that spontaneously released 01) the haafa transmitter may be exerting a feed back inhibition in the vas deferens in the resting stat phonoxybenzamine by removing this inhibition would increase the efflux of ³H-noradrenaline proluced by a single pulse. The entremaly parrow cleft in the was deference(144) was suggested as the factor responsible for the effect of apontaneously released transmitter (as a higher concentration would be attained with a marrow cleft). Although this is plausible no evilance is available at present that such phenomenan actually token place. Further, Angua and Parner (18?) observel similar potentiating affect of phenorybenza in the gill capig right atrium which has a alder elist. It ould be good that spontaneously released nor desaling on the second or the second of a sprore for the Wilner ply fast works "leverer, this is next world not he complete the the process of the strong from to Pand et al for the abs are of potential or i phentolamine of the offlue produced by stimulation in 125 Unit is was suggested that preevents, "highly ton hat permittee a of loss than a coords. this of appointer costs is to monoterinalize and an and the state of t

If negative-feedback inhibition takes place during continued release of endogenous noradrenaline at the cleft, the magnitude of the inhibition of efflux inhibition, produced by a fixed concentration of exogenous noradrenaline should decline as the synaptic concentration of coradrenaline increased by increased endogeneous transmit of therefore. Towaver, a fixed concentration of exogenous periodrenaline larveage the stimulation-twineed offlux of "therefore to "00 pulses inlinear at 7.5 mill 15 Hz to planting entont (1, 18 and 44 per cert inhibition respective) at the same three (requencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies) in the scall entotice(18.5 Hz to planting three frequencies)

· •

.

. .

Although phonogyper of and all operations repared the at mule in induced with the second sector in the second and your your $(1-\varepsilon_{1})^{2} = (1-\varepsilon_{1})^{2} = (1-\varepsilon_{1})^{2$ 网络中国人名英格兰卡莱斯人名英格兰卡尔 法法法的 计分子 化分子子 网络白垩合 网络白垩合 化分子分子 off filling and the set of the se . 1 1 . ev1 Attended with the second state of the second s · · · · alter in the lot of · • • 1 1 1 1 T

and a set of the set o

experiments would exert post-junctional effects by themselves. Utilisation of specific α -agonists and α_2 -antagonists will not overcome this confounding effect as α_2 -receptors have been discovered post-junctionally as well, during the last few years.

"alaner regards the observation of lesser pre-synaptic inhibition with increasing frequency of stimulation as being against the theory(180). It was suggested that higher eleft concentrations produced by higher frequencies of stimulation should lead to a greater effect on the pre-synaptic α_2 -receptors. However, this argument cannot be accepted as evidence against the theory as it is only a philosophical argument that regards the pre-synaptic inhibition as the dominant event at the adrenergic neural eleft. It could be argued equally well that higher (requencies of atimulation should be able to override preevanptic inhibition, as the prime function at the eleft should be the inlease of tragenittee.

Another factor that has to be considered in experiments' in solidarith processing to rodulation is the megnitudes of the increases or depresent in transmitter rises observed. In some experiments the increases in transmitter rises observed. In some experiments the increases in transmitter rises observed. In some experiments the increases in transmitter rises amount to less than alogic introduces. Although a single fold change may seem substantial a with these responses in transmitter release may not have any appreciable of it on the effector response(180). Although this by itself does not argue agricul the existence of a pre-synaptic theory if queellons the implication of the effects of pre-synaptic inhibition with respont to the cut in a response. However, it should be noted that much large the solution of the rise of the pre-synaptic inhibition

- mything - stores

In summary, although considerable evidence is available at present for the pre-synaptic α_2 -receptor theory, a number of observations remain unexplainable on the basis of this theory. The significance of the phenomenon may differ markedly not only between species and between tissues, but also with the particular experimental condition being studied. Thus, caution should be exercised in using this theory to explain experimental observations unless the existence of the phenomenon has been demonstrated in the particular tissue studied under similar experimental situations.

142

In addition to the pre-synaptic α_2 -receptors, a variety of other pre-synaptic receptors have been described at adrenergic nerve endings(15.129,171,479). Some, like the pre-synaptic α_2 =receptors, have been shown to inhibit the release of noradrenaline while others have been shown to facilitate the release of the transmitter. The inhibitory mediators include acetylcholine(187,188,189), adenosine and adenine nucleotides(190,191), histamine (H_2). 5 hydroxytryntamine(192) and some prostaglandins(121,194,195). The facilitatory substances include angiotensis 12(1) R promists(197,198) and some prostagle inter(197, 198).

METHODS

GENERAL METHODS

Lateral saphenous veins excised from anaesthetised mongrel dogs (15-25 kg body weight) of either sex were utilised in the studies. The dogs were anaesthetised with sodium pentobarbital (Sommotol) or α -chloralose (100 mg/kg body weight) and the veins excised together with the aurrounding fatty and connective tissue. Care was taken during dissection to avoid unnecessary stretching or trauma to the veins from the instruments used for the excision. The excised vein segments were immediately placed in *oxygenated, cold, modified Krebs-bicarbonate buffer solution of the following composition (mmol/1): NaCl 116.0, KCl 5.4, Cacl₂ 1.7, NaHCO₃ 22.0, NaH₂PO₄ 1.2, MgCl₂.6H₂O 1.2, CaNa₂EDTA 0.023).

Preparation of venous strips and rings(199)

The excised veins were placed in a dissecting tray containing cold Krebs-bicarbonate buffer solution and the excess fatt, and connective tissue removed while leaving the thin, nearly transparent sheath of connective dissue enrounding the tunical adventitia intact. Preserving this sheath beloed in obtaining a maximum response to transmural nerve stimulation closer to the maximum contraction produced by exogenous noradrenaline. This is possibly due to the fact that the nerves funervating the tunical media of the blood vessels course along the surface of the blood vessel before extending into the media through the adventitia. The blood vessels were then cut into either spiral strips or rings. The spiral strips were cut at an angle of approximately 45° to the long axis of the blood vessels using a pair of fine cut discention of the blood vessels using a pair of fine

and 15 - 20 mm in length. A strip cut properly tended to curl itself back into the shape of an intact blood vessel. The rings (approximately 4 mm in width) were prepared by cutting transversely across the venous segments with a pair of sharp scissors or a razor blade. During this whole procedure (of cutting strips or rings) the tissue was kept moistened with oxygenated Krebs-bicarbonate solution.

The strips and rings were suspended in tissue baths (capacity 12 ml or 22 ml) containing Krebs-bicarbonate buffer solution at a pH of 7.4. The apparatus used is shown in Fig. 11. The buffer solution was maintained at 37°C and continuously aerated with a gas mixture containing 95 per cent 0_2 and 5 per cent $C0_2$, the flow rate of which was controlled by an aerator valve. The temperature in the tissue bath fluid was maintained constant with the aid of a heater/circulater (Model No. E15, Haake Mess-Technik Gmbt U.Co., Karlshrue, Federal Republic of Germany) which circulated heated water through a water jacket incorporated in the tissue bath. Fresh Krebs-bicarbonate buffer solution was fed into the tissue bath from a reservoir which was also aerated and maintained at 37°C. The bath fluid was changed with the aid of drainage/"feeder" tubes and the bath volume was maintained constant by means of an overflow tube. Both drainage and overflow tobog ware connected to a vacuum apparatus to ensure complete drainage.

The strips were mounted between two parallel, rectangular, platinum electrodes and their lower ends attached to a momenable support, allowing fine adjustments in the lengths of the strips. The upper ends of the strips were connected to a force transducer (Model No., FT .030 Grass Instrument Co., Quincy, Mass. U.S.A.) for isometric tension recording. The cutrute of the transducers were amplified and incorded (Hodel No.



Figure 11. A diagrammatic representation of the apparatus used for the pharmacological experiments on the isolated canine saphenous vein in the present study.

2400S, Gould Inc. Cleveland, "Ohio, U.S.A.). When rings were used they were mounted on two stainless steel triangular clips and the triangular clips in turn attached to the moveable support at the lower end and the force transducer at the upper end. 5-0 silk string was used for all Transmural nerve stimulation (TNS) was applied from a attachments. stimulator via an impedance coupler(200). The electrical impulses consisted of square wave pulses: 1.0 ms duration, 10 volts. With the impedance coupler the voltage measured across the platinum electrodes was approximately equal to the voltage setting on the stimulator. The duration of the square wave pulses as well as the voltage across the electrodes were periodically monitored with the aid of an oscilloscope, to ensure that these values corresponded to the dial settings on the Before experimentation, the rings were stretched to the stimulator. optimum points of their length-tension curves using the method described by Vanhoutte and Leusen(201). This involved the determination of the length at which the response to a fixed train of electrical stimuli is a maximum (train parameters; 10 volts, 1.0 ms, 16 Hz for 10 seconds). The length of the preparation was increased (1 mm every 3 minutes) starting from a length at which the tension was approximately zero. Following 2 minutes of stabilisation at each length the preparation was stimulated With the initial increases in with the standard train of stimuli. length the response to the train of stimuli increased. The magnitude of the response reached a plateau with further increases in length and then decreased if lengthening was continued further. The optimum length was regarded as the length at which magnitude of the responses reached a The preparations were then equilibrated for a further 90 plateau. During this time the bath fluid minutes before the experiment proper-

was replaced with fresh buffer solution every 30 minutes.

Dose-response curves to all agonists were obtained in a cumulative manner, graded doses being added to give the desired concentration of the drug in the tissue bath(202). The total volume of solution added to the tissue bath during a cumulative dose-response curve was less than 4 per cent of the total volume of the bath. Following a complete doseresponse curve no other drug was tested for at least 60 minutes with repeated changes in the bath fluid during this period, to minimise any adverse effects due to desensitisation.

Stimulus-response curves to transmural nerve stimulation was obtained in a cumulative manner, by increasing the frequency two-fold at each step from 0.5 through 16 Hz (pulse duration 1.0 ms, 10 volts).

SPECIFIC METHODS

The specific protocols carried out are listed below:

Interaction Experiments: Protocol One

Protocol 1.1

Determination of the response to exogenous noradrenaline against a background contraction produced by transmural nerve stimulation (TNS)

Protocol 1.2

Determination of the response to TNS against a background contraction produced by exogenous noradrenaline.

Protocol 1.3

Determination of the response to exogenous noradrenaline against a background electrical current produced by TNS following blockade of the contractile effects of TNS with guanethidine or diltiazem.

-147

Protocol 1.4

Determination of the response to exogenous noradrenaline against

a background contraction produced by TNS and against a background contraction produced by either tyramine, methoxamine,

histamine or phenylephrine.

Protocol 1.5

Determination of the response to exogeneous noradrenaline against a background contraction produced by TNS and repetition -of the same in the presence of either propriatelol, indomethacin, aminophylline or cimetidine.

Superfusion Experiments: Protocol Two

Protocol 2.1

Determination of the effect of exogenous noradrenaline applied against a background contraction produced by TNS, on the release of tritiated noradrenaline and its metabolites produced by the TNS: experiment carried out in the presence of cocaine.

Protocol 2.2

Determination of the effect of a background exogenous noradrenaline, on the release of tritlated noradrenaline and its metabolites during TNS applied against this background: experiment carried out in the presence of cocaine.

Protocol 2.3

Column chromatographic analysis of the superfusate samples from Protocol 2.1 and Protcol 2.2 to separate the total radioactivity present in the superfusate into noradrenaline and its metabolites.

Relaxation Experiments: Protocol Three

Protcol 3.1

Determination of the effect of TNS on a vein pre-contracted with prostaglandin $F_{2\alpha}$ following blockade of the contractile response to TNS with guanethidine and phenoxybenzamine.

Protocol 3.2

Determination of the effect of the following drugs on the relaxatory response to TNS observed in Protocol 3.1: cimetidine, indomethacin, ouabain, aminophylline, cyproheptadine, tetrodo toxin, ascorbic acid, catalase.

Protocol 3.3

Determination of the effect of storage of the saphenous veins at 4°C for 9 days, on the relaxatory response to TNS observed in Protocol 3.1.

Protocol 3.4

٢

Determination of the effect of chemical sympathectomy of the saphenous vein rings using 6-hydroxydopamine, on the relaxatory response to TNS observed in Protocol 3.1

Interaction Experiments: Protocol One

In this series of experiments the interaction between transmural nerve stimulation (TNS) and exogenous noradrenaline was studied in isolated canine saphenous vein strips. After setting up of the preparation and equilibration for 90 minutes the following protocols were carried out. Only one protocol was done in each preparation. Protocol 1.1.

This was designed to compare the response to a concentration of a

149.

concentration of exogenous noradrenaline added against a background contraction produced by TNS. The protocol consisted of the following steps (Fig. 12). Firstly, a dose-response curve to noradrenaline was obtained: Step A. Tension was then allowed to return to baseline with repeated changes of the bath fluid. This was followed by a stimulus response curve to transmural nerve stimulation: Step B (approximately 60 minutes between completion of dose-response curve in step A and commencement of Step B). Then a dose of noradrenaline was added to produce a contraction between 20-80 per cent of the maximum: Step C. This was taken as the initial control. After washing and equilibration TNS was applied to produce a contraction betweeen 10-90 per cent of the control. Once this contraction reached a plateau the control dose of noradrenaline was added while maintaining the background TNS and the response determined: Step D. After washing and equilibration, the control was repeated: Step E. This was taken as the final control. Steps C. D and E of the proposal were then repeated using different magnitudes of contraction produced by buckground TNS (range: 10-90 per of the control) and different concentrations of exegencus cent noradremaline as the control (range: to produce between 20.80 per cost of the maximum contraction to noradrenaline in the preparation).

The study was designed to compare the contraction produced by a concentration of noradrenaline with the additional contraction produced by the same concentration of noradrenaline against a background of TNS. However, a direct comparison between the two would be erroneous for the following reasons. The contractions to noradrenaline in Step C and Step E (controls) and in Step D do not commence from the same le all of active tension in the vein . Active tension is provided to the same le all of active tension in the vein .

NORADRENALINE (NA) DOSE-RESPONSE CURVE CONCENTRATIONS: 10 ⁻⁷ - 10 ⁻⁵ mol/L TRANSMURAL NERVE STIMULATION (TNS) - RESPONSE CURVE STIMULUS PARAMETERS: 10V, 1 M.SEC, 0.5 - 16 HZ ADDED A DOSE OF NA TO PRODUCE A CONTRACTION BETMEEN 20-80% OF THE MAXIMUM - INITIAL CONTROL
CURVE STIMULUS PARAMETERS: 10V, 1 M.SEC, 0.5 - 15 HZ ADDED A DOSE OF NA TO PRODUCE A CONTRACTION
APPLIED TNS TO PRODUCE A CONTRACTION LESS THAN INITIAL CONTROL (RANGE 10-90%). CONTROL DOSE OF NA ADDED WHEN CONTRACTION REACHED A PLATEAU TNS MAINTAINED THROUGHOUT. EFFECT OF NA MEASURED - TEST VALUE
STER REPRATED EINAL CONTROL



٠

ç

ξ,

3188 F

**

...

151

,

Figure 12. Summary of the stops in Protocol 1.1: MAterogenous coordinate list construct nerve stimulation.

Step C and Step E while it is greater than zero by the amount of active tension produced by TNS in Step D. The magnitude of the contraction produced by a fixed amount of an agonist would decrease progressively as the active tension is increased commencing from zero. This is evident from the hyperbolic nature of the dose-response curve to agonists, i.e., a decrease in slope of the dose-response curve as one proceeds at ng the x-axis. (Note: The dose response curves to agonists is linear in the mid-range only after a logarithmic transformation of the concentration of the agonist). Thus, the amount of an agonist that is required to produce a contraction from zero to 10 per cent would be much less than the amount needed to increase the contraction from 10 to 20 per cent Conversely, the magnitude of the contraction produced by a fixed amount of an agonist would lessen as one proceeds along the weakis of the done This is identical to the situation encountered in response curve. Thus a direct comparison between the contractions Protocol 11. produced by noredignaline in thep C and thep I and that preduced in Ster D fo not possible Before a contration constant on main the magnitude of the contraction that would be produce by the outrol dege of norphrenaline componeing from one of the contest of the guilt lent to that priduced by the sull have to be leaded The yay 4 has eq Rollows Fig. 11). If the entraction proceeding the tran Dorsey V per cent of the maximum (for porndrenaline of the particular pr persion studi d) and the total contraction produced by INS and includion line and the observed contraction to the normalized line along that he cont). The extrac of entrantion to the sea ret 1. f 1stnpl' g the state of the second of the second of 317 1 Stat





Figure 13. A diagrammatic representation of the method of calculation of the expected contraction for a fixed concentration of exogenous noradrenaline (Γ_{na}) , added against a background contraction produced by transmural nervy stimulation (INS) in Frotocol 1 1. Step C, Step D and Ster E refer to the respective steps in Fratecol 1.1 (refer Fig. 12). All correctages loss be a opensed toking the maximum contraction to r^{1} a 100 ber seat.
noradrenaline dose-response curve shown in Fig. 13. The X per cent contraction produced by the background TNS was related to a notional point on the abscissa. The same concentration of noradrenaline as used in the controls was added along the abscissa starting from this point and the corresponding total contraction was read off the ordinate. This is represented by Z per cent. Thus, the <u>expected contraction</u> was taken as (Z-X per cent). Comparisons were made between the <u>observed</u> <u>contractions</u> and the <u>expected contractions</u> using the student's t-test for paired data. The dose-response curve used for calculating the <u>expected contractions</u> was obtained by pooling the data from the doseresponse curves done each day. Although this method of determining an <u>expected contraction</u> may not be without error, it was used as the best method available as a direct comparison sould be mining an expected contraction was not be without error.

*

Instant 1 2

. .

This was derived to ever the response to a frequency of transmiral very difficult of the response to the side frequency of TNS against a backgi will offertion produced by exogenous accadeonation. I all the restrict of Protocol 1.1. The protocol construct of the fullying shape fig 10. Firstly, a dese response our considerations was how Step A. This was foll will be a citized of the subject to transmiral nerve atimulation: Step 3. This a firstoon of TNS was applied to produced a contraction between 0.80 per cent of the maximum: Step 0. This was taken as the initial control. After the maximum: Step 0. This was taken as the initial control.

154

y

- STEP A NORADRENALINE (NA) DOSE-RESPONSE CURVE.
- STEP B TRANSMURAL NERVE STIMULATION (TNS)-RESPONSE CURVE STIMULUS PARAMETERS; 10V, 1 M.SEC, 0.5-16 HZ
- STEP C APPLIED TNS TO PRODUCE A CONTRACTION BETWEEN 20-80% OF THE MAXIMUM INITIAL CONTROL
- STEP D ADDED NA TO PRODUCE A CONTRACTION LESS THAN INITIAL CONTROL (RANGE 10-90%). WHEN CONTRACTION REACHED A PLATEAU, CONTROL TNS APPLIED. NA PRESENT IN BATH THROUGHOUT. EFFECT OF TNS MEASUPID - TEST VALUE



Figure 14. Summary of the stops in Protocol 1.2; NA:exogenous ore localine, TNS:transmural nerve stimulation.

of TNS was applied and the response determined while the exogenous noradrenaline remained in the tissue bath: Step D. After washing and equilibration the control was repeated: Step E. Steps C, D and E of the protocol were then repeated using different magnitudes of contraction produced by the background exogenous noradrenaline (range: 10-90 per cent of control) and different frequencies of TNS as the control (range: to produce 20-80 per cent of the maximum contraction to noradrenaline in the preparation). The frequencies of TNS and the concentrations of exogenous noradrenaline in Step D of both protocols were chosen in a manner that the total contraction in Step D would be less than the maximum obtainable with exogenous noradrenaline alone in the saphenous vein. This ensured that the total response observed in Step D was not limited by the maximum contraction attainable to noradrenaline in the preparation.

Protocol 2.2 was designed to compare the contraction produced by a frequency of TNS with the additional contraction produced by the same frequency of TNS against a background contraction produced by exogenous noradrenaline. As detailed under Protocol 1.1 a direct comparison would be erroneous as the contraction to TNS in Step C and Step E (controls) and in Step D do not commence from the same level of active tension. Thus, before a comparison could be made the magnitude of the contraction that would be produced by the control frequency of TNS commencing from an amount of active tension equivalent to that produced by exogenous noradrenaline would have to be calculated. This was done as follows (Fig. 15). If the contraction produced by TNS alone in the controls is C per cent of the maximum (for noradrenaline in the particular preparation studied) this could be related to a notional concentration

, ,



157

Figure 15. A diagrammatic representation of the method of calculation of the expected contraction for a fixed frequency of transmural nerve stimulation (TNS), applied against a background contraction produced by exogenous noradrenaline (NA) in Protocol 1.2. Step C, Step D, and Step E refer to the steps in Protocol 1.2 (refer Fig. 14). All percentages have been expressed, considering the maximum contraction to exogenous noradrenaline in the canine saphenous vein as a 100 percent.

> . .

of noradrenaline on the abscissa (represented by d mol/1). This would be the concentration of exogenous noradrenaline necessary to produce a contraction equivalent to that produced by the control frequency of TNS, and was used as a substitute for the latter in calculating the expected contractions. If the contraction produced by exogenous noradrenaline in. Step D was X per cent of the maximum (for noradrenaline in the particular preparation studied) and the total contraction produced by the exogenous noradrenaline and TNS was Y per cent the observed contraction to the TNS alone would be (Y-X per cent). The expected contraction to the same frequency of TNS commencing from a level of produced by tension equivalent to. that the active exogenous noradrenaline was calculated using the noradrenaline dose-response curve shown in Fig. 15. The X per cent contraction produced by the background noradrenaline was related notional point along the abscissa. The concentration of exogenous noradrenaline that was equivalent to the control frequency of TNS was added along the abscissa starting from this notional point and the corresponding total contraction read off the This is represented by Z per cent. ordinate. Thus, the expected contraction was taken as (Z-X per cent). Comparisons were made between the observed contractions and expected contractions using the student's t-test for paired data. The dose-response curve used for calculating was obtained by pooling the data from the the expected contractions individual dose-response curves done each day.

Results of Protocol 1.1 and Protocol 1.2 indicated that the contractions produced by exogenous noradrenaline in the isolated canine saphenous vein were "Inhibited" by background TNS as in Protocol 1.1. Therefore further experiments were performed to elucidate the mechanism

, , 158 responsible for this inhibitory phenomenon. These experimental protocols were further extensions and modifications of Protocol 1.1 as the inhibitory phenomenon was evident in this protocol only.

Protocol 1.3

This protocol was performed to determine whether the inhibitory effect of TNS on the exogenous noradrenaline contraction was due to the excitation of intramural nerves or due to the field of current per se. Steps A-E of Protocol 1.1 were done as before. Then the adrenergic neurone blocking agent guanethidine was added to the tissue bath in a concentration just sufficient to block the contractile effects of TNS at the frequency used in Step D. (Note: Complete abolition of the contractile effects of TNS up to a frequency of 32 Hz was not attempted). Guanethidine was kept in the tissue bath for a minimum of 60 minutes before further experiments were carried out and was present in the bath throughout the rest of the experiment. After the blockade of the contractile response to TNS, Steps C, D and E were repeated i.e., the two controls and Step D. In Step D, TNS, at the same frequency as used before the addition of guanethidine, was applied first. This did not produce a response but the flow of electrical current between the platinum electrodes during the current pulses would still be present. After 5 minutes of TNS the control dose of noradrenaline was added while maintaining the current and the response determined. The contraction produced by exogenous noradrenaline in Step D was compared with the mean of that produced in Step C and Step E by a student's t-test for paired Calculation of expected contractions to noradrenaline was not data. necessary as the contractions with and without guanethidine commenced from the same level of active tension (which was approximately zero in

the present instance).

Steps C, D, and E of the protocol were also repeated using the calcium channel blocker diltiazam in place of guanethidine. Diltiazem produced a differential effect on the contractions produced by TNS as compared with the contractions produced by exogenous noradrenaline in the canine saphenous vein. It was used to block the contractile effects of TNS while leaving the contractile effects of exogenous noradrenaline relatively unaltered. Again, complete blockade of TNS induced contractions up to a frequency of 32 Hz was not attempted. A concentration sufficient to block the effects of TNS at the frequency used in Step D was used. Incubations were carried out for minimum of 60 minutes with diltiazem and it was present in the bath fluid for the rest of the experiment.

Protocol 1.4

These experiments were designed to determine whether the inhibitory effect of background TNS, on the exogenous noradrenaline mediated contraction, observed in Protocol 1.1 was specific for TNS or whether it would still be evident when another agonist is substituted in place of TNS. The agonists used were:

(a) tyramine - an indirectly acting sympathomimetic agent

(b)

methoxamine-a specific α_1 agonist which is a poor substrate for uptake₁ and thus with no indirect sympathomimetic effects

(c) histamine

(d) phenylephrine

The concentration of tyramine used was kept low enough to avoid direct effects on smooth muscle which appear at higher concentrations. The concentration range which produced indirect effects only was determined in preliminary experiments where dose-response curves to tyramine were done with and without an uptake₁ inhibitor.

In Protocol 1.4, Steps A-E of Protocol 1.1 were done first. Then a dose of tyramine (or methoxamine or histamine) was added into the tissue bath to produce a contraction approximately equal in magnitude to that produced by the background TNS in Step D (Fig. 16). Once this contraction to tyramine reached a plateau, the control dose of noradrenaline was added and the response determined. The additional contraction produced by noradrenaline against a background of tyramine was compared with the additional contraction produced by noradrenaline against a background of TNS with the student's t-test for paired data. Calculation of expected contractions to noradrenaline was not necessary as the imagnitude of the background tensions (produced by TNS and tyramine) were approximately equal. In practice the triad (steps C, D and E) with (tyramine (or histamine or methoxamine) was carried out first and then the triad was repeated substituting TNS in place of tyramine as it was easier to match a contraction by TNS equal in magnitude to that produced by tyramine than vice versa.

Protocol 1.5

b)

c)

This was designed to investigate whether the inhibitory effect of TNS on the exogenous noradrenaline mediated contraction observed in Protocol 1.1 would still be present in the presence of the

a) beta-blocker propranolol (10⁻³ mol/1)

cyclo-oxygenase inhibitor indomethacin $(10^{-5} \text{ mol}/1)$

- P_1 -purinoceptor antagonist aminophylline (10⁻⁵mol/1)
- H_2 -receptor antagonist cimetidine 10^{-4} mol/1

Firstly, steps A-E of Protocol 1.1 were repeated as before. Then one of the above drugs was added to the tissue bath to produce the



162

≜NA

Figure 16. Steps of Protocol 1.4. The first part of the protocol is shown in the upper half of the figure: the control contraction to the exogenous noradrenaline (initial control), the contraction to the same concentration of noradrenaline added against a background contraction produced by transmural nerve stimulation (TNS) and the final control. The second part of the protocol is shown in the lower half of the figure: the initial control, the contraction to the same concentration of noradrenaline added against a background contraction by tyramine (of equal magnitude to the contraction produced by TNS in the first part of the study) and the final control. concentration given above. The drug was then present in the bath for the rest of the experiment. Steps C,D, and E were repeated following incubation with the drug for a minimum of 30 minutes. Only one drug was tested in one vein strip. The additional contractions produced by exogenous noradrenaline against a background of. TNS with and without each drug were compared with the student's t-test for paired data.

Superfusion Experiments: Protocol Two

In step D of Protocol 1.1 exogenous noradrenaline was added against a background contraction produced by TNS and the additional response measured. When the additional response is taken as that produced by the exogenous noradrenaline, the assumption made is that the magnitude of the contraction produced by the background TNS remains unchanged during this period. However, exogenous noradrenaline is known to produce inhibition of release of noradrenaline by (thus inhibition of the contraction by) the sympathetic nerves acting on pre-synaptic α_2 receptors(203). If such a phenomenon takes place in Step D of Protocol 1.1 this could confound the results. A similar situation could take place in Step D of Protocol 1.2 when TNS is applied against a background contraction produced by exogenous noradrenaline. Thus it was decided to investigate these two confounding effects by measuring the release of noradrenaline from the sympathetic nerves in isolated canine saphenous veins during experimental situations similar to Protocol 1,1 and Protocol 1.2.

A modification of the superfusion technique described by Vanhoutte et al(188) using tritiated noradrenaline was utilised for this study. Lateral saphenous veins were isolated from anaesthetised dogs as described under General Methods. The veins were cut into spiral strips

. / .

approximately 2-3 mm in diameter and 50-60 mm in length. These preparations were incubated in 8 ml of Krebs buffer solution containing $7-^{3}$ H-noradrenaline (specific activity 10-30 Ci/mmol) in a concentration of 10^{-6} mol/1 while aerating the solution with 95 per cent 0₂ -5 per cent CO2. This was carried out in a fumehood. After 60 minutes the strips were then transferred to a fresh solution of ³H-noradrenaline in Krebs solution and incubated for a further 60 minutes. At the end of the second period of incubation the vein strips were rinsed in 25 ml of Krebs buffer solution (without any ³H-noradrenaline) and mounted for superfusion in a glass chamber similar to that described by Hughes and Roth(204) as illustrated in Fig. 17. The strips were mounted between two platinum wires (0.032 cm in diameter) inside a funnel shaped chamber maintained at 37°C with the aid of a water jacket. The platinum wires were used as electrodes by connecting them to the stimulator-impedance coupler system (refer General Methods). The gaps between the electrodes and the strips were wide enough to allow contraction and relaxation without restraint and yet sufficiently narrow to be filled consistently by part of the superfusate retained by capillary action, thus ensuring continued electrical conductivity. The lower end of the strip was attached to a plastic holder and the upper end connected to a force transducer (Model No. FT .03C, Grass Instrument Company, Quincy, Ma. U.S.A.) for isometric tension recording. The transducer was mounted on a moveable support to allow vertical movement. The connection to the transducer was made with a wettable twine string to aid in the superfusion. The vein strip was superfused with oxygenated Krebsbicarbonate buffer solution pre-warmed to 37°C. The solution was made tog drip along the twine string on to the vein strip between the two





Figure 17. A diagrammatic representation of the apparatus used for the superfusion experiments using ³H-noradrenaline in the present study.

N .

platinum electrodes. The buffer solution cascaded over both the vein strip and the electrodes and was collected in test tubes in a fraction collector (Model No. Frac-100, Phrmacia Canada Inc. Dorval, Quebec, "Canada) through the outlet tube of the superfusion chamber. The samples were collected on a time-basis with a change of the tubes every -2 minutes. The superfusion was carried out at a constant flow rate of 3.0 ml/min with the help of a positive displacement roller pump (Model No. Miniplus 2, Gilson Medical electronics, Viliers. le.bel., France). A three-way stopcock upstream from the pump allowed rapid switching from control Krebs buffer solution to buffer solutions containing drugs whenever necessary.

After setting up of the vein strips the initial basis tension was set at 3.0g by stretching the strips (by moving the force transducer upwards). Superfusion with Krebs buffer solution was continued for 120 minutes before the experiment proper, to allow for any loosely bound \mathbb{R}^3 noradrenaline to be "washed" away and the basal efflux of radioactivity to stabilise at a steady level. The experiments (Protocol 2.1 and Protocol 2.2) were carried out in the presence of cocaine to inhibit uptake₁ as pre-synaptic α_2 inhibition has generally been demonstrated in the presence of an uptake₁ inhibitor(174). The superfusing solution was changed to Krebs buffer solution containing cocaine (3 x 10⁵ mol/1) 30 minutes before the commencement of the experiment.

Protocol 2.1

14.

In order to determine the effects of exogenous noradrenaline on the efflux of 3 H-noradrenaline during TNS, step D of Protocol 1.1 was carried out as a superfusion experiment. As exogenous noradrenaline does not change the bisal efflux of 3 H-noradrenaline significantly in

the presence of cocaine(174), Step C and Step E Protocol 1.1 (1.e., the measurement of the response to а concentration of exogenous noradrenaline alone) were not done here. The experiment proper was carried out as follows. The numbers refer to the consecutive test tube numbers in the fraction collector (each tube corresponds to a two-minute. period of collection containing 2 x 3.0 ml=6.0 ml of the superfusate). The triad of consecutive tubes used for determination of the efflux of $3_{\rm H-noradressline}$ representative for that particular period of the $3_{\rm H-noradressline}$ protocol are denoted by asterisks. These constituted the final three tubes of each period. The pooled superfusate from the same triad of tubes was used in column chromatography to separate the total radioactivity into noradrenaline per se and its different metabolites. PERIOD I: 1* 2* 3* (CONTROL)

Superfusate was collected for 6 minutes (3 tubes) as a control period for determination of the basal efflux ³H-noradrenaline.

PERIOD IL:	4 5 6 7 8* 9 * 10*	(TNS)
	TNS was applied at 2 Hz for 14 minutes with Krebs buffer solution.	while superfusing
PERIOD TIT.	11 12 13 14 15*, 16* 17*	(TNS + NA)
	Superfusing solution was changed to Krebs	buffer containing
	noradrenaline (10 ⁻⁶ moll) for the next	14 minutes while
	maintaining TNS.	· · · · · · · · · · · · · · · · · · ·
PERIOD IV:	16 19 20 21 - 22+ 23+ 24+	(TNS)

Superfusing solution was changed back to plain Krebs buffer while maintaining TNS - 14 minutes.

PERIOD V:

25 26 27 28 29 30 31* 32* 33* (CONTROL)
 TNS terminated. Superfusion was continued with plain Krebs
 buffer for the next 18 minutes.

The protocol is summarized in Fig. 18

		• •	. FOCOCOT	E.	*	- *
	2 10.				÷.	
1*		•		30		
2*	•	•		31*	· · · · · · · · · · · · · · · · · · ·	ι
3*	CONTROL	-		32*	8	
4	TNS			33*	CONTROL.	
5				END OF	EXPERIMENT	
6						• ,
7			·			-
8* .			· ·		١	
9*		· ·			р., (
10*	TNS		•			
11	TNS + NA		99 -			
12						
13			•			
] 4					c •	ħ
15*			, 1		. 1	
16*			, ,			•
17*	TNS + NA					
18	r v g				·) · · · · · · · · · · · · · · · · · ·	м <u>.</u> и.
19					* .	
20						3
21						
>?*						• ·
534						· · · · · ·
24	TNS					
25	· · · · ·					م ر ۲۰۰
6					·	•
7						
28					,	
Ú.						

Protocol 2.1

168

Fig. 18. Summary of Protocol 2.1. Numbers refer to consecutive test tube numbers in the fraction collector. Each tube represents a 2 minute period of collection of superfusate. Asterisks denote the tubes with all for submany ont column chiomalographic analysis.

Total radioactivity present in each sample collected (total of 33 tubes) was determined using a 1.0 ml aliquot of the superfusate (details given below). The superfusate collected during selected 6 minute intervals (the three consectuive test tubes indicated by asterisks) was also utilised for subsequent column chromatographic analysis to separate ⁵H-noradrenaline from its metabolites. To minimise the adsorption of tritiated noradrenaline and its metabolites on to the glass wall of these triads of test tubes, each test tube contained 0.5 ml of a stock carrier solution containing unlabelled noradrenaline and each of its five major metabolites (composition of stock solution given below). The stock carrier solution also contained sodium metables lphite, disodium ethylene diamine tetraacetic acid and hydrochloric acid as protective agents to prevent exidation of catecholamines. These tubes were kept in the refrigerator until just prior to collection of the superfusate. Following collection of the superfusate 1.0 ml was piretted out for total radioactivity measurements as explained above. 'A prection was made in the calculation of total radioactivity in orth of these samples saved for cuberqueut column chromitographic unit ste a the 6 or of or superformate in each to take can diluted on the set he allfold of of 9.5 ml of the stand or the solution. The over filling overformer Constructions (5.5) I to each control from the Database of open tetal on a provides the sport have (method on any and are set, if the to at the set of the set and kept for solum somet or it is all to that the second of

a all of congrations the sadar and for any parager remaining the second second second remaining the second se

and metabolites retained in the tissue were extracted(205) by placing each strip in a small vial containing 2.5 ml of extraction fluid (see below for composition) and agitating the contents of the vial continuously for 30 minutes. Then the strip was transferred to another vial containing the same extraction fluid. After a further 30 minutes of agitation, the strips were removed, and the two 2.5 ml extraction portions were pooled and mixed. 'A 1.0 ml aliquot from this 5.0 ml was

The amount of radioactivity present in each 2 minute (6 ml) sample collected, was expressed as a fraction of the total radioactivity present in the strip, at that particular time-referred to as Fractional release(206). Thus the radioactivity present in the last sample collected was expressed as a fraction of the sum of the total radioactivity present 1.n the tissue extraction fluid and the radioactivity present in the last sample itself. As the resultant fractional toleage values were small, each value was multiplied by 10³ for clarity, and appresend to graphs and tallon.

e.g. redisactivity present in the last sample -400 dpm/mlredisactivity present in the penultimete comple -600 dpm/mltotal radi activity in the tingue struct at the end of the experiment -600,000 dpm/mlFront colored for last sample $\frac{(400 \text{ x} 6)}{(600,000 \text{ x}^5) + (400 \text{ x} 6)}$

170

ert j

Preparation of stock carrier solution (Personal communication Lorenz RR, Vanoutte PM.)

1.

A 0.01 per cent solution of each of the following drugs in 0.01 per cent ascorbic acid was prepared by dissolving 10 mg of each drug in the ascorbic acid solution and making it up to 100 ml: noradrenaline (NA), 3,4-dihydroxyphenyl glycol (DOPEG), 3methoxy-4-hydroxymandelic acid (7MA), normetaneobrine (NMN), 3 methoxy-4-hydroxyphenyl glycol (MOFEC), and 3.5 dihydroxy mandelic acid (DOMA).

800 mg sodium metablisulphite and 800 mg di sodium thviene diamine tetra acetic acid were weighed out into a beaker and 8.0 ml of each of the solutions propared above were added in the following order, while stirring or invousiv with the aid of a magnetic stirrer.

8 ml NE

8 ml DOPEG

R m1 VMA

8 m1 NMN

8 ml MOPEC

R int parts

8 ml of 5 N hudrochlor's wild ung alled loan. The resile ung

 0.5 ± 1 of the construction as a set of the space of the test of the set of the space of the set of the set

-

magnetic stirrer and the 0.5 ml was pipetted out while stirring as the drugs are in suspension. 172

Extraction fluid:Preparation(207)

Extraction fluid used for 3 H-noradrenaline extraction from the tissue was made up as follows.

- 1. 5.5 mg Na₂EDTA and 44.3 mg of ascorbic acid were dissolved in IN acetic acid and the total volume was made up to 500 ml with IN acetic acid itself.
- 2. The resulting solution was kept at 4°C and was good for 6 months.

Radioactivity measurement

Total radioactivity, was measured in all samples collected in the fraction collector. Following the collection, the superfusate contained in each tube (6.0 ml) was mixed thoroughly with the help of a vortex mixer, and a 1.0 ml aliquot pipetted into a liquid scintilation vial. Ten ml of liquid scintilation fluid (Aquason, New England Nuclear, Canada) was added to each of these vials and mixed vigorously with the 1.0 ml of the superfusate, by shaking to form a clear fluid. Radioactivity was measured in a liquid scintilation counter (Model no. 8500, Beckman Instruments). Each vial was counted for 10 minutes or until 10,000 counts accumulated. Corrections for quenching were made with an external standard and counting efficiency was approximately 36 per cent. The cintillation vials were left for 120 minutes for dark Protocol 2.2

This protocol was carried out to determine the effects of a prior application of exogenous noradrenaline on the efflux of 3 H-noradrenaline during transmural nerve stimulation, i.e., corresponding to Protocol 1.2. Thus, Step E, Step D, and Step E of Protocol 1.2 were carried out as a superfusion experiment. Gocaine $(3 \times 10^{-5}, \text{mol}/1)$ was present in the superfusing Krebs buffer solution throughout the experiment. The experiment proper was carried out as follows. The numbers refer to the consecutive test tubes in the fraction collector. The last three tubes during each period of the protocol were selected for subsequent column chromatographic analysis as in Protocol 2.1. These test tube numbers are denoted by an asterisk.

173

PERIOD I: 1* 2* (CONTROL) Superfusate was collected for 6 minutes (three tubes) as a control period for determiantion of the basal efflux of ³Hnoradrenaline. PERIOD II: 4 5 6 7 8* 9* 10* (TNS.) TNS was applied at 3 Hz for 14 minutes while superfusing with plain Krebs buffer solution. TNS was terminated at the end of the 14th minute. (CONTROL) PERIOD ITI: 11 12 13 14 15 16 17 19 19 20 21 22 23 25 251 25+ Superfusion was continued with plath Krebs buffer solution for 31 minutes PERIOD IV: 28 29 **3**0 31 32* 33* (NA) 34* Superfusing solution was changed to Krebs buffer containing noradrenaline (3 x 10"/mol/1) for the pext 14 minutes

> . S

38 39* 40* 41* 35 36 37 (NA + TNS)TNS was applied at 3 Hz for 14 minutes while continuing the superfusion with Krebs buffer containing noradrenaline $(3.3 \times 10^{-7} \text{ mol}/1).$

1

PERIOD VI:

FERIOD V:

47 48 49 50 51 🖉 🧋 42 43 44 45 46 (CONTROL) 52 . 53 54 55 56* 57* 58* TNS was terminated and the superfusing solution changed to. plain Krebs buffer solution for the next 34 minutes as a control period.

PERIOD VII: 59 60 61 62 63* 64* 65* (TNS) TNS was applied at 3 Hz for 14 minutes while superfusing with plain Krebs buffer solution.

PERIOD VIII: 66 67 68 69 70 71 72 73* 74*²⁸ 75* (CONTROL) TNS was terminated and the superfusion continued with plain Krebs buffer solution as a control.

At the end of the experiment the total radioactivity remaining in the vein strip was extracted as explained in Protocol 2, 1. Total radioactivity was counted in each sample of superfuste collected and column a chromatohgraphic analysis was done in [#]selected samples as before. The protocol is summarised in Figure 19.

As stated above, the radioactivity present in the superfusate collected in the triad of test tubes denoted by asterisks were considered as representative for that particular period of the experiment. The last three samples during each period consitituted this which were also triad of test tubes used for the column chromatography. The mean of the radioactivities present in the three complex were used for statistical comparisons.

Tube No.	Protocol 2.2	
Tube No.	a 3 5	
1*		•
	*	- 4- 1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	, 30 (a	59 TNS
2*	31	60 .
<u>3* CONTROL</u>	32*	61
4 TNS 8	33*	62*
5	<u>34* NA</u>	63*
6	35 NA + TNS	64*
7	36 😼	65 TNS 4
8*	37	66 CONTROL
9**	38	67
<u>10* TNS</u>	39 * [≩]	68
11 CONTROL	40*	69
12	41* NA + TNS	70
13	42 CONTROL	71
14	43	72
15	44	73*
16	45	74*
17	46	75* CO TROL
18	47	e e e
19	48	
20	40	
21	5()	
22	52	
24	5 3	
₽° ' 25 *	54	
26*	5.5	
27* CONTROL	۲ 	
28 MA	57 *	
40		

Figure 19. Summary of Frotocol 2.2. Numbers rot the open to tort tube numbers in the fraction collector. Each fully a constraint to . period of collection of superfusate. Asteriak the mapping of the transmission of the complete south the second of the

.

14

۰.

4

30

۰. ۲

j., j.

.

÷

,

.

,

The efflux of 3 H-noradrenaline decays with the passage of time in most preparations in superfusion experiments. To correct for this the following procedure was used(174). Evoked release of 3 H-noradrenaline during an intervention such as TNS (or a combination of TNS and exogenous noradrenaline) was calculated by subtracting the average basal efflux of 3 H-noradrenaline during the two control periods (done before and after the intervention) from the efflux during the intervention.

Column chromatographic analysis(188,208,209)

This was used to separate the radioactive material in the superfusate into noradrenaline (NA) and its major metabolites: 3,4dihydroxyphenylglycol (DOPEG), 3,4-dihydroxymandelic acid (DOMA), normetanephrine (NMN), 3-methoxy-4-hydroxyphenylglycol (MOPEG) and 3methoxy-4-hydroxymandelic acid (VMA).

Summary: Alumina, Dowex-50 and Dowex-1 were used for the separation which was carried out in glass columns (0.5 cm in diameter). First, the catechol compounds (NA, DOPEG, DOMA) were separated from the noncatechols is adsorption of the former on alumina. The NA and DOPEG were eluted from the alumina with acetic acid, followed by DOMA which was elu ed with 0.2N and IN hydrochloric acid. NA was separated from DOPEG (in the sluted NA/DOPEG mixture) by adsorption of the former on Dowex-50 and clution later with hydrochloric acid. The non catechol compounds present in the alumina with the effluent containing MOPEG and VMA. The NMN in Dowex-50 columns with the effluent containing MOPEG and VMA. The NMN adsorbed on Doi or 50, was subsequently eluted with hydrochloric acid in ethanol. HOPEG and VMA were separated by adsorption of the latter on to Douex 1 and its colorequent elution with hydrochloric acid.

it is the petro is two) is unabled and apply attan (210)

This was carried out in a fumehood as follows.

· 1.

0

- 100 g of aluminium oxide was added to 500 ml of 2N hydrochloric acid in a 1000 ml beaker, the beaker covered with a watch glass, and heated at 90° to 100° C for 45 minutes with continuous and rapid stirring, using a magnetic stirrer-hot plate.
- 2. The beaker was removed from the heater-stirrer and the heavier particles of aluminium oxide allowed to settle for 1 1/2 minutes. The supernatant fluid (distinctly yellow in colour) was discarded along with the finer particles of aluminium oxide.
- 3. The aluminium oxide was then washed twice with fresh 250 ml portions of 2N hydrochloric acid at 70°C for 10 minutes, discarding the supernatant with the finer aluminum oxide particles each time.
- Aluminium oxide was washed with 500 ml of 2N hydrochloric acid at 50°C for 10 minutes. Supernations with the fluer alumina could lee discarded as before.
 - The alumina use the washed (stirring for 1.10 minutes) repeatedly (about 20.25 times) with fresh 200 ml postions of disifled water until the pH of the suspendion was 3.4. The market with the final particles was decauted such time.

Finally, the aluminium oride was transferred to un emporating dish and to ivated by heating it 300°C for 2 house or at 200°C overnight (A mi iff atton of the original method as recomminded by Wan outle, "... and "scheur: " principal communication) Atomics is in the scheme and "scheur: " principal communication).

· ~ '

one volume of the Dowex-50 resin was suspended in approximately three volumes of distilled water and continuously stirred for 20 minutes. The Dowex-50 was then allowed to settle' for 30 minutes and the supernatant decanted. This was repeated 4-5 times. Following the final washing the Dower-50/distilled water suspension was filtered on a Buchner funnel-Buchner flask (with d of light negative pressure in the Buchner flask) and air overnight.

Dowex=50 x 4 resin (200-400 mesh) - washing

178

- The resin was then suspended in three volumes of 2N hydrochloric 2. acid and continuously stirred for 30 minutes. The suspension was allowed to settle for 30 minutes and the supernatant discarded. This step was repeated with fresh 2N hydrochloric acid until the supernatant was almost colourless - approximately 2-3 times. (This procedure cycles the Dowex - 50 resin through a H^+ form). 3 The resin was suspended in three volumes of distilled water and stirred for 30 minutes, allowed to settle for 30 minutes, and the supernatant discarded. This was repeated once.
- 4 The Dower-50 was then suspended in three volumes of 2N sodium hydrovide and stirred for 30 minutes and the supernatant discarded (This cycles the resin through a Na⁺ form).
- The resin was then washed four times with distilled water as in 5. step three, and filtered on a Buchner funnel flask and allowed to air dry.

The mached roots was stored in a bottle at 4°C. 6.

Dower 50: an iver 1 grade cation exchange resin - AG 50W X 4, 200-400 month hydrore "io-Rad Laboratories, Richmond, california U.S.A.

1.

 \sim

Dowex-1 resin; no special washing necessary. The resin powder can be used directly in the preparation of the columns. Source; Dowex - 1: stock no. 1 x 4-400 chloride form strongly basic anion exchange resin -4 per cent cross linked - dry mesh 200-400.

Solutions used for Column Chromatography (Appendix V)

All the solutions used throughout the chromatographic procedure, including the de-ionised bi-distilled water, contained 0.1 per cent Triton X-100 unless specified otherwise. The addition of Triton X-100 not only improved the recovery of MOPEG and VMA, but made it also possible to pass effluents and eluates directly over the next column without the formation of air bubbles in its narrow part(208).

Re-pipette dispensers of different volume as indicated below, were used to facilitate delivery of the required volumes of the solutions during the chromatography procedure. As the delivered volume need not be exact, repipette dispensers of the type shown in Fig. 20 (Fisher Scientific Ltd., Toronto, Canada) were used for the purpose. When a number of samples (e.g., 6.8 samples) are analyzed simultaneous's (st is usually done) a fast delivery be omen essential making the dispenser particularly suitable.

Preparation of the Glass Columns

Glass columns used had an thernal Hameter of 0.5.4 and mere plugged with glass wool (Fig. 21). A choir store of silastic table of a attached to the follows diverse in a turn to useble a spring (come to used (a the table of the table of the turn to the table of the table of the



· •

Figure 21. The respipetto dispenses used for delivery of solutions



Figre 21. sterrammenti represe etter of a plane terra terra terra de la sectoria de la sectori

One three tier rack:

Upper: Alumina columns Middle: Dowex-50 columns Lower: Dowex-1 columns

9

The columns were mounted so that the effluent from the alumina column dropped directly into the Dowex-50 resin column, and the effluent from the latter, directly into the Dowex-1 column.

One "Wo tier rack

Prover Alumina columns (the glass columns from the top tier of

the three tier rack are transferred here).

Lower: Dower-50 Jumpa

three single ther tarks

Glass columns from the three ther and two tier racks are ultimately

Graduated cylinders (with glass stoppers, to facilitate the mixing of the solutions contained within) were labelled A,B,C,D,E, and F for collection of the effluents and eluates, containing intact noradrenaline and its five matabolites. They were numbered 1A, 2A, 3A, 8A depending on the number of samples contained simultaneously. Twenty-five ml cylinders were idequal for all i cent the A paries where fifty ml

Copy 51 Slower from tration

A margine in was rrenared by adding one volume (approximately) of discilled even 0 to present fritton X 100 to one volume of the wan of the 30 perdection and stirring continuously with

2. 1.0 ml of the Dowex-50 resin suspension was added (while stirring) to each Dowex-50 column. The resin settled at the bottom of the column above the glass wool. The dowex-50 resin should form a column about 2.5 cm in height: if the height was not adequate some more Dowex-50 suspension was added. The beaker containing he unused resin suspension was covered with parafilm and st in the refrigerator at 4°C for use over the nert for days.

1. The following were added to each column

15 ml 2N HC1

5 ml distilled water/0.1 per cent Triton & 100

15 ml sodium phosphate fulfer

5 ml distilled mater (1) pay cont Triton V 100

Each solution was added only after the previous solution had incided through the column. Let when the meniscus just reached the top of the Dowex-50 resin. Once all four solutions had drained through, the eilastic tubing attached to the lottom end of the column war. 'amped until the experiment proper was begun. It was ensured that a shall amount of fluid remained above the former for resin to sold do the of the compand.

6 Devero50 colugns are place the stand of th

I olumna from

A griggion of the second secon

- 2. 1.0 ml of the resin suspension was added to each Dowex-1 column while stirring. More resin was added, if necessary, to form a Dowex-1 plug approximately 2.5 cm tall. The balance of resin left was kept at 4°C after covering the beaker with parafilm for use over the next few days.
 - 5 ml of 0.1 per cent Triton X-100 was added and the silastic tubing at the bottom clamped once this had drained through the column.
- 4. The Dowex-1 columns were placed on the lower tier of the three tier rack.
- " ' m-chromatography separation procedure

The samples used for the column chromatography were the triads of test tubes containing the superfusate representative of each period of the superfusion experiment. The superfusate collected in the 3 tubes of each "stad were pooled together (after removing 1.0 ml from each to count for total radioactivity) and stored immediately at -23°C until the column chromatographic analysis was done. These tubes contained unlabelled NE, DOPEG, "MA, NMN, MOPEG and DOMA to minimise adsorption of the tritiated compound on to the glass. They also had disodium ethylene di the tetra acetic anid, sodium metableu'nhite and hydrochloric acid ""fective igen's 's prevent oxidation is the tritiated compounds ""fective igen's 's prevent oxidation is the tritiated compounds

While proparing "ower 50 and Dower's columns the first samples were removed from the fractor and allowed to thaw. The contouts of our sample tube were mixed with the old of a rise of the line is a settle for 5 minutes. Then if mi

was pipetted out in duplicate for determination of total radioactivity.

- 3. Then 10 ml of each sample was pipetted into plastic beakers numbered according to the samples.
- 4. Approximately 500 mg of activated alumina was added to each beaker containing 10 ml of the samples.
- 5. A suspension (alumina + sample in beaker) was prepared by continuous stirring with a stirring rod and each sample was titrated to a pH of 8.4. This was done by using 0.5 N Na₂CO₃ for coarse titration and 0.1 N Na₂CO₃ for fine adjustment of the pH. The sample was maintained at pH 8.4 for five minutes using 0.1 N Na₂CO₃ if necessary, while stirring continuously. Note: Na₂CO₃ solutions did not contain Triton X-100
- 6. 50 ml graduated cylinders labelled 1A. 2A etc., were placed under the Dowex-1 glass columns in the lower tier of three tier rack and the titrated samples and the alumina (in the plastic beakers) were added to the glass columns in the upper tier. The plastic beakers were rinsed with distilled water/0.1 per cent Triton 3-100 using a squirt bottle, to tomore fany adversal alumine from the beakers The clemps attached to the time 50 and Dowor I alumns of the three-tier tack were removed.

The alwar (1) of the comple belows through the elumina (elumn " ml of devilled water ()) per cent lifton X 100 was added to remove we alwains adhering to the glass on the upper part of the column. ()) the colutions added to the alumina column draine¹ through to the low relations in the dillector () the three through to the low relations in the dillector () the three DOPEG

\mathbf{N}	8
\sim	<u> </u>

Once the 15 ml distilled water/0.1 per cent Triton x-100 too had drained through, the alumina columns were removed from the upper tier of the three-tier rack and positioned over the Dowex-50 columns in the two-tier rack (thus the same alúmina columns now formed the upper tier of the two tier rack). 25 ml graduated cylinders labelled 1E, 2E, etc. were placed below the Dowex-50 columns in the lower tier of the two-tier rack.

9. The following were added to each column.

4 x 2 ml 0.2N acetic acid

Note: added in four stages, 2 ml at a time once the previous 2 ml had run through the column.

- 10. Alumina columns (upper tier of two-tier rack) were then removed and placed over 25 ml graduated cylinders labelled 1D, 2D, etc., in a single tier rack (refer - DOMA, Step 16).
- 11. The Dowex-50 columns (the lower tier of the two-tier rack) were rinsed with 5 ml of distilled water/0.1 per cent Triton X-100. The offluent contained DOPEG(E).
- NMN

Once all the solutions had drained through the Dowex 50 columns in the middle tier of the three-tier rack, (refer Step 7), 5 ml of distilled water/0.1 per cent Triton X-100 was added and allowed to run through.

13. These Dover 50 columns were removed and placed over 25 ml graduated cylinders labelled 1C. 2C. etc. on a single-tier tack.
NHU was eluted with 10 ml of 6N NCL/ethanol solution and collected

in the gratuated cylinders(C).

- 15. Alumina columns from the upper tier of the two tier rack were positioned over 25 ml graduated cylinders labelled 1D, 2D, etc. (refer steps 10,11).
- 16. The following solutions were added to elute DOMA:10 ml 0.2 N hydrochloric acid followed by 5 ml 1 N hydrochloric acid once the former had drained through. The solution contained in the graduated cylinders (D) contained DOMA.
- 17. The Dowex-50 columns remaining in the lower tier of two-tier rack (refer step 12) were removed and positioned over 25 and graduated cylinders labelled 1F, 2F, etc.
- 18. NA was eluted from the Dowex-50 by adding 10 ml of 2N hydrochloric acid and collected in the graduated cylinders (F).

MOPEG/VMA

DOMA

NE

- 19. Once all the solutions had drained through the Dowex-1 columns in the lower tier of the three-tier rack (refer Step 7) the columns were rinsed with 5 ml of distilled water/0.1 per cent Triton X-100. The effluent was collected in the 50 ml graduated cylinders labelled 1A, 2A, etc. This contained MOFEG.
- Freenutions and aids during the procedurat
- The solution was applied to the columns only after the provision

The resins were not allowed to dry at any time during the entire procedure. This could be considered as the most important precautionary measure during the procedure. Thus, the menisci in all the columns were observed continuously. However, one need not be frantically worried about this as a 1-2 minute delay from the time the applied fluid reaches the compound to the addition of the next solution does not cause a detectable difference in the radioactivity.

, b)

۴N

- c) A 10 ml syringe with a thin silastic tube attached to its end, was kept handy. This can be helpful when an improper fluid has been applied as one can suck out the wrong fluid immediately, in hope that the assay has not already been ruined.
- d) If one applied the strong acid first in Step 17 and then the weak acid, no obvious problem with the counts occurred.
- e) A 50 ml cylinder connected to a large bore needle with the needle piercing a rubber stopper that fits tightly at the top rim of the glass columns was kept handy. This was used to apply slight pressure above a fluid contained within a glass column if the drainage was extremely slow.
 - To improve separation of the catechol compounds that are absorbed on to the alumina in step 6 (if the separation is not satisfactory) the following procedure can be adapted. Collect the effluent from the alumina columns (upper tier of three-tier rack) in step 6, following the addition of the superfusate samples, in 25 ml graduated cylinders. Thus, the effluent is not allowed to run into the Dowex-50 and Dowex-1 columns in the middle and lower ú.

tiers of the three tier racks (This part of the experiment can be done in a single-tier rack). The effluent is then re-applied to the alumina column and the effluent allowed to run into the Dowex-

Radioactivity measurements

The volume contained in each of the graduated cylinders was measured and recorded at the end of each stage. The contents of each cylinder was then mixed well and 1.0 ml pipetted out in duplicate and the radioactivity measured in a scintilation counter (refer Superfusion Experiments for details). Each sample was counted for 10 minutes or until 10,000 counts accumulated. Unlike in the Superfusion Experiments the efficiency varied from sample to sample because of different degrees of quenching by the different solutions used in the chromatographic The efficiency varied from 30-38%. Thus, disintergrations analysis. per minute (dpm) were utilised in all calculations involved. As each sample was counted in duplicate for radioactivity, the average dpm contained in 1.0 ml of the sample was first calculated. From this value the amount of radioactivity present in the 16.5 ml $(3 \times 5.5 \text{ ml})$ of superfusate present in each of the triads of test tubes used for the column chromatographic analysis was calculated (refer Superfusion Experiments for details).

e.g. Volume of superfusate present in triad-16.5 ml -

Volume of superfusate used for column chromatography procedure $(refer step 3) \sim 10.0 \text{ m}$

a) If the final volume collected in the graduated cylinder A (containing MOPEG) is 44.1 ml, 1.0 ml from this was counted in duplicate for radioactivity. If the mean radioactivity present in 1.0 ml was 91.1 dpm the amount of radioactivity present as

189
MOPEG in 16.5 ml of the superfusate = 91.1 x 44.1 x $\frac{16.5}{10.0}$ = 6629 dpm 190

1

b) If the total radioactivity present in 1.0 ml of the superfusate (refer to column chromatography procedure, step 2) is 1801.1. The total radioactivity present in 15.0 ml of the superfusate

> = 1801.1 x 16.5 = 29718

Therefore fraction present as MOPEG = 6629/29718 = 0.223 = 22.3 per cent

The forms used for the recording of results during column chromatography are shown in Table 1 and Table 2.

UATE OF CHROMATOGRAFIED HEREE	:			1982 1982
DATE OF SUPERFUSION EXPERIMENT TRIAD NO IN SUPERFUSION EXPERIMENT	•	5	000	
TRIAD CHARACTERISTICS	:			
COLUMN NUMBER		3	0 1	
VOLUME OF SAMPLE USED FOR CHROMATOGRAPHY	:	10.	()m i	

ţ٠

• •

• • • •

, ^{...}

	======================================	TOTAL VOLUME,m1	: VOLUME : COUNTED.ml	: VIAL : NUMBER
	: ORIGINAL SAMPLE (TOTAL RADIOACTIVITY)	·	1.0 : 1.0	: 1
A	OOWEX 1 EFFLUENT (MOPEG)	44.1 :	1 O : 1.0	: 3 : 4
9B	DOWEX 1 ELUATE	14 R	1.0 1.0	: 5
3C	DOWEX 50 ELUATE (NMN)	95	1 O 1 O	: 7 : 8
30	ALUMINA ELUATE (DOMA)	; 14.5	1 0 1 0	· 9 · 10
a E	DOWEX 50 EFFLUENT (DOPEG)	13.7	1 0	11
	DOWEX 50 ELIMATE (NA)	16.3	: 1 O	13 1

Table 1: The form used for the set of the during the form

chromatogrative star in or

, · ·

ľ91

192

DATE OF CHROMATOGRAPHIC ANALYSIS 13 Oct 1982 DATE OF SUPERFUSION EXPERIMENT 11 Oct 1982 TRIAD NO. IN SUPERFUSION EXPERIMENT 5 TRIAD CHARACTERISTICS COLUMN NUMBER. 3

. .

METABOLITE FRACTION	NO	CPM	EFFICIENCY %	DPM	MEAN DPM	TOTAL VOLUME ml	TOTAL DPM/ 16 5m1	%
TOTAL RADIO ACTIVITY	1 2			· · · -	1801.1	16 5	29718	 100
MOPEG	3	• • •			91 1	 1 L L	6629	22 3
VMA	 5 6	 			·	· · · · · ·	;-	• • • •
NMN	7 8							
DOMA	9 10	••••	*****		• • • • • • •	•••••••••		
DOPEG	10 11 12					· · · · · · ·		•••
NA	13	- · -		•••••	• • • • • • •			·

•

Table 2: The form used for seconding of results during a column

chromatography erroriment

4

۰.

¥.,

Relaxation Experiments: Protocol Three

The results of the Interaction Experiments described above indicated that background TNS inhibited the contraction caused by exogenous noradrenaline in the isolated canine saphenous vein. One possible explanation for this finding is a concurrent relaxatory response produced by the transmural nerve stimulation. The present investigation was an attempt to demonstrate the existence of such a relaxatory response in the canine saphenous vein in item following blockade of the contractive response to TNS

Protocol 3.1

Lateral saphenous veine were excised from ansatheticed logs as explained under General Methods. Excess connective tissue was removed and the specimums cut into rings approximately 4 mm in width. Entreme care was taken during the excision and preparation of rings, to avoid contact of surgical instruments with the luminal surface of the blood vessel or rubbing of the opposing luminal surface of the blood vessel or rubbing of the opposing luminal surface against each other, in order to preserve the endothelium. In other rings the endothelium was machanically removed by inserting the tip of a small fissue for one into the lumen of the ring and gently rol in the tip are imen back and forth over a filt i noner for 15 months if the line insertion of the insertion with the still of the triangular static before.

To elf inste the control fir off the of TNS is the conquestion of combination of generaliding only a synchronized second to the standed bits the foll streaments. An attempt was made to block the controction figure of the second second of 32 Phys. 8 notion 193

contractile effects of TNS adequately, a combination of the two drugs First, guanethidine $(10^{-4} \text{ mol}/1)$ was introduced into the was used. bath, after the rings were set up as described above, and was present in the bath throughout the experiment. After incubation with guanethidine for 90 minutes, phenoxybenzamine (2 \times 10⁻⁵ mol/1) was added to the bath and the incubation continued for a further 30 minutes. Then the unbound phenoxybencomine present in the bath was removed by repeated changes of hath fluid with fresh Krebs solution. During the last 30 minutes of this incubation both propracial (2 \times 10⁻⁶ mol/1) and atropine (5 \times 10⁻⁶ mol/1) were also present in the tissue bath. Thus, they were added to the bath together with the phenxybenzamine at the end of the first 90 minutes of incubation with guanethidine. Atropine and propranolol were added to the bath fluid to prevent any possible muscarinic or betareceptor mediated relaxation during the subsequent stages of the These two drugs (and the guanethidine) were present in the protocol. hath fluid throughout the rest of the evperiment. In order to ensure that the concentration of propranoiol used $(2 \times 10^{-6} \text{ mol}/1)$ was queficient to provent beta recentor modiated relaxation, the response to is prevaline was test d in some rings. Isoprevaline in a concentration of a v 10⁷⁵ molt did not produce any relevantion in saphenous vein rings pre contractor dith prostagiondin For as described below. The concentration of attractive used was similar to that used by other workers . that any prosthly muscarinic relevation in smooth muscle(212).

194

At the of this period of two hours, the preparations were contracted by adding Frobs buffer solution containing prostaglandin γ_{α} (10) model). The contraction produced was approximately 80 per cent of submodule to a strainment of in the methenous veta. Once this contraction reached a plateau, the response to TNS was determined using trains of stimuli of 30 second duration at each frequency. The stimulus parameters used were as follows: square wave pulses; duration 1.0 ms, strength 10 V, frequency 1,7.4.9.16, and 32 Hz. Each train of mulses was applied following complete resource from the response to the region to aim of million the response to the region to aim of million of million and from 8-30 minutes.

Protocol 3.2

In Protocol 3.1, TNS applied after pre-contraction of the saphener vein with prestaglandin F_{2g} produced a frequency dependent relevation Protocol 3.2 was done to elucidate further, the mechanism responsible for this relevatory response to TNS. Protocol 1 to as carried of first to determine the stimulus response our is to TNC, as a control. Then one of the following trage as added to the bigane back (after scale sign the bath field with first bottles control out to the protocol g (d).

41

1. . Tool Ha shapped tubihitor totand touts (1)

" ", recentor out gonist cleatiding (10 - mat/1)

1. - Cyclo wyre and Indibiliar indows forte (1) - mol-

4. Ty receptor anta infation from the (10 - and 1).

The first of the

The other is the second structure of the second struct

1/1 - F 1

bicarbonate" solution was prepared by replacing the KCl in the Krebs-bicarbonate buffer solution used, with an equimolar concentration of NaCl.

ⁿ² Only one drug was tested in each ring. Control rings without the above drugs were run in parallel to correct for changes in the responses with time. The stimulus-response curves to TNS before and after each drug were compared by regression analysis after log transformation (to base 2.0) of the frequency values.

1 + 1 ment 3.3

As tetradetowin used in Protocol 3 7 did not abolish the relaxatory response to TNS, the neural origin of this response was investigated using cold storage of the saphenous veins. Intramutal autonomic nerves are known to degenerate when exclued blood vessels are stored at 5 (213) Some exclued apphenous veins were stored in Krebs buffer solution at 1.0 for 9 dows on described by Guimarses et al(214). Rings were monited to the classe bath following this cold storage and the reap use to intermitted trains of TNS determined after precontraction. with prestaglands. For an described in Protocol 3.1. The stimulusicanouse cruces to THE I there cold stored veins were compared with the all due to come out a done on rings prepared from the same veing on the lay foundary. Although the contractile response to TNS was and shad in the of the of verne granethidine and phenoxyberramine more spected to a movel class sings to similare the conditions in the control for a postal series to a consider a provide and a post the etimitic recent of the

Phone and

1 1 en1

1917

· · · · · · ·

1.18

'nvervetion

hydroxydopamine on the relaxatory response to TNS was investigated here. The method used for the denervation was a modification (personal communication Vanhoutte PM, Lorenz RR) of the method used by Aprigliano The rings were set up for isometric tension and Hermsmeyer(215). recording as before. Aeration of the tissue bath was stopped and 20 ml of unbuffered physiological salt solution (for electrolyte composition refer Appendix III) containing 6.0 mg of dissolved 6 hydroxydoramine $(1.46 \times 10^{-3} \text{ mol}/1)$ was added to the tissue bath in place of Krebs huffer solution. Incubation with 6-hydroxydopamine was carried out for 10 minutes. The both solution was replaced with another 20 ml of physiological salt solution containing 6-hydroxydopamine and the incubation continued for a further 10 minutes. The 6-hydroxydopamine was then removed and replaced with Krebs buffer solution and aeration with 95 per cent Ω_0 5 per cent Ω_0 recommenced. Frequent ring with fresh Krebs solution was carried out over the next 3 hours until the tension (increased by the 6-hvdroxydopamine treatment) returned to the control level. The rings were then contracted with prostaglandin $F_{2\alpha}$ (in the presence of proprovolol and atropine as before) and the atimulus-response curve to INS repeated. In some of these hydroxydopamine treated veine, guanethidine and phonosybeuramine were also appl Ed to similate the conditions in Protocol 3.1. In Protocol 3.3 stops alting 6 hvlrovvderandne troatmont stro run to parillal as compared to the stand of the state of the presences of the treatest 1. . <u>.</u> 1 ringe

1 T - E

· · ·

. .

appeared to enhance the response to TNS in Protocol 1.2, it was decided to investigate this phenomenon further. The effect of low · concentrations of exogenous noradrenaline (which by itself did not produce any increase oin tension in the saphenous vein), on the contractile response to TNS was investigated here. Saphenous vein rings were prepared for experimentation as explained in the General Methods. After an equilibration period of 90 minutes TNS was applied as 5 second trains of stimuli at 8 Hz (strength: 10V, pulse duration:0.3 ms) every 5 minutes until the contractile response to the train of TNS became stable. This stable value was regarded as the control and the stimulus parameters were maintained constant for the rest of the experiment. Next, exogenous noradrenaline was added to the tissue bath in a concentration of 9.0 \log_{10} mol/1 and the response to the trains of TNS After three to four trains of TNS the exogenous determined. noradrenaline was removed by several changes of the bath fluid, while continuing the TNS every 5 minutes. Once the contractile response to TNS reached the control value the experiment was repeated with a higher · . concentration of exogenous nor adrenaline. The concentrations of exogenous noradrenaline utilized for the protocol were as follows (in log₁₀ mol/1): -9.0, -8.5, -8.0, 7.5, 7.2.

The maximum contractile response to TNS in the presence of each concentration of exogenous non-advanceline was expressed as a percentage of the control. The experimental values at each concentration of the known exogenous non-advance and the control values were analysed as a two way analysis of variance (the control and the responses to TNS of the five concentrations of variance (control and the responses to TNS of the five concentrations of variance (control and the responses to TNS of the five concentrations of variance (control and the responses to TNS of the five concentrations of variance (constituting 6 treatment ins). When the E value is significant (p<0.05) the differences

between any two treatment groups were assessed with the least significant difference test.

STATISTICAL ANALYSIS(216,217)

For each Protocol the number of experiments reported corresponds to the number of dogs used unless specified otherwise (see Protocol 1.1 and Protocol 1.2). All data are expressed as mean ± standard error of mean (SEM). For statistical comparison of two sets of observations, the Student's t-test (paired or unpaired as appropriate) was employed; a p value of less than 0.05 was considered as a statistically significant difference between two groups of data.

The control and experimental stimulus-response curves to TNS in Protocol Three were compared as follows: A regression analysis (after log transformation to base 2.0 of the frequency values) was carried out first on each group (experimental and control) of data. The two regression lines resulting from the above were compared thereafter. The slopes of the two lines were compared initially, followed by the elevation of the lines above the x-axis (i.e, homogeneity of the Xintercept) if the slopes were found not to differ significantly. The latter procedure was carried out by an analysis of covariance.

The control and the experimental values in Protocol Four were analysed with a two-ver analysic of variable, taking each day's experiment as a fugle replicate. The control response to DNS and the responses at the 5 different concentrations of coorenous a radrenaline were regarded on 6 treatment groups. When the ' value was significant the differences between cry two treatment groups at the treatment of core of other

-199

Ę,

RESULTS

Protcol One

. Saphenous vein strips mounted for isometric tension recording, in the tissue bath, did not demonstrate any spontaneous mechanical activity. They responded to exogenous noradrenaline in a concentration dependent manner. With repeated changing of bath fluid the tension returned to approximately the same level as before the commencement of the dose-response curve. This recovery from a dose-response curve took from 30-45 minutes. TNS applied as square-wave pulses produced a frequency dependent contraction with a maximum contraction at 16-32 The maximum contraction produced by TNS in the saphenous vein Hz. strips was about 90 per cent of the maximum contraction produced by Here too the tension returned to control exogenous noradrenaline. levels following cessation of stimulation. Although this occurred even without washing, several changes of bath fluid were done during this period which varied from 15-20 minutes. Further, the bath fluid was changed every 30 minutes even when no specific intervention was carried out. In 22 preparations (strips) the basal tension was 3.2 g (mean \pm 0.18 g SEM) and the maximum tension to exogenous noradrenaline was 4.9 g (mean + 0.30 g SEM).

Fr. 1 1.1

The effect of a TNS induced background contraction, on the contraction produced by exogenous noradrenaline was examined here. The experiment was carried out using different concentrations of exogenous noradrenaline and different magnitudes of background contraction produced by TNS as explained under Methods. The concentrations of traggenous noradrenaline used ranged from 7.5 x 10^{-7} mol/l to 6.0 x 10^{-6} .

The magnitude of the control contractions produced by this mol/l. exogenous noradrenaline alone in Step C and Step E of the protocol maximum contraction for the cent of ranged from 30-90 per noradrenaline. The background contractions produced by TNS at different frequencies ranged from 5-80 per cent of the maximum contraction for The additional exogenous noradrenaline in the saphenous vein. contraction produced by exogenous noradrenaline, when added against a background contraction induced by TNS in Step D was found to be much less than the controls, i.e., Step C and Step E. Often the total contraction produced by TNS and exogenous noradrenaline in Step D was only slightly greater than the contraction produced by exogenous noradrenaline alone. On some occasions they were equal. An example from Protocol 1.1 is shown in Figure 22.

As explained under Methods, a direct comparison between the additional contraction produced by exogenous noradrenaline against a contractions produced by exogenous TNS, background of and the noradrenaline alone would be erroneous. (Thus, an expected contraction was calculated for each concentration of exogenous noradtenaline used in Protocol 1.1. depending on the magnitude of the background contraction produced by TNS (see Methods). Following this correction, the observed contractions were compared with the expected contractions. The results are summarised in Fig. 23. Seventy-eight observations were made on 12 saphenous veins (number of dogs = 12). The observed contractions in Step D were significantly less than the expected contractions with the observed/expected ratio having a mean of 82.4 per cent (SEM, 2.1 %, p<0.001). The individual data points utilised for Figure 23 are shown in Figure 24, where the observed/expected contraction ratios were



Figure 22. An example from Protocol 1.1. The additional contraction produced by exogenous noradrenaline (NA) added against a background contraction by transmural nerve stimulation (TNS), is much less than the controls (on either side); in fact, the total contraction produced by exogenous noradrenaline and TNS in the middle is not much different from the contraction produced by exogenous noradrenaline alone in the two controls.

-



Figure 23: The summary of the results from Protocol 1.1. The observed/expected contraction produced by exogenous noradrenaline (NA) added against a background contraction produced by transmural nerve stimulation (Bgrd. TNS) is significantly less than the controls.

•



Figure 24. The individual data points for the observed/expected, contractions by exogenous noroadrenaline (NA) added against a background contraction produced by transmural nerve stimulation (TNS) in Protocol 1.1, plotted against the magnitude of the background TNS contraction. The latter is expressed as a percentage of the maximum contraction produced by exogenous proadrenaline in the capine saphenous vein.

plotted against the magnitude, of the background contraction ratios produced by TNS. The latter was expressed as a percentage of the maximum contraction for exogenous noradrenaline. As can be seen, 71 of 78 data points for observed contractions were less than the expected contractions. Further, the observed/expected ratio appeared to diminish with increasing magnitude of the background contraction produced by TNS, with a significant negative correlation (r = -0.42, p(0.001). Correlation coefficient although significant, was rather low with a coefficient of determination (r^2) of 0.17.

Protocol 1.2

The effect of a exogenous noradrenaline induced background contraction, on the contraction produced by TNS was examined here. The experiment was carried out using different frequencies of TNS and different magnitudes of background contraction produced by exogenous noradrenaline as explained under Methods. The control contractions produced by TNS in Step C and Step E of the protocol ranged from 20-80per cent of the maximum contraction for exogenous noradrenaline in the vein strips (The upper limit was constrained by the fact that maximum contraction to TNS in the saphenous vein was about 90 per cent of the mayimum contraction to exogenous noradrenaline). . 'he background contractions produced by exogenous noradrenaline in Stop D, ranged from 5-75 per cent of the maximum contraction to exogenous noradrenaline. The additional contraction produced by TNS applied against the background contraction induced by exogenous noradrenalists, although less than the controls, was more than that observed in Step D of Frotocol 1.1 (which was the inciprocal of Step D in Protocol 1.2). This additional contraction was occasionally observed to be equal to the control

contraction produced by TNS alone. A representative example from Protocol.1.2 is shown in Fig. 25.

As a direct comparison between the additional contractions produced by TNS in Step D with the two control contractions in Step C and Step E was not possible, expected contractions were calculated. The results are summarised in Fig. 26. Seventy one observations were made on 12 saphenous veins (number of dogs = 12). Unlike in Protocol 1.1 where the observed contractions in Step D were significantly less than the expected contractions, the observed contractions were significantly more than the expected contractions in Protocol 1.2 with the observed/expected ratio having a mean of 132.8 per cent (SEM 3.6 %, p<0.001). The individual data points utilised for Figure 26 are shown in Figure 27 where the observed/empected contraction ratios were plotted against the magnitude of the background contraction produced by exogenous noradrenaline, the latter being expressed as a percentage of the maximum contraction for exogenois noradrenaline. As can be seen, 64 (of 71) observed contractions were greater than the expected Further, the data points appeared to be randomly contractions. distributed, the observed/expected ratio showing no significant correlation with the magnitude of the background contraction produced by exogenous noradrenaline (r=+0.06, p>0.05),

Protocol 1.3

This protocol was carried out to determine whether the inhibitory effect of TNS on the exogenous noradrenaline contraction was due to the excitation of intramural nerves or due to the field of electrical current per se.





- - *******

Figure 25. An example from Protocol 1.2. The additional contration produced by transmural herve stimulation (TNS) applied against background contraction by exogenous noradrenalize (NA) although less than the controls (on either side) is relatively mire than that the in Protocol 1.1 (refet Fig. 22) where e optimus (A act offet to herbor and fug. 1.1. The reciprocal of

ħn



4

Figure 26. The summary of the results from Protocol J-2. The observed/expected contractions produced by transmural nerve stimulation (INS) applied against a background contraction by erogenous - '----'fun (Bgrd NA) is signification by erogenous



Figure 27. The individual data points for observed/expecte contractions produced by transmural nerve stimulation (INS) suplied against a larground contraction by inconcurs foredre align (1) de col J 2. The d'served expected of electron a Pro + 1+++ 112 . . . $\mathbf{r} \mathbf{b}$ agni 10.1 " + + mkrimma NA co per r 91 · . 1 17 • + -> • 1

Steps A to E of protocol 1.1 were carried out first. This showed results similar to that observed in Protocol 1.1. Thus, TNS was found to inhibit the contraction produced by exogenous noradrenaline. The contractile response to the frequency of TNS used in Step D was blocked by the addition of guanethidine into the tissue bath. The concentration of guanethidine necessary for the blockade varied from 1×10^{-6} mol/1 to 5×10^{-6} mol/l depending on the frequency of TNS used in Step D. A representative example of Step C, Step D and Step E following this blockade is shown in Fig. 28. The contraction produced by exogenous noradrenaline in Step D was similar to the two controls (Step C, Step E) in spite of the electrical current flowing between the platinum electrodes during Step D. The contraction in Step D was compared with the mean of the two controls. The results are summarised in Fig. 29. The mean contraction produced by exogeneous noradrenaline during the pascage of an electrical current was 20 B non-cout (mean + 0.5% SER) of " P portrole (n.8, FN0.05)

A similar experiment we contribute using the calcium antagonist dilitiant bydrechl if's in place of guamethidine. The concentration of dilitiant resulted to block the contractile offsets of TNS varied from 10 to 10 f mol/1 dipenting on the frequency of TNS used in Step D. At the secondarizations the crite offsets of the evogenous noradrenative uses limitated by all for 10 m r result. As with guamethidine, the contraction produced by compared normalise plane was not codified by the results of an electrical current through the solution during Step D. The mean contraction brocked by programs noradrenalize for Step D was per cent (many 1 - 3 per cent Colling fibe controls 1 - 3, pho-05).



Figure 28. A representative example from Protocol 1.3. The two control contractions to exogenous noradrenalice clong (on either side) and the contraction to the same concentration of noradrenaline added while applying TNS (middle) are shown. All three responses were elicited in the pression of guine bid he in the formation to block the contraction of guine bid he in the formation to block the contraction of guine bid he in the formation to block the contraction of guine bid he in the formation to block the contraction of guine bid he in the formation to block the contraction of guine bid he in the formation to block the

: ٢



Figure 29. Summary of results from Protocol 1.3. The contraction produced by exogenous noradrenaline (y-axis) in the controls is compared with the contraction produced by the same concentration of noradrenaline added against a background of transmural nerve stimulation (TNS) with the contractile effect of TNS blocked with guanethiding (n=8, p>0.05).

Protocol 1.4

These experiments were designed to determine whether the inhibitory effect of background TNS, on the exogenous noradrenaline mediated contraction, observed in Protocol 1.1 was specific for TNS or whether it would still be evident when another agonist is substituted in place of " TNS.

213

The response to exogenous noradrenaline added against a background contraction produced by TNS was compared with the response to the same concentration of exogenous noradrenaline added against a background contraction produced by tyramine (or methoxamine or histamine or phenylephrine). In this protocol, the additional response to exogenous noradrenaline was expressed as a percentage of the mean of the two controls i.e., Step C and Step E. The background contraction produced by TNS was matched (as close as possible) to that produced by tyramine to enable a direct comparison of the additional responses to be made. All four compounds, tyramine, methoxamine, histamine and phenylephrine produced a concentration dependent contraction in saphenous vein strips/rings. Dose-response curves to each of these drugs were performed in a preliminary set of experiments to get an idea of the concentration range to be used in the protocol proper-Tyramine, methoxamine and phenylephrine produced complete dose-response curves at a concentration range from 10^{-7} to 10^{-4} mol/1. In the case of tyramine, the dose-response curves were repeated in the presence of conside (10^{-5}) mol (1) to determine the concentration range in which its direct effects (on the smooth muscle) could be avoided. In the presence of consine, tyramine produced no appreciable contractions up to a concentration of 6 x 10 mol/1. In the protocol resper the concentration of tyramius used

was kept below this to avoid direct effects on smooth muscle. Saphenous veins were far less responsive to histamine compared with the adrenergic agonists described above. It often produced no contractions until a concentration of 10^{-4} mol/1 was reached.

οτφία στο το το το Αναφία, αταφία τα αταφία τα αταφία Αναφία τα παγία τα από τα α A representative example from the experiments using tyramine hydrochloride is shown in Fig. 30. The additional contraction produced by exogenous noradrenaline against a background contraction induced by tyramine, was much greater than that produced against a background contraction induced by TNS. This difference was apparent with different magnitudes of background contraction as well as different concentrations The pooled results are summarised in Fig. of exogenous noradrenaline. contraction produced by the exogenous 31 where the additional noradrenaline is expressed as a percentage of the controls. The additional contraction against a background of TNS was 67.1 per cent (mean \pm 2.5% SEM) of the control, while the additional contraction against a background of tyramine was 86.5 per cent (mean \pm 3.4% SEM) of the control (n=16, p<0.001).

In the experiments using phenylephrine hydrochloride as the background agonist, the drug was used in a concentration range from 1 x 10^{-6} mol/1 to 4 x 10^{-6} mol/1. The pooled results of these experiments are summarised in Fig. 32. The additional contraction produced by exogenous noradrenaline against a background of TNS was 56.2 per cent (mean ± 3.3%, SEM) of the control while the additional contraction against a background of phenylephrine was 68.3 per cent (mean ± 3.4% SEM) of the control. The additional contraction against a background of phenylephrine was significantly greater than that against a background of TNS (p<0.001, n-11).



▲ 2³5×10⁶ Nor

Enil

A

Figure 30. A representative example from Protocol 1.4 with tyramine hydrochloride (Tyr) as the background agonist. The response to exogenous noradrenaline (Nor) added against a background contraction by transmural nerve stimulation (TNS) (Upper middle), the response to the same concentration of noradrenaline added against a background contraction by tyramine (lower middle) and the four control contractions to exogenous noradrenaline alone are shown. The additional contraction produced by exogenous noradrenaline against a background contraction by tyramine is much more than that produced against a background contraction by TNS.



Figure 31. Results from Protocol 1.4 (background agonists:tyramine, histamine and methoxamine). The additional contraction produced by exogenous noradrenaline against a background contraction by transmural nerve stimulation (TNS) is compared with the additional contraction to the same concentration of noradrenaline against a background contraction by tyramine (left), histamine (middle) and methoxamine (right). The additional contraction to noradrenaline is expressed as a percentage of the control. Control refers to the contraction produced by exogenous noradrenaline alone, i.e., with no background agonist or TNS. Tyr: tyramine, Hist:histamine, Meth:methoxamine.

 \hat{r}^{i}

····· ,



Figure 32. Summary of results from Protocol 1.4 (background agonist: phenylephrine). The additional contraction produced by exogenous noradrenaline against a background contraction by transmural nerve stimulation (TNS) is compared with the additional contraction to the same concentration of noradrenaline against a background contraction by phenylephrine. The additional contraction to noradrenaline is expressed as a percentage of the control. Control refers to the contraction produced by noradrenaline alone, i.e., with no background agonist or TNS. PE: phenylephrine.

į.

In the experiments using methoxamine hydrochloride. as. the -background agonist, the drug was used in a concentration range from 1 x 10^{-6} mol/1 to 6 x 10^{-6} mol/1. The pooled results of these experiments are summarised in Fig. 31. The additional contraction produced by exogenous noradrenaline against a background of TNS was 60.8 per cent (mean \pm 5.4% SEM) of the control while the additional contraction against a background of methoxamine was 84.3 per cent (mean \pm 5.6% SEM) of the control. The additional contraction against a background methoxamine was significantly greater than that against a background of TNS (p <0.001, n=8).

In the experiments using histamine dihydrochloride as the background agonist, the drug was used in a concentration range from 5 x 10^{-5} mol/l to 4 x 10^{-4} mol/l. The pooled results of these experiments are summarised in Fig. 31. The additional contraction produced by exogenous noradrenaline against a background of TNS was 59.6 per cent (mean \pm 3.2% SEM) while the additional contraction against a background of histamine was 91.7 per cent (mean \pm 2.6% SEM). The additional contraction greater than that against a background of TNS (p<0.001, n=12).

Protocol 1.5

This protocol was designed to investigate the mechanism responsible for the inhibition observed in Protocol 1.1, utilising "antagonists" to some possible mediators of the inhibitory phenomenon.

The effect of propranolol, indomethacin, aminophylline and cimetidine on the inhibition of the exogenous noradrenaline mediated contraction by background TNS was investigated here. Thus, Step C, Step D and Step E of Protocol 1.1 were carried out first and then repeated in

the presence of one of the above mentioned drugs. The additional contractions produced by exogenous noradrenaline against a background of TNS with and without the drug were compared. Each was expressed as a percentage of the mean of the two controls, i.e, Step C and Step E.

In the experiments using the β blocker propranolol it was used in a concentration of 10^{-5} mol/1. This concentration of propranolol was sufficient to block the relaxant effects of the beta-agonist isoprenaline (following α -blockade) in the canine saphenous vein. This was established in preliminary experiments. At this concentration, the contractile response to exogenous noradrenaline was increased by up to 10 per cent in some preparations. The pooled results from the experiments using propranalol are summarised in Fig. 33. The additional contraction produced by exogenous noradrenaline was 66.3 per cent (mean \pm 4.6%, SEM) of the control in the absence of propranolol, and 58.4 per cent (mean \pm 3.0% SEM) in the presence of the drug. The latter value was significantly less than the former (0.05 > p>0.01, n=10).

In the experiments using indomethacin, the drug was used in a concentration of 10^{-5} mol/l. At this concentration it did not modify appreciably the magnitude of the contractions produced by exogenous noradrenaline or TNS. The pooled results of these experiments are summarised in Fig. 34. The additional contraction produced by exogenous noradrenaline was 67.8 per cent (mean $\pm 1.9\%$ SEM) of the control in the absence of indomethacin and 65.0 per cent (mean $\pm 2.2\%$ SFM) of the control in the significantly different from each other (p 0.05, p 11).

In the experiments using the P_1 -purinomeptor entagenest, aminophylline, the drug was used in a concentration of 10⁻⁵ mol⁽¹⁾. At

2.19



Figure 33. Results from Protocol 1.5 (propranolol). The additional contraction produced by exogenous noradrenaline against a background contraction by transmural nerve stimulation (TNS) with and without propranalol (10⁻⁵ mol/l) are shown. The additional contraction to noradrenaline is expressed as a percentage of the control. Control refers to the contraction produced by noradrenaline along. i.e., with no background TNS. Prop:Programolol

• •



221

. 1

Figure 34. Results from Protocol 1.5 (indomethacin and cimetidine). The additional contraction produced by exogenous noradrenaline against a background contraction by transmural nerve stimulation (TNS) with and without (a) indomethacin 10^{-5} mol/1 (left) and (b) cimetidine 10^{-5} mol (right) are shown. The additional contraction to cradrentline is expressed as a percentage of the control. Contraction to the contraction produced by poradrenaline alon.



222

Figure 35. Results from Protocol 1.5 (aminophylline). The additional contraction produced by exogenous noradrenaline against a hackground contraction by transmural nerve stimulation (TNS) with and without aminophylline (10^{-5} mol/1) are shown. The additional contraction to exogenous noradrenaline is expressed as a percentage of the control. Cont of refers to the contraction produced by contraction for a long. (a.

0

8

D

this concentration aminophylline did not significantly affect the contractile responses to exogenous noradrenaline or TNS. As with indomethacin, the additional contraction produced by exogenous noradrenaline against a background of TNS did not appear to be modified by the presence of aminophylline. The pooled results of these experiments are summarised in Fig. 35. The additional contraction produced by exogenous noradrenaline was 57.8 per sent (mean \pm 4.6% SF⁽¹⁾) of the control in the absence of eminophylline and 55 is per cent (mean \pm 3.1%, SEM) of the control in the presence of the drug. These two significantly different from each other (p^{ND, 05}, n.9).

In the parentments using elmetidine the drug was used in a con-entration of 10 5 mol/1. The presence of eimetidine did not cause 10 any approximate alteration in the magnitude of the minaetile recommento DNS or exogenous predienaline. The pooled results from these cimetidine experiments are summarised in Fig. 3'. The additional contraction produce he ever nois noradinaline was 65.2 per cont (near + 2.6% SFM) of the cont of the minaetile recommendation of 66 per cent (mean + 2.5 the of the minaetile recommendation of 66 per these two mations is a summarised in real

Frotacol 1wo

53

This process as a ried out to determine theorem (1) are scaped x_2 inhibition takes place during the copy instal conditions will be x_2 inhibitions takes place during the copy instal conditions will be x_2 inhibitions there are symplet for a labor of x_2 is the difficure of examples of the copy o

Protocol 2.1

Step D of Protocol 1.1 was carried out as a superfusion experiment in this part of the study. The radioactivity measurements in the superfusate have been expressed below as fractional release per 2 minute period of collection. All values have been multiplied by 10^3 for the sake of clarity as the fractional release is almost always a number with two to three decimal places. Thue, the fractional release values given helow have to be multiplied by 10^{-3} in order to obtain the true fractional release.

Iwo hours after the commencement of the superfusion, the experiment proper was bebun. At this time the basal efflux of tritiated compounds had reached a steady level. In fine coveriments the total radioactivity the superfugate at this time was In 1.58 (mean \pm 0.10 SEM). Unmetabolised ³H-moralise constituted only 4.3 per cent of this The deaminated compound DOPEG (3,4-dihydroxytotal radioactivity. phenylglycel) formed 34.3 per cent of this fraction and the orthomethylate' deaminated compound MOPEG (3-methoxy, 4-hydroxyphenylglycol) 20.2 per cent: DOMA (3,4-dihydroxymandelic acid) and VMA (vanilly) mandelie acid) constituted approximately 17 per cent each and NMN (normal another the) only 4 per cent. Thus, doaminated compounds (DOFEG, DOMA) formed approximately 51 per cent of the basal efflux, the pmothylated deputated compainds (MOPEG, VMA) approximately 38 per cent. and o m thilstod compounds (IMMI) approximately 4 per cent. When TNS was aprilat at > 117 surtained outraction resulted. This was accompanied indicactivity, ³H-moradrenaline and the for ener total 1 1 be total fractional release amounting to 4.97 (mean ± m tal litting sith. sumptainitable " normationaline constituted 26.2 per

cent of the total radioactivity (fractional release, mean 1.3 \pm 0.12 SEM) with MOPEG, VMA, NMN and DOMA constituting approximately 23.3, 7.7, 18.3 and 10.3 per cent respectively. Fig. 36 shows a representative example from Protocol 2.1 and the summary data for total radioactivity and the different metabolite fractions during the experiments (n=5). After 14 minutes of TNS, exogenous noradrenaline was introduced into the superfusing fluid in a concentration of 10^{-6} mol/l while maintaining the TNS at 2 Hz for 14 minutes. This resulted in a further increase of tension accompanied by a decrease in the efflux of total radioactivity ³H-noradrenaline. The fractional release of total as well as radioactivity was 2.60 (mean \pm 0.33 SEM) while that of ³H-noradrenaline was 0.36 (mean \pm 0.06 SEM) during this period. At the end of the 14 minutes the exogenous noradrenaline was removed from the superfusing fluid while maintaining the INS. This resulted in an increase in the fractional release of total radioactivity (mean 4.75 + 0.84 SEM) and 3Hnoradienaline (mean 1.11 t 0.11 SEM) to levels close to that observed prior to the introduction of exogenous poradrenaline. After another 14 minutes the TNS was terminated with a resultant decrease in the efflux of tritiated compounds to basal levels. At the end of this price the frantional release of theal radio tive i was 1 is (mean + 0 12, OPH) and that of ³H-coradiovaline was 0.06 (mean + 0.01 GPM) which comparable to the basal offline at the commencement of the experience

Thus, the introduction of exogenous noradronaline into the superineing fluid during TNS, resulted in a degreese in the efflue of total indicactivity or cell as ³Henored resulted induced by the TNS. The efflue produced by TNS is press in a spanna ar adductivity with the more description of spanna ar adductivity with the more description.


Figure 36. Summary of results from Protocol 2.1 The effect of exogenous noradrenaline added against a background of transmural nerve atimulation (TNS), on the total radioactivity of superfusate and efflux of ³H-noradrenaline and metabolites produced by TNS (bottom) (refer Table 5 for standard errors of the mean) An example from Protocol 2.1 is shown also. Changes in tension (top) and total radioactivity in the superfusate (middle) during the experiment are shown. All radioactivity measurements are expressed as fractional release x 10³. NA:unmetabolised noradrenaline, Deaminated:deaminated metabolites (DOMA and DOPEC), OMDA:ormethylated deaminated metabolites (MOPEG and VMA), "Maintermetanephrine. Total radioactivity in the superfusate.

after the introduction of exogenous noradrenaline (NA) using a paired ttest. The results are summarised below as fractional release (mean \pm SEM, n=5).

Total radioactivity

TNS only 4.85 ± 0.84 TNS + NA 2.60 ± 0.33, Ratio 0.558

p<0.001

³H-noradrenaline

TNS only 1.20 ± 0.11

p<0.01

TNS + NA 0.36 ± 0.06

Ratio 0.300

Evoked release was also calculated for each period of stimulation during the protocol (evoked release is the efflux during a period of intervention minus the basal efflux, i.e., the increase in efflux produced by the intervention itself). Comparisons made using evoked release values (instead of the total release as given above) too demonstrated the inhibition of the efflux of total radioactivity and ³Hnoradrenaline by exogenous noradrenaline.

The results of this protocol indicated that exogenous noradrenaline significantly inhibited the efflux of total radioactivity as well as ³H-noradrenaline produced by TNS. All the date from the column chromatographic analysis during the present protocol are summatized in Table 3.

DOPEG	0.54±0.02 (34.3)	0.56±0.10 (11.4)	0.48±0.06 (18.3)	0.56±0.11 (11.9)	0.37±0.04 (25.0)
VHOQ	0.26±0.01 0.54±0.02 (16.7) (34.3)	0.51±0.03 (10.3)	0.28±0.07 、(10.8)	0.43±0.03 (9.1)	0.18±0.01 (11.8)
NHN	0.06±0.01 (4.0)	0.91±0.11	0.32±0.03 (12.1)	0.73±0.09 (15.4)	0.07±0.07 (4.6)
VHA	0.28±0.03 (18.0)	0.38±0.02 (7.7)	0.38±0.03 (14.7)	0.54±0.05 (11.4)	0.41±0.05 (27.5)
MOPEG	0.32±0.02 (20.2)	1.16±0.05 (23.3)	0.66±0.01 (25.5)	1.20±0.05 (25.3)	0.34±0.01 (23.0)
W	0.07±0.01	1.30 1 0.12 (26.2)	0.36±0.06 ' (14.0)	1.11±0.11 (23.4)	0.06±0.01 (4.0)
TOTAL RADIOACTIVITY	1.58±0.10*	4.97±0.82	2.60±0.33	4.75±0.84	1.48±0.19
XYPER IMENTAL PER IOD	I CONTROL	SNL 11	VN +	.V TNS	V CONTROL
X	1	11	11	N .	>

Summary of results of column chromatographic analysis of samples from Protocol 2,1 TNS: Transmural nerve stimulation, 2 Hz; NA: Exogenous noradrenaline, 10⁻⁶ mol/1; Table 3:

٢

*Mean fractional release ± SEM; ** per cent of total radioactivity; ***Recovery.of

radioactivity for all samples during the analysis = 96.5 per cent (mean ± 0.64% SEM).

Protocol 2.2

Step C, Step D and Step E of Protocol 1.2 were carried out here as a superfusion experiment. The protocol proper was begun two hours after the commencement of the superfusion when the basal efflux of the tritiated material had come down to a steady, low level. The fractional release of total radioactivity at this time was 2.06 (mean \pm 0.20 The percentages of the different metabolite fractions contained SEM). in the total radioactivity were similar to the values obtained (at the commencement of the experiment proper) in Protocol 2.1; NA: 4.2 per cent (fractional release: mean 0.09 ± 0.01 SEM) MOPEG: 18.1 per cent VMA: 16.3 per cent, NMN: 3.5 per cent, DOMA: 18.2 per cent, DOPEG: 35.1 per cent. A representative example from the present protocol is shown in Fig. 37 together with the summary data for total radioactivity and the metabolite fractions during the experiments (n=5). After the control period, TNS was applied at 3 Hz. This produced an increase in the fractional release of total radioactivity (mean 5.49 \pm 0.80 SEM) as well as 3 H-noradrenaline (mean 1.44 ± 0.08, SEM). The latter constituted approximately 26.3 per cent of the total radioactivity during this TNS was terminated after 14 minutes and a control period period. followed, during which the efflux of tritiated compounds returned to ³H-(total radioactivity: mean 1.65 ± 0.08 SEM. basal levels noradrenaline 4.1 per cent of total). Exogenous noradrenaline in a concentration of 3.3 x 10^{-7} mol/1 was introduced into the superfusing solution after this control period, for 14 minutes. This resulted in an increase in tension but it did not change the total efflux or the relative proportions of the metabolite fractions to any appreciable The total fractional release during this period was 1.45 (mean extent.



Figure 37. Summary of results from Protocol 2.2. The effect of background exogenous noradrenaline on the total radioactivity of the superfusate and efflux of ³H-noradrenaline and metabolites (bottom) (refer Table 6 for standard errors of the mean). An example from Protocol 2.2 is shown also. Changes in tension (top) and total radioactivity in the superfusate (middle) during the experiment are shown. All radioactivity measurements are expressed as fractional release x 10^3 . NA:unmetabolised noradrenaline, Deaminated:deaminated metabolites (DOMA and DOPEG), OMDA:o-methylated deaminated metabolites (MOPEG and VMA), TNS:transmural nerve stimulation, NMN:normetanephrine, TOTAL:total radioactivity in the superfusate.

 \pm 0.07 SEM) and the ³H-noradrenaline release was 0.07 (mean \pm 0.01 SEM). At the end of 14 minutes TNS was applied (at the same frequency as used before) while continuing the superfusion with fluid containing This produced a further increase in tension exogenous noradrenaline. accompanied by an increase in the efflux of tritiated compounds. The total radioactivity present in the superfusate during this period was 3.81 (mean \pm 0.89, SEM) and the 3μ -noradrenaline content 0.89 (mean \pm 0.10 SEM: 23.4 per cent of total). The relative proportions of the different metabolites were similar to that observed during application of TNS in the absence of a background contraction produced by exogenous noradrenaline. At the end of this period (14 minutes) TNS was terminated and the superfusing solution changed back to one without exogenous noradrenaline. The efflux of tritiated material fell to basal levels during this period: total radioactive mean 1.39 ± 0.08 SEM 3 H-noradrenaline:mean 0.06 ± 0.01 SEM. TNS applied after this and control period produced an increase in tension accompanied by an enhanced efflux of tritiated material. The total efflux of radioactivity during this period was 4.86 (mean \pm 0.77 SEM) and the 3 Hnoradrenaline efflux 1.31 (mean ± 0.05 SEM, 26.9 per cent of total). TNS was terminated at the end of 14 minutes resulting in a fall of the efflux of tritiated material to basal levels.

³н--During this protocol, the total both efflux and the noradrenaline efflux produced by TNS in the presence of exogenous noradrenaline was less than that produced in its absence. The means of the efflux values during the two periods of TNS (in the absence of exogenous noradrenaline) were compared with the efflux produced by TNS in the presence of exogenous noradrenaline by a paired t-test. The results are summarised below as fractional release (mean + SEM, n=5).

Total radioactivity

TNS only 5.07 ± 0.79 NA + MS 3.81 ± 0.89 Ratio 0.73

³H-noradrenaline

TNS only 1. 5 0.06 NA + TNS 0.89 ± 0.10 Ratio 0.6

Comparison of the same using evoked fractional release i.e., efflux during an intervention minus the basal efflux) demonstrated similar results. All the data from the column chromatographic analysis are summarised in Table 4.

p<0.05

p<0.05

Therefore, in both Protocol 2.1 and Protocol 2.2 significant inhibition of the TNS induced efflux of tritiated compounds including unmetabolised ³H-noradrenaline occurred in the presence of exogenous noradrenaline in the superfusing solution. Evoked release of tritiated compounds during TNS in the presence of exogenous noradrenaline was expressed as a percentage of that in the absence of the exogenous drug (evoked fractional release was utilised as it reflects the efflux This was done for each individual induced by TNS more accurately). These percentages (which reflect both protocols. experiment in inversely the degree of inhibition produced by exogenous noradrenaline) from the two protocols were compared by a students t-test for unpaired data. The mean percentage for Protocol 2.1 was 32.9 per cent (SEM 2.7%) and for Protocol 2.2 was 57.6 per cent (SEM 9.9%). These two values were significantly different from each other (p<0.05).

***Recovery of radioactivity for all samples during the amalysis = 95.7 per cent S: Transmural nerve stimulation, 3 Hz; Na:Exogenous noradrenaline 3.3x10⁻⁷ aol/i: *Mean fractional release ± SEM; ** per cent of total radioactivity; rean ± U.58% SEM)

ummary of results of column chromatographic analysis of samples from Protocol 2.2 ble 4 :

, t

١ , ,

V ITY NA HOPEG VHA NHN DOMA DOPEG	* 0.09±0.01 0.37±0.03 0.33±0.06 0.07±0.01 0.38±0.02 0.72±0.06 (4.2) (18.1) (16.2) (3.5) (18.2) (35.1)	1.44±0.08 1.19±0.07 0.42±0.05 0.80±0.04 0.62±0.04 0.69±0.12 (26.3) (21.6) (7.6) (14.5) (11.3) (12.5)	0.07±0.02 0.29±0.02 0.35±0.06 0.08±0.02 0.29±0.02 0.49±0.04 (4.1) (17.4) (21.4) (4.9) (17.3) (29.7)	0.07±0.01 0.25±0.01 0.27±0.06 0.06±0.01 0.24±0.02 0.51±0.04 (4.61) (16.9) (18.8) (4.2) (16.5) (35.3)	1,89±0.10 0,92±0.05 0.35±0.05 0.49±0.04 0.45±0.05 0.59±0.10 23.4) (24.2) (9.2) (12.9) (11.8) (15.4)).06±0.01 0.27±0.02 0.30±0.06 0.05±0.01 0.21±0.02 0.45±0.05 4.1) (19.2) (21.3) (3.4) (14.9) (32.1)	3140.05 1.3240.16 0.3540.03 0.7440.04 0.5040.04 0.5040.07
 OTAL ADIOACTIVITY	06±0.20 [*] 0.	. 49±0.80		45±0.07 0.	rf 68 1 -0-88	.139±0.08	86±0. 77
IXPERIMENTAL PERIOD	: control	SNL I.	I CONTROL	¥¥ .	Vr.	CONTROL	SN1 -

]

.....

Protocol Three

A possible explanation for the inhibition of the exogenous noratirenaline contraction by TNS in Protocol 1.1 is a concurrent relaxatory response by TNS. The present protocol was an attempt to demonstrate the existence of such a relaxatory response in the canine saphenous veins, following blockade of the contractile response to TNS.

Saphenous vein rings responded to transmural nerve stimulation with frequency dependent contractions which were maximum at a frequency of When the stimulus-response curves were repeated (in 16-32 Hz. preliminary experiments) following incubation of the vein rings with tetrodotoxin (10^{-6} mol/l) for 30 minutes, the contractile response to TNS was almost completely abolished (Fig. 38). The maximum contraction at 32 Hz in the presence of tetrodotoxin was 3.4 per cent (mean \pm 1.7 per cent SEM, n=5) of the control value. This confirmed that the contractile response to whe TNS (applied as square-wave pulses 1.0 ms duration, 10 V strength) was due to the activation of intramural nerves. In the experiment proper, blockade of this contractile response TNS was achieved, by to a combination of guanethidine and phenoxybenzamine, Following the addition of guanethidine into the tissue bath an increase in tension was observed in the vein rings. The magnitude of this increase varied considerably from preparation to Some preparations demonstrated intermittent contractions preparation. superimposed on the increased tone. At the end of 90 minutes when phenoxybenzamine (2 x 10^{-5} mol/1) was added, the tension returned to the basal value and remained relatively stable thereafter. After another 30 minutes TNS was applied at a frequency of 32 Hz to ensure adequate sympathetic blockade. At this time the response to TNS was almost "ompletely aNolished in most usin rings.





Figure 38: The effect of tetrointoxin $(10^{-6} + 1/1)$ of the introduction to the maximumal in the initial set of the s

Protocol 3.1

The basal tension in the vein rings was 2.95 g (mean \pm 0.19 g SEN) at the commencement of the experiment proper (n=17). ... Following the blockade of the contractile response to TNS, the rings were made to contract by the addition of prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/1). The contraction reached a plateau after 10-15 mintues. The tension at this time was 17.53 g (mean ± 1.27 g SEM, n-17). TNS applied at this stage as intermittent trains of stimuli (30) s duration) at each frequency (1-32 Hz), elicited a frequency dependent relaxation in the rings (Fig. 39A). Relaxation commenced within a few seconds of the commencement of INS. Complete recovery from the relaxation (following cessation of TNS) occurred within 5-10 mintues. although in some instances, especially at higher frequencies, it took considerably longer (up to 30 minutes). Further, at the higher frequencies a second, delayed, relaxation was sometimes observed when the TNS was tarminated at the end of 30 Thus, the reaconse consisted of two chases: an initial aeconda. transient relaxation and a second, delayed, relaxation with slow recovery (Fig. 39B). The second phase (when theerved) was sometimes of greater magnitude then the initial phase. The magnitude of the relevation, in general, write consider bly from tissue to theme. The relexation was observed on 's guilt 's preparations in which the contractile response to INS was not completely shallshed by the guanethid (ne phenoxybenzemine is a treatment All the data from the frequency response curves or summarised in Fig. 40. The meximum refination observed was 3 3 o (means + 0.18 g SEM, n=17).

The oder to determine the out in relexation observed with modiated





Figure 39. Response of canine applendue vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10 5 mol/1), to transmural nerve stimulation (TNS) following sympathetic and muscarinic blockade using guanethidine (10 10 mol/1), pheno(vhentamine (2010 mol/1) propranolol (7910 5 mol/1) and atropine (5 × 10 6 mol/1). The applied as intermittent 30 second train of stimuli at each frequency. As demonstrating a frequency dependent relaxation to intermittent trains of grimuli. B: example of a biphasi response to TNS at 32 Hz demonstration as initial transfect plan for and a second delay d after relays

a true and a grant to a state of the second



Figure 40. The relationship of the magnitude of the relaxation to transmural nerve stimulation (TNS) to frequency of TNS, in canine saphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/l) following sympathetic and muscarinic blockade. TNS:trains of 30 second limit in the anch frequency, pulse duration: 1.0 ms, strength:10 V.

238

· · · ·

.

mechanically denuded of endothelium, (as described under Methods) with parallel controls. Confirmation of the preservation of the endothelium in the control rings and the removal of the endothelium by the mechanical denudation was obtained by the use of scanning electron microscopy. The rings were removed at the end of the experiments and carefully cut open to be fixed and processed for scanning electron microscopy of their luminal surfaces. Two electron-micrographs from control and de-endothelialised rings are shown in Fig. 41. As can be seen, the ring mechanically denuded of the endothelium showed no evidence of intact endothelium. On the other hand, the endothelium was preserved in 80-90 per cent of the luminal surface in control rings.

The optimum basal tension for the rings mechanically denuded of endothelium (mean; 2.60g \pm 0.16 g SEM, n=12) was not significantly different from the controls (mean; 2.75g \pm 0.18g SEM, n=12) (p>0.05). Further, the response of these rings to prostaglandin $F_{2\alpha}$ (maximum tension: mean; 17.90g \pm 1.33g SEM) was also not significantly different from the controls (maximum tension: mean 18.77 g \pm 1.45g SEM) (p>0.05, n=12). TNS applied to these rings elicited a frequency dependent relaxation. The data are summarised in Fig. 42. The maximum relaxation observed was 3.50 g (mean \pm 0.42g SEM, n=12) in the de-endothelialised rings and 3.40g (mean \pm 0.21g SEM, n=12) in the control rings. The stimulus-response curves compared by regression analysis with analysis of covariance (See under statistical Methods) did not show a significant difference (p>0.05).

When TNS was applied continuously, in a "cumulative" fashion at frequencies of 1 through 32 Hz (instead of intermittent trains of stimuli for 30 seconds at each frequency) the pattern of relevation was

Figure 41: A. Scanning electron micrograph of the intimal surface of a saphenous vein ring fixed and processed at the end of an experiment. Approximately 80 per cent of the intimal surface has endothelial cells. White bars at the bottom = $10 \ \mu\text{m}$. Magnification:1.42 x 10^3 . B. Scanning electron micrograph of the intimal surface of a vein ring mechanically denuded of the endothelium. White bars = $100 \ \mu\text{m}$. Magnification:1.43 x 10^2 . Note: absence of endothelial cells.

A

B







242

Figure 42. The effect of mechanical de-endothelialisation on the relaxatory response to transmural nerve stimulation in canine saphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/1) following sympathetic and muscarinic blockade. TNS:trains of 30 second duration at each frequency.

inconsistent, with a definite frequency dependent relaxation being observed only occasionally (Fig. 43). All rings showed a relaxatory response at 1 Hz. When the frequency of stimulation was increased, there was no further relaxation from 2-8 Hz in most rings. In some preparations the tension returned towards the baseline during these intermediate frequencies. At higher frequencies of stimulation (16-32 Hz) most preparations responded with further relaxations. Nevertheless, the maximum relaxation observed at 32 Hz with this mode of stimulation appeared to be less than that observed during a 30 second train of a stimuli at 32 Hz. Following continuous stimulation, some preparations showed a delayed relaxation, the recovery from which was prolonged, occasionality taking up to 30 minutes. It was also incomplete at times. Since the responses elicited by intermittent trains of stimuli (30 second) were more reproducible than those elicited by continuous stimulation, all drug effects were tested against intermittent trains of stimuli.

Protocol 3.2

The effect of different drugs on the relaxatory response to TNS was investigated here. A stimulus-response curve to TNS (applied as intermittent 30 trains) was carried out first and then repeated in the presence of the drug.

TNS applied in the presence of the fast Na⁺ channel inhibitor tetrodotoxin (10^{-6} mol/l) elicited a frequency dependent relaxation. The maximum relaxation in the presence of tetrodotoxin was 2.62 g (mean \pm 0.20 g SEM, n=6) and in its absence was 2.77 g (mean \pm 0.25 g SEM, n=6). The results are summarised in Fig. 44. The stimulus-response curves with and without tetrodotoxin were not significantly different



Figure 43. Response of canine saphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/l), to transmural nerve stimulation (TNS) following sympathetic and muscarinic blockade. Stimulus-response curves to TNS done "cumulatively" using continuous stimulation (in contrast to the protocol shown in Fig. 39). A. TNS at 0.5 Hz produced a relaxation which was reversed partially with an increase in the frequency of stimulation (i.e. 1 to 8 Hz). However at the higher frequencies (16 to 32 Hz) the relaxation was evident again. B. Example of a more definite frequency dependent relaxation with continuous TNS:such a response was seldom observed with this mode of stimulation.



Figure 44. Effect of tetrodotoxin on the relaxatory response to transmural nerve stimulation (TNS) in canine saphenous vein rings precontracted with prostaglandin $F_{2\alpha}$ following sympathetic and muscarinic blockade. TNS:trains of 30 second duration at each frequency.

from each other (p>0.05). As mentioned before, this concentration of tetrodotoxin almost completely abolished the contractile response to TNS (prior to application of guanethidine and phenoxybenzamine) in the saphenous vein.

Incubation with the H₂-receptor antagonist cimetidine in a concentration of 10^{-4} mol/1 did not antagonise the relaxatory response to TNS. The maximum relaxation in the presence of cimetidine was 2.88 g (mean \pm 0.53 g SEM, n=5) and the maximum relaxation in the control preparations was 2.92 g (mean \pm 0.47 g SEM, n=5). The pooled data from the stimulus-response curves are shown in Fig. 45. There was no significant difference between the two curves (\dot{p} >0.05, n=5).

Indomethacin in a concentration of 10^{-5} mol/1 failed to modify the relaxatory response to TNS. The maximum relaxation with and without indomethacin were 2.82 g (mean \pm 0.40 g SEM) and 3.23 g (mean \pm 0.38 g SEM) respectively (n=6). The pooled data shown in Fig. 45 demonstrated no significant difference between the two stimulus-response curves (p>0.05, n=6).

In the experiments with the P_2 -receptor antagonist aminophylline the drug was used in a concentration of 10^{-5} mol/1. At this concentration it had no effect on the contraction produced by the prostaglandin $F_{2\alpha}$. The maximum relaxation to TNS in the presence of aminophylline was 2.82 g (mean \pm 0.49 g SEM, n=6) and that in its absence was 3.08 g (mean \pm 0.39 g SEM, n=6). The pooled data are shown in Fig. 46. There was no significant difference between the stimulusresponse curves with and without aminophylline (p>0.05, n=6).

The introduction of ouabain into the tissue-bath fluid in a concentration of 2 x 10^{-4} mol/1 regulted in a gradual increase of the



Figure 45. Effect of cimetidine (left) and indomethacin (right) on the relaxatory response to transmural nerve stimulation (TNS) in canine saphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/1) following sympathetic and muscarinic blockade. TNS:trains of 30 second duration at each frequency. The drug concentrations are given in mol/1.





Figure 46. Effect of aminophylline (10^{-5} mol/l) on the relaxatory response to transmural nerve stimulation (TNS) in canine saphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10^{-5} mol/l) following sympathetic and muscarinic blockade. TNS:trains of 30 second duration at each frequency.

active tension produced by the prostaglandin $F_{2\alpha}$ in some rings. This increase varied from 0-15 per cent of the active tension prior to the introduction of the drug. The relaxatory response to TNS was almost abolished in the presence of ouabain (the effect appeared to be similar regardless of the presence or absence of increased active tension, induced by ouabain). The maximum relaxation in the presence of the drug was 0.55 g (mean \pm 0.09 g SEM, n=6) and that in its absence 2.73 g (mean \pm 0.24 g SEM, n=6). The decrease in the response was much more evident at the lower frequencies with practically to relaxation being observed up to (and including) a frequency of 8 Hz (Fig. 47). The two stimulus response curves were significantly different from each other (p<0.001, n=6).

In the experiments using "zero-K⁺ Krebs buffer" solution a marked decrease in active tension was observed when the normal Krebs buffer solution in the tiskue bath was replaced by the above solution, in onite of the continued presence of of prostaglandin $F_{2\alpha}$ in the bath. The active tension fell to 15-35 per cent of the control value (mean; 23.75 per cent) after 30 minutes of incubation in the zero K⁺ Krebs buffer. TNS applied at the end of this period failed to produce any significant relaxation. The maximum relaxation observed in zero K⁺ Trabs buffer was 0.26g (mean \pm 0.13g SEM, n=5) while that in the curvels was $^{-}.08g$ (mean \pm 0.3g SEM, n=5). The pooled stimulum response curves was zignificantly different from each other (Fig. 47, r $^{-0.1}$, n=5).

A possible role for free radicale in the observed relaxatory response to TNS was investigated by using accordinated (10^{-4} mol/l) and the enzyme catalase (50 ug/ml). Ascordinated is a non-life according to form the life spectformuly of the life spectformula of t

249

3,



.)

Figure 4". Effect of ourbain $(2 \times 10^{-4} \text{ mol/l})$ and zero-K⁺ "Krebs" buffer solution on the relaxatory response to transmural nerve stimulation (TNS) in canine suphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/l) following sympathetic and muscarinic blockade. TNS: peroxide radicals. The two drugs by themselves did not alter the contractile responses to prostaglandin $F_{2\alpha}$. TNS applied in the presence of these drugs produced a frequency dependent relaxation as before. The maximum relaxation observed in the presence of ascorbic acid was 2.53 g (mean \pm 0.19 g SEM, n=7) while that in its absence was 2.31 g (mean \pm 0.25 g SEM, n=7). The stimulus-response curves constructed from the pooled data were not significantly different from each other (Fig. 48, p>0.05, n=7). The maximum relaxation to TNS in the presence of catalase and in the controls were 2.98 g (mean \pm 0.43 g SEM) and 3.00 g (mean \pm 0.52 g SEM) respectively (n=6). The pooled data did not show any significant difference between the stimulus-response inves with and without catalase (Fig. 48, p>0.05, n=6).

Protocol 3.3

ja)

Q

The effect of storage of the isolated saphenous veins at 4°C for 9 days ("cold storage") in Krebs buffer solution, on the relaxatory response to TNS observed above was investigated in this protocol. The contractile response to the prostaglandin $F_{2\alpha}$ in these veins following cold storage was approximately 30-50 per cent of that observed in fresh rings taken from the name dogs. When isoprenaline (3 x 10⁻⁶ mol/1) was applied to these rings pre-contracted with prostaglandin $F_{2\alpha}$ (in the absence of the β-blocker, progranolol), the rings responded with relaxations; the tension produced by prostaglandin $F_{2\alpha}$ was 6.43 g (mean \pm 0.67 g SEM) while the magnitude of the relaxation to isoprenaline was 6.37 g (mean \pm 0.066 g SEM) (n=3). These results indicated that the rings of saphenous veins, even after cold storage for 9 days were capable of contraction and relaxation in response to TNS (applied to the relaxation in response to TNS the relaxation to isoprenous



Figure 48. Effect of ascorbic acid (left) and catalase (right) on the relaxatory response to transmural nerve stimulation (TNS) in canine' saphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/1) following sympathetic and muscarinic blockade., TNS: trains of 30 second duration at each frequency.

.

rings following pre-contraction with prostaglandind $F_{2\alpha}$) was almost abolished in these veins. Experiments done from rings taken from the same veins on the day of excision (i.e., prior to cold storage) served as controls. The maximum relaxation observed in the venous rings following cold storage was 0.46 g (mean \pm 0.23 g SEM) and that in the control rings was 3.12 g (mean \pm 0.62 g SEM) (n=5). The pooled data are summarised in Fig. 49. The pooled stimulus-response curve in the venous rings following cold storage was significantly different from that in the controls (p<0.01, n=5). In some (three preparations) of these rings phenoxy enzamine and guanethidine too were applied to simulate the experimental conditions in the "fresh" rings. These rings too failed to show any relaxation to transmural nerve stimulation.

Protocol 3.4

The effect of acute chemical sympathetic denervation using 6-hydroxydopamine, on the relaxatory response to TNS was investigated in this protocol. There was an immediate increase in tension when the 6-hydroxydopamine was applied. This reached a peak in 4-5 minutes and gradually declined thereafter. With the second application of 6-hydroxydopamine after 10 minutes, a rise in tension was observed again. The tension returned to the basal level over the next. 1-3 hours. The contractile response to TNS (at 32 Hz) was 10-20 per cent of the control value, 2 hours after the application of 6-hydroxydopamine, in most veins. At the end of 4 hours the contractile response was less than 5 per cent in almost all veins studied. Thus, the investigation of the TNS induced relaxation was carried out approximately 4 hours after the application of the 6-hydroxydopamine. Frequent rinsing of the tissue bath solution was carried out during this period. Rings (taken



Figure 49. Effect of cold-storage of the isolated canine saphenous veins at 4°C for 9 days on the relaxatory response to transmural merve stimulation (TNS) in the vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/1) following sympathetic and muscarinic blockade. TNS:trains of 30 second duration at each frequency.

.

from the same vein) without the application of 6-hydroxydopamine were used as controls. Guanethidine and phenoxybenzamine were used in these rings to block the contractile response to TNS. TNS applied following pre-contraction with prostaglandin $F_{2\alpha}$ elicited a frequency dependent relaxation in the 6-hydroxydopamine treated rings. The maximum relaxation observed was 3.2 g (mean \pm 0.41 g SEM) in these rings. The maximum relaxation observed in the control rings was 2.88 g (mean \pm 0.69g SEM, n=4). The pooled data are summarised in Fig. 50. The pooled stimulus-response curves with and without 6-hydroxydopamine treatment were not significantly different from each other (p>0.05, n=4).

Protocol Four

Five second grains of TNS applied at a frequency of 8 Hz (10 V, 0.3 ms) produced mansient contractions with complete recovery in 60-120 With repeated trains of TNS (every 5 minutes) the response seconds. became stable after 5-10 applications. The response at this time had a mean of 2.61 g \pm 0.40 g SEM (n=10). Low concentrations of noradrenaline added at this point produced a concentration dependent potentiation of the contractile response to the train of TNS (Fig. 51). This potentiation was evident within 30 seconds of the addition of the exogenous noradrenaline, and reached a maximum within 5-10 minutes. The potentiation of the response to TNS was still present even 30 minutes after the addition of exogenous noradrenaline. After removal of the noradrenaline from the tissue bath, the potentiation disappeared and the response to the train of TNS returned to the control value within 10-15 The highest concentration of exogenous noradrenaline utilised minutes. for this protocol was $-7.2 \log_{10} \text{ mol/l}$. At this concentration, noradrenaline produced a small contraction in some vein rings' (mean



256

Figure 50. The effect of chemical sympathetic denervation with 6-hydroxydopamine (6-OHDA) on the relaxatory response to transmural nerve stimulation (TNS) in canine saphenous vein rings pre-contracted with prostaglandin $F_{2\alpha}$ (10⁻⁵ mol/1) following sympathetic and muscarinic blockade. TNS:trains of 30 second duration at each frequency.



Figure 51. A representative example from Protocol Four. Trains of transmural nerve stimulation (TNS) were applied approximately every 5 minutes. Dots represent the points of addition of exogenous noradrenaline (NA) to the tissue bath cumulatively. A concentration dependent potentiation of the response to the train of TNS by exogenous noradrenaline NA is shown. W:rinsing of the tissue bath with tresh Krebs buffer solution.

0.34 g \pm 0.15 g SEM, n=10). The response to the train of TNS at this concentration of noradrenaline was 246.2 per cent (mean \pm 36.9 SEM, n=10) of the control. The responses to TNS at the different concentrations of background exogenous noradrenaline are summarised in Fig. 52. The results of the present protocol indicated a potentiation of the response to TNS by low background concentrations of exogenous noradrenaline. Identical results were obtained when the exogenous noradrenaline was added cumulatively and the response determined at each stage instead of discreet concentrations with rinses between as explained in the Methods.



Figure 52. Summary of results from Protocol Four. A concentration dependent potentiation of the response to the by exogenous noradrenaline (NA) is shown. TNS:transmural nerve stimulation. LSD:least significance difference test.

259

÷.

DISCUSSION

260

The results described in the preceeding chapter can be summarised as follows:

Interaction Experiments

Protocol 1.1: The observed contraction to exogenous noradrenaline when added against a background contraction produced by TNS was significantly less than the calculated expected contraction. Further the observed contraction/expected contraction ratio had a. significant negative correlation with the magnitude of the background contraction produced by TNS.

Protocol 1.2: The observed contraction to TNS against a background contraction produced by exogenous noradrenaline was, on the other hand, significantly more than the calculated expected contraction. The observed contraction/expected contraction ratio had no significant correlation with the magnitude of the background contraction produced by exogenous noradrenaline.

Protocol 1.3: The inhibition of the exogenous noradrenaline induced contraction by background TNS was not evident when this background TNS contraction was blocked by using either guanethidine or diltiazem while maintaining the electrical current.

Protocol 1.4: The additional contraction produced by exogenous nroadrenaline against a background contraction produced by either tyramine, methoxamine, histamine or phenylephrine was significantly more than the contraction produced against a background contraction of similar magnitude produced by TNS.

Protocol 1.5: The inhibition of the exogenous noradrenaline induced contraction by background TNS was not significantly decreased by the

presence of either propranolol, indomenacin, aminophylline or cimetidine in the tissue bath during the protocol.

Superfusion experiments

Protocol 2.1: In this superfused saphenous vein preparation, exogenous noradrenaline was applied against a background contraction produced by TNS (as in Protocol 1.1) to examine the release of 3 H-noradrenaline induced by TNS. It was found that the exogenous noradrenaline significantly depressed the amount of 3 H-noradrenaline as well as the total radioactivity (i.e., 3 H-noradrenaline and its metabolites) in the superfusate.

Protocol 2.2: In this protocol TNS was applied against a background contraction produced by exogenous noradrenaline (as in Protocol 1.2). The presence of a background of exogenous noradrenaline decreased the amount of 3 H-noradrenaline as well as the total radioactivity present in the superfusate during TNS, as compared with that present during the application of TNS by itself.

Relaxation experiments

Following blockade of the contractile response of Protocol 3.1: saphenous vein rings to TNS (using guanethidine and phenoxybenzamine), application of TNS on a pre-contracted vein ring elicited a frequencyconsistent dependent relaxation. The response was more and more evident with the frequency-dependency reproducible, and intermittent trains of stimuli (at different frequencies) as compared continuous "cumulative" stimulation. Mechánical dewith endothelialisation of the vein rings had no significant effect on the relaxatory response.

Protocol 3.2: Tetrodotoxin, cimetidine, indomethacin, aminophyline,
ascorbic acid and catalase had no significant effect on the relaxatory response to TNS observed in Protocol 3.1. However, the response was almost abolished by ouabain and zero-K⁺ Krebs buffer solution. Protocol 3.3: When Protocol 3.1 was carried out following storage of the excised saphenous veins at 4°C for 9 days in Krebs buffer the relaxatory response to TNS was no longer evident. Protocol 3.4: Chemical sympathectomy using 6-hydroxydopamine had no significant effect on the relaxatory response to TNS observed in Protocol 3.1.

Calculation of expected contraction in Protocol One

error.

In protocol 1.1 the effect of a background TNS induced contraction, the on contractile response exogenous noradrenaline to investigated. It was found that the observed contraction was 82.4 per cent (mean) of the expected contraction. Further, the observed contraction/expected contraction ratio showed a significant negative/ correlation with the magnitude of the background TNS contraction. The calculation of the expected contractions was an essential part of the analysis in this protocol for reasons outlined in the Methods section. However, both the inhibition of exogenous noradrenaline contraction by background TNS (as compared with the controls) and the significant negative correlation between the observed contraction/expected contraction ratio were present even when the analysis was carried out with the unmodified data. In fact, both these findings were much more unmodified data. Thus, calculation of the expected evident with the contractions did not qualitatively alter the results. Nevertheless, calculation of the expected contractions as outlined may not be without

First, the method does not take into consideration the changes in the response of the preparations with the passage of time during the experiment. This occurs as the dose-response curve carried out at the commencement of the experiment was utilised in the calculation of the expected contractions. Preliminary experiments (not reported in the thesis) demonstrated that the maximum contraction to exogenous noradrenaline decreased by less than 15 per cent over an 8-10 hour period during an experiment. However, if the maximum contraction of each dose-response is taken as a 100 per cent, the percentage change ar any one concentration of noradrenaline was less than 5 per cent except at the lowest part of the dose-response curve.

A second error associated with the calculation of the expected contractions may result from equating contractions TNS, to to concentrations of, exogenous noradrenaline. Although the contractile response to TNS is also dependent on noradrenaline (released at the nerve endings), the relative contribution of α_1 and α_2 receptors to the contractile response to endogenous noradrenaline (released by TNS) may be different to the relative contribution of α_1 and α_2 receptors in the case of exogenous noradrenaline (see below). Therefore equating contractions to TNS, to a concentration of exogenous noradrenaline may not be quite accurate. Although a quantitative estimate for the errors resulting from the two factors cannot be calculated, these errors are observed ' difference the produced thé in to unlikely have contraction/expected contraction ratios between Protocol 1.1 and Protocol 1.2.

The "inhibitory" effect of background TNS on the contraction produced by exgenous noradrenaline observed in Protocol 1.1 was

obtained using a wide range of concentrations of the exogenous drug and a wide range of frequencies of stimulation with TNS. Thus, the present study, unlike most other similar studies, (on the interaction between exogenous noradrenaline and TNS) provides data on this interaction spanning the whole range of the dose-respone curves rather than on a few selected points. Only the upper 10-15 per cent of the dose-response curves were excluded as it was necessary to avoid any limitation on the contraction produced during the experiment by the maximum contraction attainable with each preparation.

In Protocol 1.2 the "reciprocal of protocol 1.1" was carried out. Thus the response to TNS was measured against a background contraction produced by exogenous noradrenaline. The expected contractions were calculated to compensate for the different levels of active background tension produced by exogenous noradrenaline during the protocol (Fig. 14). This yielded an unexpected result in that the additional contraction produced by TNS against a, background of exogenous noradrenaline was found to be significantly more than the expected contractions. Further, there was no significant correlation between the observed contraction/expected contraction.

The results obtained in Protocol 1.1 where background TNS was found. to inhibit the contraction produced by exogenous noradrenaline could be explained on the basis of three hypotheses.

1. Pre-synaptic α_2 -inhibition(5,15) may account for the results: During Step D of Protocol 1.1 exogenous noradrenaline was added to the tissue bath against a background contraction induced by TNS. The response to the exogenous noradrenaline was measured as the

additional contraction produced at this point. This assumes that the contraction produced by TNS remained unaltered during this period. However, it is possible that the exogenous noradrenaline added may in fact depress the TNS contraction by acting on presynaptic aggreceptors. This would result from an inhibition of the release of endogenous noradrenaline at the adrenergic nerve endings leading to a decreased concentration of endogenous noradrenaline at the synaptic cleft and thus a decreased contractile response. Such a pre-synaptic effect would lead to an under-estimation of the contractile reeponse to exogenous noradrenaling during Step " and thus to the conclusion that hackground Inhilita 1111 1 110 contraction by exogenous neradranalina

The elevationeous release of a relayatory (vasodilator) substance by me may account for the results. If an inhibitory transmitter is record by DIC is addition to the endogenous noradrenaline (e) of traducing the star tile reapone to TNS), this could lead as alled for at the structure connecto the exogenous observative and the target to the Rel I to the Flague bath against If auch a coloratory to comfitter ware . 1 1 1.0 1. Inner 1 . . . 11 the strate 17 fan fiom two posthla 1 , 1) 1 1 g . . . 1.4 the from a appriate grandflater marve **P** • • • • • • 1 a g 1 the stress of constructs nerves to the 1 1 re molt , techning of transmissi nerve structure in with the theory of or of a fold of ourront the second reaction of any a to the second second from the . . .

the adrenergic nerves (as long as their thresholds for activation are comparable). On the other hand, the inhibitory transmitter may exist together with noradrenaline in the adrenergic nerve ending as a co-transmitter(219,220). Adenosine triphosphate, an inhibitory trrusmitter in some non-edrenergic, non-cholipergic ('purinergic') nerves. is believed to be present together with noradrenaline in adrenergic nerve endings(210). There is suggestive evidence for the evidence for other transmitters such as sometostatin and enlaphalin as an induced in adrenergic neurones(218).

١

An interaction between post-synaptic of and post synaptic of 7 receptor effects more account for the results. If, as postulated in the Introduction, endogenous noradronaline released by TNS acts relatively more on the functional receptors and the exogenous noradrenalize relativity more in the extra junctional ra receptors. Interaction between the effects of in and appreciation stimulation could take place. Place a, a discretions have our here finentigated to do the annumber extent 160 - 5 fra house in his review, angreate that a tribunch that is a star recept to oth typps and inter or " " " " " Not to at 1 . . . ********** · + 11.0 and the second mm + + 1 10 1 1 1 1 1 1 1 1 1 1 4 m re ten es mennesset of the free free states and the second states for the second second to thene 1 .. The Co יידרי גידער יידרי 1 1.11

The presence of both α_1 and α_2 post-synaptic receptors have been demonstrated in the canine saphenous vein (used in the present study) by De Mey and Vanhoutte(12,223). In this study, yohimbine (an α -receptor antagonist with a greater affinity for the a_2 sub-type) was found to cause a parallel shift to the right of the dose-response curves to noradrenaline in isolated saphenous vein rings (also in splenic artery, femoral artery and femoral vein) although the slope of the Schild plot was found to be significantly less than unity (0.69). Prazosin, a relatively selective α_1 -antagonist was also found to cause a shift of the dose-respone curves to noradrenaline to the right although it was not a true parallel shift as the maximum was depressed at some concentrations. The presence of both α_1 and α_2 receptors was demonstrated by using relatively selective α_1 -agonists (phenylephrine, methoxamine) as well as α_2 -agonists (clonidine, tramazoline) in this st…dv. The authors concluded that exogenous poradrenaline acted on both To and C, post-synaptic receptors in producing contraction of the saphenous with rings with the lower concentrations of the amines acting mainly on the ap recentors. Unfortunately, the effects of selective antagonists with TMS indical succession to the complementary was not names at in this sight

In more ψ_n is the presence of both α_1 and α_2 poet exception perceptors and the action of exercisic northrenalfs on both types of inceptors are been demonstrated in the canfue sephenous mein, an interaction atoms the transfer of registers remeter a powerble Influence of sympathetic nerve activity upon the sensitivity to exogenous noradrenaline

Review of the literature relating to TNS/exogenous noradrenaline interaction revealed relatively few studies. For instance, subsensitivity to exogenous noradrenaline associated with electrical stimulation was reported by Rapoport and Bevan in the isolated ear arteries of the rabbits(224).

Arterial rings taken from rabbits sacrificed by stunning followed by rapid exsanguination (a procedure accompanied by a large increase in sympathetic tone) were found to be less sensitive to noradrenaling than arteries taken from rabbits sacrificed by a lethal dose of pentobarbital (a procedure accompanied by decreased sympathetic activity) in this study. This subsensitivity lasted for at least 6 hours. Further, the response to noradrenaline $(4.6 \times 10^{-8} \text{ mol/l})$ was tested following eight 2 minute periods of TNS over a period of 2.5 hours (pulse duration 0.3 ms. frequency 16 Hz, supramaximal voltage) and compared to the response following periodic applications of histamine (2 x 10 6 mol/1). The response to noradrenaline following prior stimulation with TNS was significantly less than that following prior treatment with biotosine Nowever, there were no significant difference between the former of the response to controle which were hopt or allenged during the 2.2 hour period of pertodic TNS application. It outputs on high that sympathetic activity is the top adaptivity to the action In roaged " oth miscle to noradiens? ne

A similar de roos in err netver en 's sub-maximal secter et tons f ear arteries (225,226). A voltage of 15-30 V across the electrodes in the presence of tetrodotoxin to block action potentials was utilised to effect direct stimulation of the muscle. There was no difference in response to KCl (26.4 mmol/1) or the maximum contractile responses to noradrenaline, histamine or 5-hydroxytryptamine. Direct electrical stimulation failed to alter the response of <u>rabbit saphenous veins</u> to noradrenaline or histamine in this same study. Bevan and Bapoport concluded that the subsensitivity observed in the ear arteries⁷ was likely to be due to a change in the smooth muscle cells, beyond the receptors, as the decreased sensitivity was relatively non-specific, affecting noradrenaline, histamine and 5-hydroxytryptamine.

Unlike in Protocol 1.1 and Protocol 1.2 where the effects of a combination of TNS and exogenous nroadrenaline were investigated, the studies discussed above dealt with the effects of prior effectrical stimulation (transmural and direct) on the respone to a subsequent application of exogenous noradrenaline. However, they are mentioned here as they represent the only studies in the recent ligerature which . have shown a subsensitivity to noradrenaline produced by electrical stimulation in vascular smooth muscle. On the other hand, a supersensitivty to exogenous noradrenaline induced by continuous nerve stimulation. following pre-treatment with reserpine in the nictitating membrane of the spinal cat was shown by Trendelenberg(227). Following pre-treatment with reservine, 3 mg/kg intraperitoneally (24% hours prior) to the experiment), continuous pre-ganglionic stimulation was found to whift the dust response curve to (1) noradrenaline by a factor of 4.2. illus above to constrivity disappeared rapidly following the termination

269

.

of the nerve stimulation. Thus, this short-term pre-treatment with reserpine did not alter the sensitivity of the nicitiating membrane to noradrenaline in the unstimulated state. The findings in this study were explained on the basis of noradrenaline taken up by uptake₁ into the adrenergic nerve endings being immediately released during nerve stimulation (but not in the unstimulated state) leading to the supersensitivity.

In summary, a survey of the <u>literature</u> did not offer an adequate . and satisfactory explanation for the findings in Protocol 1.1 which demonstrated an inhibition of the response to exogenous noradrenaline by background TNS.

Influence of exogenous noradrenaline upon the sensitivity to transmural nerve stimulation

Protocol 1.2 where background exogenous The findings in noradrenaline appeared to facilitate the response to TNS are even more puzźling. Hope, Law et al. (228) investigated the effects of noradrenaline and adrenaline on the vasoconstrictor response to TNS and on the efflux of ³H-noradrenaline produced by TNS in the isolated rabit ear artery(228). TNS was applied as a train of invelses at 5 Hz for 30 seconds (pulse duration 1.0 mg, supramagimal voltage) Exogenous noradrenaline was used in three concentrations: 0.05, 0.5 and 5.0 Superfusion, with 0.5 and 5.0 μ mol/1 concentrations of 1. mol/1. noradrenaline resulted in a marked decrease in the additional contractile response to the train Superfueton of TNS. noradrenaline at 0.05 pmo1/1 decreased the response 2 ITT approximately 8" per cour of the control value This was account hv in the offlux of derreagen the 1.91.01 the Erfar St. a. 1

noradrenaline and its metabolites: no separation carried out). Similar results were obtained with adrenaline. The authors concluded that the vasoconstrictor response to TNS was decreased by exogenous noradrenaling and adrenaline as a result of pre-junctional inhibition and by other mechanisms desensitisation of the post-junctional such the **as** The evidence for the latter was the fact that the responses receptors. to TNS were reduced even after cessation of goradrenalize administration at a time when the transmitter release had returned to the control value or even exceeded it. As no allowance was made for the increased background active tension produced by the exogenous noradrenaline the interpretation of the decreased response to TNS during exogenous noradrenaline administration is made difficult. However. the subsensitivity to TNS following cessation of administration of noradrenaline when the tension had returned to basal levels appears more conclusive.

A study carried out by Su(229) however, demonstrated different regults. The effects of low concentrations of noradrenaline. adrenaline, phenylephrine, methoxamine, naphazoline and oxymetazoline, on the vasoconstrictor response to a 40 second train of TNS (8 Hz, 0.3 ms duration) were investigated in a variety of rabbit blood vessels in , this study. The α -agonists mentioned above were used in concentrations which did not cause any contractile effects on their own. The response to TNS in mesenteric arterial rings were markedly potentiated by subthreshold concentrations of the above drugs. This effect was well maintained for at least 30 minutes and was concentration dependent. A large part of this potentiating effect developed within 5 seconds. The some concentrations of poradrenaline which potentiated the contractile

271

response to TNS, decreased the stimulation-induced efflux of ³н́– noradrenaline in this preparation. This potentiating effect was not by uptake1 inhibition (cocaine), puptake7 inhibition prevented (metanephrine) or beta-blockade (propranolol). The sabbit ear artery, the saphenous artery and the pulmonary artery did not show this potentiation of TNS induced \sim contraction by α -agonists. In fact, inhibition was produced by some agonists. The rabbit brachial artery demonstrated a slight potentiation. The authors concluded that a postsynaptic mechanism is probably involved in this potentiation (in the mesenteric artery) as the pre-synaptic actions of these agonists lead to no effect/decrease in the release of ³H-noradrenaline in this A similar potentiation of the response of the smooth preparation. muscle to TNS was shown in the guinea-pig vas deferens by Sjostrand and Swedin(230). A number of agonists including noradrenaline, adrenaline, acetylcholine, bradykinin, histamine and substance P in concentrations that had no effects on thier own, enhanced the responses to TNS by up to 100 per cent. These drugs also enhanced the responses to direct electrical stimulation of muscle following prior (6.12 days before) surgical deveryation or ofter blockade if action potentiale to the intramural with tetredotoxis. NPT 98 This indicated that the enhancement was likely to be a result of a pool avantic effect of the agonists. These studies also combastas the relation sector oversimplification that a use syn atter inhibition. catecholamines would invariably homper the transitest a process whole. A prientiation similar to that whom is a ~h 1 could account for the cohanced regionar to orogenein manteresters there is the

In Protocol Four of the present study the effect of low concentrations (no contractile effects on their own) of exogenous noradrenaline on the responsento a train of TNS was investigated. This demonstrated a potentiation of the response to TNS similar to that observed in the above mentioned studies. The maximum response observed amounted to, 246.2 per cent of the control value. This phenomenon, if present under the conditions employed in Protocol 1.2, could account for the higher observed contractions to TNS (as compared with the expected contractions) when applied against a background contraction produced by exogenous noradrenaline. There is no reason to believe that the postulated post-synaptic potentiating mechanism would not function when the concentration of noradrenaline used produced a contraction on its own (i.e. above threshold). However, an increase in tension produced by the background noradrenaline would tend to mask the potentiation because of the added confounding factor of a decreased response to a fixed stimulus as one proceeds up the dose-response curve.

The findings in Protocol Four demonstrated also, that pre-synaptic γ_2 -inhibition of TNS by exogenous noradrenaline is unlikely to be of any biological significance during conditions similar to those employed in this protocol in the canino saphenous vein. The mechanism responsible for the potentiation of the response to TNS is not clear at present.

influence of electrical current on the response to exogenous noradrenaline

In Frotocol 113, it was found that inhibition of the exogenous noradrenaline contraction by TNS (observed in Protocol 1.1) was not evident, when the response to exogenous noradtenaline was measured following blockade of the background TNS induced contraction with responsibilities, while maintaining the electrical current. Guanethidine,

an adrenergic neurone blocking agent, depletes the stores of noradrenaline from adrenergic nerve endings and also prevents the release of the transmitter. However, it has no effect on the passage of the pulses of current between the two electrodes in the tissue when 'TNS' is applied. Thus, in this protocol, what is measured is the effect of background pulses of electrical current (strength 10 V, duration 1.0 ms, different frequencies) on the response to exogenous noradrenaline. No significant effect was demonstrated. As stated before, guanethidine was used in this protocol at the lowest concentration that blocked the effect of TNS at the frequency used in the first part of the protocol, i.e., where the effect of exogenous noradrenaline was measured against a background contraction by TNS. This protocol' demonstrated that the inhibition of the exogenous noradrenaline contraction was not due to the field of current per se, acting directly on the vascular smooth muscle or by causing oxidation of the exogenous peredrenaline in the tissue bath. This finding was confirmed in the experiments using the calcium channel antagonist dilitizion to place of guenethidine to block the contractile effects of An interacting observation during the latter experimente was TNS. that diltty zem in a concentration of 10 5 to 10 6 mol/1 was found to block the constructive of TNS (at 1-4 Hz) completely while leaving the magnitude of the contraction produced by exogenous noradrenaline about 70-80 per cent of its control value. nt Thie Ca21 difforential offact of channel autagenists on the contraction The concered with that produced by exogenous produced by nor transtore to the. and r eap nous vers was also shown to Canherite

the in the present study

. .

ŵ

demonstrated with both diltiazem and verapamil in the above study.

Ð

In Protocol 1.4 it was demonstrated that the additional contraction produced by exogenous noradrenaline against a background of TNS was significantly less than that produced against a background contraction of similar magnitude produced by tyramine, phenylephrine, methoxamine or histamine. Thus, the inhibition of the exogenous noradrenaline contraction produced by background TNS (Protocol 1.1) was not present with pround contractions produced by the above named agonists. In fact potentiation of the exogenous noradrenaline contraction was present in the case of tyramine, methoxamine and histamine where the mean magnitude of the contractions in the presence of background agonist contractions were over 80 per cent of the control contractions by noradrenaline alone with some values even exceeding the controls.

Contraction of smooth muscle by tyramine is mediated by two pharmacological actions: (1) an indirect sympathomimetic effect which leads to the release of endogenous noradrenaline from the adrenergic nerve endings present in the tissue (2) a direct effect on smooth muscle leading to contraction. The indirect sympathomimetic effect is usually seen at lower concentrations of tyramine as compared with the latter effect. In exerting this indirect effect, tyramine is first taken up into the adrenergic nerve endings by uptake1. This tyramine then leads to the displacement of endogenous noradrenaline within the synaptic regicles. Some of the noradrenaline displaced into the neuroplasm comes out of the nerve endings and produces the contraction of the smooth miscle while the balance is metabolised by intraneuronal enzymes. [n the present study, the contractile effects of the concentrations of tyramine used ((6 x 10-5 mol/1) were complexely blocked by the uptake,

inhibitor cocaine (this was established in preliminary experiments - see Results). Thus, the effects would have been produced predominantly by its indirect effects via the release of endogenous noradrenaline. Nevertheless, the inhibitory effect on the exogenous noradrenaline produced by TNS (which also releases endogenous noradrenaline) was not observed with tyramine. This could be explained in three ways.

Firstly, although both tyramine and TNS lead to the release of endogenous noradrenaline, the two mechanisms of release are not identical. Tyramine, as explained above, leads to the displacement of endogenous noradrenaline from the storage vesicles and its subsequent "leakage" into the synaptic cleft. TNS, on the other hand, leads to the release of endogenous noradrenaline via exocytosis of the storage Here noradrenaline is released into the synaptic cleft vesicles. together with other vesicular contents such as dopamine- β -hydroxylase and ATP. If the inhibitory effect of background TNS, on the exogenous noradrenaline contraction observed in Protocol 1.1 was due to the simultaneous release of an inhibitory co-transmitter, this inhibition observed with tyramine which roleases on/logenous would not be Secondly, if the inhibitivy offect noradrenaline by displacement. observed in Protocol 1.1 was due to the release of an inhibitory transmitter released from a new edienergic nerve (present in the scale) together with adremergic nerves) is INS, this release would not wall take place in the case of tyrumine whose actions on the advance gin using ending depends on a specific intake process. Thus, it is no likely to act on a new advence give nerve. Thirdly, the results shell be urplated come titles between ex genoue actualiant and transfor and a the contraction to added agoing

tyramine(148,232). As tyramine has a higher affinity for uptake₁ than noradrenaline it would act as a competitive inhibitor of uptake₁ for noradrenaline. This would lead to a higher blophase concentration of noradrenaline and therefore a potentiation of its effects. Although this third possibility is known to occur and is likely to play some part in the results observed with tyramine, the other two possibilities cannot be excluded on the findings presented to this stage..

 α_1/α_2 receptor interaction

An interaction between the effects of α_1 receptors and α_2 receptors was considered as a probable explanation for the inhibition of exogenous noradrenaline contraction by background TNS. This postulated interaction should still occur in the tyramine experiments where the endogenous noradrenaline released at the adrenergic nerve endings by tyramine would act relatively more on the intrasynaptic (probably α_1) receptors just as in the case with TNS. However, the inhibitory effect nor evident under these circumstances. was Thus, the tyramine experiments do not support α_1/α_2 receptor interaction as a probable explanation for the findings in Protocol 1.1. Further evidence in favour of the above conclusion is provided by the experiments using phonylephrine and methoxamine. Both these drugs have been considered as relatively specific α_1 advenoceptor agonists(16,233) with phenylephrine possesing a minor indirect sympathomimetic action in addition. In these experiments the effect of a padrenoceptor stimulation on the contraction of exogenous noradronaline was tested. No inhibition of the contraction by evegenous peradronaline was evident under these circumstances. Thus is consistion between α_1 and α_4 remains stimulation appears to be an

ค

unlikely explantation for the inhibitory effect on erogenous noradrenaline contraction by INS, in Protocol 1.1.

The experiments using methoxamine help shed further light on the potentiation observed with tyramine. Methoxamine is an extremely poor substrate for the untake, process(153). Thus, unlike tyramine, it would not compete with the 'exogenous noradrenaline for the uptake. Nevertheless, the inbihition of exogenous noradrenaline contraction was not observed against a background contraction produced by methoxamine where the results were very similar to those observed with background tyramine. Thus, competition for optake, between tyramine and exogenous noradrenaline may not be the sole avaianation of the lack of inbihition of the exogenous noradrenaline may not be the sole avaianation of the lack of inbihition produced by tramine.

The encodmental finitors up to this point out two possibilities as explanations for the findings in trained 1.1. The star avaptic receptor tabiliton 1 the area of a work earling remains a ponallitity, "one of the orperimontal's to see 1. (with tyran ne. phonelophetic, mothing in and his gut a source the tree needs industion synchreaters. To first the an only using transfer any a many the constraint of a set of a set of the set of the TRAM A STAR OF THE THE the straight straight 110 terrend to solution of the tractic generic Although a release of neralional sector sectories is a birth to possible of this high one control, the island to the substance is selen in general ingeneral conditionally a base have to indified the last of the con-(in the reaction of the that the second of the second second second Sorth 1 2 K+

278

 $\sqrt{2}$

exogenous noradrenaline did not produce any inhibition of the release of 3 H-noradrenaline produced by tyramine in the above study. In Protocol 1.4 the inhibition of exogenous noradrenaline contraction by background TNS was not observed with a background of tyramine. This may have been due to the missence of pre-synaptic α_{2} -inhibition under the latter figure tances.

The second possibile explanation for the findings in Protocol 1.1 is simultaneous release of an inhibitory transmitter. This may appear a contransmitter or from a different intramural nerve-The erroriments with gnanethidine which blocks the release from adrenergic norme endings are more in favour of a contransmitter than a separate herve. This is because no inhibition was observed under these circumstances. As guarethidine is not likely to prevent release of transmitter from a non-adrenergic nerve, the inhibition should have been still evident if the inhibitory transmitter was released from such a nerve. The experiments using diltiazem appear to support the preavoiable hypothesis rather than the inhibitory transmitter hypothesis. As dilligrow blocks the contractile effects of TNS and presumably not the rolonge of a adremative, the inhibitory effects of a released nac d typemitter bould be a been evilent under these circumstances: but no ' th'thop of the exogenous noradrenaline contraction was observed. To ever it a known that Ca²¹ entagoniets may block the throad flow of "n" long at the narve endings which trigger the release of the course namillers (17) . (Though this could cours at higher concentrations then that such recessary to block the smooth muscle channel, and a settern at the neuronal membrane membrare rolean the transmitter campet pr vontine

Investigation of the possible mediators of the inhibition observed in Protocol 1.1

An attempt was made to identify the postulated relaxatory substance in Protocol 1.5 by using pharmacological antagonists against possible relexatory mediators. Firstly, a beta-receptor mediated relaxation by the noradrenaline released by TNS was investigated by repeating Protocol 1.1 in the presence of propranalol. The presence of β receptors in the smooth muscle of the canine saphenous weing was shown by Guimarnes er al(234,235,236). Further, noradrenaline was shown to be effective as an agonist at these ("ecoptors, although much meaker than both adrenaline and isoprenaline In these investigations propranolol wis used to a concentration of 5 x 10^{17} to block the β receptor mediated effects. In Frotocol 1.5 of the present study propranalal used in a concentration of 10 5 mol/1 did not aboligh the inhibition of the Engennue noradrenaling mediated contraction by background TNS. In fact, the exogenous noredrenaline medicied contraction against a trolground of TNS was significantly loss (thus the inhibition mole' is the presure of programatol. Thus, a β -receptor methated effect to sufficive decount for the findings in Frotocol 1 1

When the had been shown to produce playettes in valuate shouth muscles by enting in histaming is action a count if the muscle of 1 membrine (237,238). With each of the as the smooth point of the histories in the blood count up 11 for a concent up 11 for a concent up 11 point (1, 139). We can be to concent for the muscle of the mut numbries in the count of the block of the mut numbries in the count of the block of the count of the coun

the product of the second by methods

 H_2 -receptor antagonist cimetidine. The presence of cimetidine did not significantly alter the response to exogenous noradrenaline against a background of TNS. Therefore, it is unlikely that the inhibition of exogenous noradrenaline contraction observed in Protocol 1.1 is mediated by a H_2 -receptor.

During the last decade the existence of purinergic nerves in nonvascular as well as vascular smooth muscle has been documented(72,241). Adenosine triphosphate (ATP) and/or adenosine are believed to be the neurotrepemitters in these purinergic nerves. Purinoceptors are of two F₁ purinoceptors are most sensitive to adenosine types. and progressively less sensitive to adenosine monophosphate (AMP), adenosine diphosphare (ADF) and ATF On the other hand, the Porpurinoceptors are most sensitive to ATF and progressively less sensitive to ADP, AMP and adappendence. Both ATP and adenosine have been shown to cause relaxation in canine saphenous veins pre contracted with noradrenaline(211). Further. ATP is known to be present in adrenergic nerve endings together with noredrenaline in the synaptic vesicles(130). Thus, it is possible that the inhibit ry effect on the exogenous noradrenaline contraction by THS was mediated by a purinergic substance. Aminophylline was used as a r_1 puringceptor antagonist in the present study. The inhibition of the exogenous perchanaline contraction by the was still evident under these elycumetances. "owever, the effects of a F2 purinceptor antegonist was not investigated in the resent study. Thus, the role for a purinergic substance as a possible mediator for the inhibitory effect on the ogenous poradronaline contraction, cannot be ruled out with the Table evidence in the present study.

Prograduate state to from areabidante rold via the cyclo-

·281

oxygenase pathway are present in most mammalian tissues, although they are not stored to any significant extent except in the seminal fluid(242). Prostaglandins are believed to be primarily local or tissue hormones that have their effects, at or near to the site of synthesis. They exert a vasodepressor or a vasoconstrictor action on vascular smooth muscle depending on the particular prostaglandin involved, the type of blood vessel and the species tested(243). Prostaglandin- E_1 (PGE1) has been shown to reduce the pressor response to intravenous catecholamines in the rabbit and rat vasculature(194). Further. adrenergic nerve stimulation induces synthesis and release of prostaglandins in many tissues(244). Such an effect could theoretically occur during the application of TNS in Protocol 1.1. resulting in the synthesis and release of an inhibitory prostaglandin. Thus, the effect of the cyclo-oxygenase inhibitor indomethacin on the response to exogenous noradrenaline against a background of TNS was tested in the present study. As the inhibitory effect of TNS on the exogenous noradrenaline was still evident under these circumstances an arachidonic acid metabolite of the cyclo oxygeness pathway is unlikely to be the mediator for this inhibitory effect. Now on, the participation of a metabolite of the liver canase pathwar of a solitude gold met bolism cannot be another that the second second of the second second second second second second second second second 11. Kygenne

1

In summary the experiments leng in Protocol 1.5 very not fruitful in the jde tiffication of the postulated inhibitory transmitter. However, the results observed muggested that the inhibitory effort is unlikely to be madiated by a Disceptor is However, a

en properties of the map about the second second

Role of pre-synaptic α_2 -inhibition in the findings in Protocol 1.1

In Protocol Two the role of pre-synaptic α_2 -receptor mediated inhibition during the experimental conditions of Protocols 1.1 and 1.2 were investigated using ³H-noradrenaline. The experiments were carried out in the presence of the uptake, inhibitor cocaine hydrochloride. In Protocol 2.1 exogenous noradrenaline added against a background TNS contraction, significantly decreased the total radioactivity as well as intact ³H-noradrenaline fraction in the superfusate. the Total radioactivity was decreased to 55.8 per cent (mean) of the value during the application of TNS only (Period II and Period IV of the protocol), and ³H-noradrenaline to 30.0 per cent (mean). In Protocol 2.2, the total radioactivity and the ³H-noradrenaline in the superfusate during the period where TNS was applied against a background of exogenous poradrenaline was significantly less than that during the periods of The total radioactivity was 73.0 per cent application of TNS only. (mean) of its value during TNS alone and ³H-noradrenaline 66.0 per cent of its value.

If it is assumed (as is usually done) that the tritiated noradrenaline in the superfusate is a satisfactory measure of the amount of noradrenaline released at the adrenergic nerve ending, there is ovidence for pre-synaptic inhibition by the exogenous noradrenaline added during Protocol 2.1 and Protocol 2.2 (which were similar to Protocol J.1 and Protocol 1.2 respectively). Further, the degree of inhibition appears to be greater in Protocol 2.1 (and thus presumably in Protocol J.1), with the degrease in total radioactivity and the 3 Hnoradrenaline amounting to 44.2 per cent (100-55.8) and 70 per cent (100-30) restrictively as compared with 3° per cent (100-73) and 34.0 per cent (100-66) respectively in Protocol 2.2 (and thus presumably in Protocol 1.2). The TNS induced contraction was smaller in magnitude than the exogenous noradrenaline contraction in Protocol 2.1 and vice versa in Protocol 2.2. Thus, TNS was applied at a higher frequency in Protocol 2.2. The degree of pre-synaptic inhibition is known to be inversely related to the frequency of stimulation of the adrenergic nerves(129,186). Thus the difference in the degree of inhibition observed in the two protocols is not unexpected. Protocol 2.2 which utilised a higher frequency of stimulation demonstrated a lower degree of inhibition as expected.

Superficially this difference in the degree of inhibition may seem to account for the different results observed in Protocol+ 1.1 and Protocol 1.2. In Protocol 1.1 background TNS was found to inhibit the contraction by exogenous noradrenaline. In Protocol 1.2 a background exogenous noradrenaline was found to enhance the contraction by TNS. However, Protocol 2.1 and Protocol 2.2 were carried out only at a single frequency (in each case) of TNS and a single concentration of exogenous noradrenaline. This tended to highlight the difference in the degree of inhibition in the two protocols. Protocol 1.1 and Protocol 1.2, on the other hand, were carried out using a multitude of concentrations of exogenous noradrenaline and a multitude of frequencies of TNS. The difference between the two protocols was entirent throughout the range of TNS employed (refer Fig. 24 and Fig. 27). This makes it unlikely that the difference in the degree of pre-synaptic inhibition would account for the difference observed between Fratocol 1.1 and Protocol 1.2.

Inhibition of the THE induced release of "H noradrenative by

284

a

observed in the Protocol 2.1 and Protocol 2.2 of the present study), has been demonstrated by Lorenz et al(174). In this study the effect of exogenous noradrenaline on the release of tritiated material produced by TNS was investigated also in the absence of an uptakel inhibitor. The addition of noradrenaline (1.2 x 10^{-6} mol/l) to the superfusing fluid increased the tension (by 3.7 g mean $\pm 0.6 \text{ g}$ SEM), the total radioactivity of the superfusate, the efflux of ³H-noradrenaline, its metabolites deaminated and normetanephrine. When this same concentration of ³H-noradrenaline was added against a background contraction by TNS (2 Hz, 9 V, 2.0 ms) the further increase in tension was minimal (<0.2 g) (Fig. 53). This was similar to the results observed in Protocol 1.1 of the present study. Further, with the addition of exogenous noradrenaline in the above study, there was no decrease in the total radioactivity and the ³H-noradrenaline in the The only significant effect was an augmentation of the superfusate. efflux of deaminated metabolites of ³H-noradrenaline. Thus, in the absence of uptake inhibition (the condition present in Protocol 1.1 and Protocol 1.2 of the present study), pre-synaptic inhibition by exogenous noradrenaline could not be demonstrated in the canine saphenous vein. Investigations using canine abdominal sorts, superior mesenteric artery, splenic artery and the splenic capsule too showed a similar inability to demonstrate pressynaptic inhibition in the absence of uptake, blockade in the above study. Only the canine portal vein showed an inhibition of the efflux of ³H-noradrenaline by exogenous noradrenaline in the absence of cocaine.

The results of the above study were explained on the basis of possible displacement by exogenous noradrenaline, of the

Figure 53. Effect of norodrenaline on tension, total radioactivity (superfusate and efflux of ³H-noradrenaline and metabolites during electrical stimulation of comine saphenous vein strips before (a) and after (b) containe (3 x 10^{-5} mol/1). The asterisks indicate times at which superfusate was analysed chromatographically to determine to els of ³H noradrenaline and metabolites. Values shown are mean \pm SEM (Reproduced with permission from S Karger AC; Basel, Blood "assola 10⁻²). 16:113-125).



• Cocoine, 3 k 10⁻⁵M Electric Stimulation, 9 V, 2 Hz ES

noradrenaline within the adrenergic nerve terminal obscuring the demonstration of the pre-synaptic inhibition(174). This was supported by the observation that during TNS, exogenous noradrenaline augmented the efflux of deaminated metabolites in the absence of cocaine but not in its presence. As mentioned before, exogenous noradrenaline (by itself) also produced an increase in the efflux of total radioactivity as well as ³H-noradrenaline. The latter finding makes it difficult to draw a firm conclusion about the pre-synaptic effects of exogenous NA However, whether the noradrenaline comes out of the during TNS. adrenergic nerve terminal by displacement or exocytosis would not make a difference to i contractile effects as long as it comes out in the form intact (unmetabolised) noradrenaline. Thus, tyramine, which acts predominantly by causing displacement of noradrenaling from adrenergic nerve terminals, is still able to cause a powerful contractile effect in a vascular smooth muscle. In the above study, although the added exogénous noradrenaline may have produced an increased efflux of deaminated metabolites the amount of unmetabolised ³H-noradrenaline in the superfusate remained unchanged (Fig. 51). Thus, the contraction produced by TNS should have remained unchanged during the period of addition of exogenous noradrenaline. In spite of this the additional contraction produced by exogenous noradrenaline was less than 0.2 g as mentioned before.

In summary, no evidence for pre-synaptic inhibition by exogenous noradrenaline was available in the canine saphenous vein in the absence of uptake₁ inhibition although one cannot definitely exclude such a phenomenon in this experimental situation. This finding makes it difficult to conclude that pre-synaptic α_2 -inhibition is the mechanism

responsible for the inhibitory effect of TNS, upon the contraction produced by exogenous noradrenaline observed in Protocol 1.1. Another observation also argues against the pre-synaptic inhibition mechanism. Pre-synaptic inhibition is known to decrease with increasing frequency Thus, if pre-synaptic inhibition were to of stimulation(129,186). account for inhibition of the contraction of exogenous noradrenaline in Protocol 1.1, one would expect this latter inhibition to be less with increasing frequency of TNS. However, the observed/expected contraction ratio showed a significant tendency to decrease with increasing magnitude of background TNS in Protocol 1.1 (Fig. 24). The significant contraction ratio а negative observed/expected had correlation (-0.42) with the magnitude of the background TNS. Although this correlation coefficient is relatively low, it is emphasised that one would expect a positive correlation coefficient if pre-synaptic inhibition were to account for the findings in Protocol 1.1.

Relaxation to TNS

As pre-synaptic α_2 inhibition did not seem likely to account for the inhibition of exogenous noradrenaline contraction by TNS in Protocol 1.1, another possibility that may account for this finding. The concurrent release of an inhibitory neurotransmitter by TNS had to be considered. This was investigated in Protocol 3:1 following blockade of the contractile response to TNS. The adrovergic neurone blocker guanethidine (10⁻⁴ mol/1) and the r antagonist phonoxybenzamine, had to be used in combination to effect a blockade of the contractile response to TNS up to a frequency of 32 Hz. Although both the above in gs tore used in relatively bigh concurrations, this the uneroidable as the contractive performance of a contraction to the contraction as the

contraction of about 90 per cent of the maximum contraction to exogenous noradrenaline. The vein rings pre-contracted with prostaglandin $F_{2\alpha}$ following this blockade demonstrated a frequency dependent relaxation to TNS applied as intermittent trains of stimuli. As the experiments were carried out in the presence of both atropine and propranolol, the relaxation could be described as non-adrenergic, non-cholinergic. in nature. The relaxatory response demonstrated a slow recovery and a biphasic response especially at the higher frequencies.

λ.,

Relaxatory responses to TNS, similar to those observed in the present study, have demonstrated been in certain cerebral arteries(245,246,247) and canine coronary arteries in vitro(248). For instance, Bevan et al(246) described a neurally mediated vasodilatation in cat cerebral and extra-cerebral arteries. The relaxation observed was biphasic in nature with an atropine (5 x 10^{-7} mol/1) sensitive component followed by a delayed atropine resistant component. In the present investigation, although a biphasic response was observed occasionally (especially at the higher frequencies of stimulation), both components of the response were found to be resistant to atropine at a concentration of 5 x 10^{-6} mol/l. Ito and Takeda(212) described a relaxatory response to TNS in the cat tracheal smooth muscle following an increase of tone produced by 5-hydroxytryptamine. This relaxatory response was reduced by propranolol (2 x 10^{-6} mol/1) but a significant part of the response was non-adrenergic and non-cholinergic in nature. The authors remarked on a slow recovery from the relaxation produced by the transmural nerve stimulation, similar to that observed in the present study.

Mechanism responsible for the relaxatory response to TNS

TNS leads to contraction or relaxation in smooth muscle by the activation of intramural nerves. This is achieved by induction of action potentials in these nerves with a resultant release of neurotransmitter from the nerve endings. Tetrodotoxin, a specific fast Na⁺ channel blocker has been used as a test for the neural origin of in isolated smooth relaxatory or contractile responses to TNS muscle(249), since it is believed to block the action potentials in the intramural nerves. This phenomenon was investigated in Protocol 3.2. Tetrodotoxin. in a concentration of 10^{-6} mol/1, did not significantly affect the relaxatory response to TNS observed in Protocol 3.1. However, this same concentration of totroliticatin almost completely abolished the contractile response to TNS (up to a frequency of 32 Hz), prior to guanethidine and phenoxybenzamine treatment in the present study. In most of the studies cited above(245,246,247) tetrod towin abolished the relaxatory responses to TNS. Thus, the inability to block the relaxation by tetrodotoxin in the present study could be considered as evidence against a neural origin for this relaxation. However, recent evidence indirates the existence of tetrodotoxin resistant nerve mammal' au potentials and near transmitter relation in action perves(250, 251, 252). Cardebo at all253) also have described a relaxatory response to The which was resistant to " "redetasty in isclated robbit and col, corobial a decoronary acception. Thus it could he argued that there offects are modified by the set with strict totrodotoring peofstant intromast nerses

Another heat which has been continued alton to establish the neuron

storage of isolated tissue at $4^{\circ}C_{\sim}$ in a buffer solution for 9-10 Prolonged cold storage has been shown to cause slow days(254). degeneration of the intramural nerves with irreversible loss of ability to store and release neurotransmitter in isolated smooth muscle(15,213). In Protocol 3.3 of the present study after a period of nine days in storage at 4°C, TNS produced no contractile responses in the saphenous veins, indicating a "denervation" of the sympathetic However, these preparations responded to prostaglandin $F_{2\alpha}$ and nerves. isoprenaline in the conventional manner with contraction and to relaxation respectively although the contractile response was significantly less than the controls. In such veins the relaxatory responses to TNS were abolished also. These findings indicate that a neural mechanism may indeed be involved in the observed relaxatory "egronse.

A second potential mechanism which may mediate this relaxatory response is a direct (non-neurogenic) effect on the smooth muscle cells by the electrical impulses per se. Such an explanation was offered by Pooke et al(248) for the relaxatory response to transmural nerve etimulation observed in the canine coronary arteries since this response was resistant to proluged cold storage as well as to tetrodotoxin. Although the collition of the relaxatory response following cold storage in the present alors and suggests a neurally mediated response, it does not exclude a direct relaxatory offect on the smooth muscle because cold storage also could have had a specific effect upon a hypothetical mechanism responsible for such a direct relaxatory effect.

The of origin of the postulated inhibitory neurotransmitter

it and following non-strangmitter released by the TNS we

responsible for the relaxation observed in Protocol 3.1, it could theoretically originate from two sites (1) from an as yet, unidentified inhibitory intramural nerve (2) from the synaptic vesicles of adrenergic nerves where 1t mav exist as a contrantitor together with noradrenaline The latter possibility does not seem likely as the relaxation to TNS in Protocol 3.1 was demonstrated following guanethidine treatment which would block the colorse of noradrenaling from sympathetic nerves. He ortholess, this was further investigated by hydroxydopamine to produce a chemi al everythectome in the üsing venoue Inga 6 hidrow topomine treatment feaulin the deprioration in advenergin inves such as itoplasmic childring, with al changer ghaonin NE denne interventelos jand mitechnolitat certifing demonstrated by /pitcliano stat in the portal scin and conductantes of the rat(215-255) there thing a much stilling i hours of the treatment the the true of the unitarity that a prostille cotransmittor or out terr her with porglossiller. In the does no vestelos would in rolens i ha op INC fillout op and transmust after t hydroxydop mtre Topoyor the comment would be in the ly a plfpot a non-stranger to terrange to age on 6 hydromater after sectors only of a little to the second of the tener of the tener of the tener of the second the second the tener of the second the tener of the second the tener of the second tener of tener parone of mentor by product mutato from any lips be a to structor uptake Inform the mail a top a sport of Alexander a and experimental to the second of the start fragment for the of the off is a NONCE . When a loss of faith hydroxydopoole treated information, in sponser 1 frage in the started of the line it appears with the "enr e to the second contract to define with considering thet ... · • ʻ 1 11 G the second se

292

· · · ·

observed in the present study. Further support for this conclusion is provided by the inability of tetrodotoxin to significantly affect the relaxatory response to TNS in Protocol 3.2. As tetrodotoxin is known to block the release of noradrenaline from adrenergic nerve endings (by blocking the fast Na^+ channel of the action potential) it would also block the simultaneous release of a co-transmitter.

Possible mediators of the relaxatory response to TNS

Ø

Free radicale have been implicated as a possible mediator of relaxatory responses in vascular smooth muscle(257). A free radical is defined as any atom, group of atoms, or molecule in a particular state with one unpaired electron accupying an outer orbital ("biradical" - two unpaired (lectrong)(258). These radicals have been implicated in a variety of disease states in recent years(258) and have also been shown to produce relaxation in some blod vessels(259). Hyperoxia is one of several factors that are believed to precipitate the generation of free radicale in living tiggue(259). In ignlated emouth muscle experiments the tissue have in expension with a minimum of 95 per cent 0_2 and 5 per cent Cop. This stature has how shown to produce "2 tensions of around 640 mm lg in the thrang bath colution(260). The passage of pulses of current through such a hyperosic medium (during TWS) could be regarded a colorital inform in the generation of these free radicale. Thus, fine indiate can be considered as a potential mediator of the relevators ply opening observed in Frotecol 3.1 of the present study. The possible tote for these radicals are generally investigated by using druge that function as free radiant enorongers (250). However, this is made determine by the enterbrie is variant of free radicate and a relate of relation de water in top exectifie echaempere (259)

Protocol 3.2 of the present study the effect of two such scavengers on the relaxatory response to TNS was investigated. The two drugs used were catalase (a scavenger of hydrogen peroxide and peroxide radicals) and ascorbic acid (a non-specific scavenger)(258,259). These two drugs did not significantly alter the relaxatory response to TNS. Absence of an effect by catalase suggests that hydrogen peroxide is unlikely to plav a role in the relaxation observed in the present study. The conclusion to be drawn from the lack of effect of ascorbic acid is more difficult.

In addition in Frotocol 3.2 the effect of cimeridine (H₂-receptor antagonist), indomethacin (cyclo-oxygenase inhibitor), amipophylline (P₁-purinoceptor antagonist) on the relaxatory response to TNS was investigated. None of these drugs had a significant effect on the response to TNS. Thus, it is unlikely that the relaxation to TMS observed in Protocol 3.1 is madiated by a H₂-receptor stimulation, a P₁-purinoceptor atimulation or by an arachidonic acid metabolite of the system of the system and a significant effect of the system of the system.

The Na⁴/K⁴ Albace inhibitor onabain abolished the relaxatory responses observed in the present study. Since a relatively high concentration of the drug (2 x 10⁻⁴ mol/1) was necessary for ithis blockade a non-specific effect of onabain invalated to the membrane Na⁴/K⁴ ATPace inhibition has to be considered. However nerve P⁴ Freise buffer polation which inhibits the Na⁴/K⁴ pump, also abolished the relaxatory isopones to 70° Thus it appears 197211 to 11 to 1

infinence of mode of MR on the relexation

An increasing enderstand charaction was the fact that

consistent relaxation than continuous stimulation of the isolated venous ring. This is in contrast to the sympathetically mediated contractile effects of transmural nerve stimulation in the isolated canine saphenous vein where continuous stimulation produces a consistent, frequency dependent response(261). Although a frequency dependent uniform response to continuous transmural nerve stimulation is usual in the classical autonomic nerves, a significantly enhanced response to intermittent bursts of stimulation has been described by Edwards and Bloom in the gastro-intestinal tracts of calves(262). This response was associated with the release of a hombesin-like-peptide from intramural nerves from the gastro-intestinal tract. The unusual finding with intermittent transmural nerve stimulation in the present study could be explained also by an enhanced release of such a neurotransmitter.

In summary, the findings of Protocol Three supported the hypothesis that TNS produces a non-adrenergic, non-cholinergic relaxation in the canine saphenous vein pre-contracted with prostaglandin $F_2\alpha$. This relaxation, although residuant to tetrodotoxin, was abolished by cold storage of the veins for 9 days: It is therefore suggested that this relaxatory response could be mediated by an as yet unidentified, tetrodotoxin resistent relaxatory nerve although a direct effect of TNS on the smooth muscle cannot be definitely excluded. Even if the latter mechanism more responsible for the relaxation, it would still be potentially important in isolated smooth muscle experiments as it is likely to occur during TNS in such experiments.

Role of TNS induced relaxation in Protocol 1.1 Finall, one has to consider the role played by this TNS induced

noradrenaline contraction by background TNS observed in Protocol 1.1. Although TNS induced relaxation in Protocol 3.1 was demonstrated following the blockade of the contractile response, there is no reason to believe that the same phenomenon would not occur in the absence of such blockade. However, the relaxatory response would not be apparent because of the bigger magnitude of the contractile effect. Thus, this TNS induced relaxation could theoretically account for the inhibition of the exogenous noradreanline contraction by background TNS observed in Protocol 1.1. Both effects were not antagonised by propranolol, cimetidine, aminophylline, or indomethacin. The inhibitory effect observed in Protocol 1.1 was present with a background contraction with TNS only; other agonists substituted in place of TNS failed to produce any inhibition. Tyramine, which produces its contractile effects by releasing endogenous noradrenaline too did not produce inhibition indicating that the inhibitory effect was dependent on TNS rather than on endogenous noradrenaline. In addition, the inhibitory effect, observed in Protocol 1.1 was found to be significantly more (or the observed/expected contraction ratio less) with increasing magnitude of background TNS. This relationship, as mentioned before, is against presyneptic inhibition being the mechanism responsible for the findings of From (1.1.1.) However, the above relationship of 1d to explaine ' on the hasis of the findings of Frotonni 3.1 whore the manifolds of the nelawatory effect was dependent on the frequency of INS.

Although the abave evidence is suggestive, some facts remain unexplained at the present time. (1) The findings in Prot col 1.2 where a background exemption and trensling contraction was found to pot ordate the contraction by US. Is not contraction was found to pot ordate

hypothesis. The "relaxatory effect" produced by the TNS when applied against a background of exogenous noradrenaline should have limited the total contraction. (2) The inhibitory effect on exogenous noradrenaline contraction was not observed in Protocol 1.3 where the contraction produced by TNS was blocked using guanethidine or diltiazem while maintaining the electrical current. Why the relaxatory effect to TNS observed in Protocol 3.1 did not limit the contraction by exogenous noradrenaline is not clear.

These discrepancies may be explainable on the basis of a number of confounding mechanisms operating during Protocol 1.1 and Protocol 1.2 where the interaction between exogenous noradrenaline and TNS was studied. (1) The possible effects of pre-synaptic α_2 -inhibition when exogenous noradrenaline is applied against a background of TNS and vice versa. (2) The possible potentiation by exogenous noradrenaline of the contractile effects of TNS as observed by Su(229) and Sjöstrand et al(230), and demonstrated in the canine saphenous vein in Protocol Four of the present study. (3) The effects due to the relaxatory response to TNS observed in Protocol 3.1.

The findings and conclusions from the present investigation can be summarised as follows.

- 1. A background TNS contraction inhibited the contraction produced by exogenous noradrenaline in canine saphenous veins.
- ? This inhibitory effect appeared to be greater with increasing magnitude of background TNS.
- 3. The inhibitory effect was specific for background TNS, as it was not observed with background contractions produced by tyramine (an indirectly acting sympathomimetic agent), methoxamine, phenylephrine and histamine.
- The inhibitory effect was not antagonised by propranalol, cimetidine, aminophylline or indomethacin. Thus, the inhibitory effect is unlikely to be mediated by a β -receptor, a H₂-receptor, a P₁-purinoceptor or a prostaglandin metabolite of the cyclo-oxygenase pathway.
- 5. Protocol 1.1 and Protocol 1.2 carried out in a superfused that 'exogenous Protocol demonstrated preparation (in Two) inhibited the TNS induced efflux of total noradrenaline radioactivity and intact ³H-noradrenaline during both protocols.
- 6. Following sympathetic blockade, TNS elicited a frequency dependent relaxation. This relaxation was observed in the presence of propranolol and atropine, i.e., it was non-adrenergic and noncholinergic in nature.
- 7. This relaxatory response to TNS was resistant to tetrodotoxin but was abolished following cold storage of the saphenous veins. The relaxatory response was not affected by chemical sympathectomy with 6-hydroxydopamine.
- 8. The relaxatory response was not blocked by cimetidine, aminophylline, indomethacin, ascorbic acid or catalase. It was abolished by ouabain and zero K^4 Krebs buffer solution suggesting that a functional Na^4/K^4 membrane pump was necessary for the observed relaxation.
- ⁹. Thus, in the canine saphenous vein TNS elicits a non-adrenergic, non-cholinergic relaxation, possibly mediated by a tetrodotoxin resistant nerve.

10. This non-adrenergic, non-cholinergic relaxation to TNS can be put forward as a possible explanation for the inhibition of exogenous noradrenaline mediated contraction by background TNS observed in Protocol 1.1. However, other factors are also likely to be responsible for the findings in Protocol 1.1 and Protocol 1.2. 11. The contractile response to TNS in canine saphenous veins was potentiated by background exogenous noradrenaline induced contractions.

(sub-threshold) concentrations of exogenous noradrenaline 12. Low elicited a potentiation of the responses to trains of TNS' in the canine saphenous vein. If this pehnomenon is present with higher concentrations of exogenous noradrenaline, it could account for the findings in Protocol 1.2. i.e. the enhancement of the response to TNS by background contractions produced by exogenous noradrenaline. 13. There appears to be a significant interaction between the effects of exogenous noradrenaline the effects and of transmural nerve stimulation in isolated canine saphenous veins.

Considerations for the future

A number of findings from the present study merit further investigation. Firstly, no definite conclusion was arrived at, as to the origin of the relaxatory response observed in Protocol Three. Definite proof for a non-adrenergic, non-cholinergic intramural nerve, resistant to tetrodotoxin could be obtained if the relaxatory response could be demonstrated while stimulating the nerve trunk supplying the blood vessel proximally instead of applying transmural nerve, stimulation. The canine lateral spahenous vein derives its adrenergic finervation from the upper lumbar roots via the lumbar sympathetic trunk. If the postulated relaxatory innervation also takes the same anatomical pathway stimulation at a proximal site may indeed be

Thus, in experiment could be carried out in vivo on an possible. isolated, perfused, saphenous vein segment to simulate Protocol Three. The contractile response to the stimulation of the lumbar sympathetic guanethidine would have to be blocked using and nerves phenoxybenzamine. Following this blockade the vein could be contracted with prostaglandin $F_{2}\alpha$ and the response to lumbar sympathetic nerve stimulation elicited with the venous segment perfused at constanct flow with Krebs buffer solution while measuring the hydrostatic pressure The biggest problem in this protocol would be the inside the vein. isolation of the perfused venous segment from the rest of the circulation without damaging its innervation tø any significant This isolation from the rest of the circulation would be extent. essential to avoid any secondary passive changes in pressure from the concomitant arteriolar constriction during the lumbar sympathetic stimulation.

Another approach that can be utilised to determine the origin of the relaxatory response in Protocol Three would be the morphological identification of non-adrenergic, non-cholinergic nerves in the saphenous vein. Electron-microscopy as well as immuno-histochemistry can be adapted for this purpose. The latter method using antibodies to vasoactive intestinal polypeptide and substance P has demonstrated the presence of these substances in nerves in a number of blood vessels.

The second question that has to be resolved is the mechanism behind the potentiation of the response to TNS by exogenous noradrenaline in Protocol Four. Recent experiments (in this laboratory) have demonstrated a similar potentiation of the response to TNS with

methoxamine (selective α_1 -receptor agonist), cloudine (relatively

selective α_2 -agonist) and adrenaline. The potentiation produced by the sub-threshold concentrations of noradrenaline and adrenaline may suggest a role for circulating catecholamines in vivo in enhancing the responses to sympathetic nerve activity. At present circulating catecholamines are not believed to exert any direct action on vascular smooth muscle as their plasma concentrations are well below the threshold for activation of vascular smooth muscle in vitro. Thus the investigation of the findings in Protocol Four, with a view to elucidating the mechanism behind the potentiation would be useful.

Thirdly, the ability of the calcium antagonist, diltiazem to block the contractile response to TNS while leaving the response to exogenous noradrenaline relatively unaltered is also interesting and deserves further investigation.

In summary, the results obtained in the present study may pave the way for a number of diverse investigations.

BIBLIOGRAPHY

Weiner N. Norepinephrine, epinephrine and the sympathomimetic amines. In: Gilman AG, Goodman LS, Gilman A, eds., The pharmacological basis of therapeutics. New York: Macmillan Publishing Co., Inc., 1980; 138-175.

 Paterson G. The response to transmural stimulation of isolated arterial strips and its modification by drugs. J Pharm Pharmacol 1965; 17:341-349.

Shepherd JT, Vanhoutte PM. The human cardiovascular system: Facts and concepts. New York; Raven Press, 1979; 180-207.

ż.

4.

Starke K, Wagner J, Schumann HJ. Adrenergic neurone-blockade by clonidine: comparison with guanethidine and local anaesthetics. Arch Int Pharmacodyn Ther 1972; 195:291-308.

5. Langer SZ. Presynaptic regulation of carecholamine release. Biochem Pharmacol 1974; 23:1793-1800.

6. Berthelsen S, Pettinger WA. A functional basis for classification of α -adrenergic receptors. Life Sci 1977; 21:595-606.

Bentley SM, Drew GM, Whiting SB. Evidence for two distinct types of post-synaptic a-adrenoceptor. Br J Pharmacol 1977; 61:116P-117P.

Drew GM, Whiting SB. Evidence for two distinct types of post Synaptic gradrenoceptors in vascular smooth muscle in vivo - Br J Pharmacol 1979: 67:207-215.

 Docherty JR, MacDonald A, McGrath JC. Further sub-classification of α-adrenoceptors in the cardiovascular system, vas deferens and anococcygeus muscle of the rat. Br J Pharmacol 1979: 67.421P-422P.

- Timmermans PBMWM, Kwa HY, Van Zwieten PA. Possible sub-division of post synaptic α-adrenoceptors mediating pressor responses in the pithed rat. Naunyn Schmiedebergs Arch Pharmacol 1979; 310:189-193.
- 11. Timmermans PBMWM; Van Zwieten PA. Vasoconstriction mediated by postsynaptic α_2 -adrenoceptor stimulation. Naunyn Schmiedebefgs Arch Pharmacol 1980; 313:17-20.
- 12. De Mey J, Vanhoutte PM. Uneven distribution of post junctional alpha₁- and alpha₂-like adrenoceptors in canine arterial and venous smooth muscle. Circ Res 1981; 48:875-884.
- Hirst GDS, Neild TO. Evidence for two populations of excitatory receptors for noradrenaline on arteriolar smooth muscle. Nature 1980; 283:767-768.
- 14. Hirst, GDS, Neild TO. Localisation of specialised noradrenaline receptors at neuromuscular junctions on arterioles of the guineapig. J Physiol (Lond) 1981; 313:343-350.
- 15. Vanhoutte PM, Verbeuren TJ, Webb RC. Local modulation of adrenergic neuroeffector interaction in the blood vessel wall. Physiol Rev 1981; 61:151-247.
- 16. McGrath JC. Evidence for more than one type of post-junctional α-adrenoceptor. Biochem Pharmacol 1982; 31:467-484.
- 17. Osswald W, Guimaraes S, Coimbra A. The termination of action of catecholamines in the isolated venous tissue of the dog. Naunyn Schmiedebergs Arch Pharmacol 1971; 269;15-31.
- 18. Coimbra A, Ribeiro-Silva A, Osswald W. Fine structural and autoradiographic study of the adrenergic innervation of the dog lateral saphenous vein. Blood vessels 1974; 11:128-144.

- 19. Caro CG, Pedley TJ, Seed WA. Mechanics of the circulation. In: Guyton AC, ed.; Gardiovascular Physiology. London: Medical and Technical Publishers, Chap. 1.
- Rhodin JAG, Architecture of the vessel wall. In: Bohr DF, Somlyo AP, Sparks HV. eds. The cardiovascular system, volume II, Vascular smooth muscle. Bethesda: American Physiological Society, 1980; 1-31. (Pappenheimer JR; Forster RF, Mommaerts WFHM, Bullock TH. eds. Handbook of physiology, Section 2).
- 21. Wolinsky H, Glagov S. Structural basis for the static mechanical properties of the aortic media. Circ Res 1964: 14:400-413.
- 22. Furchgott RF, Bhadrakom S. Reactions of strips of rabbit aorta ' epinephrine, isopropylarterenol, sodium nitrite and other drugs J Pharmacol Exp Ther 1953; 8:129-143.
- 23. Herlihy JT. Helically cut mescular strip preparation: geometrical considerations. Am J Physical 1980: 238:H107 H109.

المرابي بالمناج الجارك فالأرفع فيكبه

- 24. Rhodin JAG. The ultrastructure of mammalian arterioles and recapillary sphincters. J Ultrastruc Res 1957: 18:181-223.
- 25. Teleo CH, Glagov S, Kolslev BF. Special structural features of the rat portal uning Anat Pool 1970, 166,509,560.
 - 26. McConnell JG. Roddie, IC. A comparison of the belaviour of ' longitudinal and circular smooth muscle in the second of vetra. J. Physical (Lond) 1970 - 207-821 CMR.
 - 27. Keele CA, Neil E, Joels N. Capillary circulation. It: Samson Wright's applied physiology. Dith ad Oxforts Contribution for Press, 1982; 81-82.
 - 29. Hartshorne DJ. Biochemical basis (rentraction () in the amouth muscle () of () or () 100 () 100 ()

- 29. Webb RC, Bohr DF. Regulation of vascular tone, molecular mechanisms. Prog Cardiovasc Dis 1981; 24:213-242.
- 30. Movsesian MA. Calcium physiology in smooth muscle. Prog Cardiovasc Dis 1982; 25:211-224.
- Somlyo AP, Somlyo AV. Vascular smooth muscle I. Normal structure, pathology, blochemistry and biophysics. Pharmacol Rev 1968; 20:197-272.
- 3? Somiyo AV. Ultrastructure of vascular smooth muscle. In: Bohr DF. Somiyo AV, Sparks HV. eds. The cardiovascular system, volume II, Vascular smooth muscle. Betheeda: American Physiological Society. 1980:33-67. (Pappenheimer TR, Forster RF, Mommaerts WFHM, Bullock TH, eds. Handbook of Physiology).
- 33. Basar E, Weiss C. Vasculature and Circulation. "Amsterdam: Flaevier/North-Holland Biomedical Press, 1981:71-96.
- 36. Devine CE, Somlyo AV, Somlyo AP. Sarcoplasmic reticulum and excitation contraction coupling in mammalian smooth muscles. J Cell Biol 1972: 52:690-718.
- Forbes MS, Rennels ML, Nelson E. Caveolar systems and sarcorinsmic reticulum in coronary smooth muscle colls of the mouse. J Ultractivict Res 1020-67-125-130

Phodin 'AG. Fine structure of vescular walls in mammale. With special reference to smooth muscle component. Physical Rev 1967 47:48 81.

Devine CE, S mino AV, Somiyo AP. Surceplasmic retirulum and mitoch udria as attin accumulating it. in grooth muscle

- 38. Somlyo AV, Somlyo AP. Strontium accumulation by sarcoplasmic reticulum and mitochondria in vascular smooth msucle. Science 1971; 174. 955-958.
- 39. Somiyo AP, Chaldakov C. Burnstock G et al. Structural characteristics, mechanisms of contraction. innervation and proliferation of smooth muscle cells. In: Wolf S, Werthessen MT eds. The smooth muscle of the artery. New York: Plenum Press. 1975; 1-80 (Back N, DiLuz'o NR, Halpern B et al. eds. Advances in experimental medicine and Mology; Vol 57).
- 10. Vallieres J, Scarpa A, Somivo AP. Sub cellular fractions of smooth muscle: isolation. substrate utilisation and Ca²⁺ transpor by main pulmonary artery and mesenteric into mileobondria. Arch Biochem Bi phys 1275: 170:659-669.
- 51 Somlyo AF, Somlyo AV, Shuman H. Electron probe analysis of vascular on oth muncle. composition of mit devices of the state sytemizer. Contrast. 2009, 01:316–335.
- Burneto (*) Structure Flemouth muscle and the fondingstion. I Bulbrie R. Priding M., Jones AW, Comits T. etc., Local of Direct Areas of this bered that 100 p. 1.62

306

.

- 46. Fisher BA, Bazby RM. Reorientation of myofilaments during contraction of a vertebrate smooth muscle. Am J Physiol 1977; 232:C5-C14.
- 47. Craig R, Megerman J. Assembly of smooth muscle myosin in to side-
- 48. Murphy RA. Mechanics of vascular smooth muscle. In: Bohr DF, Somlyo AV, Sparks HV. eds. The cardiovascular system, volume II, Vascular smooth muscle. Bethesda: American Physiological Society, 1980; 325-351. (Pappenheimer JR, Forster RF, Mommaerts WFHM, Bullock TH. eds. Handbook of Physiology).
- 49. Murphy RA. Filament organisation and contractile function in vertebrate smooth muscle. Ann Rev Physiol 1979; 41:737-748.
- Rosenbluth J. Smooth muscle: an ultrastructural basis for the dynamics of its contraction. Science 1965; 148:1337-1339.
- Huxley AF, Miedergerke R. Structural changes in muscle during contraction. Nature 1954; 173:971-973.

Huxley H, Hanson J. Changes in the cross striations of muscle during contraction and stretch and their structural futerpretation - Pature 1001: 173:973 976.

Hartsborne MJ, Gorecka A. Biochemistry of the contractile protring of smooth muscle. In: Bohr DF, Somlyo AV, Sparks HV, edg. The cardiovectular system, volume II. Vascular smooth muscle Bethesda: Ameri an "hysiologi al Scolety, 1980:93-120 (Fappenh imer JR, Forster DI Theman and UNUM Buillock TH. edu Hantle C. Fhysiology)

⁴ Neering LR, Morgan KG. Use of acquorin to study excitationcontraction "ounling in more than the muscle Peture 1980: 288.005 [97]

- 55. Morgan JP, Morgan KG. Vascular smooth muscle: the first recorded Ca²⁺ transients. Pfluegers Arch 1982; 395:75-77.
- 56. Blinks JR, Prendergast FG, Allen DG. Photoproteins as biological calcium indicators. Pharmacol Rev 1976; 28:1-93.
 - 57. Walsh MP, Bridenbaugh R, Kerrick WGL, Hartshorne DJ. Gizzard Ca²⁺ - independent myosin light chain kinase: evidence in favour of the phosphorylation theory. Fed Proc 1983; 42:45-50.
 - 58. Ebashi S, Kodama A. A new protein factor promoting aggregation of tropomyosin. J Biochem 1965; 58:107-108.
 - 50. Stull IT, Sanford CF. Differences in skeletal, cardiac and smooth muscle contractile element regulation by calcium. In: Weiss GB ed. New perspectives on calcium antagonists. Bethesda: American Physiological Society, 1981; 35-46. (Fishman AP, Berne PM. Morgan HE, eds. Clinical Physiology Series).
 - 60. Walters M, Marston SB. Fhosphorylation of the calcium idnregulated thin filaments from vascular smooth muscle. A new regulatory mechanism? Biochem J 1981; 127:127-139.
 - (1) Lehman W, Szent Gvorgvi AG. Regulation of muscle entraction. Distribution of actin control and myorin control in the autout bingdom - 1 Cen Thysics 1275:66:1000.
 - Seller "P. Fhosphorslatic dependent regulation of themical monority of Mit Charles 1911, 206 and a 50
 - 63 Hartshorne DJ, Mrwa U. Pogulation of second and the second sec

Torochiof A. S. tab ine S. S. Girardy, hehe to S. Sanothio,

65. Cheung WY. Calmodulin. Sci Am 1982; 246:62-70.

- 66. Walsh MP. Calmodulin dependent myosin light chain kinases. Cell Calcium 1981; 2:333-352.
- 67. Ebashi S, Toyo-oka T, Nonomura Y. *Gizzard troponin: J Biochem 1975; 78:859-861.
- 68. Ebashi S, Nonomura Y, Nakamura S, Nakasone H, Kohama K. Regulatory mechanism in smooth muscle: actin-linked regulation. Fed Proc 1982; 41:2863-2867.
- 69. Nonomura Y, Ebashi S. Calcium regulatory mechanism in vertebrate smooth muscle. Biomed Res 1980; 1:1-14.
- 70 Robinson BF, Collier JG. Vascular smooth muscle: correlations between basic properties and responses of human blood vessels. Br Med Bull 1979: 35:305-312.
- 71. Owman C, Edvinsson L, Nielson KC. Autonomic neuroceptor mechanisms in brain vessels. Blood vessels 1974; 11:2-31.
- 77. Campbell, G. Gibbons IL. Nonadrenergic, noncholinergic transmission in the autonomic nervous system: Purinergic nerves. In: Kalsner S. ed. Trends in autonomic pharmacology; vol T. Beltimore Munich: Urban & Schwarzenberg, 1979; 103-144.
- 73 Bevan MA, Bevan RD, Duckles SP. Adrenergic regulation of vascular smooth muscle. In: Bohr DF, Somlyo AP, Sparks HV, eds. The cardiovascular system, vol II, Vascular smooth muscle. Bethesda; American Physiological Society, 1980; 515-566. (Pappenheimer JR, Forster RF, Mommaerts WFHM, Bullock TH, eds. Handbook of Ebygiology)
- 74. Burnstock G. Autonomic innervation and transmission. Br Med Bull 1979: 35:255-262.

75. Osswald W, Guimaraes S. Adrenergic mechanisms in blood vessels: Morphological and pharmacological aspects. Rev Physiol Biochem Pharmacol 1983; 96:53-122.

سور با مان ب

t • ...

- Bancroft FW. Venomotor nerves of the hind-limb. Am J Physiol 76. 1898; 1:477-485.
- 77. Donegan JF. The physiology of the veins. J Physiol (Lond) 1921; 55:226-245.
- 78. Webb-Peploe MM. Cutaneous venoconstrictor response to local cooling in the dog. Circ Res 1969; 24;607-615.
- 79. Keele CA, Neil E, Joels N. Membrane transport. In: Samson Wright's applied physiology. 13th ed. Oxford: Oxford University Press. 1982:7-20.
- 80. Harder DR. Membrane electrical activation of arterial smooth muscle. In: Crass MF III, Barnes CD, eds. Vascular smooth muscle: Metabolic, ionic and contractile mechanisms. New York: Academic Press, 1982; 71-97 (Barnes CD, ed. Research topics in physiology).
- Johansson B, Somlyo AP. Electrophysiology and excitation-81. contraction coupling. In: Bohr DF, Somlyo AP, Sparks HV, ede. The cardiovascular system, vol II. Vascular smooth muscle-Bethesda: American Physiological Society, 1980; 301-323 (Pappenheimer JR, Forster RF, Mommaerts WFHM, Bullack TH, eds. Handbook of Fhysiology).
- Fleming WW. The electrogenic Na⁺/K⁺ -pump in smooth muscle: 92. physiologic and pharmacologic significance. Ann Rev Pharmacol Toxicol 1980; 20:129-149.

83. Brading AF. Maintenance of ionic composition. Br Med Bull 1979;

35:227-234.

. . . .

84. Holman ME, Neild TO. Membrane properties. Br Med Bull 1979; 35:235-241.

85. Brender D; Strong CG. Shepherd JT. Effects of acetylstrophanthidin on isolated veins of the dog. Circ Res 1970; 26:647-655.

- 311

- 86. Keatinge WR. Ionic requirements for arterial action potential. J Physiol (Lond) 1968; 194:169-182.
- 87. Brading AF. How do drugs initiate contraction in smooth muscle. Trends in Pharmacol Sci 1981; 2:201-205.
- 88. Cuthbert AW, Sutter MC. The effect of drugs on the relation between action potential discharge and tension in a mammalian vein. Br J Pharmacol 1965; 25:592-601.
- 89. Biamino G, Johansson B. Effects of calcium and sodium on contracture tension in the smooth muscle of the rat portal vein. Pfluegers Arch 1970; 321:143-158.
- 90. Uvelius B, Johansson B. Relation between extracellular potassium ion concentration and contracture force after abolition of spike discharge in isolated rat portal vein. Blood Vessels 1974; 11:120-127.
- 91. Somlyo AV, Somlyo AP. Electromechanical and pharmacomechanical coupling in vascular smooth muscle. J Pharmacol Exp Ther 1968; 159:129-145.
- 92. Farley JM, Miles PR. Role of depolarization in acetylcholine induced contractions of dog trachealis muscle. J Pharmacol Exp Ther 1977; 201:199-205.
- 93. Kitamura K, Kuriyama H. Effects of acetylcholine on the smooth muscle cell of isolated main coronary artery of the guinea pig.

J Physiol (Lond) 1979; 293:119-133. 94. Ilen JC, Bukoski RD. Current status of vascular smooth muscle subcellular calcium regulation. In: Crass MF III, Barnes CD, eds. Vascular smooth muscle: Metabolic, ionic and conractile , mechanisms. New York: Academic Press, 1982: 99-134 (Barnes CD, ed. Research Topics in Physiology).

- 95. Brading AF, Widdicombe JH. The use of lanthanum to estimate the numbers of extracellular cation-exchanging sites in the guinea-pig taenia coli, and its effects on transmembrane monovalent ion movements J Physiol (Lond) 1977; 266:255-273.
- 96. Langer GA., The structure and function of the myocardial cell surface. Am J Physiol 1978; 235:H461-H468.
- 97. Bolton TB. Mechanisms of action of transmitters and other substances on smooth muscle. Physical Rev 1979; 59:606-718.
- 98. Hurwitz L, McGuffee LF, Little SA, Blumberg H. Evidence for two distinct types of potassium-activated calcium channels in an intestinal smooth muscle. J Pharmacol Exp Ther 1980, 214:574-580.
- 99. Van Bremmen C, Aaronson P; Loutzenhiser R, Meisheri K. Calcium fluxes in isolated rabbit aorta and guinea pig taenia coli. Fed Proc 1982; 41:2891-2897.
- 100. Weiss GB. Sites of action of calcium antagonists in vascular smooth muscle. In: Weiss GB ed. New perspectives on calcium antagonists. Bethesda: American Physiological Society, 1981; 83-94 (Fishman AP, Berne RM, Morgan HE, eds. Clinical Physiology Series).
- 101. Meisheri KD, Hwang O, Van Breemen C. Evidence for two separate Ca²⁺ pathways in smooth muscle plasmalemma. J Memb Biol 1981; 59:19-25.

et et e a lorra alla

313

102. Van Breemen C, Farinas BR, Gerba P, McNaughton ED. Excitationcontraction coupling in rabbit aorta. Studied by the lanthanum method for measuring cellular calcium influx. Circ Res 1972; 30:44-54.

- - -

104.

106.

103. Van Breemen Ç, McNaughton E. The separation of cell membrane calcium transport from extracellular calcium exchange in vascular . smooth muscle. Biochem Biophys Res Commun 1970; 39:567-574.

Godfraind T, Miller RC, Socrates Lima J. Effects of yohimbine, rauwolscine and corynanthine on contractions and calcium, fluxes induced by depolarization and prostaglandin $F_{2\alpha}$ in rat aorta. Br J Pharmacol 1983; 80:115-121.

105. Scatchard G. The attractions of proteins for small molecules and ions. Ann NY Acad Sci 1949; 51:660-671.

Hudgins PM, Weiss GB. Effects of Ba, Sr and stimulatory agents on Ca^{2+} movements and contraction in vascular smooth muscle. Fed. Proc 1969; 28:541.

107. Hudgins PM, Weiss GB. Differential effects on calcium removal upon vascular smooth muscle contraction induced by norepinephrine, histamine and potassium. J Pharmacol Exp Ther 1968; 159:91-97.

108. Karaki H, Kubota H, Uraakawa N. Mobilisation of stored calcium for phasic contraction induced by norepinephrine in rabbit aorta. Eur J Pharmacol 1979; 56:237-245.

109: Deth R, Casteels R. A study of releasable Ca fractions in smooth muscle cells of the rabbit forta. J Gen Physiol 1977; 69:401-416.

110. Deth R, Van Breemen C. Agonist induced release of intracellular Ca^{2+} in the rabbit arota. J Memb Biol 1977; 30:363-380.

111. Somlyo AP, Somlyo AV, Shuman H, Endo M. Calcium and monovalent. ions in smooth muscle. Fed Proc 1982; 41:2883-2890.
112. Suzuki S, Sugi S. Ultrastructural and physiological studies on the longitudinal body wall muscle of Dolabella auricularia. Localisation of intracellular calcium and its translocation during mechanical activity. J Cell Biol 1978; 79:467-478.
113. Sugi H, Daimon T. Translocation of intracellularly stored calcium during the contraction-relaxation cycle in guinea-pig taenia coli. Nature 1977; 269:436-438.

الأسرية مسيدة بيبا مردير المراجع والمراجع الأس

- 114. Jones AW. Content anf fluxes of electrolytes. In: Bohr DF, Somlyo AP, Sparks HV, eds. The cardiovascular system, vol II, Vascular smooth muscle. Bethesda: American Physiological Society, 1980; 253-299. (Pappenheimer JR, Forster RF, Mommaerts WFHM, Bullock TH. eds. Handbook of Physiology).
- 115. Blaustein MP. Sodium ions, calcium ions, blood pressure regulation and hypertension: a reassessment and a hypothesis. Am J Physiol 1977; 232:C165-C173.
- 116. Brading AF. Calcium-induced increase in membrane permeability in the guinea pig taenia coli: evidence for involvement of a sodiumcalcium exchange mechanism. J Physiol (Lond) 1978; 275:65-84.
- 117. Schatzmann HJ. Dependence on calcium concentration and stoichiometry of the calcium pump in human red cells. J Physiol (Lond) 1973; 235:551-569.
- 118. Kramer GL, Hardman JG. Cyclic nucleotides and blood vessel contraction. In: Bohr DF, Somlyo AP, Sparks HV, eds. The cardiovascular system, vol II, Vascular smooth muscle. Bethesda: American Physiological Society, 1980; 179-199. (Pappenheimer 'P. Forster RF, Mommaerts WFHM. Bullock TH. ede. Handbook of Physiology).

والمراجع المراجع المراجع والمراجع والمراجع Hardman JG. Cyclic nucleotides and smooth muscle contraction: 119. some conceptual and experimental considerations. 'In: Bulbring E, Brading AF, Jones AW, Tomita T. eds. Smooth muscle: an assessment of current knowledge. Austin: University of Texas Press, 1981; 249-262.

الأحياء ويتبر والمترين والمرك

315

- Adelstein RS, Conti MA, Hathaway DR, Klee CB. Phosphorylation of 120. smooth muscle myosin light chain kinase by the catalytic sub unit of adenosine 3':5'-monophosphate dependent protein kinase. J Biol Chem 1978; 253:8347-8350. •
- 121. Fitzpatrick DF, Szentivanyi A. Stimulation of calcium uptake into aortic microsomes by cyclic AMP and cyclic AMP-dependent protein kinase. Nauyn Schmiedeberg Arch Pharmacol 1977; 298:255-257.
- 122. Andersson R, Nilsson K. Cyclic AMP and calcium in relaxation in intestinal smooth muscle. Nature New Biol 1972; 238:119-120.
- Kukovetz WR, Poch G, Holzmann S. Cyclic nucleotides and 123. relaxation in vascular smooth muscle. In: Vanboutte PM, Leusen I, eds. Vasodilatation. New York: Raven Press, 1981; 339-353.
- Collins GA, Sutter MC. Quantitative aspects of cyclic AMP and 124. relaxation in the rabbit anterior mesenteric-portal vein. Can J Physiol Pharmacol 1975; 53:989-997.
- 125. Diamond J. Role of cyclic nucleotides in control of smooth muscle contraction. In: George WJ, Ignarro LJ, eds. Advances in cyclic nucleotide research, Vol 9. New York: Raven Press, 1978; 327-340.
- 126. Verma SC, McNeill JH. Isoproterenol-induced relaxation, phosphorylase activation and cyclic adenosine monophosphate levels in the polarized and depolarized rat uterus. J Pharmacol Exp Ther 1976: 198:539--547.

•

 127. Katsuki S, Arnold WP, Murad F. Effects of sodium nitroprusside, nitroglycerine and sodium azide on levels of cyclic nucleotides
 and mechanical activity of various tissues. J Cyclic Nucleotide Res 1977; 3:239-247.

- 128. Schultz KD, Schultz K, Schultz G. Södium nitroprusside and other smooth muscle relaxants increase cyclic GMP levels in rat ductus deferens. Nature 1977; 265:750-751.
- 129. Westfall TC. Local regulation of adrenergic neurotransmission. Physiol Rev 1977; 57:659-728.
- 130. Mayer SE. Neurohumoral transmission and the autonomic nervous system. In: Gilman AG, Goodman LS, Gilman A, eds. The pharmacological basis of therapeutics. New York: Macmillan Publishing Co., Inc., 1980; 56-90.
- 131. Kopin IJ, Silberstein SD. Axons of sympathetic neurons: transport of enzymes in vivo and properties of axonal sprouts in vitro. Pharmacol Rev 1972; 24:245-254.
- 132. Klein RL, Lagercrantz H. Insights into the functional role of noradrenergic vesicles. In: Klein RL, Lagercrantz H, Zimmermann H, eds. Neurotransmitter vesicles. London: Academic Press, 1982; 219-239.
- 133. Nelson DL, Molinoff PB. Distribution and properties of adrenergic storage vesicles in nerve terminals. J Pharmacel Function 1976: 196:346-359.
- 134. Sparks HV. Effect of local metabolic factors on vascular smooth muscle. In: Bohr DF, Somlyo AP, Sparks HV, eds. The cardiovascular system, vol II, Vascular smooth muscle. Bethesda: American Physiological Society, 1980; 475-513 (Pappenheimer IR. Forster RF, Mommerts WFHM, Bullock TV eds., Mandbook of Physiology).

- 135. Gershon MD. Effects of tetrodotoxin on innervated smooth muscle preparations. Br J Pharmacol Chemother 1967; 29:259-279.
 - 136. Smith, AD, Winkler H. Fundamental mechanisms in the release of catecholamines. In: Blaschko H, Muscholl E., eds. Handbook of experimental pharmacology, vol XXXIII, Catecholamines. New York: Springer-Verlag, 1972; 538-617.
 - 137. Gothert M, Nawroth P, Neumeyer H. Inhibitory effects of verapamil, prenylamine and D600 on Ca²⁺-dependent noradrenaline release from the sympathetic nerves of isolasted hearts. Naunyn Schmiedebergs Arch Pharmacol 1979; 310:11-19.
 - Zimmerman H. Vesicle re-cycling and transmitter sclease. Neuroscience 1979; 4:1773-1804.

a. . .

- 139. Basbaum CB, Heuser JE. Morphological studies of stimulated adrenergic axon varicosities in the mouse vas deferens. J Cell Biol 1979; 80:310-325.
- 140. Smith AD. Proteins of vesicles from sympathetic axons: chemistry, immuno-reactivity, and release upon stimulation. Neurosci Res Prog Bull 1979; 8:377-382.
- 141. Klein RL. Chemical composition of the large noradrenergic vesicles In: Klein RL, Lagercrantz H, Zimmermann H, eds. Neurotransmitter vesicles. London: Academic Press, 1982; 133-174.
- 14" Kopin IJ, Breese GR, Krauss KR, Weise VK. Selective release of newly synthesised norepinephrine from the cat spleen during sympathetic nerve stimulation. J Pharmacol Exp Ther 1968; 161:271-278.
- 143. Hughes J, Roth RH. Variation in noradrenaline output with changes in stimulus frequency and train length: role of different noradrenaline pools. Br J Pharmacol 1974; 51:373-381.

. 317

144. Bevan JA. Some functional consequences of variation in adrenergic synaptic cleft width and in nerve density and distribution. Fed Proc 1977; 36:2439-2443. 318

- 145. Ljung B, Bevan JA, Pegram BL, Purdy RE, Su M. Vasomotor nerve control of isolated arteries and veins. Acta Physiol Scand 1975; 94:506-516.
- 146. Bevan JA, Su C. Variations of intra- and perisynaptic adrenergic transmitter concentrations with width of synaptic cleft in vascular tissue. J Pharmacol Exp Ther 1974; 190:30-38.
- 147. Vanhoutte PM, Coen EP, De Ridder WJ. Verbeuren TJ. Evoked release of endogenous norepinephrine in the canine saphenous vein. Inhibition by acetylcholine. Circ Res 1979; 45:608-614.
- 148. Iverson LL. Catecholamine uptake processes. Br Mad Bull 1973: 29:130-135.
- 149. Iverson LL. Role of transmitter uptake mechanisms in synaptic neurotransmission. Br J Phrmacol 1971: 41:571-591
- 150. Lorenz RR, Vanhoutte PM. Inhibition of adrenergic neurotransmission in isolated veins of the dog by potentium tone. I Physical (Lond) 1975: 246:479-500.
- Palaic D, Panieset JC. Inhibition of the produced line of the function of the produced line of the function of the product of
- 152 Shore PA. Transport and starships of biogenity interest the Pa. Pharmachi 1972; 12-209 226
- 101 Westfall Die Supersensitivity of smooth mus le Int Pulbring F, Breding AF Jours AW, Tomita I educ Fronch muscle; on a gest of mirror in the second se

5

154. De la Lande IS, Frewin D, Waterson JG. The influence of sympathetic innervation on vascular sensitivity to noradrenaline. Br J Pharmacol Chemother 1967; 31:82-93.

- 1

- 155. Gillespie JS. Uptake of noradrenaline by smooth muscle. Br Med Bull 1973; 29:136-141.
- 156. Tipton KF. Biochemical aspects of monoamine oxidase. Br Med Bull 1973; 29:116-119.
- 157. Sharman DF. The catabolism of catecholamines:recent studies. Br Med Bull 1973; 29:110-115.
- De la Lande IS Adrenergic mechanisms in the rabbit ear artery. Blood Vessels 1975: 12:137-160.
- 159. Muldoon SM, Vanhoutte PM, Tyce GM. Norepinephrine metabolism in canine saphenous vein: prevalence of glycol metabolites. Am J Fhysiol 1978; 234:H235-H243.
- 160. Powis G. Binding of catecholamines to connective tissue and the effect upon the responses of blood vessels to noradrenaline and to nerve stimulation. J Physiol (Lond) 1973; 234:145-162.
- "" Kalsner S. The lack of effect of oxytetracycline on responses to sympathetic perve stimulation and catecholamines in vascular tissue. " I Pharmonol 1976: 58:261-266
- Brandao F, Guimaraes S. Institution of endogenous noradronaling by electrical stimulation in titut of demonstrations and plana Vescule 1774: 11:45-54
 - De Paiva MO, Ouimaraes S. A comparative study of the uptake and metabolism of momentaling and ad enaline by the isolated saphenous vett of the local line of the holdshire of the metabolism.

164. Starke K. Taube HD, Borowski E. Pre-synaptic receptor systems in catecholaminergic transmission. Biochem Fharmacol 1977: 26:259 268.

Ţ

- 165. Paton WDM, Thompson JW. The mechanism of action of adrenaline on the superior cervical ganglion of the cat. Proc Jut Congr Physics Sci 19th, 1953; 664-665.
- 166. Brown GL, Gillespie JS. The output of sympathetic "ransmitter from the spleen of the cat. J Thystol (1997; 1987; 1987; 100
- 167. Farnebo LO, Hambarger B. Drug induced charges f the release of ³H-noradrenaline from field estimulated relifits. Br 1915 - 1 1971: 43:97-106
- Kirpekar SM. Puig M Effect of flow-stop on noradrenaline release from normal spleens and spleens treated with cocaine, observation in the or observation. Br J Flormanni 1971: 40-350 060.
- 112 Langer SL, Adver E. Energ A, Stefano FJE. The role of the receptor in regulating noradrenaline overflow by nerstimulation. Free Jun Congr Physick Sci. 25th, 1211-2005.
- Starke P. Influence of verecepto stim laste such that is the second star back of the star star schedules. 71: 19:420.
- 1981: 21-7 W
- Pismukes K, De Börr AA, Mulder AH. On the mocha ism of the second second stated medulation of ³H nerodicostics is in the second secon

Starke F. Endo T. Taube HD. Relative present presentation notenol of a democeptor agonists in the ability lineary

- 174. Lorenz RR, Vanhoutte PM, Shepherd JT. Interaction between neuronal amine uptake and prejunctional alpha-adrenergic receptor activation in smooth muscle from canine blood vessels and spleen. Blood Vessels 1979:16:113-125.
- 175. Starke K, Montel H. Influence of drugs with affinity for αadrenoceptors on noradrenaline release by potassium, tyramine and dimethylphenyl-piperazinium. Eur J Pharmacol 1974; 27:273-280.
- Westfall TC, Leighton HJ. Effect of decentralisation on presynaptic receptor regulation of NE release. Pharmacologist 1976; 18.208.
- Vanhoutte PM, Collis MG, Janssens WJ, Verbeuren TJ. Calcium dependence of pre-junctional inhibitory effects of adenosine and acetylcholine on adrenergic neurotransmission in canine saphenous veine. Eur / Fharmacol 1981; 72:189-198.
- 179 Sanders KM, Ross G. Inhibition of in vivo neural vasoconstriction by exogenous catecholamines. Blood Vessels 1975; 12:13-20.
- 179, Westfall TC. Evidence that noradrenergic transmitter release is regulated by pre-synaptic receptors. Fed Proc 1984; 43:1352-1357.
- Kalaner S. The pre-synaptic receptor controversy. Trends in Flarmancel Sci 1982; 3:11-16, 18-21.
- Rand MI. McCullach MW, Story DF. Fredback modulation of coreduction transmission Trends in Fharmacol Sci 1982; 3:8-11, 16 10
- Angus 34, Former Pf. Evidence against pre-synaptic α-adrenoceptor mod lation of a site compatientic transmission. Nature 1980;

321

- 183. Kalsner S. Single pulse stimulation of guinea-pig vas deferens and the pre-synaptic receptor hypothesis. Br J Pharmacol 1979; 66:343-349.
- 184. Kalsner S, Chan CC. Adrenergic antagonists and the presynaptic receptor hypothesis in vascular tissue. J Pharmacol Exp Ther 1979; 211:257-264.
- 185. Dubocovich ML, Langer SZ, Massingham R. Lack of correlation between pre-synaptic inhibition of noradrenaline release and end organ responses during nerve stimulation. Br J Pharmacol 1980; 69:81-90.
- 186. Langer SZ. Pre-synaptic receptors and their role in the regulation of transmitter release. Br J Pharmacol 1977; 60.481-497.
- 187. Vanhoutte PM, Levy MN. Prejunctional cholinergic modulation of adrenergic neurotransmission in the cardiovascular system. Am J Physiol 1980; 238:H275-H281
- 188. Vanhoutte PM, Lorenz RR, Tyce GM. Inhibition of norepinephrine-³H release from sympathetic nerve endings in veing by acetylcholine. J Pharmacol Exp Ther 1973; 195:386-394.
- ind. Shepherd JT, Lorenz RR, Tyce GM, Canhoutte PM. A etvicholine inhibition of transmitter release from adrenergic nerve cerminal modiated by muscerinic recentors. For these 1979, 17 101-124
- 100 Verhaeghe RH, Vanhoutte PM, Shepherd II Inhibition of symmathetic neurotransmission in comine blood vessels by all users and adomine nucleatides - Circ Pes 1017: 40-208 215
- The Mey T, Rurnstock G, Venhoutte PM. Modulation of the evoked release of non-adrebalize in caning suphemous vein vic pre-ivmentic reprocession of the two controls of the controls of the control of the two controls of two controls of the two controls of two controls

322

 ~ 1

ş

405.

- 192. McGrath MA, Shepherd JT. Histamine and 5-hydroxytryptamineinhibition of transmitter release mediated by H₂- and 5hydroxytryptamine receptors. Fed Proc 1978; 37:195-198.
- 193. Malik KU, Ryan P, McGiff SC. Modification by prostaglandin E_1 and E_2 , indomethacin and arachidonic acid of the vasoconstrictor responses of the isolated perfused rabbit and rat mesenteric arteries to adrenergic stimuli. Circ Res 1976; 39:163-168.
- 194. Horton EW. Prostaglandins at adrenergic nerve endings. Br Med Bull 1973; 29:148-151.
- 195. Malik KU. Prostaglandins-modulation of adrenergic nervous system. Fed Proc 1978; 37:203-207.
- 196. Zimmerman BG. Actions of angiotensin on adrenergic nerve endings. Fed Proc 1978; 37:199-202.
- 197. Westfall TC, Peach MJ, Tittermary V. Enhancement of the electrically induced release of norepinephrine from the rat portal vein: mediation by β_2 -adrenoceptors. Eur J Pharmacol 1979; 58:67-74.
- Majewski H, Rand MJ, Tung LH. Activation of pre-junctional β adrenoceptors in rat atria by adrenaline applied exogenously or released as a contransmitter. Br J Pharmacol 1981; 73:669-679.
- 199. Furchgett RF. Spiral-cut strips of rabbit aorta for in vitro studies of responses of arterial smooth muscle. In: Bruner HD, Root WS, Lands AM, eds. Methods in medical research, Vol 8, Chicago: The Year Book Publishers, Inc., 1960: 177-186.
- 700. Duckles SP, Silverman RW., Transmural nerve stimulation of blood vessels in vitro: a critical examination. Blood Vessels 1980; 17:53-57

- 201. Vanhoutte P, Leusen L. The reactivity of isolated venous preparations to electrical stimulation. Pfluegers Arch 1969; 306:341-353.
- 202. Van Rossum JM, Hurkmans JAThM, Wolters CJJ. Cumulative doseresponse curves. II. Technique for the making of dose response curves in isolated organs and the evaluation of drug parameters. Arch Int Pharmacodyn Ther 1963; 143:299-330.
- 203. Shepherd JT, Vanhoutte PM. Local modulation of adrenergic neurotransmission. Circulation 1981; 64:655-666.
- 204. Hughes J, Roth RH. Evidence that angiotension enhances transmitter release during sympathetic nerve stimulation. Br J Pharmacol 1971; 41:239-255.
- 205. Lorenz RR, Powis DA, Vanhoutte PM, Shepherd JT. The effects of acetylstrophanthidin and ouabain on the sympathetic adrenergic neuroeffector junction in canine vascular smooth muscle. Circ Res 1980; 47:845-854.
- 206. Cohen RA, Shepherd JT, Vanhoutte PM. Pre-junctional and postjunctional actions of endogenous norepinephrine at the sympathetic neuroeffector junction in canine coronary arteries. Circ Res 1983; 52:16-25.
- 207. Levin JA. Paper chromatographic assay of (³H) norepinephrine and its five major metabolites. Anal Biochem 1973; 51:42-60.
- Verbeuren TJ, Coen E, Vanhoutte PM. Determination of ³H norepinephrine and its metabolites in superfusate from isolated blood vessels. Arch Int Pharmacodyn Ther 1977; 227:171-74.
- 209 Graefe KH, Stefano FJE, Langer SZ. Preferential metabolism of (*) ³H-norepinephrine through the deaminated glycol in the tot vas deference. Biochem Pharmacol 1973; 22:1147-1160

- 210. Anton AH, Sayre DF. A study of the factors affecting the aluminum oxide-trihydroxyindole procedure for the analysis of catecholamines. J Pharmacol Exp Ther 1962; 138:360-375.
- 211. De Mey JG, Vanhoutte PM. Heterogeneous behavior of the canine arterial and venous wall-importance of the endothelium. Circ Res 1982; 51:439-447.
- 212. Ito Y, Takeda K. Non-adrenergic inhibitory nerves and putative transmitters in the smooth muscle of the cat trachea. J Physiol (Lond) 1982; 330:497-511.
- 213. Vanhoutte PM. Physical factors of regulation. In: Bohr DF, Somlyo AV, Sparks HV. eds. The cardiovascular system, volume II, Vascular smooth muscle. Bethesda. American Physiological Society, 1980; 449. (Pappenheimer JR, Forster RF, Mommaerts WFHM, Bullock TH. eds. Handbook of Physiology).
- 214. Guimaraes S, Osswald W, Cardoso W, Branco D. The effects of cocaine and denervation on the sensitivity to noradrenaline, its uptake and the termination of its actions in isolated venous tissue. Naunyn Schmiedebergs Arch Pharmacol 1971; 271:262-273.
- ?15. Aprigliano 0, Hermsmeyer K. In vitro denervation of the protal vein and caudual artery of the rat. J Pharmacol Exp Ther 1976; 198:568-577.
- 216. Sokal RR, Rohlf FJ. Biometry: 2nd ed., San Francisco. WH Freeman and Company, 1981:170-177, 499-530.
- 217. Snedecor GW, WG Cochran. Statistical Methods: 7th ed. Ames, Iowa. Towa State University Press, 1980:385-388.
- 218. Burnstock G. Neurotransmitters and trophic factors in the autonomic nervous system. J Physiol (Lond) 1981; 313:1-35.

- 219. Burnstock G. Do some nerve cells release more than one transmitter? Neuroscience 1976; 1:239-248.
- 220. Viveros OH, Diliberto EJ, Daniels AJ. Biochemical and functional evidence for the co-secretion of multiple messengers from single and multiple compartments. Fed Proc 1983; 42:2923-2928.
- 221. Flavahan NA, McGrath JC. An analysis of α_1 and α_2 -adrenoceptor mediated pressor effects of adrenaline. Br J Pharmacol 1981; 72:519P.
- 222. Flavahan NA, McGrath JC. Blockade by yohimibine of prazosinresistant pressor effects of adrenaline in the pithed rat. Br J Pharmacol 1980; 69:355-357.
- 223. De Mey JG, Vanhoutte PM. Differences in pharmacological properties of post junctional alpha-adrenergic receptors among arteries and veins. Arch Int Pharmacodyn Ther 1980; 244:328-329.
- 224. Rapoport R, Bevan JA. Acute stress reduces the sensitivity of the vasculature to sympathetic control. Experientia 1979; 35:1609-1611.
- 225. Rapoport RM, Bevan JA. Vascular smooth muscle contraction induced by direct electrical stimulation and its subsequent effect on reactivity to agonists. J Pharmacol Exp Ther 1981; 218:375-381.
- 226. Rapoport RM, Bevan JA. Mechanism of electrical stimulationinduced subsensitivity of vascular smooth muscle. J Pharmacol Exp Ther 1981; 218:382-389.
- 227. Trendelenberg U. Supersensitivity to norepinephrine induced by continuous nerve stimulation. J Pharmacol Exp Ther 1966; 151:95-102.

- 228. Hope W, Law M, McCulloch MW, Rand MJ, Story DF. Effects of some catecholamines on noradrenergic transmission in the rabbit ear artery. Clin Exp Pharmacol Physiol 1976; 3:15-28.
- 229. Su C. Potentiative effects of alpha agonistic sympathomimetic amines on vasoconstriction by adrenergic nerve stimulation. J Pharmacol Exp Ther 1980; 215:377-381.
- 230. Sjostrand NO, Swedin G. On the mechanism of the enhancement by smooth muscle stimulants of the motor responses of the guinea-pig vas deferens to nerve stimulation. Acta Physiol Scand 1974; 90:513-521.
- 231. Vanhoutte PM, Rimele TJ. (Personal Communication).
- 232. Muscholl E. Autonomic nervous system: newer mechanisms of adrenergic blockade. Ann Rev Pharmacol 1966; 6:107-128.
- 233. Starke K. α-adrenoceptor subclassification. Rev Physiol Biochem Pharmacol. 1981; 88:199-236.

. 6

- 234. Guimaraes S. Further study of the adrenoceptors of the saphenous vein of the dog: influence of factors which interfere with the concentrations of agonists at the receptor level. Eur J Pharmacol 1975: 34:9-19.
- 235. Guimaraes S, Osswald W. Adrenergic receptors in the veins of the dog. Eur J Pharmacol 1969; 5:133-140.
- 236. Guimaraes S, Paiva MQ. Two distinct adrenoceptor-biophases in the vasculature: one for α and the other for β -agonists. Naunyn Schmiedebergs Arch Pharmacol 1981; 316:195-199.
- 237. Brody MJ. Histamine and vascular smooth muscle. In: Bevan JA, Godfraind T, Maxwell RA, Vanhoutte PM, eds. Vascular neuroeffector

mechanisms. New York: Raven Press, 1980; 82-87.

- 238. Tucker A, Weir EK, Reeves JT, Grover RF. Histamine H₁- and H₂receptors in pulmonary and systemic vasculature of the dog. Am Physiol 1975; 229:1008-1013.
- 239. Ryan, MJ, Brody MJ. Neurogenic and vascular stores of histamine in the dog. J Pharmacol Exp Ther 1972; 181:83-91.
- 240. Ryan MJ, Brody, MJ. Distribution of histamine in the canine autonomic nervous system. J Pharmacol Exp Ther 1970; 174:123-132.
- 241. Burnstock G. Cholinergic and purinergic regulation of blood vessels. In: Bohr DF, Somlyo AV, Sparks HV. eds; The cardiovascular system. Volume II. Vascular smooth muscle. Bethesda: American Physiological Society, 1980: 567-612 (Pappenheimer JR, Forster RF, Mommaerts WFHM, Bullock TH, eds. Handbook of Physiology).

242. McGiff JC. Prostaglandins, prostacyclin and thromboxanes. Ann Rev Pharmacol Toxicol 1981; 21:479-509.

- 243. Altura BM, Altura BT. Vascular smooth muscle and prostaglandins. Fed Proc 1976; 35:2360-2366.
- 244. Messina EJ, Weiner R, Kaley G. Prostaglandins and local circulatory control. Fed Proc 1976; 35:2367-2375.

· .

- 245. Lee TJF, Su C, Bevan JA. Non-sympathetic dilator innervation of cat cerebral arteries. Experientia 1975; 31:1424-26.
- 246. Bevan JA, Buga GM, Jope CA, Jope RS, Moritoki H. Further evidence for a muscarinic component to the neural vasodilator innervation of cerebral and cranial extracerebral arteries of the cat. Circ Res 1982; 51:421-29.

- 247. Winquist RJ, Webb RC, Bohr DF. Relaxation to transmural nerve stimulation and exogenously added norepinephrine in porcine comprai vessels. Circ Res 1982; 51:769-776.
- 248. Rooke T, Cohen RA, Verbeuren TJ, Vanhoutte PM. Non-neurogenic inhibitory effect of electrical impulses in isolated canine coronary arterias. Eur J Pharmacol 1982; 80:251-254.
- 249. Duckles SP. Transmural electrical stimulation: Distinguing between activation of nerves and smooth muscle. In: Barge JA, Godfraind T, Maxwell RA, Vanhoutte PM, eds. Vascular neuroeffector mechanisms. New York: Raven Press, 1980; 33-35.
- 250. Blaschke E, Uvnas B. No effect of tetrodotoxin on catecholamine release from the perfused Cat adrenal gland. Acta Physiol Scand 1981; 113:267-269.
- 251. Hirst GDS, Spence I. Calcium action potentials in mammalian peripheral neurones. Nature 1973; 243:54-56.
- 252. McAfee DA, Yarowsky PJ. Calcium dependent potentials in the mammalian sympathetic neurone. J Physiol (Lond) 1979; 290:507-523.
- 253. Hardebo JE, Hanko J, Owman C. (Abstract). Characterisation of relaxant responses to transmural stimulation in isolated blood vessels. 1983; 20:195-196.
 - 254. Lee TJF, Hume WR, Su C, Bevan JA. Neurogenic vasodilatation of cat cerebral arteries. Circ Res 1978; 42:535-542.
 - 255. Aprigliano O, Rybarczyk KE, Hermsmeyer K, Van Orden LS III. Adrenergic neuronal degeneration induced in portal vein and caudal artery by 6-hydroxydopamine in vitro. J Pharmacol Exp Ther 1976; 198:578-588.

256. Kostrzewa RM, Jacobwitz DM. Pharmacological actions of 6hydroxydopamine. Pharmacol Rev 1974; 26:199-288.

- 257. Furchgott RF, Zawadski JV, Cherry PD. Role of endothelium in the vasodilator response to acetylcholine. In: Vanhoutte PM, Leusen I, eds. Vasodilatation. New York, Raven Press, 1981; 49-66.
- 258. Del Maestro RF. An approach to free radicals in medicine and biology. Acta Physiol Scand (Suppl) 1980; suppl 492:153-168.
- 259. Rosenblum WI. Effects of free radical generation on mouse pial arterioles: probable role of hydroxyl radicals. Am J Physiol 1983; 245:H139-H142.
- 260. De Mey JG, Vanhoutte PM. Interaction between Na⁺, K⁺ exchanges and the direct inhibitory effect of acetylcholine on canine femoral arteries. Circ Res 1980; 46:826-836.
- 261. Vanhoutte PM, Clement D, Leusen I. The reactivity of isolated veins to electrical stimulation. Arch Int Physiol Biochem 1967; 75:641-657.

ford.

262. Edwards AV, Bloom SR (Abstract). The importance of the pattern of stimulation in relation to autonomic responses. Proceedings of the International Union of Physiological Sciences (20th Congress). 1983; 15:291

Appendix I

Drugs and Chemicals

The Krebs buffer solution used, had the following composition (mmol/1) NaCl 116.0, KCl 5.4, CaCl₂ 1.2, NaHCO₃ 22.0, NaH₂PO₄ 1.2, MgCl₂ .6H₂O 1.2, glucose 10.1, CaNa₂EDTA 0.023.

Sources of drugs used

 aluminum oxide (Woelm Pharma-GmbH & Co., Eschwege, West Germany)

2. aminophylline: MW 420.4 (Abbot Laboratories Ltd., Montreal, Canada).

3. ascorbic acid: MW 176.1 (Sigma Chemical Co., St. Louis, U.S.A.)

4. atropine: MW 289.4 (Sigma Chemical Co., St. Louis, U.S.A.)
5. catalase (Sigma Chemical Co., St. Louis, U.S.A.)

6. cimetidine hydrochloride: MW 288.8 (Sigma Chemical Co., St. Louis, U.S.A.).

7. cocaine hydrochloride: MW 339.8 (Sigma Chemical Co., St. Louis, U.S.A.).

8. 3.4-dihydroxymandelic acid (DOMA):MW 184.1 (Sigma Chemical Co., St. Louis, U.S.A.)

9. 3.4 dihydroxyphenylglycol (DOPEG):MW 170.2 (Sigma Chemical Co., St. Louis, U.S.A.).

10 diltiazem hydrochloride: MW 451.0 (Sigma Chemical Co., St. Louis, U.S.A.).

11.	di sodium ethylene diamine tetra-acetic acid:MW 336.2 (Sigma
	Chemical Co., St. Louis, U.S.A.).
12.	Dowex-1 (Sigma Chemical Co., St. Louis, U.S.A.).
13.	Dowex-50 (Bio-Rad Laboratories, Richmond, California, U.S.A.).
, ¹⁴ .	Guanethidine sulphate:MW 294.4 (Ciba-Geigy Canada Ltd.).
.15.	Histamine dihydrochloride:MW 184.1 (Sigma Chemical Co., St
	Louis, U.S.A.).
16,	6-hydroxydopamine hydrochloxide MP 205.7 (Sigma Chemical Co.,
	St. Louis, U.S.A.).
17.	Indométhacin hydrochloride MU 357.8 (Sigme Chemical Co., St
	Louis, U.S.A.).
18.	Methoxamine bydrochloride (vagoual):MW 247.7 (Burrougha
	Wellcome Ltd., Montreal, Conada)
19.	3. Methoxy-4hydroxymandelt antid (VMA) Hull 198 2 (Stamp Chroning)
	Co., St. Louis, U.S.A.).
,	3 Methows (hydroxypteryless) (pere or so is in the contract of the
	Co., St. Loute - C.C. S.
۰ ،	Noradr neline (free ways) (WW 319.) (free best of
	Louis, U.S.A.),
••	7 ³ A normaline (specific activity 10 10 ()
	Nuclear, Boston U.S.A.).
C 1	Novi tar philne budrocht ille (11) - 11 - 11

.

332

9

w'

د. م

****** *	
24.	Ouabain (octahydrate):MW 728.6 (Sigma Chemical Co., St. Louis,
	U.S.A.).
25.	Octyl phenoxy polethoxyethanol (Triton X-100) (Sigma Chemical
	Co., St. Louis, U.S.A.).
26.	Pentobarbital sodium (Somnotol): (M.T.C. Pharmaceuticals,
	Hamilton, Canada).
?-	Phenoxybenzamine hydrochloride:NW 340.3 (Smith Kline and French Canada Ltd.
28.	Phenylephrine hydrochloride:MW 203." (Sigma Chemical Co., St.
	Louis, U.S.A.).
· 0	Propranolol hydrochloride:MW 295.8 (Sigma Chemical Co., St. & Louis, U.S.A.).
) (* .	Prostaglandin $F_{2\alpha}$: (Prostin $F_{2\alpha}$, Dinoprost tromethamine) MW
	475.6 (Upjohn Company of Canada, Don Mills, Canada
1 1	Tetrodotoxin:MW 319 (Sigme Chemical Co., St. Louts, U.S.A.).
	Tvramine hydrochlovite: 173.6 (Sigma Chemical o, Sr. Louis
יח.	a concentrated stock solutions (10 4 mol(1 in most cases) of the
	propared in distilled wates. 1.0 mg of acdium metableulphire
	d to each 10 ml of the stock solutions to minimise oxidation in
he cas	e of easily oxidisable druge. Suitable dilutions of the drugs
	pared in Trabs buffer solutions each day.

•

stant is 2 100pt aligned to the tissue bath to produce the desired g .

· • · · · • •

concentration in the bath fluid. Continuous bubbling of the tissue bath fluid with 95 per cent 0_2 -5 per cent $C0_2$ ensured that this desired concentration in the fluid was achieved almost instantaneously. The highest concentration of the drugs during dose-response curves were maintained in the tissue bath for as short a duration as possible to minimise desensitization. Drugs were removed from the bath fluid by repeated rinsing of the bath with fresh Krebs buffer solution. When a drug had to be kept in the tissue bath of a fixed concentration throughout an experiment (α_2 , propranolol and atropine in Protocol 3.1) the drugs were added to the reservoir of Krebs solution feeding the baths. This prevented the epsated additions of small slippoir of the drugs that would chere is not thus $\pi^2\eta^2$ is all drugs that would chere is not thus $\pi^2\eta^2$ is all drugs that would chere is not thus $\pi^2\eta^2$ is all

Appendix II

Solutions used for column chromatographic analysis 2 litres 2N hydrochloric acid 1. 332 ml of 12N HC1 400 ml of 0.5% Triton X-100 1268 ml H₂0 2. 1 litre sodium phosphate buffer pH 6.5 6.6 g KH₂PO₄ 2.67 g Na₂HPO₄ 1 g Na₂EDTA Make up to 11 with 0.1% Triton x 100: pH checked and titrated to 6.5 if necessary. 1 litre 6N HC1/ethanol 250 ml .12N HC1 500 ml ethanol 240 ml H₂0 (no Triton X-100) 1 litre 0.2N hydrochloric acid 100 ml 2N HC1 200 ml 0.5% Triton X-100 700 ml H20 'itre 1 N hydrochloric arid 500 ml 2N HCL 200 ml 0.5% Triton X-100 300 ml Har 1 litre 0 2N noetic sold 200 ml l N scetis acid 200 ml 0.57 Triton X-100 700 ml H_0 7. ' 'itre 0.1% Triton X-100 200 ml 0.5% Triton X-100

800 ml H20

Appendix III

Solutions used during chemical sympathectomy with 6-hydroxydopamine

Composition of unbuffered physiological salt solution used (mmol/l): NaCl 136.7, MgCl₂ 2.1, KCl 2.7, CaCl₂ 1.8, glueose 5.4, CaNa₂EDTA 0.023.

The pH of the solution was adjusted to 4.9 using reduced glutathione. 6.0 mg of 6-bydroxydopamine was dissolved in 20 ml (1.46 x 10^{-3} mol/1). of this physiological salt solution just prior to use on each day.

Appendix IV

The definitions of some frequently used terms in Pharmacology are given below. The definitions do not exactly correspond to the correct meaning of these terms in the English language, nevertheless, these definitions have become irrevocably embedded in Pharmacological literature.

Washing, rinsing: the replacement of the bath fluid with fresh Krebs buffer solution containing no drugs (unless specified otherwise).

Dose:

•

the aliquot of a drug added into the tissue bath to produce a desired concentration in the bath fluid following the dilution of the drug in the bath.

Doserreponse curve:

refers to the effect to an agonist (y axis) plotted against the concentration of the agonist (x axis) i.e. concentration effect curve.

Incubation:

Exposing a tissue to a desired concentration of a drug for a specified length of time. This is commonly used in reference to antagonists which are kept in contact with a tissue for a length of time before repetition of the agonists effects.