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TITLE: XIth International Symposium on Cholinergic Mechanisms - Function and Dysfunction

PRINCIPAL INVESTIGATOR: Israel Silman, Ph.D.

CONTRACTING ORGANIZATION: The Weizmann Institute of Science
Rehovot 76100 Israel

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PREPARED FOR: U.S. Army Medical Research and Materiel Command
Fort Detrick, Maryland 21702-5012

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NSN 7540-01-280-5500
XIth INTERNATIONAL SYMPOSIUM ON CHOLINERGIC MECHANISMS-FUNCTION AND DYSFUNCTION & 2nd MISRAHI SYMPOSIUM ON NEUROBIOLOGY

St. Moritz, Switzerland, May 5 - 9, 2002

PROGRAM AND ABSTRACTS
XIth INTERNATIONAL SYMPOSIUM ON CHOLINERGIC MECHANISMS-FUNCTION AND DYSFUNCTION & 2nd MISRAHI SYMPOSIUM ON NEUROBIOLOGY
St. Moritz, Switzerland
May 5 - 9, 2002

ORGANIZED BY

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<table>
<thead>
<tr>
<th>TABLE OF CONTENTS</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Committees</td>
<td>3</td>
</tr>
<tr>
<td>Acknowledgements</td>
<td>4</td>
</tr>
<tr>
<td>General Information</td>
<td>6</td>
</tr>
<tr>
<td>Social Events</td>
<td>8</td>
</tr>
<tr>
<td>Accompanying Persons' Program</td>
<td>8</td>
</tr>
<tr>
<td>Scientific Program</td>
<td></td>
</tr>
<tr>
<td>Monday, May 6, 2002 – Opening</td>
<td>11</td>
</tr>
<tr>
<td>Tuesday, May 7, 2002</td>
<td>13</td>
</tr>
<tr>
<td>Wednesday, May 8, 2002</td>
<td>15</td>
</tr>
<tr>
<td>Thursday, May 9, 2002</td>
<td>17</td>
</tr>
<tr>
<td>Posters</td>
<td></td>
</tr>
<tr>
<td>Monday, May 6, 2002</td>
<td>21</td>
</tr>
<tr>
<td>Tuesday, May 7, 2002</td>
<td>24</td>
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<tr>
<td>Wednesday, May 8, 2002</td>
<td>27</td>
</tr>
<tr>
<td>Thursday, May 9, 2002</td>
<td>30</td>
</tr>
<tr>
<td>Abstracts</td>
<td></td>
</tr>
<tr>
<td>Poster Abstracts</td>
<td>33</td>
</tr>
<tr>
<td>Index</td>
<td></td>
</tr>
</tbody>
</table>
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(Alphabetical)

ACCOMMODATION
Kenes Tours, the official Symposium travel agent, will operate a hospitality desk during the Symposium.

BADGE
On registering you will receive your Symposium kit in which you will find your name badge. You are kindly requested to wear your badge during all sessions and events.

LANGUAGE
The official language of the Symposium is English.

LIST OF PARTICIPANTS
A list of participants who have registered prior to the Symposium is displayed on the bulletin board. Please amend/add your name and address to the list.

LOCATION
Laudinella Hotel, Address: CH-7500, St. Moritz, Switzerland
Tel: +41 81 836 0000; Fax: +41 81 836 0001
E-mail: info@laudinella.ch
Website: www.Laudinella.ch

LUNCHES
An organized lunch is arranged for all participants and accompanying persons staying at the Laudinella hotel, on Monday, Tuesday, Wednesday and Thursday. Participants staying at the Hotel Loffler will also receive lunch vouchers for the Laudinella hotel.

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Posters will be on display on Monday, Tuesday, Wednesday and Thursday and will change every day. Please check your poster scheduling in the program. Your poster should be displayed as per the Board No. in the Program. Posters are to be mounted from 07:45-08:30 on the day your poster is scheduled. Please remove your poster at the conclusion of sessions on your assigned day. The Organizing Committee will not be responsible for posters which are not removed on time. Poster presenters are requested to be present at their poster board during coffee breaks.

REGISTRATION

REGISTRATION/INFORMATION/SECRETARIAT DESKS
Registration desks will be situated at the Laudinella Hotel as follows:
Sunday, May 5 from 16:00 – 20:00.
The desks will reopen on Monday, May 6 at 07:30 and will stay open throughout the sessions on Tuesday, Wednesday and Thursday.

SPEAKERS READY ROOM (Oral Presentations)
A speakers’ ready room will be provided for speakers and will be available every day 30 minutes prior to sessions.

Audiovisual Equipment

1. Overhead projection (projection of transparencies). If using transparencies, please inform the technician in the Speakers’ Ready Room at least 30 minutes before the start of your session.

2. Slide projection (projection of 35mm slides). If using slides, you are requested to load them in a slides’ tray and check them in the Speakers’ Ready Room at least 30 minutes prior to the start of your session.

3. If using a Powerpoint (or any other computer) presentation, please note you need to bring it on a floppy disk (3.5”/1.44 MB) or on a CD (no ZIP disk !!) and load it on the computer in the session hall, at least 30 minutes before the start of the session (during breaks).

You may supply your own laptop computer as a back-up. If using a MacIntosh laptop computer, please confirm that it has a VGA socket for external signal and come to check it in the session hall, at least 30 minutes before the start of the session (during breaks).

4. If using video, please ensure that it is VHS multi-system format. Please check it in the session hall, at least 30 minutes before the start of the session (during breaks).
SOCIAL EVENTS

Sunday, May 5, 2002

19:30  Get Together Reception

(included in the registration fees of participants and accompanying persons).

Wednesday, May 8, 2002

21:00  Farewell Dinner

(included in the registration fees of participants and accompanying persons)

ACCOMPANYING PERSONS' PROGRAM

All registered accompanying persons are invited to the Get Together Reception on Sunday and the Farewell Dinner on Wednesday, as well as to the following half-day tour:

Tuesday, May 7, 2002

PONTRESINA

Enjoy a scenic drive from St. Moritz to Pontresina where you will visit the interesting Alpine Museum of mountaineering, hiking and the history of alpinism.

Afterwards, take a leisurely walk around this typically swiss and picturesque village whilst enjoying the breathtaking surrounding natural scenery. Return to St. Moritz will be by horse-drawn sleigh - promised to be an unforgettable experience!

Departure for the tour from the hotel lobby at 09:00 and approximate time of return is 13:00.
XIIth INTERNATIONAL
SYMPOSIUM ON
CHOLINERGIC MECHANISMS-
FUNCTION AND DYSFUNCTION
&
2nd MISRAHI SYMPOSIUM ON
NEUROBIOLOGY

SCIENTIFIC PROGRAM
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MONDAY, MAY 6, 2002

08:00 – 08:30
WELCOME AND GREETINGS

08:30 – 09:20
PLENARY SESSION 1: HEILBRONN LECTURE
Chairperson: V.P. Whittaker, UK

08:30 ACETYLCHOLINE RECEPTORS: PROBING FUNCTIONALLY SIGNIFICANT STRUCTURAL CHANGES WITH SITE-DIRECTED REACTIONS
A. Karlin, USA

09:20 - 10:40
SESSION 1
Chairperson: F.J. Barrantes, Argentina

09:20 THE CRYSTAL STRUCTURE OF MOLLUSCAN AChBP REVEALS THE LIGAND BINDING DOMAIN OF THE NICOTINIC ACETYLCHOLINE RECEPTOR

09:40 THE BINDING SITE OF THE ACETYLCHOLINE RECEPTOR: FROM SYNTHETIC PEPTIDES TO SOLUTION AND CRYSTAL STRUCTURE

10:00 STRUCTURAL STUDIES ON THE NICOTINIC ACETYLCHOLINE RECEPTOR
Y. Paas, France

10:20 MOLECULAR BASIS OF THE SELECTIVITY OF NEUROTOXINS TOWARDS nAChR SUBTYPES
D. Servent, C. Gaillard, B. Gilquin, S. Antil-Delbeke, P.J. Corringer, J.P. Changeux, A. Menez, France

10:40 Coffee Break and Poster Viewing

11:10 – 12:50
SESSION 2
Chairperson: D. Bertrand, Switzerland

11:10 ALLOSTERISM OF THE NICOTINIC ACETYLCHOLINE RECEPTOR
F. Hucho, G.M. Bixel, M. Krauss, Germany

11:30 STRUCTURE AND DYNAMICS OF ACETYLCHOLINE RECEPTOR AND ITS LIPID MICROENVIRONMENT: MOLECULE TO CELL
S. Antollini, J. Baier, M. Blanton, I. Bonini, B. De los Santos, M.C. Gallegos, I. Garbus, M.F. Pediconi, M. Prieto, A.M. Roccham, J. Wenz, F.J. Barrantes, Argentina

11:50 SITE-DIRECTED REACTIVE PROBES FOR STRUCTURAL AND FUNCTIONAL INVESTIGATION OF CHOLINERGIC PROTEINS
F. Kotzyba-Hibert, S. Lourdew, C. Che, T. Grutter, A. Specht, A. Mourot, M. Goeldner, France

12:10 NMR STRUCTURE OF ALPHA-BUNGAROTOXIN IN COMPLEX WITH AN ACHR
ALPHA-SUBUNIT PEPTIDE REVEALS THE BASIS FOR SPECIES SPECIFIC RESISTANCE TO THE TOXIN AND HOW ALPHA-NEUROTOXINS INHIBIT ACETYLCHOLINE BINDING TO THE RECEPTOR

12:30 ACTIVATION INHIBITION AND UPREGULATION OF THE HUMAN NEURONAL NICOTINIC ALPHA4/BETA2 RECEPTOR BY A PARTIAL AGONIST
V. Itier, D.C. Bertrand, Switzerland

12:50 Lunch Break and Poster Viewing
MONDAY, MAY 6, 2002

14:45 - 16:25

SESSION 3
Chairperson: S.C. Froehner, USA

14:45 ORGANIZATION OF CYTOSKELETON OF MUSCLE FIBERS BY MUSCLE ACTIVITY AND AGRIN
G. Bezakova, T.L. Lomo, Switzerland, Norway

15:05 THE ROLE OF PSY1 NUCLEOTIDE RECEPTOR IN THE FORMATION OF NEUROMUSCULAR
Junctions
K.W.K. Tsim, E.A. Barnard, Hong Kong, UK

15:25 THE DYSTROPHIN COMPLEX - A SCAFFOLD FOR SIGNALING PROTEINS AT SYNAPSES
M.E. Adams, S.C. Froehner, USA

15:45 AGRIN BLOCKADE IMPAIRS LATE BUT NOT INITIAL STAGES OF FUNCTIONAL INNERVATION OF
HUMAN MUSCLE IN VITRO
T. Mars, K. Mis, M.P. King, A.F. Miranda, Z. Grubic, Slovenia, USA

16:05 ORGANIZING ACETYLCHOLINESTERASE MOLECULES AT THE NEUROMUSCULAR SYNAPSE
R.L. Rotundo, S.G. Rossi, J.M. Quintero, L.M. Kimbell, USA

16:25 Coffee Break and Poster Viewing

17:00 - 18:20

SESSION 4
Chairperson: J. Massoulie, France

17:00 THE FUNCTIONAL HETERO-OLIGOMERIC FORMS OF CHOLINESTERASES
J. Massoulie, France

17:20 TRANSCRIPTIONAL AND POST-TRANSCRIPTIONAL EVENTS CONTROLLING EXPRESSION OF
ACETYLCHOLINESTERASE IN DEVELOPING AND ADULT MUSCLES
B.J. Jasmin, L.M. Angus, G. Belanger, J. Deschenes, F. Nassralia, Canada

17:40 DENSITY AND LOCALIZATION OF ACETYLCHOLINESTERASE IN VERTEBRATE NEUROMUSCULAR
JUNCTIONS
L. Anglister, Israel

18:00 FOUR ACETYLCHOLINESTERASE GENES IN THE NEMATODE CAENORHABDITIS ELEGANS
D. Combes, Y. Fedon, J-P. Toutant, M. Arpagaus, France

20:30 - 21:50

SESSION 5
Chairperson: H. Soreq, Israel

20:30 THE MOLECULAR NEUROBIOLOGY OF ACETYLCHOLINESTERASE VARIANTS: FROM STRESSFUL
INSULTS TO ANTISENSE INTERVENTION
H. Soreq, D. Glick, Israel

20:50 TRANSCRIPTIONAL CONTROL OF THE CHOLINERGIC GENE LOCUS (CGL): A MOSAIC MODEL FOR
REGULATION OF THE CHOLINERGIC PHENOTYPE
L.E. Eiden, B. Schuetz, M. Goeddes, C. Depboylu, M.K-H. Schafer, E. Weihe, USA, Germany

21:10 REGULATION OF CHOLINERGIC GENE EXPRESSION BY NRSF/REST
M. Shimojo, L.B. Hersh, USA

21:30 NEUROTRANSMITTER PHENOTYPE SWITCH IN DEVELOPING NEURONS – GENOMIC AND
PROTEOMIC APPROACHES
M. Linial, Y. Bledi, Y. Bogoch, Israel
Tuesday, May 7, 2002

08:30 - 09:20

Plenary Session 2: Couteaux Lecture

Chairperson: S. Tsuji, France

08:30 Wanderings in and about active site gorges and subunit interfaces
P. Taylor, B. Molles, J. Shi, S. Camp, Z. Radic, USA

09:20 - 10:40

Session 6

Chairperson: J.M. Gonzalez-Ros, Spain

09:20 Regulation of neurotransmitter release: calcium and ion channels

09:40 Macromolecular architecture of active zone material and its role in synaptic transmission
U.J. McMahan, USA

10:00 Persistent and transient inhibition of acetylcholine release from motor terminals by botulinum toxin A and E are due to their cleavage products (SNAP-25 1-197 and 1-180) having different life-times
J.O. Dolly, G.O. Lisk, F.A. Meunier, N. Mohammed, P.G.P. Foran, UK

10:20 Mediatorphore no longer an artefact
M. Israel, Y. Dunant, Switzerland, France

10:40 Coffee Break and Poster Viewing

11:10 - 12:50

Session 7

Chairperson: I. Parnas, Israel

11:10 Muscarinic presynaptic receptors are involved in the control of the time course of neurotransmitter release in frog and mouse neuromuscular junctions
I. Parnas, I. Slutsky, H. Parnas, Israel

11:30 Structural and functional conservation of snare complexes
D. Fasshauer, W. Antonin, M. Margittai, S. Pabst, R. Jahn, Germany

11:50 Genetic dissection of synaptic function in Drosophila
T.L. Schwarz, USA

12:10 Genetic regulation of cholinergic neurotransmitter phenotypes
P.M. Salvaterra, M-H. Lee, S. Song, USA

12:30 Signaling pathways that regulate the cholinergic gene locus expression
B. Berse, I. Lopez-Coviella, T. Mellott, B.E. Slack, M.T. Follett, R.S. Thies, L. Li, J.K. Blusztajn, USA

12:50 Lunch Break and Poster Viewing
TUESDAY, MAY 7, 2002

14:45 - 16:25

SESSION 8

Chairperson: **P. Marchot, France**

14:45 COMPARATIVE STRUCTURAL STUDIES ON CONJUGATES OF *TORPEDO CALIFORNICA* AND HUMAN ACETYLCHOLINESTERASES WITH ORGANOPHOSPHATE NERVE AGENTS

15:05 CRYSTAL STRUCTURE OF RECOMBINANT HUMAN BUTYRYLCHOLINESTERASE: NEW INSIGHTS INTO THE CATALYTIC MECHANISMS OF CHOLINESTERASES
Y. Nicolet, F. Nachon, P. Masson, O. Lockridge, J-C. Fontecilla-Camps, *France, USA*

15:25 INTRINSIC TRYPHTOPHAN FLUORESCENCE OF CHOLINESTERASES: DIRECT, NON-PERTURBING MONITORING OF ENZYME-LIGAND INTERACTIONS
Z. Radic, E. Kim, P. Taylor, *USA*

15:45 SURPRISING FINDINGS FROM THE FUNCTIONAL ANALYSIS OF HUMAN ACETYLCHOLINESTERASE ADDUCTS OF ALZHEIMER'S DISEASE DRUGS
A. Ordentlich, C. Kronman, D. Barak, N. Ariel, D. Kaplan, B. Velan, A. Shaffer, *Israel*

16:05 UNFOLDING AND FOLDING OF *TORPEDO CALIFORNICA* ACETYLCHOLINESTERASE

16:25 *Coffee Break and Poster Viewing*

17:00 - 18:20

SESSION 9

Chairperson: **P.Y. Masson, France**

17:00 SCAVENGER PROTECTION AGAINST ORGANOPHOSPHATES BY CHOLINESTERASES
B.P. Doctor, A. Saxena, M.T. Clark, Y. Rosenberg, D.M. Maxwell, D.E. Lenz, Y. Ashani, *USA, Israel*

17:20 A COMPLEX ARRAY OF POST-TRANSLATION MODIFICATIONS DETERMINES THE CIRCULATORY LONGEVITY OF ACETYLCHOLINESTERASE IN A HIERARCHICAL MANNER
C. Kronman, T. Chitlaru, A. Ordentlich, B. Velan, A. Shaffer, *Israel*

17:40 PRESYNAPTIC INHIBITION OF CENTRAL ACETYLCHOLINE RELEASE WITH A1 LIGANDS: PREVENTION OF CHOLINERGIC CRISIS
H.P.M. van Helden, T.J.H. Bueters, B. Groen, M. Danhof, A.P. Ijzerman, *The Netherlands*

18:00 POLYURETHANE IMMOBILIZED ENZYMES: OP SENSING AND DECONTAMINATING MATRIXES
WEDNESDAY, MAY 8, 2002

08:30 - 09:20

PLENARY SESSION 3: SALPETER LECTURE
Chairperson: U.J. McMahen, USA

08:30 CHEMICAL KINETICS PARAMETERS AND RECEPTOR DEGRADATION RATES AT THE NEUROMUSCULAR JUNCTION
E.E. Salpeter, USA

09:20 - 10:40

SESSION 10
Chairperson: S. Fuchs, Israel

09:20 CONGENITAL MYASTHENIC SYNDROMES (CMS): MULTIPLE MOLECULAR TARGETS AT THE NEUROMUSCULAR JUNCTION
A.G. Engel, K. Ohno, S.M. Sine, USA

09:40 ANTIBODIES TO ACETYLCHOLINE RECEPTORS AND MUSK IN MYASTHENIA GRAVIS AND RELATED DISORDERS
A. Vincent, UK

10:00 INHIBITORY EFFECTS OF MUSCARINIC RECEPTOR AUTOANTIBODIES ON PARASYMPATHETIC NEUROTRANSMISSION IN SJÖGREN’S SYNDROME
S.A. Waterman, S. Lester, T.P. Gordon, M. Rischmueller, Australia

10:20 IMMUNOTHERAPY OF MYASTHENIA GRAVIS: ANTIGEN-SPECIFIC MUCOSAL TOLERANCE AND ANTAGONISTS OF KEY CYTOKINES AND COSTIMULATORY FACTORS
M.C. Souroujon, S-H. Im, S. Fuchs, Israel

10:40 THE ALPHA7 NACHR L250T MUTATION IN MICE: A MODEL FOR EPILEPSY IN MEN
A. Orr-Urtreger, R.A. Sack, M. Kedmi, A. Harmelin, Z. Gil, Israel

Coffee Break and Poster Viewing

11:10 - 12:50

SESSION 11
Chairperson: N.C. Inestrosa, Chile

11:10 NEURODEGENERATIVE PROCESSES IN ALZHEIMER’S DISEASE
N.C. Inestrosa, G.V. De Ferrari, J.L. Garrido, A. Alvarez, M. Bronfman, Chile

11:30 ACETYLCHOLINESTERASE FACILITATES AMYLOID DEPOSITION IN A MOUSE MODEL OF ALZHEIMER’S DISEASE
T. Rees, P. Hammond, S. Younkin, H. Soreq, S. Brimijoin, USA, Israel

11:50 ACETYLCHOLINE HYDROLYSIS AT THE MAMMALIAN SKELETAL NEUROMUSCULAR JUNCTION: MORE THAN ONE ENZYME
E. Krejci, J. Minic, J. Molgo, France

12:10 MOLECULAR AND FUNCTIONAL DIVERSITY IN NICOTINIC ACETYLCHOLINE RECEPTOR GENE FAMILIES OF C. ELEGANS AND D. MELANOGASTER
D.B. Sattelle, UK

12:30 GENETIC DISSECTION OF AN ACETYLCHOLINE RECEPTOR INVOLVED IN NEURONAL DEGENERATION
M. Treinin, S. Halevi, L. Yassin, Israel

12:50 Lunch Break and Poster Viewing
SESSION 12

Chairperson: V.I. Tsetlin, Russia

14:15 NEURONAL NICOTINIC RECEPTORS, ALLOSTERIC POTENTIATING LIGANDS (apls), AND ENDOGENOUS METABOLITES: IMPLICATIONS FOR TREATMENT OF ALZHEIMER'S DISEASE (AD)
E.X. Albuquerque, M.D. Santos, M. Alkondon, E.K. Moon, A. Maelicke, USA, Brazil, Germany

14:35 THE RATIONALE FOR USING GALANTAMINE TO TREAT DIFFERENT DEMENTIA TYPES
A. Maelicke, S. Lilienfeld, C. Grantham, Germany, USA, Belgium

14:55 PREVENTION OF APOPTOSIS BY GALANTAMINE: A NOVEL THERAPEUTIC STRATEGY FOR ALZHEIMER'S DISEASE
M. Garcia-Lopez, Spain

15:15 PRECLINICAL STUDIES OF GALANTAMINE USING A FORM OF ASSOCIATIVE LEARNING SEVERELY IMPAIRED IN ALZHEIMER'S DISEASE
D. Woodruff-Pak, USA

15:35 A COMMON AGONIST AND POTENTIATOR FOR ALPHA7 NICOTINIC AND 5-HT3 SEROTONIN RECEPTORS
R. Zwart, L. Broad, C. Felthouse, K. Pearson, G. McPhie, E. Sher, UK

15:55 Coffee Break and Poster Viewing

SESSION 13

Chairperson: U.Z. Littauer, Israel

16:30 MOLECULAR BIOLOGY OF ALZHEIMER'S DISEASE – THERAPEUTIC PERSPECTIVES
R. Nitsch, Switzerland

16:50 BETA-AMYLOIDS, TAU HYPERPHOSPHORYLATION AND COGNITION ARE BENEFICIALLY AFFECTED BY M1 MUSCARINIC AGONISTS - PERSPECTIVES IN ALZHEIMER'S DISEASE TREATMENT

17:10 CHOLINERGIC DEFICITS AND NON-COGNITIVE BEHAVIOURAL CHANGES IN PATIENTS WITH DEMENTIA
P. Francis, C.P.L-H. Chen, M.M. Esiri, J. Keene, UK

17:30 CROSS-TALK BETWEEN APOLIPOPROTEIN E THEAMYLOID PRECURSOR PROTEIN AND BRAIN INFLAMMATION
D.M. Michaelson, S. Meilin, G. Ophir, Y. Ezra, L. Oron, S.M. Beni, E. Shohami, Israel
THURSDAY, MAY 9, 2002

08:30 - 09:20

PLENARY SESSION 4: BRZIN LECTURE
Chairperson:  E. Reiner, Croatia

08:30  ACETYLCHOLINESTERASE REGULATION IN SKELETAL MUSCLES
J. Sketelj, N. Crne-Finderle, P. Pregelj, Slovenia

09:20 - 10:40

SESSION 14
Chairperson:  U. Drews, Germany

09:20  CHOLINERGIC CORTICAL TERMINATIONS ESTABLISH CLASSICAL SYNAPSES AND UNDERGO AGE-RELATED ATROPHY
A.C. Cuello, P. Turrini, M.A. Casu, T.P. Wong, Y. De Koninck, A. Ribeiro Da Silva, Canada

09:40  DEVELOPMENT OF CHOLINERGIC PROJECTIONS TO CORTEX: POSSIBLE ROLE OF NEUROTROPHINS IN TARGET SELECTION.
R. Robertson, J. Yu, USA

10:00  ACTIVATION OF THE CHOLINERGIC SYSTEM DURING COGNITIVE PROCESSES
G. Pepeu, M.G Giovannini, M. Pazzagli, J. Cangioli, M.B. Passani, Italy

10:20  PRECLINICAL AND CLINICAL STUDIES ON THE ROLE OF MUSCARINIC RECEPTORS IN THE PHARMACOTHERAPY OF SCHIZOPHRENIA
F.P. Bymaster, A. Shekhar, K.W. Perry, K. Rasmussen, D. McKinzie, C.C. Felder, USA

10:40  Coffee Break and Poster Viewing

11:10 - 12:50

SESSION 15
Chairperson:  K. Loffelholz, Germany

11:10  CENTRAL CHOLINERGIC NEURONS IN CULTURE: REGULATION OF SURVIVAL AND FUNCTION
M. Segal, N. Landman, V. Greenberger, Israel

11:30  HOW IS THE BRAIN SUPPLIED WITH CHOLINE, BUT PROTECTED AGAINST EXCESS CHOLINE?
K. Loffelholz, J. Klein, Germany

11:50  MEASURING CEREBRAL ACETYLCHOLINE ESTERASE ACTIVITY IN ALZHEIMER DEMENTIA BY PET FUNCTIONAL PARAMETRIC IMAGING
K. Herholz, G. Zundorf, B. Bauer, S. Weisenbach, W-D. Heiss, Germany

12:10  BLOOD-BRAIN BARRIER DISRUPTION IS ASSOCIATED WITH ABNORMAL CORTICAL THETA RHYTHM GENERATION: THE POTENTIAL INVOLVEMENT OF ACETYLCHOLINESTERASE
E. Aviv, I. Shelef, H. Golan, A. Korn, O. Tomkins, L. Pavlovsky, A. Friedman, Israel

12:30  CONTROL OF ACETYLCHOLINE RELEASE UNDER STIMULATORY CONDITIONS BY ITS BIOSYNTHETIC PRECURSORS: GLUCOSE AND CHOLINE
J. Klein, S. Kopf, K. Loffelholz, Germany

12:50  Lunch Break and Poster Viewing
THURSDAY, MAY 9, 2002

14:45 - 16:25

SESSION 16

Chairperson: A. Enz, Switzerland

14:45  TREATMENT OF DEMENTIA WITH CHOLINESTERASE INHIBITORS
A.D. Korczyn, Israel

15:05  CHOLINESTERASE INHIBITORS STABILIZE COGNITIVE DECLINE IN ALZHEIMER'S DISEASE
E. Giacobini, Switzerland

15:25  GENDER DIFFERENCES IN THE ACTIONS OF CHOLINESTERASE INHIBITORS
M. Weinstock, R-H. Wang, Israel

15:45  Amyloid Precursor Protein Processing Properties of the Novel Neuroprotective Cholinesterase Monoamine Oxidase Inhibitor, TV326 and Its Optical Isomer, TV3279
M.B.H. Youdim, M. Phalach-Yogev, O. Bar-Am, M. Weinstock, T. Amit, Israel

16:05  Novel Bifunctional Compounds Eliciting Cholinergic and Anti-Inflammatory Activity for the Treatment of CNS Impairments
G. Amitai, R. Adani, I. Rabinovitz, G. Sod-Moriah, H. Meshulam, Israel

16:25  Coffee Break and Poster Viewing

17:00 - 18:20

SESSION 17

Chairperson: S. Tucek, Czech Republic

17:00  Roles of External Loops of Muscarinic Receptors in Interactions Between N-Methylscopolamine and Allosteric Modulators
A. Krejci, S. Tucek, Czech Republic

17:20  Structure and Activation of Muscarinic Acetylcholine Receptors
E.C. Hulme, Z-L. Lu, M.S. Bee, C.A.M. Curtis, UK

17:40  Generation and Analysis of Muscarinic Acetylcholine Receptor Knockout Mice
A. Duttaroy, M. Yamada, J. Gomez, W. Zhang, R. Makita, T. Miyakawa, F. Bymaster, C. Felder, C. Deng, J. Wess, USA, Japan

18:00  Closing Remarks
A.G. Karczmar, USA
XIth INTERNATIONAL SYMPOSIUM ON CHOLINERGIC MECHANISMS-FUNCTION AND DYSFUNCTION & 2nd MISRAHI SYMPOSIUM ON NEUROBIOLOGY

POSTERS
MONDAY, MAY 6, 2002

LIST OF POSTERS 1

1  EFFECT OF THYROID HORMONE ON ACETYLCHOLINESTERASE MRNA LEVELS IN THE SLOW SOLEUS AND FAST EDL MUSCLE OF THE RAT
   P. Pregelj, J. Sketelj, Slovenia

2  ACETYLCHOLINESTERASE mRNA EXPRESSION IN RAT SPINAL CORD
   K. Mis, E. Davidson, H. Park, M.P. King, T. Mars, Z. Grubic, Slovenia, USA

3  NEUROMUSCULAR JUNCTION FORMED IN CO-CULTURE WITH EMBRYONIC SPINAL CORD IS ACCOMPANIED BY CO-DIFFERENTIATION OF NEURONAL AND GLIAL CELLS
   T. Mars, K.J. Yu, X. Tang, A.F. Miranda, Z. Grubic, F. Cambi, M.P. King, Slovenia, USA

4  FUNCTIONAL EXPRESSION AND STOICHIOMETRY OF THE NOVEL HUMAN ALPHA9ALPHA10 HETEROMERIC NICOTINIC ACETYLCHOLINE RECEPTOR

5  LARGE SCALE EXPRESSION OF THE EXTRACELLULAR AND CYTOPLASMIC DOMAINS OF THE DROSOPHILA ADHESION PROTEIN, GLIOTACTIN

6  STRUCTURAL REORGANIZATION OF THE ACETYLCHOLINE BINDING SITE OF TORPEDO NICOTINIC RECEPTOR REVEALED BY DYNAMIC PHOTOAFFINITY LABELING
   T. Grutter, F. Kotzyba-Hibert, S. Bertrand, D. Bertrand, M. Goeldner, Switzerland

7  COEXPRESSION OF ALPHA10 AND ALPHA9 NICOTINIC ACETYLCHOLINE RECEPTORS IN RAT DORSAL ROOT GANGLION NEURONS
   K.S. Lips, U. Pfeil, R.V. Haberberger, W. Kummer, Germany

8  MAPPING THE ACETYLCHOLINE BINDING SITES OF TORPEDO NICOTINIC RECEPTOR USING PHOTOAFFINITY LABELING: PAST, PRESENT AND FUTURE
   F. Kotzyba-Hibert, A. Mourot, T. Grutter, M. Goeldner, France

9  NFkB REGULATES THE ACTIVITY OF HUMAN ACETYLCHOLINESTERASE PROMOTER IN MUSCLE

10 THE INITIAL BINDING OF ACETYLCHOLINESTERASE AND PERLECAN OCCURS INSIDE THE CELL PRIOR TO EXTERNALIZATION
    S.G. Rossi, R.L. Rotundo, USA

11 ROLE OF MDX NERVE AND MUSCLE IN REGULATING NEUROMUSCULAR JUNCTION PROPERTIES: A STUDY USING MUSCLE TRANSPLANTS
    A.R. Durrant, M. Szabo, L. Anglister, M.M. Salpeter, USA, Israel

12 CRYSTAL STRUCTURE OF THE TETRAMERIZATION DOMAIN OF ACETYLCHOLINESTERASE AT 2.3A RESOLUTION
    M. Harel, H. Dvir, S. Bon, W.Q. Liu, C. Garbay, J.L. Sussman, J. Massoulie, I. Silman, Israel, France

13 SCALING UP OF PRODUCTION, PURIFICATION AND REFOLDING OF A CHIMERIC THREE-FINGERED TOXIN WITH SPECIFICITY FOR ACETYLCHOLINESTERASE

14 ROLE OF SPONTANEOUS MUTATIONS OF NEURONAL NICOTINIC RECEPTORS IN ADNFLE
    D. Bertrand, I. Favre, H. Phillips, S. Bertrand, S.F. Berkovic, J.C. Mulley, Switzerland, Australia

15 EXPRESSION OF COLQ AT THE NEUROMUSCULAR JUNCTION
    C. Legay, France
<table>
<thead>
<tr>
<th>No.</th>
<th>Title</th>
<th>Authors</th>
</tr>
</thead>
<tbody>
<tr>
<td>17</td>
<td>CLONING AND CHARACTERIZATION OF ACETYLCHOLINESTERASE GENE IN CHICKEN</td>
<td>X. Zhang, K.W.K. Tsim, D.C.C. Wan, Hong Kong</td>
</tr>
<tr>
<td>18</td>
<td>CANDOXIN A NEW SNAKE TOXIN SPECIFIC FOR THE ALPHA7 nAChR</td>
<td>E. Charpantier, S. Nirthanan, P. Gopalakrishnakone, M.C.E. Gwee, H.E. Khoo, L.S. Cheah, R. Manjunatha Kini, D. Bertrand, Switzerland, Singapore</td>
</tr>
<tr>
<td>19</td>
<td>MUSCARINIC RECEPTOR REGULATION OF EVOKE ACETYLCHOLINE RELEASE IS AFFECTED BY ACETYLCHOLINESTERASE INACTIVATION AT THE MOUSE NEUROMUSCULAR JUNCTION</td>
<td>J. Minic, J. Molgo, E. Krejci, France</td>
</tr>
<tr>
<td>21</td>
<td>NICOTINIC ACETYLCHOLINE RECEPTOR AND MUSK ARE CLUSTERED IN C2C12 CELLS VIA LIPIDIC RAFTS</td>
<td>F. Stetzkowskil-Marden, S. Marchand, J. Cartaud, France</td>
</tr>
<tr>
<td>22</td>
<td>IDENTIFICATION OF SPECIES DIFFERENCES IN THE PHARMACOLOGY OF THE ALPHA-7 NICOTINIC RECEPTOR USING THE ANTAGONIST RADIOLIGAND [3H]-METHYLLYCACONITINE</td>
<td>N. Crawford, K. Finlayson, J. Sharkey, J.S. Kelly, UK</td>
</tr>
<tr>
<td>23</td>
<td>ACETYLCHOLINESTERASE IS REQUIRED FOR NEURONAL AND MUSCULAR DEVELOPMENT IN ZEBRAFISH</td>
<td>M. Behra, X. Cousin, C. Bertrand, J.L. Vonesch, A. Chatonnet, U. Strehle, France</td>
</tr>
<tr>
<td>24</td>
<td>THE 14.3.3 GAMMA PROTEIN IS PART OF THE MUSK SIGNALING COMPLEX AT THE NEUROMUSCULAR JUNCTION</td>
<td>L. Strochlic, A. Cartaud, M. Recouveure, R. Graihe, J.P. Changeux, J. Cartaud, France</td>
</tr>
<tr>
<td>25</td>
<td>CONSTRUCTION AND CHARACTERISATION OF A CHIMERIC HUMAN ALPHA 7 NICOTINIC ACETYLCHOLINE / MOUSE 5HT3 RECEPTOR</td>
<td>P.J. Craig, R. Zwart, S. Bose, R.E. Beattie, E.A. Folly, L.R. Johnson, E. Bell, N.M. Evans, S.G. Volsen, E. Sher, N.S. Miller, L.M. Broad, UK</td>
</tr>
<tr>
<td>26</td>
<td>ALPHA-CONOTOXINS PnIA AND A10L-PnIA STABILISE DIFFERENT STATES OF THE CHICK NEURONAL ALPHA 7 ACETYLCHOLINE RECEPTOR</td>
<td>R.C. Hogg, S. Bertrand, P.F. Allowood, D.J. Adams, D.C. Bertrand, Australia, Switzerland</td>
</tr>
<tr>
<td>28</td>
<td>RAPSYN ESCORTS THE NICOTINIC ACETYLCHOLINE RECEPTOR ALONG THE EXOCYTIC PATHWAY VIA THE ASSOCIATION WITH LIPID RAFTS</td>
<td>S. Marchand, A. Devillers-Thiry, S. Pons, J-P. Changeux, J. Cartaud, France</td>
</tr>
<tr>
<td>29</td>
<td>IDENTIFICATION AND CHARACTERIZATION OF A DIVERSE FAMILY OF NEUROTOXIN-LIKE PEPTIDES FROM THE SOUTH AMERICAN CORAL SNAKE</td>
<td>T. Kubo, G. Baptista, X. Yang, S. Kobayashi, M. Takeda, A. Prieto-Da-Silva, T. Yamane, Japan, Brazil</td>
</tr>
<tr>
<td>30</td>
<td>MODELS OF THE EXTRACELLULAR DOMAIN OF THE NICOTINIC RECEPTORS AND OF AGONIST AND CA++ BINDING SITES</td>
<td>N. Le Novere, T. Guettier, J-P. Changeux, France</td>
</tr>
<tr>
<td>31</td>
<td>ORIGIN OF ACETYLCHOLINESTERASE IN THE DEVELOPING NEUROMUSCULAR JUNCTION</td>
<td>M. Jevsek, T. Mars, Z. Grubic, Slovenia</td>
</tr>
</tbody>
</table>
LIST OF POSTERS 1

32 PROBING THE BINDING INTERFACE BETWEEN THE NICOTINIC ACETYLCHOLINE RECEPTOR AND A SHORT ALPHA-NEUROTOXIN THROUGH RECEPTOR-BIOTINYLTEOXIN-STREPTAVIDIN TERNARY COMPLEXES
F. Teixeira, A. Menez, P. Kessler, France

33 THE AGONIST BIPHASIC DOSE-RESPONSE CURVE OF THE HUMAN ALPHA4BETA2 RECEPTOR BECOMES MONOPHASIC IN THE PRESENCE OF PKC INHIBITORS OR IN LOW LEVELS OF EXTRACELLULAR CALCIUM IONS
I Bermudez, L.M. Houlihan, UK

34 NFkB REGULATES THE ACTIVITY OF HUMAN ACETYLCHOLINESTASE PROMOTER IN MUSCLE

35 MOLECULAR CLONING OF NICOTINIC ACETYLCHOLINE RECEPTOR SUBUNIT GENES FROM THE PEACH-POTATO APHID, MYZUS PERSICAE
M. Kirwan, Y. Huang, M.S. Williamson, A.L. Devonshire, J.D. Windass, S. Dunbar, S.J. Lansdell, N.S. Millar, UK

36 NEW ESSENTIAL RESIDUES IN CHOLINESTERASE ACYL POCKET
S.N. Moralev, Russia

37 VARIABILITY OF SUBSTRATE SPECIFICITY IN CHOLINESTERASES OF VERTEBRATES AND INVERTEBRATES
E.V. Rozengart, S.N. Moralev, Russia
TUESDAY, MAY 7, 2002

LIST OF POSTERS 2

1. 2-AMINOPERIMIDINE IS AN EFFECTOR OF CHOLINESTERASES
   Y. Shalitin, D. Segal, D. Gur, *Israel*

2. PERIPHERAL BINDING OF ETHOPROPAZINE TO HORSE SERUM BUTYRYLCHOLINESTERASE
   E. Reiner, G. Sinko, A. Stuglin, V. Simeon-Rudolf, *Croatia*

3. SOME CONSIDERATIONS AS TO THE MOLECULAR MECHANISM OF CHOLINESTERASE CATALYSIS
   N.B. Brovtsyna, E.V. Rozengart, A.A. Suvorov, S.N. Moralev, *Russia*

4. PHOSPHONYLATION OF ACETYLCHOLINESTERASE AND THE PROPENSITY FOR REACTIVATION
   ANALYZED BY CHIRALITY AND MUTAGENESIS
   Z. Kovarik, Z. Radic, H.A. Berman, P. Taylor, *Croatia, USA*

5. THE FIRST TWO NATURALLY OCCURRING ACTIVATORS / REACTIVATORS OF
   ACETYLCHOLINESTERASE
   R. Gupta, S.S. Thakur, *India*

6. KINETICS OF INTERACTION OF ETHOPROPAZINE ENANTIOMERS WITH
   BUTYRYLCHOLINESTERASE AND ACETYLCHOLINESTERASE
   G. Sinko, Z. Radic, V. Simeon-Rudolf, E. Reiner, P. Taylor, *Croatia, USA*

7. REACTIVATION STUDY INDICATES THAT THE ORIENTATIONS OF HI-6 MAY DIFFER IN
   REACTivating ACETYLCHOLINESTERASE INHIBITED WITH ORGANOPHOSPHATES AND
   ORGANOPHOSPHONATES
   C. Luo, A. Saxena, H. Leader, Z. Radic, D.M. Maxwell, P. Taylor, B.P. Doctor, *USA*

8. KINETIC AND X-RAY CRYSTALLOGRAPHY STUDIES ON THE INTERACTION OF CHOLINESTERASES
   WITH THE ANTI-ALZHEIMER DRUG RIVASTIGMINE

9. ATTEMPTS TO ENGINEER AN ENZYME-MIMIC OF BUTYRYLCHOLINESTERASE BY SUBSTITUTION
   OF THE SIX DIVERGENT AROMATIC AMINO ACIDS IN THE ACTIVE CENTER OF
   ACETYLCHOLINESTERASE
   D. Kaplan, A. Ordentlich, D. Barak, N. Ariel, C. Kronman, B. Velan, A. Shafferman, *Israel*

10. MALDI-TOF/MS ANALYSIS OF ACETYLCHOLINESTERASE-LIGAND CONJUGATES: A TOOL FOR
    RESOLUTION OF MECHANISTIC PATHWAYS
    E. Elhanani, A. Ordentlich, O. Dgany, D. Kaplan, Y. Segall, R. Barak, B. Velan,
    A. Shafferman, *Israel*

11. INFLUENCE OF WATER ON THE FUNCTION OF ACETYLCHOLINESTERASE
    R.H. Henchman, K. Tai, T. Shen, J.A. McCammon, *USA*

12. THE FLUCTUATING SYNAPSE
    K. Kaufmann, Germany

13. X-RAY STRUCTURE OF TORPEDO ACHE COMPLEXED WITH BIFUNCTIONAL LIGANDS RELATED
    TO HUPA: NOVEL DRUGS FOR TREATMENT OF ALZHEIMER'S DISEASE
    D.W. Wong, H.M. Greenblatt, D. Shaya, P.R. Carlier, Y.-P. Pang, Y.-F. Han, I. Silman,
    J.L. Sussman, *Israel, USA, Hong Kong*

14. CRYSTALLIZATION AND DETERMINATION OF THE X-RAY STRUCTURE OF HUMAN ACHE
    H. Dvir, G. Kryger, J.L. Johnson, T.L. Rosenberry, I. Silman, J.L. Sussman, *Israel, USA*

15. 3D STRUCTURE OF TORPEDO CALIFORNICA ACETYLCHOLINESTERASE COMPLEXED WITH
    HUPRINE X
    J.L. Sussman, *Israel, Spain, USA*
TUESDAY, MAY 7, 2002 (continued)

LIST OF POSTERS 2

16 LIGAND INDUCED CONFORMATIONAL CHANGES IN THE OMEGA LOOP OF ACETYLCOLINESTERASE REVEALED BY FLUORESCENCE SPECTROSCOPY
J. Shi, Z. Radic, A. Boyd, P. Taylor, USA

bis-ACTING GALANTHAMINE DERIVATIVES AS IMPROVED DRUGS IN THE SYMPTOMATIC TREATMENT OF ALZHEIMER'S DISEASE
H.M. Greenblatt, C. Guillou, B. Badet, C. Thal, I. Silman, J.L. Sussman, Israel, France

18 COMPARISON OF TWO REACTION SCHEMES FOR THE HYDROLYSIS OF ACETYLTHIOCHOLINE BY BUTYRYLCHOLINESTERASE
V. Simeon-Rudolf, G. Sinko, A. Stuglin, J. Stojan, M. Golicnik, E. Reiner, Croatia, Slovenia

19 QUANTAL ACETYLCHOLINE RELEASE THROUGH MEDIATOPHORE PROTEOLIPID OVER-EXPRESSED IN NEUROBLASTIC CELLS
A. Bloc, J. Falk-Vairant, M. Malo, M. Israel, Y. Dunant, Switzerland, France

20 STIMULATION OF NICOTINIC RECEPTORS INDIRECTLY INCREASES ACETYLCHOLINE RELEASE IN RAT STRIATUM
V. Dolezal, V. Zemlickova, S. Tucek, Czech Republic

21 CAPILLARY ZONE ELECTROPHORESIS DETECTS UNWANTED CHOLINESTERASE-BOUND HIDDEN LIGANDS THAT MODULATE ENZYME CONFORMATIONAL STABILITY
D. Rochu, F. Renault, C. Bon, P. Masson, France

22 CRYSTAL STRUCTURE OF TORPEDO CALIFORNICA ACETYLCHOLINESTERASE WITH A NOVEL GALANTHAMINE DERIVATIVE: IMPLICATIONS FOR THE DESIGN OF NEW ANTI-ALZHEIMER DRUGS
M.C. Siotto, C. Bartolucci, D. Lamba, Italy

23 X-RAY STRUCTURE OF SOMAN-AGED HUMAN BUTYRYLCHOLINESTERASE
F. Nachon, Y. Nicolet, P. Masson, J-C. Fontecilla-Camps, O. Lockridge, France, USA

24 STUDIES ON DYNAMICAL TRANSITIONS IN CHOLINESTERASES

25 TETANIC FADE IS REVEALED BY BLOCKING PRESYNAPTIC NICOTINIC RECEPTORS CONTAINING ALFA4BETA2 AND ALFA3BETA2 SUBUNITS AFTER REDUCING THE SAFETY FACTOR OF NEUROMUSCULAR TRANSMISSION
M. Faria, L. Oliveira, M.A. Timoteo, M.G.B. Lobo, P. Correia-de-Sa, Portugal

26 MOLECULAR CHARACTERISATION OF ACETYLCHOLINESTERASE FROM THE PEACH-POTATO APHID MYZUS PERSICAE(SULZ.)
M.C. Andrews, C.G. Bass, M.S. Williamson, G.D. Moores, UK

27 PECULIARITIES OF KINETIC BEHAVIOUR OF FISH S/ABRAMIS BALLERUS/ BLOOD SERUM CHOLINESTERASE
V.D. Tonkopii, Russia

28 STRUCTURAL INSIGHTS INTO THE INTERACTIONS AT THE ACETYLCHOLINESTERASE PERIPHERAL ANIONIC SITE
Y. Bourne, P. Taylor, H.A. Berman, Z. Radic, P. Marchot, France, USA

29 LETHAL EFFECTS OF HEAD-TO-TAIL 3-ALKYLPYRIDINIUM POLYMERS ISOLATED FROM THE MARINE SPONGE RANIERA SARAI: ACHE INHIBITION OR UNSPECIFIC BINDING TO SERUM PROTEINS?
M. Bunc, K. Sepcic, A. Rotter, T. Turk, A. Vidmar, D. Suput, Slovenia

30 PIPERONYL BUTOXIDE: A SPECIFIC INHIBITOR OF INSECTICIDE RESISTANT ACETYLCHOLINESTERASE
R.V. Gunning, Australia

31 SIGNIFICANCE OF PARAMETERS BETWEEN VARIOUS KINETIC SCHEMES FOR CHOLINESTERASES
J. Stojan, M. Golicnik, Slovenia
LIST OF POSTERS 2

32 STUDIES OF ACETYLCHOLINESTERASE FROM THE PEACH-POTATO APHID, *MYZUS PERSICAE* (SULZ.)
N. Javed, M.S. Williamson, A.L. Devonshire, R.C. Viner, T. Lewis, G.D. Moorses, UK

33 EXPLORING THE ACHE GORGE WITH GALANTHAMINE
G. Fels, E. Linnemann, E. Luttmann, C. Pilger, Germany

34 HYSTERESIS IN BUTYRYLCHOLINESTERASE CATALYSIS: EVIDENCE FOR SUBSTRATE-INDUCED
CONVERSION OF THE ENZYME FROM LATENT TO OPERATIVE FORM
P-Y. Masson, M-T. Froment, F. Nachon, L.M. Schöpfer, France, USA

35 A CALCIUM-PROTON ANTIPORT IN CHOLINERGIC AND GLUTAMATERGIC SYNAPTIC VESICLES
M. Cordeiro, V. Bancila, A. Bloc, Y. Dunant, Switzerland

36 OXACHEIN, A NOVEL POTENT INHIBITOR OF ACETYLCHOLINESTERASE FROM A PLANT—*OXALIS CORNICULATA* L.
R. Gupta, A. Gupta, India

37 MECHANISM OF ACETYLCHOLINESTERASE INHIBITION BY FASCICULIN: A 5 NANOSECOND
MOLECULAR DYNAMICS SIMULATION
K. Tai, T. Shen, R.H. Henchman, Y. Bourne, P. Marchot, J.A. McCammon, USA, France

38 PROF. RENE COUTEAUX AND HIS PRESYNAPTIC ‘ACTIVE ZONE’
S. Tsuji, France

26
LIST OF POSTERS 3

1. MITOGEN-ACTIVATED PROTEIN KINASE KINASE INHIBITS CILIARY NEUROTROPHIC FACTOR-ACTIVATED CHOLINE ACETYLTRANSFERASE GENE EXPRESSION
   T. Mellott, I. Lopez-Coviella, J.K. Blusztajn, B. Berse, USA

2. EFFECTS OF METHOMYLL ON SPELENS AND APOPTOSIS

3. PLANT CHOLINESTERASE ACTIVITY AS A BIOSENSOR FOR TOXINS IN THE ENVIRONMENT
   V.V. Roshchina, Russia

4. EFFECTS OF CARBAMATE INSECTICIDES ON RAT NEURONAL ALPHA4BETA4 NICOTINIC RECEPTORS AND RAT BRAIN ACETYLCHOLINESTERASE
   C.J.G.M. Smulders, T.J.H. Bueters, H.P.M. Vijverberg, The Netherlands

5. SITE-SPECIFIC ANALYSIS OF GLYCAN STRUCTURES ON PLASMA-DERIVED HUMAN (Hu) AND HORSE (Eq) BUTYRYLCHOLINESTERASES (BchE)
   G.E. Garcia, D.R. Moorad-Doctor, G. Lockridge, C.B. Millard, C.A. Broomfield, USA

6. RECOVERY FROM DESENSITIZATION OF A NEURONAL NICOTINIC RECEPTOR
   S. Voytenko, R.J. Lukas, R. Gruener, USA

7. MECHANISM AND STRUCTURAL REQUIREMENTS OF XANOMELINE WASH-RESISTANT BINDING TO M1 MUSCARINIC RECEPTORS
   J. Jakubik, E. El-Fakahany, S. Tucek, Czech Republic, USA

8. IS THE G-PROTEIN-COUPLED M2 MUSCARINIC RECEPTOR A VOLTAGE SENSOR?
   O. Tour, N. Dascal, Y. Ben Chaim, P. Parnas, H. Parnas, USA, Israel

9. SOME BASIC RULES GOVERNING OLIGOSACCHARIDE-DEPENDENT CIRCULATORY RESIDENCE OF GLYCOPROTEINS ARE REVEALED BY MALDI-TOF MAPPING OF THE MULTIPLE N-GLYCANS ASSOCIATED WITH RECOMBINANT BOVINE ACETYLCHOLINESTERASE
   C. Kronman, T. Chitlaur, N. Seliger, S. Lazar, A. Lazar, L. Zilberstein, B. Velan, A. Shafferman, Israel

10. EFFECT OF POST-TRANSLATION MODIFICATIONS OF HUMAN ACETYLCHOLINESTERASE ON ITS CIRCULATORY RESIDENCE
    T. Chitlaur, C. Kronman, S. Lazar, N. Seliger, B. Velan, A. Shafferman, Israel

11. CHANGES IN NEURONAL CHOLINERGIC RECEPTOR BINDING SITES AT DIFFERENT AGES IN TRANSGENIC MICE OVEREXPRESSING HUMAN ACETYLCHOLINESTERASE
    M.M. Svedberg, A-L. Svensson, I. Bednar, A. Nordberg, Sweden

12. TRANSGENIC OVEREXPRESSION OF READTHROUGH ACETYLCHOLINESTERASE (ACHE-R): DISTRIBUTION OF ACHE-R AND CFOS IN BRAIN IN RELATION TO BEHAVIOR

13. STUDY ON THE MECHANISM OF BLOCKADE OF ACETYLCHOLINE RELEASE BY SNAKE PRESYNAPTIC PLA2 NEUROTOXINS ON NERVE TERMINALS
    O. Rossetto, M. Rigoni, P. Caccini, C. Montecucco, Italy

14. 'READTHROUGH' ACETYLCHOLINESTERASE FORMS NEURONAL COMPLEXES WITH PKC BETA II AND ITS WD CARRIER RACK1
    E.H. Sklan, K.R. Birikh, S. Shoham, H. Soreq, Israel

15. THE ROLE OF READTHROUGH ACETYLCHOLINESTERASE IN THE PATHOPHYSIOLOGY OF MYASTHENIA GRAVIS
    T. Evron, Y. Hamra, N. Boneva, S. Seidman, T. Brenner, H. Soreq, Israel
16. CHRONIC ACETYLCHOLINESTERASE OVEREXPRESSION INDUCES MULTILEVELED ABERRATIONS IN NEUROMUSCULAR PHYSIOLOGY
N. Farchi, H. Soreq, B. Hochner, Israel

17. EXPRESSION OF THE CHOLINERGIC GENE LOCUS IN THE TRACHEAL EPITHELIUM OF THE RAT
U. Pfeil, L. Eberling, K.S. Lips, R.V. Haberberger, W. Kummer, Germany

18. MUSCARINIC RECEPTORS AND TRP-CHANNELS IN PRIMARY SENSORY NEURONS OF THE RAT
R. Haberberger, S. Wiegand, M. Kress, Germany

19. DIVERSE MOLECULAR MECHANISMS UNDERLYING CONGENITAL MYASTHENIC SYNDROMES
R.G. Webster, R. Croxen, S. Brownlow, M. Brydson, S. Haslam, C. Young, C. Slater, J. Newsom-Davis, A. Vincent, D. Beeson, UK

20. ACETYLCHOLINESTERASE KNOCKOUT MICE HAVE INCREASED SENSITIVITY TO SCOPOLAMINE AND ATROPINE
A. Hrabovska, O. Lockridge, E. Duyssen, USA, Slovak Republic

21. DOWNREGULATION OF MUSCARINIC RECEPTORS IN MICE DEFICIENT IN ACETYLCHOLINESTERASE
B. Li, E.G. Duyssen, O. Lockridge, USA

22. TARGETING OF THE HUMAN VESICULAR ACETYLCHOLINE TRANSPORTER TO CHOLINERGIC SUBDIVISIONS IN TRANSGENIC MICE
B. Schuetz, E. Weihe, L.E. Eiden, Germany, USA

23. THE DIURNAL ACTIVITY OF ACETYLCHOLINESTERASE INHIBITORS
B.M. Davies, USA

24. THE MUSCARINIC M1 RECEPTOR AS A THERAPEUTIC TARGET FOR COGNITIVE DEFICITS: PRECLINICAL PHARMACOLOGY AND KNOCKOUT MOUSE STUDIES
C.C. Felder, K.S. Gannon, F.P. Bymaster, A. Porter, D.L. McKinzie, J. Wess, N.M. Nathanson, UK, USA

25. RESCUE OF THE ACETYLCHOLINESTERASE KNOCKOUT MOUSE BY FEEDING A LIQUID DIET; PHENOTYPE OF THE ADULT ACETYLCHOLINESTERASE DEFICIENT MOUSE
E.G. Duyssen, J.A. Stribley, D. Fry, S. Hinrichs, O. Lockridge, USA

26. ROLE OF MUSCARINIC RECEPTORS IN THE ACTIVATION OF THE SUBICULO-ACCUMBENS PROJECTION
S.N. Mitchell, S. Moss, A. Sharott, UK

27. BRAIN PENETRATION AND BEHAVIOURAL PROPERTIES OF A POTENT ALPHA 7 NICOTINIC ACETYLCHOLINE RECEPTOR AGONIST IN THE RAT

28. FINE-TUNING MODULATION OF NEURONAL MUSCARINIC M1 (FACILITATORY) AND M2 (INHIBITORY) RECEPTORS ACTIVATION BY ADENOSINE AT THE RAT NEUROMUSCULAR JUNCTION
L. Oliveira, M.A. Timoteo, P. Correia-de-Sa, Portugal

29. IDENTIFICATION OF SIGNALING PROTEINS DOWNSTREAM OF THE TYROSINE KINASE MUSK IN CLUSTERING OF ACETYLCHOLINE RECEPTORS
R. Willmann, P. Mittaud, C. Fuhrer, Switzerland

30. DRAMATIC DEPLETION OF CELL SURFACE ACETYLCHOLINE MUSCARINIC RECEPTORS M2R DUE TO LIMITED DELIVERY FROM INTRACYTOPLASMIC STORES IN NEURONS OF ACETYLCHOLINESTERASE (ACHE) DEFICIENT MICE.
V. Bernard, C. Brana, I. Liste, O. Lockridge, B. Bloch, France, USA

31. CHEMICAL MODIFICATION OF RECOMBINANT HUMAN ACETYLCHOLINESTERASE BY POLYETHYLENE GLYCOL GENERATES AN ENZYME WITH EXCEPTIONAL CIRCULATORY LONGEVITY
O. Cohen, C. Kronman, T. Chililaru, S. Lazar, N. Seliger, D. Kaplan, A. Ordentlich, B. Velan, A. Shafferman, Israel
WEDNESDAY, MAY 8, 2002 (continued)

LIST OF POSTERS 3

32  HUPERZINE A AND DONEPEZIL ATTENUATE STAUROSPORINE-INDUCED APOPTOSIS IN RAT CORTICAL NEURONS VIA BCL-2 AND BAX REGULATION AND INHIBITION ON CASPASE-3
H.Y. Zhang, X.C. Tang, China

33  INSECT GROWTH REGULATORS INHIBIT ACETYLCHELINESTERASE ACTIVITY IN B-BIOTYPE BEMISIA TABACI IN AUSTRALIA
E.L.A. Cottage, R.V. Gunning, Australia
THURSDAY, MAY 9, 2002

LIST OF POSTERS 4

1  BIOCHEMICAL CHARACTERISATION OF MICE TRANSGENIC FOR A MUTATION IN AMYLOID PRECURSOR PROTEIN (APP) KNOWN TO CAUSE FAMILIAL ALZHEIMER'S DISEASE
   P.T. Francis, K.L. Matthews, K.E. Heslop, P.F. Chapman, UK

2  SYMPATHETIC SUPERIOR CERVICAL GANGLIA (S.C.G.) OF CAT CHOLINERGIC RELAY OF HYPOTHALAMIC-STIMULATED ORGAN-SPECIFIC VASCULAR CHANGES - RELEVANCY TO NORMAL AND TO CLINICAL DYSAUTONOMIC FUNCTION
   B. Blum, J. Israeli, Israel

3  CATECHOLAMINE INDUCED CYTOTOXICITY AND ITS PROTECTION, TAURINE AND ANALOGUES: SOME MORE THOUGHTS
   R.C. Gupta, India

4  THE EFFECT OF TYROSINE ON COGNITIVE FUNCTION IN ANIMAL MODELS FOR ANOREXIA NERVOSA
   D. Ben Shushan, Y. Avraham, S. Hao, S. Mendelson, E.M. Berry, Israel

5  THE EFFECT OF DIET RESTRICTION, SEPARATION STRESS AND TYROSINE ADMINISTRATION ON THE CHOLINERGIC SYSTEM IN MICE
   S. Hao, Y. Avraham, S. Mendelson, E.M. Berry, Israel

6  EFFECTS OF LITHIUM CHLORIDE ON MEMORY PERFORMANCES OF MICE IN ELEVATED PLUS-MAZE TEST
   P. Yamanturk, L. Eroglu, Turkey

7  EFFECTS OF 7-NITROINDAZOLE ON MEMORY PERFORMANCES OF RATS TRAINED FOR THREE-PANEL RUNWAY TASK: HIPPOCAMPAL CHOLINERGIC ENZYME ACTIVITIES
   P. Yamanturk, Y. Uluirceri, S. Bekpinar, H. Kayuncuoglu, Turkey

8  ACETYLCHOLINE AND NO-MEDIATED CGMP SYNTHESIS IN THE RAT BRAIN
   W.C.G. Van Staveren, M. Markerink-van Ittersum, H.W.M. Steinbusch, J. De Vente, The Netherlands

9  HYDROCORTISONE AFFECTS THE DENSITIES OF CARDIAC MUSCARINIC AND ADRENERGIC RECEPTORS
   J. Myslivecek, J. Rincen, S. Tucek, Czech Republic

10 EEG EVALUATION OF HUPERZINE A, A REVERSIBLE CHOLINESTERASE INHIBITOR
    S.L. Hale, H. Ved, A. Williams, B.P. Doctor, F. Tortella, USA

11 HUPERZINE A AND CHOLINESTERASE INHIBITORS: GLUTAMATE AND BENZODIAZEPINE RECEPTOR INTERACTIONS
    S.V. Nigam, B.P. Doctor, H.S. Ved, R.K. Gordon, USA

12 CALCIUM MOBILISATION AND CELLULAR CONTRACTION OF EMBRYONIC LENS VESICLE AND NEURAL TUBE ON MUSCARINIC CHOLINERGIC STIMULATION
    U. Dreus, M. Oppitz, G. Schriek, Germany

13 CHOLINERGIC MODULATION OF CHEMOTAXIS IN HUMAN MELANOMA CELLS
    A. Boss, S. Noda, M. Sailer, M. Oppitz, U. Dreus, Germany

14 COMBINED ANDROGEN-DONEPEZIL TREATMENT IN POST-STROKE REHABILITATION
    J.W. Crayton, L.M. Konopka, A.G. Karczmar, USA

15 LOCALISATION OF THE HIGH-AFFINITY CHOLINE TRANSPORTER-1 IN RAT SKELETAL MUSCLE AND SPINAL CORD
    K.S. Lips, U. Pfeil, R.V. Haberberger, W. Kummer, Germany

16 CHOLINERGIC-GLUTAMATERGIC INTERACTIONS IN HIPPOCAMPAL NEURONS: POSSIBLE ROLE IN THE NORMAL AND DISEASED HIPPOCAMPUS
    L. Pavlovsky, A. Friedman, Israel

17 HUPERZINE A, A PROMISING ANTI-ALZHEIMER'S AGENT, REDUCES STAUROSPORINE-INDUCED APOPTOSIS IN NG108-15 CELLS
    Y-F. Han, X-Q. Xiao, D-C. Wu, Y. Gao, W-L. Ho, N.T-K. Lee, Y. Fu, K.W.K. Tsim, Hong Kong
THURSDAY, MAY 9, 2002 (continued)

LIST OF POSTERS 4

18 DOPAMINE RELEASE FROM RAT STRIATAL SLICES IN VITRO AND FUNCTIONAL EFFECTS IN 6-OHDA TREATED RATS IN VIVO ARE MEDIATED BY BETA2 CONTAINING NICOTINIC ACETYLCOLINE RECEPTORS

19 THE EFFECTS OF GALANTAMINE IN PATIENTS WITH REFRACTORY SCHIZOPHRENIA RECEIVING RISPERIDONE
J.P. McEvoy, T.B. Allen, USA

20 CEREBRAL METABOLIC ACTIVATION WITH CHOLINESTERASE INHIBITOR THERAPY IN ALZHEIMER'S DISEASE
M. Mega, I.D. Dinov, M. Manese, J. Felix, S.M. O'Connor, J.L. Cummings, A.W. Toga. USA

21 THE ROLE OF LIPID PEROXIDATION IN THE MECHANISM OF NEUROTOXICITY OF ORGANOPHOSPHATES
V.D. Tonkopili, Russia

22 RESCUE OF THE NEURODEGENERATIVE PHENOTYPE IN AD11 ANTI-NGF MICE
S. Capsoni, S. Giannotta, A. Cattaneo. Italy

23 ACUTE CHOLINERGIC RESCUE OF SYNAPTIC PLASTICITY IN THE NEURODEGENERATING CORTEX OF ANTI-NERVE GROWTH FACTOR MICE
E. Pesavento, S. Capsoni, L Domenici, A. Cattaneo. Italy

24 THE EFFECT OF NICOTINE ON EXPRESSION OF NICOTINIC RECEPTORS IN THE BRAIN OF PATIENTS WITH ALZHEIMER'S DISEASE

25 A PEPTIDE FROM THE C-TERMINAL OLIGOMERISATION DOMAIN OF HUMAN SYNAPTIC (T-FORM) ACETYLCOLINESTERASE FORMS CLASSICAL AMYLOID FIBRILS
M.G. Cottingham, M.S. Hollinshead, D.J.T. Vaux. UK

26 NICOTINE INDUCES GLUTAMATE RELEASE FROM HIPPOCAMPAL MOSSY FIBRES SYNAPTOSONES
V. Bancila, A. Bloc, Y. Dunant. Switzerland

27 NICOTINIC CHOLINERGIC ACTIVATION OF MAGNOCELLULAR ENDOCRINE NEURONS OF THE HYPOTHALAMUS
M. Zaninetti, E. Tribollet, D. Bertrand, R. Ogier, M. Raggenbass. Switzerland

28 UNDERSTANDING THE DUAL MODE OF ACTION OF REMINYL(R) USING A VIRTUAL SYNAPTIC CLEFT
M. Lazarewicz, A. Spiros, L. Finkel, R. Carr, H. Geerts. USA

29 ALTERED ACTIVITY OF CHOLINERGIC ENZYMES IN MUSCLES AND BRAIN OF THE OBESE-DIABETIC (OB/OB) MOUSE
M.C. Lintern, L. Cooke, H. Scriven, M.E. Smith. UK

30 EFFECT OF PYRIDOSTIGMINE ADMINISTRATION ON ACETYLCOLINESTERASE AND CHOLINEACETYLTRANSFERASE ACTIVITY IN THE GUINEA-PIG STRIATUM AND CEREBELLUM
M.E. Smith, M.C. Lintern, C.J. Brewer, J.R. Wetherell. UK

31 NICOTINIC BETA4 RECEPTOR MEDIATED ACETYLCOLINE RELEASE FROM RAT INTERPEDUNCULAR NUCLEUS
F.A. Jones, L.R. Johnson, N. Evans, S. Bose, P.J. Craig, S.G. Volsen, I.A. Pullar. UK

32 CALCIUM CONDUCTANCE AND CHOLINE SENSITIVITY OF SLOW CHANNEL SYNDROME ACETYLCOLINE RECEPTOR MUTANTS
I. Spreadbury, R. Webster, D. Beeson, A. Vincent. UK

33 BEYOND THE USUAL SUSPECTS. A CHOLINERGIC ROUTE FOR PANIC ATTACKS
M. Battaglia, A. Ogliari, C. Maffei. Italy
THURSDAY, MAY 9, 2002 (continued)

LIST OF POSTERS 4

34 SELECTIVE HISTOCHEMICAL STAINING OF PERINEURONAL ACETYLCHOLINESTERASE (AChE) IN THE LIVING ENTERIC NERVOUS SYSTEM (ENS) OF RAT AND GUINEA-PIG
S. Tsuji, R. Nakatomi, H. Tsuchiya, I. Motelica-Heino, K. Hirai, Y. Katayama, K. Ishii, T. Hashikawa, France, Japan

35 NICOTINIC ACETYLCHOLINE RECEPTOR alpha5 SUBUNITS MODULATE OXOTREMORINE-INDUCED SALIVATION AND TREMOR
N. Wang, A. Orr-Urtreger, J. Chapman, R. Rabinowitz, A.D. Korczyn, Israel

36 AUTONOMIC FUNCTION OF NEURONAL NICOTINIC ACETYLCHOLINE RECEPTORS alpha5 SUBUNITS
N. Wang, A. Orr-Urtreger, J. Chapman, R. Rabinowitz, R. Nachman, A.D. Korczyn, Israel

37 DEFICIENCY OF beta NICOTINIC ACETYLCHOLINE RECEPTOR SUBUNITS CAUSES AUTONOMIC CARDiac AND INTESTINAL DYSFUNCTIONS
N. Wang, A. Orr-Urtreger, J. Chapman, R. Rabinowitz, R. Nachman, A.D. Korczyn, Israel

38 ALPHA7 ACETYLCHOLINE RECEPTOR IN SCHIZOPHRENIA: DECREASED mRNA LEVELS IN PERIPHERAL BLOOD LYMPHOCYTES
O. Perl, T. Ilani, R.D. Strous, S. Fuchs, Israel

39 USE OF THE MORPHING GRAPHICS TECHNIQUE TO VISUALIZE CONFORMATIONAL DIFFERENCES BETWEEN ACHEs FROM DIFFERENT SPECIES AND INHIBITOR-INDUCED CONFORMATIONAL CHANGES
T. Zeev-Ben-Mordehai, I. Silman, J.L. Sussman, Israel
IN MEMORIAM EDITH HEILBRONN
Victor P. Whittaker
Wolfson College, Cambridge, UK

Edith Heilbronn died on May 11, 1999. She was the main instigator of the series of International Symposia on Cholinergic Mechanisms of which this is the 11th; this is also the first since her death. The Organizing Committee have therefore felt it appropriate that her contribution to cholinergic mechanisms should be honoured at this meeting.

Edith was born in 1925, the only child of a German-Jewish couple, members of the large Jewish community of Fürth, Northern Bavaria. In 1938 the family sought sanctuary in Sweden, where her father had business connections. She started her career as a technician in the Werner-Green Institute but by attending evening classes she obtained her fil lic from Stockholm University. In 1957 she joined the scientific staff of the Swedish Defence Research Establishment. She rose steadily, becoming chief of an enlarged Biochemistry Section within the Department of Chemistry, meanwhile obtaining her doctorate from Uppsala University. She worked on the OP nerve gases, then on the isolation of the nicotinic acetylcholine receptor and the induction of an experimental form of myasthenia gravis by injecting it into rabbits.

By now well-known internationally, Edith felt the need to bring together others working in the field of cholinergic function. In February 1970 she organized the first ISCM in Skokloster, an old castle near Uppsala. Few who were present will forget the snowy landscape, the outdoor barbecue, the abundantly flowing glogg and above all the exciting exchange of scientific work among most of the leading groups, all testimony to Edith’s flair for organization. In 1979 Edith created a Department of Neurochemistry and Neurotoxicology within the Natural Science Faculty of Stockholm University. The stream of students from home and abroad who passed through the department did much to establish the scientific study of these subjects in Sweden and abroad.

ACETYLCHOLINE RECEPTORS: PROBING FUNCTIONALLY SIGNIFICANT STRUCTURAL CHANGES WITH SITE-DIRECTED REACTIONS
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Both in the ACh binding sites and in the cation-conducting channel of the nicotinic ACh receptor, changes in the reactivity of either native cysteines or of cysteines substituted for native residues have been correlated with changes in functional state. Changes in reaction rates of target cysteines reflect changes in the reagent's path to the target and changes in the local environment of the target. We have dissociated these two kinds of influence on the reaction rates to determine the position of the gate in the channel in both the resting state and the desensitized state and to determine the intrinsic electrostatic potential in the binding site and in the channel. Widespread changes in the reactivity of substituted cysteines in the channel lining reflect an extensive pathway of propagation of the perturbation from the ACh-binding sites to the region of the channel gate.

THE FUNCTIONAL ORGANIZATION OF BRAIN NICOTINIC RECEPTORS AT THE AMINOACID LEVEL: ACTIVATION, DESENSITIZATION AND UP-REGULATION
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The nicotinic receptor (nAChR) pentamer of 300KD carries all the structural elements which account for channel activation, desensitization, and up-regulation by acetylcholine (Corringer et al., 2000). The data obtained by affinity labeling, site-directed mutagenesis, and X-Ray crystallography (with the snail acetylcholine binding protein by Smit & Sixma groups) (Breje et al., 2001) about the relevant sites and conformational transitions are reviewed (Grutter & Changeux, 2001). The relative contributions of the 6 distinct loops of the ACh binding site located at the boundary between subunits, together with that of the M2 segment and M1-M2 loop from the ion channel are discussed. Structural elements contributing to desensitization and up-regulation are presented. The data are interpreted in terms of a multi-states allosteric model (Edelstein et al., 1997, Changeux and Edelstein, 1998).

References

THE CRYSTALL STRUCTURE OF MOLLUSCAN ACHBP REVEALS THE LIGAND BINDING DOMAIN OF THE NICOTINIC ACETYLCHOLINE RECEPTOR
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We have solved the crystal structure of Acetylcholine binding protein (AChBP), a homolog of a ligand binding domain of the nicotinic acetylcholine receptor (nAChR). AChBP is a glia-produced water-soluble protein from Lymnaea stagnalis, which is involved in synaptic modulation. Its homopentamer has sequence and pharmacological similarity to the nAChR ligand-binding domain, but it lacks a transmembrane domain. The nAChRs belong to the superfamily of pentameric ligand gated ion-channels, or Cys-loop receptors, which mediate rapid chemical transmission of signals. These allosteric transmembrane proteins include the nicotinic acetylcholine, the serotonin 5HT3, GABA A, GABAC and glycine receptors.

The crystal structure shows that in the AChBP protomers have an immunoglobulin-like topology. Ligand-binding sites are located at each of five subunit interfaces and contain residues contributed by biochemically determined 'loops' A to F. The subunit interfaces are highly variable within the ion-channel family, whereas the conserved residues are stabilizing the protomer fold. This AChBP structure is relevant for the development of drugs against e.g. Alzheimer's disease and nicotine addiction.
THE BINDING SITE OF ACETYLCHOLINE RECEPTOR: FROM SYNTHETIC PEPTIDES TO SOLUTION AND CRYSTAL STRUCTURE
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Our group has been employing short synthetic peptides, encompassing sequences from the acetylcholine receptor (AChR) α-subunit for the analysis of the binding site of AChR. A 13-mer peptide mimotope, with similar structural motifs to the AChR binding region, was selected by α-bungarotoxin (α-BTX) from a phage-display peptide library. The solution structure of a complex between this library-lead peptide and α-BTX was solved by NMR spectroscopy, indicating that the bound peptide adopts an almost globular conformation. Based on this NMR study and on structure-function analysis of the AChR binding site, additional 56 peptides, resulting from systematic residue replacement in the lead peptide, one or more replacements at a time, were designed and characterized. Of these, four binding peptides, designated high affinity peptides (HAPs), homologous to the binding region of AChR, inhabiting the binding of α-BTX to AChR with IC50 of 2 nM. The solution and crystal structures of complexes of α-BTX with HAP, were solved, demonstrating that the HAP fits snugly to α-BTX and adopts a β-hairpin conformation. The X-ray structures of the bound HAP and the homologous loop of the acetylcholine binding protein (AChBP), are remarkably similar. Their superposition results in a model indicating that α-BTX wraps around the receptor binding-site loop, and in addition, binds tightly at the interface of two of the receptor subunits, where it inserts a finger into the ligand-binding site. Our proposed model explains the strong antagonistic activity of α-BTX, and accommodates many of the biochemical data on the mode of interaction of α-BTX with AChR.

ALLOSTERISM OF THE NICOTINIC ACETYLCHOLINE RECEPTOR
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The nicotinic acetylcholine receptor is an allosteric protein. Agonist binding and channel opening as well as interactions with non-competitive inhibitors (NCIs) are cooperative processes. It has been proposed that the symmetry model (Monod-Wyman-Changeux Model, MWC) describes the receptor’s allostery best. This model postulates: i. a simple axis of symmetry and ii. a preformed equilibrium of functional receptor states, in this case the resting (channel closed), the active (channel open) and the desensitized (channel closed, high agonist affinity) states shifted by ligand binding. Experimentally, it is difficult to discriminate between this and alternative (e.g. the induced fit model) models, because the methods of investigation may affect the equilibrium between states. One prediction of the MWC model is preferential binding of agonists to the activated and desensitized states, and of antagonists to the resting state. We developed a method to “freeze” the equilibrium by covalent cross-linking and to fix acetylcholine receptors either in the resting or in the desensitized state. Binding studies performed with these fixed-state receptors are not compatible with the MWC model.

MOLECULAR BASIS OF THE SELECTIVITY OF NEUROTOXINS TOWARD αACHR SUBTYPES
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Snake alpha-neurotoxins block nicotinic acetylcholine receptors (AChR) from peripheral and neuronal tissues, with high affinity (10^-10^-14 M), alpha-Cbtx, from Naja kaouthia, like other long chain neurotoxins, can block muscular-type and alpha7 neuronal receptors. The physiological significance of this property is unclear but the molecular elements associated with this dual activity are identified. Thus a core of toxin residues, at the tip of the toxin central loop, bind to both receptor subtypes, whereas other residues interact to one subtype only. This situation suggests that the binding core binds to receptor residues that are present in both receptor subtypes whereas the subtype-specific toxin residues interact with differential receptor residues. To understand the molecular basis of this selectivity, we performed complementary double mutant cycle experiments between alpha-Cbtx and the neuronal AChR. These data were used to assist a docking calculation between the toxin and a structural model of alpha7 AChR, derived from the recently solved 3D structure of a soluble homopentameric homologue of the extra-cellular domain of AChR. This analysis provides a rational basis for selectivity of protein antagonists that block AChRs.

STRUCTURE AND DYNAMICS OF ACETYLCHOLINE RECEPTOR AND ITS LIPID MICROENVIRONMENT: MOLECULE TO CELL
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Two approaches to identify the occurrence of lipid sites in the membrane-bound nicotinic acetylcholine receptor (AChR) will be described: a) Förster-type resonance energy transfer (FRET) and fluorescence studies using the probe Laurdan; b) single-channel recordings of AChR mutated in relevant residues of transmembrane (TM) domains. Changes in FRET efficiency induced by fatty acids, phospholipid and cholesterol led to the identification of discrete sites for these lipids on the AChR protein. Spectroscopy studies of N-(1-pyrenyl)maleimide (PM)-labelled intact Torpedo AChR protein and TM peptides reconstituted into liposomes have served to explore their topography relative to the bilayer. From spin label quenching of PM-labelled Cys residues in αM1, αM4, γM1 and γM4 we could reach the conclusion that they all lie in a shallow position. For M4 segments, this is compatible with a linear α-helical structure, but not so for M1, for which “classical” models locate Cys residues at the centre of the hydrophobic stretch. The TM topography of M1 can be rationalized on the basis of the presence of non-helical structure, and/or of kinks attributable to the occurrence of the evolutionarily conserved proline residues. The latter is a striking feature of M1 in the AChR and in fact in all members of the rapid lipid-gated ion channel superfamly.

The effect of lipids on lipid domain ("raft") formation and AChR stabilization is currently being characterized using fluorescence methods in vitro and in living cells. Cellular studies are aimed at understanding how lipid domains relate to AChR targeting/stabilization at the cell surface. Towards this goal, cells expressing AChR are interrogated with fluorescence microscopy in combination with various lipid modification procedures and correlated with Laurdan generalized polarization (GP) studies. Changes in cell-surface fluorescence of alpha-bungarotoxin are observed upon cholesterol and sphingolipid modification, and correlated with modification of AChR targeting to the cell-surface and with changes in the physical state of the plasmalemma in various mammalian AChR-expressing cells.

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SITE-DIRECTED REACTIVE PROBES FOR STRUCTURAL AND FUNCTIONAL INVESTIGATION OF CHOLINERGIC PROTEINS

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The recently solved crystal structure of a glutamate-derivatized acetylcholine binding protein (ACHBP) [1] will prompt homology modelling of related ligand-gated ion channel proteins and allow the docking of receptor modulators. These models, however, will require biochemical and pharmacological studies for their validation.

We have successfully used photoaffinity probes such as [3H] DDF (p-Dimethylamino benzene Diazonium Fluoroborate) and [3H] DCTA (Diazotized Cylohexylamidopyryl Tris(ethyl) Ammonium), to characterize the acetylcholine binding sites on both Torpedo AChR and AChE [2,3] and are presently developing an alternative approach to obtain similar information on recombinant receptors. The method proposes an extension of the SCAM methodology [4] using cysteine-reactive site-directed affinity ligands which react irreversibly with the engineered cysteines [5]. The formation of a specific covalent bond, between selected Cys mutants and high-affinity site-directed labels, demonstrates the specific interaction between the ligand analog and the mutant protein. Such studies will permit accurate ligand receptor interaction studies, by defining precise anchoring points between the receptor and the ligand. Specifically, they will allow the delineation of reliable pharmacophores. The interaction of epibatidine with alpha7 and alpha4beta2 receptors is being investigated using this approach (Collaboration with D. Bertrand - CMU Geneva). The synthesis and the pharmacological properties of reactive epibatidine derivatives will be described. To gain a dynamic structural insight on functional proteins we also investigated the photochemistry of cholinergic enzyme activities for potential time-resolved crystallographic studies on CEs [6]. The photochemistry of AChE and BuChE activities was demonstrated using either caged enzyme substrates or by caging directly the catalytic serine of the enzymes.


NMR STRUCTURE OF α-BUNGAROTOXIN IN COMPLEX WITH AN ACHR α-SUBUNIT Peptide REVEALS THE BASIS FOR SPECIES SPECIFIC RESISTANCE TO THE TOXIN AND HOW α-NEUROTOXINS INHIBIT ACETYLCHOLINE BINDING TO THE RECEPTOR

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The α-subunit of the acetylcholine receptor (αAChR) contains the major binding site for the snake venom derived antagonist α-bungarotoxin (α-BTX). We solved the three-dimensional structure of an αAChR-peptide (residues 182-202) in complex with α-BTX using 2D 1H-NMR spectroscopy. The bound AChR-peptide adopts a β-hairpin conformation, which associates to the toxin through a novel intermolecular β-sheet, that manifests both hydrophobic and electrostatic interactions resulting in high affinity. This structure correlates the observed changes in toxin binding affinity with mutagenesis studies and with the naturally occurring mutations of αAChR in different animal species and α-subunit types. Based on the structure of a homologous molluscan acetylcholine binding protein, a model of the extracellular domain of the AChR was constructed. Two α-BTXs were docked to this model with the assistance of our NMR data. Lined with aromatic residues, the acetylcholine binding-sites at the interface of α- and γα-subunits are occupied by the side-chain of toxin residue Arg-36. This arginine which is conserved amongst all α-neurotoxins sterically prevents acetylcholine from binding to the receptor. These findings coincide with previous mutagenesis studies and illustrate the inhibition mechanism of AChR by α-neurotoxins.

ACTIVATION INHIBITION AND UPREGULATION OF THE HUMAN NEURONAL NICOTINIC α4/β2 RECEPTOR BY A PARTIAL AGONIST

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With the expression in cell lines of human neuronal nicotinic receptor it became possible to investigate the properties of these receptors using whole cell recording and very fast drug application. Moreover, these preparations opened the possibilities to better examine the effects of prolonged exposure to a given compound and therefore mimic conditions occurring in the brain. Examination of the activation and inhibition dose-response curve unveiled that human α4/β2 receptors display a high and a low affinity state. Prolonged exposure to nicotine (8 hours) caused a marked displacement of the high versus low affinity ratio, together with an increase of the maximal evoked current. To get a further understanding in this basic mechanism we have examined the effects caused by a partial agonist. Accurate determination of both the activation and inhibition dose-response curve unveiled that human α4/β2 receptors display a high and a low affinity state. Prolonged exposure to nicotine (8 hours) caused a marked displacement of the high versus low affinity ratio, together with an increase of the maximal evoked current. To get a further understanding in this basic mechanism we have examined the effects caused by a partial agonist. Accurate determination of both the activation and inhibition dose-response curve unveiled that human α4/β2 receptors display a high and a low affinity state. Prolonged exposure to nicotine (8 hours) caused a marked displacement of the high versus low affinity ratio, together with an increase of the maximal evoked current. To get a further understanding in this basic mechanism we have examined the effects caused by a partial agonist. Accurate determination of both the activation and inhibition dose-response curve unveiled that human α4/β2 receptors display a high and a low affinity state. Prolonged exposure to nicotine (8 hours) caused a marked displacement of the high versus low affinity ratio, together with an increase of the maximal evoked current. To get a further understanding in this basic mechanism we have examined the effects caused by a partial agonist. Accurate determination of both the activation and inhibition dose-response curve unveiled that human α4/β2 receptors display a high and a low affinity state.
THE ROLE OF P2Y1 NUCLEOTIDE RECEPTOR IN THE FORMATION OF NEUROMUSCULAR JUNCTIONS

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In vertebrate neuromuscular junctions, adenosine 5'-triphosphate (ATP) is stored at the motor nerve terminals and is co-released with acetylcholine during neural stimulation. Several lines of evidence suggest that the synaptic ATP can act as a synaptic-organizing factor at the neuromuscular junctions, mediated by metabotropic P2Y1 receptors. P2Y1 receptor mRNA in chicken muscle is very abundant before hatching and again increases in the adult. The P2Y1 receptor protein is known to be restricted to the neuromuscular junctions and co-localized with AChRs in adult muscle, but not in the chick embryo. In chicks after hatching, this P2Y1 localization develops over about 3 weeks. Denervation or crush of the motor nerve (in chicken or rat) caused decrease in the muscle P2Y1 transcript, which was restored on regeneration. The activation of P2Y1 receptor by adenosine nucleotides stimulated the accumulation of inositol phosphates and intracellular Ca2+ mobilization in cultured chick myotubes. The receptor activation leads to an increase in the expression of transcripts encoding AChR alpha-subunits and AChE. In addition, the expression of P2Y2 and P2Y4 receptors were also restricted at the neuromuscular junctions. These results provide evidence for a novel function of ATP in directing the gene expression of post-synaptic functional effectors.

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THE DYSTROPHIN COMPLEX – A SCAFFOLD FOR SIGNALING PROTEINS AT SYNAPSES

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One important function of the dystrophin complex in muscle is to link the extracellular matrix (via dystroglycan) to the actin cytoskeleton, thus providing stability to the membrane during contraction. Recent evidence, however, suggests that the dystrophin complex is also a signaling complex. Two classes of dystrophin-associated proteins, the dystrobrevins and the syntrophins, form a scaffold for numerous signaling proteins, including ion channels, kinases and nNOS. Five syntrophins, named α, β1, β2, γ1 and γ2, are known. Each is a PDZ containing protein. In skeletal muscle of mice lacking α-syntrophin, nNOS and the water channel, aquaporin-4, are absent from the membrane. Furthermore, the neuromuscular junctions are abnormal, with reduced levels of acetylcholine receptor and acetylcholinesterase. The localization of nNOS and aquaporin-4 can be restored by genetic rescue expressing α-syntrophin. However, a modified form of α-syntrophin lacking the PDZ domain is not able to restore either nNOS or aquaporin-4 to the membrane. Thus, in vivo, both of these proteins require the α-syntrophin PDZ domain. In contrast to muscle, in which aquaporin-4 levels are greatly reduced, the levels in brain are unaffected by the absence of α-syntrophin. However, immunoelectron microscopy reveals that aquaporin-4 is mislocalized and resides mostly on intracellular organelles. Thus, α-syntrophin appears to be involved in regulating the expression, trafficking, and/or localization of certain membrane proteins.

AGRIN BLOCKADE IMPAIRS LATE BUT NOT INITIAL STAGES OF FUNCTIONAL INNERRATION OF HUMAN MUSCLE IN VITRO

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Agrin is a nerve-derived factor which promotes clustering of components of neuromuscular junctions (NMJs) by the activation of Muscle specific kinase (MuSK). Unlike MuSK deficiency which prevents NMJ formation completely, agrin deficiency permits some degree of synaptogenesis of NMJs. In order to provide more differentiated insight into the role of agrin in the NMJ formation, we followed functional maturation of NMJs in the in vitro innervated human muscle under the conditions of agrin blockade. Agrin was blocked by Agr33, a monoclonal antibody demonstrated to efficiently reduce AChR clustering. In the control experiments Agr33 was replaced by Agr86, a monoclonal antibody with little effect on AChR clustering. First and relatively sparse contractions are normally observed between days 7 and 10 of co-culture. Agr33 as well as Agr86 had no effect in this initial stage of muscle innervation. However, Agr33-, but not Agr86- treated co-cultures failed to increase the number of contraction units during the subsequent 5 days when this number significantly increased in the untreated co-cultures. Our results demonstrate that a small population of NMJs reach contraction-competent level relatively early even in the absence of functional agrin, while most of the primarily established neuromuscular contacts need full agrin activity to reach contraction-competent level.

ORGANIZING ACETYLCHOLINESTERASE MOLECULES AT THE NEUROMUSCULAR SYNAPSE

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The major form of acetylcholinesterase (ACHE) at the vertebrate neuromuscular junction (NMJ) is the asymmetric (A12) form consisting of three tetramers of catalytic subunits attached to a collagen-like tail. These molecules are locally synthesized and assembled, and, following externalization, become tightly attached to the synaptic basal lamina. The appearance of ACHE on the cell surface is locally controlled by signals generated at the overlying plasma membrane. To study this local control of ACHE expression, sealed chambers were used to isolate regions of individual myotubes, and subsequent treatment with agents that increased or blocked membrane depolarization. Blocking membrane depolarization resulted in upregulation of ACHE, whereas signals generated in response to membrane depolarization downregulate the enzyme and increased A12 ACHE assembly. In situ hybridization, RNase protection, and nuclear run on studies suggested that local control occurs at the level of transcription. However, once synthesized, the enzyme is externalized where it accumulates on the extracellular matrix. Several lines of evidence including in vitro and in culture binding studies, as well as studies using genetically modified mice, indicate that the heparan sulfate proteoglycan perlecian is responsible for localizing ACHE on the synaptic basal lamina. Thus, while local regulatory events are responsible for producing sufficient ACHE molecules, interactions with extracellular matrix components are responsible for the final localization.
THE FUNCTIONAL HETERO-OLIGOMERIC FORMS OF CHOLINESTERASES

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The acetylcholinesterase gene of vertebrates may generate several splice variants which possess the same catalytic domain but differ by their C-terminal regions. However, a single type of catalytic subunit (type T, or AChEt) exists in all vertebrate classes, and represents the AChE species which is expressed in the nervous tissue and muscles of mammals. Butyrylcholinesterase (BChE) subunits possess an homologous C-terminal region (T peptide). The T peptide consists of 40-41 residues, which include a cysteine at position 4 of the C-terminus, and a series of seven aromatic residues, including three tryptophans, which are fully conserved in all vertebrates. The T peptide can form an amphipathic α-helix with an hydrophobic face covered by these residues. AChE and BChE T subunits can form homo-oligomers, and also associate with anchoring proteins, generating asymmetric forms and membrane-bound tetramers. Our group has cloned a specific collagen, ColQ, which represents the "tail" of asymmetric forms, and more recently a transmembrane protein, PRIMA, which represents the hydrophobic anchor of membrane-bound tetramers. We characterized a proline-rich motif of ColQ, the PRAD ("proline-rich attachment domain"), which is responsible for association with cholinesterases; PRIMA also possesses a proline-rich motif which differs somewhat from the PRAD of ColQ, but clearly serves the same function. We further showed that the interaction relies on the C-terminal T peptide of cholinesterases, which may thus be considered as an autonomous "tryptophan (W) amphipathic tetramerization" (WAT) domain. Mutagenesis studies showed that the prolines of PRAD and the aromatic residues of WAT play a critical role in this interaction, and that only cysteines located near the C-terminus of the T peptide could form disulfide bonds with the cysteines located at the N-terminus of another subunit. In collaboration with some of the groups of H. Sihlan and J. Sussman, we have observed that the PRAD of ColQ forms a complex with four WATs: the four parallel α-helical T peptides form a cylinder around the PRAD, organized as a polypeptide II helix and oriented in the opposite direction; the aromatic side chains are stacked with prolines and the charged residues are located on the outside, and form acidic-basic pairs, within each T peptide. The assembly of PRAD and WATs organizes tetramers of AChEt and BChE subunits which are anchored respectively in the basal lamina of neuromuscular junctions, and in the cell membranes, particularly in the central nervous system: ColQ and PRIMA therefore condition the functional localization of cholinesterases.

TRANSCRIPTIONAL AND POST-TRANSCRIPTIONAL EVENTS CONTROLLING EXPRESSION OF ACETYLCHELINOSTERASE IN DEVELOPING AND ADULT MUSCLES

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Despite recent progress in our understanding of the biosynthetic events regulating AChE in muscle, our knowledge of the specific molecular mechanisms remains fragmentary. Recently, we began to examine this issue using several experimental systems. Our findings showed that the synaptic accumulation of AChE mRNAs results from a preferential activation of the AChE gene in myonuclei close to the neuromuscular junction. Mutation and deletion studies further demonstrated the critical role of the first intron, particularly that of an intronic N-box motif, in regulating the synaptic expression of the AChE gene. Additional studies revealed that this N-box, along with an adjacent E-box, are also critical for increasing transcription of AChE during early myogenesis. In these studies, we noted that the sustained elevation of AChE mRNAs in myotubes also involves post-transcriptional events. Interestingly, post-transcriptional mechanisms also play a key role in regulating the abundance of AChE transcripts in fast vs slow muscles, and in denervated muscles. Yet, the contribution of transcriptional and post-transcriptional events under these experimental conditions appeared subject to developmental influences. Experiments performed using the rat superior cervical ganglia, indicate that these two types of mechanisms play a similar role in regulating AChE in neurons. Thus, regulation of AChE mRNAs in excitable cells depends on the complex interplay between transcriptional and post-transcriptional mechanisms whose relative importance are influenced by the state of differentiation and maturation of the cholinergic synapses.

ACETYLCHELINE HYDROLYSIS AT THE MAMMALIAN SKELETAL NEUROMUSCULAR JUNCTION: MORE THAN ONE ENZYME

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Two enzymes hydrolyze acetylcholine (ACH) in mammals, acetylcholinesterase (AChE) and butyrylcholinesterase (BChE). These two enzymes share a noteworthy molecular polymorphism. These forms exist as a result of two catalytic subunits, each encoded by one gene, AChE and BChE, and of two anchors, ColQ and PRIMA. Each anchor has two functions 1) organization of an enzyme tetramer; 2) targeting of the tetramer to the extracellular domain. ColQ anchors tetramers in the basal lamina and PRIMA anchors tetramers on the cell surface. At the neuromuscular junction (NMJ), ACh actions appear multiple since each of the cells that organize the NMJ are sensitive to ACh: 1) the muscle endplate is activated by quantal ACh release, 2) the prejunctural motor nerve terminals generate fasciculation and back-firing after AChE inhibition and 3) the Schwann cell response to ACh release. To understand this complexity, we started genetic approaches to remove cholinesterases oligomers. In the absence of ColQ, the mice live but are affected by a myasthenia, like humans affected by mutations in ColQ. We have shown that in ColQ-/- mice AChE is absent, but BChE is still present in the telogral domain, probably anchored by PRIMA. To evaluate the functional consequence of the absence of AChE and the presence of BChE, we quantified nerve-evoked muscle contraction at different frequencies of nerve stimulation. The muscle failed to maintain a tetanus during repetitive stimulations (higher than 30 Hz). At 10 Hz of nerve stimulations, the contraction of the ColQ mutant is very similar to the control, but the mutant is very sensitive to BChE inhibition, while the control is less affected. This indicates that BChE controls the level of ACh during repetitive stimulations. However, BChE does not play the same role as AChE because miniature Endplate Potentials (mEPPs) and EPPs due to spontaneous and evoked release are not modified after BChE inhibition. This clearly suggests that the ACh level is controlled differently depending on the localization and/or the nature of the enzyme.

DENSITY AND LOCALIZATION OF ACETYLCHELINOSTERASE IN VERTEBRATE NEUROMUSCULAR JUNCTIONS

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The proper function of synaptic acetylcholinesterase (AChE) is determined by its concentration and position in the synaptic cleft. The present work describes the use of derivatives of fasciculin (Fas2), a polypeptide anticholinesterase toxin, as probes for determination of AChE densities at cholinergic synapses and for high resolution EM localization of AChE in synaptic clefts. Comparative studies of AChE densities and distribution at neuromuscular junctions (nms) were done by quantitative EM-autoradiography of muscles labeled with radioiodinated Fas2. These studies revealed several-fold differences in junctional AChE site densities in muscles of different vertebrates. Junctional AChE sites/μm² in mouse sternomastoid and lizard intercostal endplates were 4 and 2.5-fold higher than in frog cutaneous pectoris nms, respectively. AChE sites were distributed over the primary clefts and the full depth of the junctional clefts. Localization studies with nanogold-conjugates of Fas2 revealed that gold-labeled AChE sites were distributed over the basal lamina in the primary cleft and the postjunctional folds. Quantitative data analysis demonstrates that AChE sites are almost exclusively located on the basal lamina rather than pre- or postsynaptic membranes and are distributed in the primary cleft and full depth of the postjunctional folds. This localization pattern of AChE assures the hydrolysis of ACh bouncing off receptors. and eliminates its unnecessary re-binding.

FOUR ACYTLCHOLINESTERASE GENES IN THE NEMATODE CAENORHABDITIS ELEGANS

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Several genes encode acetylcholinesterase (ACHE, EC 3.1.1.7) in nematodes. In Caenorhabditis elegans we have cloned and sequenced the four ace genes; studied their genomic organization and their tissue-specific expression using GFP reporter constructs transfect in vivo. Two genes, ace-1 and ace-2, encode two major ACHEs with different pharmacological properties and tissue repartition. ace-1 is expressed in muscle cells (1.2) and ace-2 is mainly expressed in neurons (3). One likely hypothesis is that both ace-1 and ace-2 contribute (pre- and post-synaptically) to the neuromuscular ACHE explaining that single mutants in ace-1 or ace-2 have no alteration in locomotion. Interestingly, ace-1 has a C-terminus homologous to that of the T variant of vertebrate ACHE, whereas ace-2 possesses a C-terminal signal for glycosylation as the H variant in vertebrates. ace-3 represents a minor proportion of the total ACHE activity in C. elegans and is expressed in a few cells (pharyngeal muscle cells and CAN cells). ace-3 is highly resistant to the usual inhibitors of ACHE. ace-4 is found a few hundred base pairs upstream of ace-3 on chromosome II. Both genes are organized as an operon and are first transcribed as a bicistronic messenger. However, no enzyme corresponding to ace-4 was found. 1. Arpagaus et al., 1994, J Biol Chem 269, 9957-65. 2. Culetto et al., 1999, J Mol Biol 290. 951-66. 3. Combes et al., 2000, J Mol Biol 300, 727-42.

WANDERINGS IN AND ABOUT ACTIVE SITE GORGES AND SUBUNIT INTERFACES

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A wide variety of natural toxins that inhibit motor activity have evolved from both plant and animal species for purposes of predation or protection from predation. The unique specificities of these toxins permit their use to not only distinguish subtypes of receptors or other targets, but also probe the structures of individual binding sites within a single oligomeric receptor. In addition, these toxins and related ligands provide information on the conformational dynamics of the nAChR and ACHE not available in crystal structures. We show here the structural bases for the Conus magus toxin's, V-conotoxin-MI, preferential affinity for the V+ subunit interface over V and V' interfaces; the Viper toxin's, Waglerin-1, preference for the V', interface and the krait toxin, Naja mossambica mossambica I, preference for the V+ and V' subunits over V'. Distinct structural determinants can be defined indicating that discrete regions, formed from distant portions of the linear sequence form the binding site. In fact, these regions show the appropriate proximity relationships in the recently crystallized soluble acetylcholine binding protein from snail. Through the use of pairwise mutations in the receptor and the respective toxins and thermodynamic mutant cycle analysis, it has been possible to determine proximity relationships of residues on the receptor and toxin, orient the toxin within the binding sites, and delineate individual residue contributions to the binding energetics. A second 3-fingered toxin, fasciculin, is a high affinity inhibitor of ACHE. Through cysteine substitution mutagenesis and selective fluorophore conjugation, changes in residue microenvironments, solvent exposure and sequential motion of individual side chains have revealed that fasciculin and small ligand binding induces distinctive allosteric changes in the omega loop (Cys 69-Cys 96) of mouse ACHE. Hence these physicochemical parameters provide an essential link between molecular dynamics computations and the static crystal structures.

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REGULATION OF NEUROTRANSMITTER RELEASE: CALCIUM AND ION CHANNELS

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Release of transmitter from the cholinergic presynaptic nerve terminals is one of the key processes in neuronal communication. It is a highly controlled process and is regulated by a large number of intrinsic and extrinsic factors. Transmitter release regulation is one of the main determinants of synaptic plasticity. It also serves as an important target for the action of drugs and hormones.

Three aspects of the regulation of transmitter release will be discussed: ion channels in the presynaptic nerve terminal surface membrane, the post-fusion control of transmitter release and the calcium dynamics in presynaptic boutons.

Ion channels at the nerve terminal. The main regulator of quantal transmitter release is the intracellular calcium ion concentration. This in turn is regulated by the calcium ions that enter through the surface membrane and the calcium ions released from intracellular stores. These processes are controlled by several hundred of different ion channel molecules that affect directly and indirectly the calcium flux.

The post-fusion control of transmitter release. The cholinergic synaptic vesicle of Torpedo electrotonic neuron contains acetylcholine at a very high concentration. Most of the acetylcholine is bound to an intravesicular ion exchange matrix. Entry of cations into the vesicles can displace the transmitter and facilitate its release. We propose that the non-selective ion channels in the vesicle membrane can thus exert a post fusion regulation of transmitter release.

Calcium dynamics in nerve terminals and sea urchin eggs. Four aspects of calcium dynamics in the presynaptic nerve terminals will be discussed in view of their possible role in transmitter release: calcium oscillations, calcium waves, calcium sparks and calcium noise.

MACROMOLECULAR ARCHITECTURE OF ACTIVE ZONE MATERIAL AND ITS ROLE IN SYNAPTIC TRANSMISSION

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The position of active zone material at the nervous system's synapses, next to the synaptic vesicles docked at the presynaptic plasma membrane and to calcium channels within the membrane---both of which are directly involved in neurotransmitter release during impulse transmission---has raised questions as to the material's relationship to the vesicles and channels and to its function. My colleagues and I have examined for the first time the intricate arrangement and associations of structural components of this compact, proteinaceous organelle. Our approach was to use electron microscope tomography to generate 3D reconstructions of tissue sections from a model synapse, the frog's neuromuscular junction, and then to apply segmentation and surface rendering schemes for characterizing specific structures within the active zone material. Our findings lead to the hypothesis that the active zone material helps dock synaptic vesicles and anchor calcium channels and that the architecture of the material provides for both a particular spatial relationship and a structural linkage between the vesicles and channels. The structural linkage may well include proteins that mediate the calcium-triggered exocytosis of neurotransmitter by synaptic vesicles during synaptic impulse transmission.
MUSCARINIC PRESYNAPTIC RECEPTORS ARE INVOLVED IN THE CONTROL OF THE TIME COURSE OF NEUROTRANSMITTER RELEASE IN FROG AND MOUSE NEUROMUSCULAR JUNCTIONS

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Neurotransmitter release (amount and kinetics) in fast synapses is considered to be controlled by [Ca^2+]. Recent studies suggest that the action potential plays a direct role in determining the kinetics of release by a mechanism which involves presynaptic autoreceptors. Perfusion of a frog nmj with either the M2/M4 muscarinic antagonist, methoctramine, or with exogenous AChE, prolongs the kinetics of ACh release. This effect is reversed by muscarine. Methoctramine does not affect either the excitatory nerve terminal current or the presynaptic Ca^2+ currents. These results support a novel hypothesis according to which depolarization initiates release by relieving a tonic block produced by the M2 receptor and rebinding of ACh to the M2 receptor terminates release.

Phrenic-diaphragm preparation of knockout mice without functional M2 receptors were compared with those of wild type mice. Experimental manipulations that affected [Ca^2+]i greatly changed the amount, but not the kinetics of release in wild-type mice. In contrast, in the mutant mice, treatments that affected [Ca^2+]i affected both the amount and also the kinetics of release. The behavior of release in M2-Ko mice follows predictions from the Ca2+ hypothesis, while the behavior in wild type mice follows predictions of the Ca2+-voltage hypothesis.

NEUROTRANSMITTER PHENOTYPE SWITCH IN DEVELOPING NEURONS – GENOMIC AND PROTEOMIC APPROACHES

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P19 cells are embryonal carcinoma cells that serve as a model for studying differentiation processes including commitment to cell lineage. We have studied P19 cells following activation of neuronal and glial differentiation. The potential of these cells to mature and efficiently release neurotransmitter (NT) was established in our lab. We discovered that several variables, most notably, cell density and various neurotrophic factors affects neuronal maturation, survival and most surprisingly, the choice of NT phenotype. We showed that P19 mature to functional cholinergic neurons but also to glutamatergic neurons according to culture manipulation protocols. Those changes are associated with changes in gene expression. A large-scale holistic view on gene expression was obtained by DNA chip technology. Several predicted but some novel signaling cascades are activated a result of manipulating the culture by neurotrphins and by elevating the cell density. We have observed that NT phenotype acquisition in P19 cells is mediated by cell-cell contact and hypothesized that cell surface proteins to be essential for consolidation of NT phenotype switch. We have used proteomics approach to focus on the relevant membranous proteins and applied comparative 1D and 2D analyses to isolate specific target proteins. Membraneous proteins were isolated from P19 cells that were maintained in varying culturing conditions that support cholinergic vs. glutamatergic phenotypes. Membranes were collected at different time windows following neuronal induction. Differentially expressing proteins were excised and sent to Mass-spectroscopy analysis. The analysis is based on MALDI and Electrospray mass spectrometry (ES-MS). Currently, over 30 proteins were successfully analyzed by such methodology. Using 2D gels we were able to improve detection level and could identify relatively low expressing proteins. Some proteins were identified multiple times in independent experiments. Most intriguing proteins are a variant of Drebin - a putative dendritic-shaping molecule; Prohibitin that signifies post-mitotic cells; and several cytoskeletal-signaling molecules. Surprisingly, we observed marked changes in expression of a large group of heat shock proteins and their regulators along maturation of the neurons. The importance of a global unbiased proteomics view on developing neurons will be discussed.
GENETIC DISSECTION OF SYNAPTIC FUNCTION
IN DROSOPHILA
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Many questions of synaptic cell biology can be addressed by the study of
mutations in model systems. This lab has used Drosophila genetics to
elucidate the function of synaptic proteins. Mutations in neuronal-
synaptobrevin, a vesicular SNARE protein, demonstrate distinct mechanistic
requirements for minis and evoked release. By substituting related SNAREs,
we have tested the specificity of trafficking and the hypothesis that SNAREs
may uniquely target an individual vesicle class to the appropriate target
membrane. Mutations of the C2A domain of synaptotagmin have also been
examined and demonstrate that calcium-binding by this domain is not
essential for calcium-dependent transmission. To facilitate the identification
of
new components of the axon and terminal, we have devised a novel method
to screen for synaptic defects. By genetic manipulations, flies are screened
that are heterozygous for mutations but whose eyes are completely
homozygous for the mutation. Synaptic mutations are then recovered by
screening for blind flies with characteristic electrophysiological defects. A
gene called milton was isolated in this manner and is important for
intracellular transport of organelles to synaptic terminals. Intracellular
transport is thought to require adapter proteins for molecular motors in order
to identify and bind specific cargos, determine destinations, and anchor
cargoes after transport. Milton appears to link kinesin to mitochondria.
Mitochondria are completely absent from the milton photoreceptor nerve
terminal and axon, but present and apparently functional in the cell body.
Milton is present on mitochondria and is associated with kinesin heavy
chain. Milton contains significant homology to the Huntingtin-binding
domain of Huntington-associated protein 1 (HAP1). We propose that Milton
is a mitochondrion-specific kinesin adapter protein required for axonal
transport of mitochondria.

REGULATION OF CHOLINERGIC GENE EXPRESSION BY
NRSF/REST
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The cholinergic gene locus contains the genes for both the biosynthetic
enzyme choline acetyltransferase and the vesicular acetylcholine transporter.
Within the 5' region of the gene is found the 21 base-pair NRSF/REST
sequence to which the transcriptional repressor Neuron Restrictive Silencer
Factor (NRSF)/REST silencing factor (REST) binds and silences the gene.
Although the cholinergic gene is active in PC12 cells, in a mutant PC12 cell
line lacking protein kinase A the cholinergic gene is repressed. Although
both wild type and mutant PC12 cells express NRSF/REST, only wild type
cells express a neuron specific truncated form called REST4. We have
found REST4 can form a hetero-oligomer with NRSF/REST and block its
binding to the NRSE. We have found that REST4 can be trafficked to the
nucleus and that this requires the participation of zinc finger domains. We
have uncovered signals in REST4 (and presumably NRSF/REST) that are
responsible for targeting to the nucleus, for entry into the nucleus, and for
release into the nucleus. These will be described.

COMPARATIVE STRUCTURAL STUDIES ON CONJUGATES OF
TORDOCA CALIFORNICA AND HUMAN
ACETYLCOLINESTERASES WITH ORGANOPHOSPHATE NERVE
AGENTS
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Detailed understanding of structure-function relationships in the inhibition of
ACHE by OP nerve agents is a prerequisite for developing prophylactic and
therapeutic approaches for treating nerve-agent intoxication. Knowledge of the
3D structures of native AChE and of relevant OP conjugates is essential for
this. Our determination of the 3D structure of Torpedo californica AChE,
followed by solution of the 3D structures of its 'aged' conjugates with soman,
and both its 'aged' and 'non-aged' conjugates with VX, and of its
conjugate with tabun has furthered this objective.
The 'aged' conjugates of TaChE; with sarin and soman display essentially
identical 3D structures, thus providing structural models for the transition state
during the deacylation step with acetylcholine as substrate.
Comparison of the native TaChE structure and of the 'non-aged' and 'aged'
conjugates with VX reveals reversible movement of the catalytic histidine.
This movement involves a shift in hydrogen bonding from E327 to E199,
which has been postulated to participate in substrate and OP reactions.
The 3D structures of the rhAChE/FAS-II complex with DFP, sarin and VX
have been determined. Significant differences were observed relative to the
corresponding TaChE conjugates. It is premature to assign these differences
to the bound FAS-II, which makes direct contact with the backbone of
residues contributing to the acyl pocket. In or to inherent species differences
in plasticity of the acyl pocket. Such differences may also affect relative
susceptibilities of electric organ and mammalian AChEs to other OP
inhibitors, such as PMSF and the anti-Alzheimer drug, rivastigmine.
CRYSTAL STRUCTURE OF RECOMBINANT HUMAN BUTYRYLCHOLINESTERASE: NEW INSIGHTS INTO THE CATALYTIC MECHANISMS OF CHOLINESTERASES

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Acetylcholinesterase (AChE) and butyrylcholinesterase (BChE) are two related enzymes with different substrate and inhibitor specificities. BChE hydrolyzes a large variety of ester-containing drugs such as cocaine and scavenges organophosphorous and carbamate toxic esters. Mutants of BChE capable of hydrolyzing organophosphates have been designed. However, their activity needs to be improved to be of operational interest for prophylaxis and/or treatment of nerve agent poisoning, and decontamination purposes.

Unlike AChE, no X-ray structure of BChE is known, mainly because of the high glycosylation content of natural BChE, preventing crystal growth. Therefore, most structure studies of BChE have relied on homology models built from the Torpedo californica AChE structure. Although these models were instrumental in understanding some aspects of the AChE and BChE specificity differences, they are not accurate enough as templates for the rational design of mutants with particular catalytic features.

We recently crystallized a recombinant monomeric low-glycosylated form of human butyrylcholinesterase. The structure was solved at 2.6 Å resolution by molecular replacement using the Torpedo californica AChE structure as a starting model. Both enzyme structures are very similar. However, the active site of BChE presents an unexpected feature which may change the current interpretation of the molecular mechanisms of cholinesterases.

SURPRISING FINDINGS FROM THE FUNCTIONAL ANALYSIS OF HUMAN ACETYLCHOLINESTERASE ADDUCTS OF ALZHEIMER'S DISEASE DRUGS

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Determination of the 3D-structure of acetylcholinesterase (AChE) of Torpedo californica over a decade ago and more recently that of human enzyme together with extensive targeted mutagenesis of the mammalian AChEs led to a fine mapping of the multiple functional subsites within the active center of the enzyme. Our library of single and multiple human AChE mutants defining the various subsites was used to kinetically analyze interactions with various AChE inhibitors including tacrine (Cognex), huperzine-A, rivastigmine (Excelon), physostigmine, pyridostigmine, E2020 (Aricept, Donepezil) and galanthamine which are considered or currently in use for the treatment of Alzheimer's disease (AD). Such functional analysis characterized the key domains within the active center that are essential for accommodation of these prototypic inhibitors. Furthermore it allowed defining major structural features of the individual inhibitors that determine affinity and specificity for the enzyme. Some important and unexpected interactions were revealed by the functional analysis that could have not been anticipated from the 3D-structure of inhibitor-AChE complexes. These findings emphasize the importance of complementing the structural data with functional characterization of biological target molecules. Thus it appears that screening of lead compounds with a library of human AChE mutants may be a very useful and cost effective way for structure-based design and development of new therapeutics for AD.

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INTRINSIC TRYPTOPHAN FLUORESCENCE OF CHOLINESTERASES: DIRECT, NON-PERTURBING MONITORING OF ENZYME-LIGAND INTERACTIONS

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Intrinsic fluorescence of acetylcholinesterases (AChE, EC 3.1.1.7) and butyrylcholinesterases (BuChE, EC 3.1.1.8) was investigated to monitor enzyme-ligand interactions. The 10-13 tryptophans of cholinesterases emit fluorescence in 330 - 340 nm range upon excitation with UV light. Stern-Volmer analysis of collisional quenching of mouse AChE fluorescence by NaN, in the absence and presence of inhibitory peptide fasciculin 2, indicates that tryptophans of the AChE active center gorge contribute disproportionally to the overall fluorescence. Binding of non-fluorescent ligands, non-absorbing in 330-340 nm range and incapable of resonance energy transfer, such as decamethonium, BW286c51, edrophonium, ethopropazine, acetylcholine and choline, quench AChE and BuChE fluorescence. However, binding of fasciculin 2 and carbamoylation of the active site increase mouse AChE fluorescence intensity about 15%, thus suggesting conformational change involving tryptophans in the interaction. The rate of the conformational change appears faster than equilibration of reversible enzyme-inhibitor complexes thus allowing monitoring of inhibitor association and dissociation reactions in the millisecond time frame. Use of catalytically inactive mouse AChE S203A mutant allows for direct titration of AChE with acetylcholine revealing two binding sites for the substrate, one with Kd of ~80 uM and the other one of ~30 nM. In addition, measurements of rates of quenching of intrinsic AChE fluorescence upon association of edrophonium with mouse wild-type and H447L mutant AChE demonstrated a pH dependence consistent with the protonation state of H447 of the catalytic triad. In conclusion, monitoring of intrinsic tryptophan fluorescence of AChEs and BuChEs, is a sensitive and non-perturbing method of studying interactions of these enzymes with variety of nonfluorescent ligands. (Supported by grants from the NIH and DAMD).

UNFOLDING AND FOLDING OF TORPEDO CALIFORNICA ACETYLCHOLINESTERASE


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Chemical modification of Cys231 in TaAChE by various sulfhydryl reagents results in its conversion to one of two principal states. One of these states, produced by disulfides and by alkylating agents, is a stable state which displays typical features of a partially unfolded molten globule (MG) state, based on CD, intrinsic fluorescence and ANS binding. The second state, produced by mercury derivatives and the natural thiosulfate, allicin, which has spectroscopic characteristics very close to that of the native (N) state is metastable: at room temperature it converts spontaneously with a half-life of ~1.5 h to the MG state. We named this state quasi-native (Nq). Denaturation of TaAChE in the Nq state by glutathione or cysteine causes rapid release of the bound reagent, and concomitant recovery of most of the enzymatic activity. In contrast, similar denaturation of the MG enzyme produces no detectable recovery of enzymic activity. Transition to the MG state of N TaAChE, as well as of enzyme in the Nq state, is greatly accelerated in the presence of phosphatidylcholine liposomes. Introduction of osmyltes (glycerol, sucrose, tetramethyl-N-oxide), as well as of certain divalent cations (Mg2+, Ca2+, Mn2+) retards transition of both the N and Nq states to the MG state. The mechanisms underlying transition of the N and Nq states to the MG state both in the absence and presence of liposomes, as well as the stabilisation induced by chemical chaperons, will be discussed according to the following scheme: N ↔ Nq ↔ MG.
SCAVENGER PROTECTION AGAINST ORGANOPHOSPHATES
BY CHOLINESTERASES

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The failure of current pharmaceutical approaches to provide complete protection against toxic organophosphates and other OP compounds has led to the development of enzyme scavengers to reduce their toxicity. Among the enzymes that hold promise as scavengers, significant advances have been made using cholinesterases (ChEs). These enzymes were found to be effective bioscavengers against a variety of OPs in rodents as well as in non-human primates. Pretreatment of theseus monkeys with fetal bovine serum (FBS) ChE or horse/human serum butyrylcholinesterase (BChE) protected them against 5 LD50 of soman challenge. Monkeys pre-treated with ChEs were devoid of any behavioral incapacitation after such soman challenges. Bioscavengers can afford protection against mortality, as well as all of the adverse physiological/behavioral effects of nerve agent exposure. The use of bioscavengers would provide a capability for extended protection against a wide spectrum of nerve agents and eliminate the need for extensive post-exposure therapy. We have recently isolated and purified several homologous quantities of human serum BChE from Cohn fraction IV and have evaluated its use as a bioscavenger for safety and efficacy. We plan to examine human BChE for its lack of an autoimmune response. Preliminary results with purified monkey serum BChE, when re-administered into monkeys twice at one month interval, indicated that most of the enzyme remained in circulation after both injections. Also no antibody was detected against homologous BChE after the repeated administration of monkey enzyme. The mean retention time of this homologous BChE was approximately two weeks. These results bode well for the use of plasma-derived human BChE as a pretreatment drug for humans.

A COMPLEX ARRAY OF POST-TRANSLATION MODIFICATIONS DETERMINES THE CIRCULATORY LONGEVITY OF ACETYLCHOLINESTERASE IN A HIERARCHICAL MANNER

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The post-translation modifications of the mature forms of acetylcholinesterase (AChE) include processing at the glycosylation and enzyme subunit assembly level. While these modifications do not appear to affect catalysis, they have a remarkable impact on the pharmacokinetic behavior of the enzyme which may limit its use as a therapeutic bioscavenger. Biochemical and genetic intervention in several post-translation processes of AChE were carried out. The various AChE glycopolypeptides generated were subjected to subunit assembly determination, extensive MALDI-TOF structure analysis of their N-Glycans, and pharmacokinetic profiling. Accordingly, disialylated forms of AChE displayed a very rapid removal rate (mean residence time, MRT= ~3 min) regardless of their subunit-assembly state, while partially sialylated forms displayed a longer residence time (MRT= 60-100 min) which could be further improved by their conversion into tetramers (MRT= ~200 min). Fully sialylated AChE exhibited an enhanced mean residence time (MRT= 400 min) as compared to the partially sialylated forms, which could be further extended by tetramerization (MRT= 740-1340 min). Unraveling this hierarchical linkage between post-translation modifications and circulatory retention of AChEs, allowed us to generate recombinant products indistinguishable from the native long-lived plasma-derived enzyme. These findings may be extended to other glycoproteins with pharmaceutical potential.

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PRESYNAPTIC INHIBITION OF CENTRAL ACETYLCHOLINE RELEASE WITH A, LIGANDS: PREVENTION OF CHOLINERGIC CRISIS

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The objective of this study is to explore a new strategy to counteract organophosphate poisoning via administration of receptor-mediated inhibition of acetylcholine (ACh) release.

I.v. administration of the A1 full agonist N2-cyclopentyladenosine (CPA) 1 min after a s.c. soman, sarin or tabun poisoning (1.5 – 2LD50), resulted in (i) prevention/ postponement of chewing, salivation, convulsive activity, and respiratory distress, (ii) improvement of 24 h survival, (iii) low levels of extracellular ACh in the brain. Since partial A1 agonists have only limited cardiovascular effects compared to full agonists, the effectiveness of these partial agonists in attenuating the central evoked ACh release was examined in two ways:

(1) Perfusion striatal neuronal symptomatics from which [1H]-ACh release was evoked by 4-aminopyridine in the presence or absence of A1 receptor agonists.

(2) Inverse brain microdialysis with a dialysis probe in the corpus striatum.

In the striatal symptomatics, CPA caused a dose-dependent inhibition (IC50 = 42 ± 11 %) of the evoked [1H]-ACh release. 3'-deoxy-CPA (3DCPA), 8-propylamino-CPA (8PCPA) and 8-butyramino-CPA (8BCPA) attenuated the release to a similar extent, although at 10-fold higher dosages. Microdialysis experiments demonstrated similar results on the inhibition of ACh release by CPA and its analogues, although higher IC50 values were obtained. In conclusion, CPA, 3DPCPA and the 8-alkyl analogues reduced the ACh release in the striatum area both enrothropic and enrothropic in a concentration-dependent manner and appeared to be as efficacious as the full agonist CPA, suggesting that central inhibition of ACh release may be obtained without the above-mentioned side effects. The present results encourage further investigation of the therapeutic potential of these low efficacy agonists for the treatment of organophosphate poisoning.

MEASURING CEREBRAL ACETYLCHOLINE ESTERASE ACTIVITY IN ALZHEIMER DEMENTIA BY PET FUNCTIONAL PARAMETRIC IMAGING

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Background and Purpose: In-vivo measurement of the expression of acetylcholine esterase (AChE) is of primary interest in dementia because of the crucial role of the cholinergic system in memory and attention. 11C-labeled N-methyl-4-piperidine-acetate (11C-MP4A) is a tracer that is highly specific for AChE and has kinetic properties that are favorable for measuring cortical AChE activity. Using this tracer, we describe the results of a new method for standardized parametric images in Alzheimer dementia (AD).

Methods: We studied eleven patients with probable AD (NINCDS-ADRD criteria, mean age 62.8 years, mean mini mental status score 21.2) and six normal control subjects. Subjects were examined with dynamic PET (Siemens/CTI EXACT) in 3D-mode over 60 minutes after injection of 550-740 MBq 11C-MP4A. A dynamic sequence of 6x30 seconds, 2x60 seconds, 2x150 seconds, 10x300 seconds scans was used after acquisition of a 10 min transmission scan with 3 Ge-68 rods. Images were normalized to stereotactic space. Parametric images of tracer hydrolysis by AChE (kS) were generated by a recently developed voxel-wise dynamic curve fitting procedure using the putamen as a reference tissue. To localize the regions that demonstrate a significant change of AChE activity two sample t-tests were performed voxel by voxel using SPM99.

Results: Mean cortical kS values of late onset subjects (kS = 0.072 ± 0.0009/min) and early onset AD subjects (kS = 0.0686 ± 0.00072/min) were significantly reduced compared to controls (kS = 0.1019 ± 0.0055 min). The corresponding significance levels were p = 4.6 10-4 and p = 1.2 10-6, respectively. The most significant reduction of AChE activity was located in inferior temporal cortex.

Conclusion: Noninvasive measurement of local cerebral AChE activity with PET is possible and can be applied in a clinical setting. Severe reduction of AChE activity is present in mild to moderate AD of early and late onset. This technique has the potential to contribute to differential diagnosis of dementia, to measure cerebral AChE inhibition by drugs, and to identify individuals with particularly severe cholinergic deficits who may benefit most from treatment with cholinergic agents.

References:
BLOOD-BRAIN BARRIER DISRUPTION IS ASSOCIATED WITH
ABNORMAL CORTICAL THETA RHYTHM GENERATION: THE
POTENTIAL INVOLVEMENT OF ACETYLCHOLINESTERASE

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Numerous pathological diseases of the central nervous system have been reported to involve perturbation of the integrity of the blood-brain barrier (BBB). Although the mechanisms underlying BBB disruption and the possible risk-factors are still unknown, previous studies point to possible AChE involvement. We have developed a fast technique for the analysis of computerized tomography (CT) images, thus enabling the screening of patients for perturbation of the BBB. Our data show frequent, focal or diffuse BBB disruption in a wide range of neurological disorders. Clinical data, blood and CSF biochemical analyses pointed to general stress-associated mechanisms as the common denominator, rather than a single disease process. Moreover, the radiological data suggest that BBB dysfunction may persist for at least several weeks. To explore the possible effects of BBB disruption on cortical activity, patients with focal BBB disruption where examined with high density 128 channel electroencephalography. Abnormally high power theta activity was noted in all patients with BBB disruption. In 9 out of 13 patients examined, low-resolution electrotomography (LORETA), localized the abnormal slow wave origin in the same brain region as the anatomical BBB lesion. In a separate animal model, electrophysiological recordings in mouse brain slices suggested cholinergic-mediated enhanced glutamatergic activity as an important contributor to abnormal cortical activity. Our data suggests that persistent BBB disruption may underlie abnormal brain activity and resultant neurological disorders.

CHEMICAL KINETICS PARAMETERS AND RECEPTOR
DEGRADATION RATES AT THE NEUROMUSCULAR JUNCTION

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Two projects will be described which were started by Mika Salpeter before her tragic and untimely death. One project presents an exhaustive catalog of modeling results for miniature endplate currents (mepcs) for many choices of numerical values for chemical kinetics (and other) input parameters. The catalog will be available to the e-mail reader. In relation to Myasthenia Gravis, one can follow changes in mepc risetime and amplitude as cleith width and AChR site density are varied separately or together. The second project concerns acetylcholine receptor (AChR) site densities (SD) at the neuromuscular junction during two transitional periods, denervation and reinervation. EM autoradiography was used. The simplest theoretical model predicts an upward transient in SD during denervation and a downward transient during reinervation. The former is observed, but not the latter.

Degradation experiments suggest that a form of AChR with a 3.5 day half-life is inserted during a five day reinervation period (faster than intact AChR; slower than embryonic AChR). It is inserted at a large enough rate to keep SD constant, so that mepc amplitude can already go to the full adult value during this transition period.

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ANTIBODIES TO ACETYLCHOLINE RECEPTORS AND MUSK IN
MYASTHENIA GRAVIS AND RELATED DISORDERS

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Antibodies to acetylcholine receptors (AChR) cause the autoimmune disease, myasthenia gravis (MG), which results in muscle weakness and fatigue. About 15% of patients with classical MG do not have antibodies to the muscle AChR. In vitro their serum antibodies appear to act by disturbing an intracellular signalling pathway that leads to AChR phosphorylation. In collaboration with Dr Werner Hoch (Max Planck Institute for Developmental Biology) we showed that 70% of MG patients without AChR antibodies, have antibodies to the muscle specific kinase, MuSK (Hoch et al Nat Med 7, 365, 2001). These antibodies can inhibit agrin-induced, MuSK-dependent, AChR clustering in vitro. Thus these antibodies are potentially pathogenic and are likely to reduce the number of AChRs or alter the structural integrity of the neuromuscular junction. A small proportion of MG women transfer a transient MG to their babies, so called neonatal MG. Very occasionally, the baby is born with severe multiple joint contractures. Studies with Dr David Beeson, using adult and fetal AChR expressed in Xenopus oocytes, demonstrated the presence in these mothers of antibodies that specifically inhibit the fetal isoform of the AChR. Moreover, if the human antibodies to fetal AChR are injected into pregnant mice, the mouse fetuses become paralysed and develop joint contractures. Antibodies to voltage-gated calcium channels and voltage-gated potassium channels are involved in other neurological disorders. These antibodies can also affect the autonomic and central nervous system causing a range of different symptoms including autonomic dysfunction, cerebellar ataxia, or sleep and memory disorders. Autoantibodies to specific ion channels at cholinergic synapses are responsible for a range of different neurological disorders.

-11-
INHIBITORY EFFECTS OF MUSCARINIC RECEPTOR AUTOANTIBODIES ON PARASYMPATHETIC NEUROTRANSMISSION IN SJÖGREN'S SYNDROME

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Sjögren's syndrome (SS) is an autoimmune disorder characterized by dry eyes and mouth (sicca syndrome) and lymphocytic infiltration of lacrimal and salivary glands. Abnormalities of parasympathetic neurotransmission may contribute to the glandular dysfunction. We developed a functional assay to investigate autoantibody-mediated effects on parasympathetic neurotransmission.

Serum and purified IgG was obtained from patients with primary and secondary SS and controls. Contraction of isolated murine bladder strips in response to stimulation of M1-muscarinic receptors by carbachol or by endogenous acetylcholine released from postganglionic parasympathetic nerves was measured before and after the addition of patient serum or IgG. Sera from 5/9 patients with primary SS and 6/6 patients with secondary SS inhibited carbachol-evoked bladder contraction by approximately 50%, as well as the action of neuronally released acetylcholine at M1-muscarinic receptors. Sera from 7/8 healthy individuals and 8/8 disease controls had no effect. Anti-muscarinic receptor activity was localised in the IgG fraction. The autoantibodies were associated with bladder symptoms and other autonomic features.

Conclusion: Autoantibodies that act as antagonists at M1-muscarinic receptors on smooth muscle occur in patients with primary and secondary SS. These autoantibodies appear to contribute to sicca symptoms and may explain associated features of autonomic dysfunction. Development of screening assays will enable study of their broader clinical relevance.

IMMUNOTHERAPY OF MYASTHENIA GRAVIS: ANTIGEN-SPECIFIC MUCOSAL TOLERANCE AND ANTAGONISTS OF KEY CYTOKINES AND COSTIMULATORY FACTORS

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Developing immunotherapeutic approaches for myasthenia gravis has been our goal for many years. Mucosal administration of recombinant fragments corresponding to the human acetylcholine receptor (AChR) α-subunit suppresses ongoing experimental autoimmune myasthenia gravis (EAMG) in rats. Treated animals exhibit a Th1 to Th2/Th3 shift in their cytokine profile and down-regulation of costimulatory factors. In severely affected rats, this antigen-specific approach may need to be supported by direct modulation of key cytokines and costimulatory factors known to be involved in the pathogenesis of EAMG. To address this question, myasthenic rats were injected by antibodies either to the proinflammatory cytokine IL-18 or to the costimulatory factor CD40L. These treatments act via different mechanisms but both lead to alleviation of clinical symptoms even when given at the chronic phase of EAMG. Both impaired AChR-specific Th1 cell differentiation with no effect on Th2-type responses, and treatment by anti-IL-18 antibodies led to elevation of the Th3-type suppressive cytokine, TGF-β and to generation of regulatory cells. Anti-CD40L administration led to a significant decrease in humoral responses, whereas anti-IL-18 treatment affected mainly cellular responses to AChR. The most significant suppressive effect of both treatments was observed 2-3 weeks after initiation of treatment and was later diminished, implying that blockade of either IL-18 or CD40L may not be sufficient to suppress chronic myasthenia. We therefore suggest that antagonists to key cytokines and/or costimulatory factors be used in conjunction with antigen-specific treatments such as mucosal administration of AChR recombinant fragments.

THE ALPHA7 NACHR L250T MUTATION IN MICE: A MODEL FOR EPILEPSY IN MEN

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Alpha7 nicotinic acetylcholine receptors (nACHRs) are sparsely distributed throughout the peripheral and central nervous systems. Several studies have suggested that central alpha7 nicotinic receptors may influence sensitivity to nicotine-induced seizures in mice. In order to investigate the role alpha7 in seizure, we tested heterozygous mice with a threonine for leucine substitution at position 250 (+/L250T) within the channel domain of the alpha7 subunit. This mutation was previously shown to cause a partial gain of function in this receptor subtype (i.e., increased current amplitude and agonist affinity and decreased desensitization of the receptor). We demonstrated that administration of low doses of nicotine to alpha7+/L250T mice significantly increased the sensitivity to nicotine-induced seizures. EEG recordings and spectral analysis of these mutants showed high amplitude rhythmic activity that was in correlation with the behavioral changes induced by nicotine. Pretreatment with the alpha7 nicotinic receptor antagonist methyllycaconitine (MLA) inhibited the seizures induced by nicotine. Our data suggest that the alpha7 subunit is involved in seizure generation. Since the alpha7 L250T mutation resembles the human alpha4 and beta2 nACHR mutations found in autosomal dominant nocturnal frontal lobe epilepsy, the +/L250T animals may serve as an important tool to study the mechanism and treatment of human epilepsy.

THE MOLECULAR NEUROBIOLOGY OF ACETYLCOLINERESTASE VARIANTS: FROM STRESSFUL INSULTS TO ANTISENSE INTERVENTION

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The early discovery of the acetylcholine hydrolyzing enzyme, acetylcholinesterase (AChE) and its function in terminating cholinergic neurotransmission made it the focus of intense research for much of the past century. More recent studies on the complexity of AChE gene regulation and the accumulating evidence for some of the long-suspected "non-classical" actions of this protein call for exploring the molecular neurobiology of AChE splicing variants. To this end, we have combined transgenic animal models with genomic and cell culture studies aimed at revealing the enzymatic and morphogenic roles of 3-alternative splicing products of the human AChE gene under normal and disease conditions. There are three, not one AChE variants, each with a different C-terminal peptide, capacity for multimeric assembly and potentially distinct non-classical function(s). Both stressful stimuli and exposure to AChE inhibitors induce transcriptional activation and shift alternative splicing toward overproduction of the normally rare "readthrough" variant, AChE-R. This may be beneficial for suppressing the initial insult, but detrimental under long-term conditions. Partially 2'-O-methylated oligodeoxynucleotides capable of ameliorating AChE-R overproduction emerge as promising research tools for proving the putative involvement of AChE-R in such detrimental consequences, as well as potential therapeutic agents. Specific advantages of such agents stem from their degree of variant selectivity, which enables maintenance of cholinergic neurotransmission while preventing the stress-induced imbalance among AChE variants.
ACETYLCHOLINESTERASE FACILITATES AMYLOID DEPOSITION IN A MOUSE MODEL OF ALZHEIMER'S DISEASE

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Alzheimer's type amyloid plaques contain acetylcholinesterase (AChE), among other proteins. In vitro, AChE associates with beta-amyloid (A-beta) and hastens formation of insoluble amyloid fibrils in a manner that might promote plaque formation (Alvarez et al., 1998). To determine if this effect is clinically relevant for Alzheimer's disease, we investigated Tg2576 mice, which overexpress human amyloid precursor protein (hAPP) and develop plaques at about 9 months (Hsiao et al., 1996). Tg2576 mice were crossed with mice harboring a human AChE transgene (Brieri et al., 1996) to yield F1 hybrids that overexpress both hAChE and hAPP in brain. By 6 months, the cerebral cortex of the hybrid mice demonstrated plaques reacting with thioflavin S and with antibodies to A-beta 1-40 and 1-42. Plaque burden at this stage was modest (~125 plaques per hemisphere) and quantitative ELISA showed very little SDS-insoluble, formic acid extractable A-beta, but levels of SDS-soluble A-betawere about 25% above control. By 9 months of age, plaques had become larger and more numerous, while formic acid extractable A-beta 1-40 and 1-42 had reached levels double those in age-matched controls. These results suggest that direct interactions with AChE can promote amyloid deposition in brain, initially as an atypically loose, detergent soluble matrix. Therefore, drugs designed to target protein-protein interactions between AChE and A-beta should be tested for potential to retard progression of Alzheimer's disease.

NEURODEGENERATIVE PROCESSES IN ALZHEIMER'S DISEASE

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Alzheimer's disease (AD) is a progressive dementia paralleled by selective neuronal death, which is probably caused by the cytotoxic effect of the amyloid beta-peptide (Abeta). We have previously shown that the small-plaque component acetylcholinesterase (AChE) induces amyloid fibril formation, forms macromolecular complexes (Abeta-AChE) being incorporated into the growing fibrils, and that such complexes increased amyloid neurotoxicity. Therefore, we have searched for molecular changes induced by Abeta-AChE complexes, both in neuronal cells in vitro and in rats injected in the dorsal hippocampus with Abeta/Abeta-AChE complexes, as an in vivo model of the disease. Here, we will discuss that Abeta/Abeta-AChE neurotoxicity results in the destabilization of endogenous levels of key components of the Wnt signal transduction pathway. Moreover, activation of this signaling cascade either with lithium, protein kinase C (PKC) agonists or with conditioned media containing the Wnt-3a ligand, induce survival of post-mitotic neurons against neurotoxicity and rescue the deficit in spatial learning induced by such cytotoxic agents. Given that it has been recently observed that nicotine receptors (vz. alpha7AR), which may signal through PKC isoenzymes, binds Abeta with nanomolar affinities, we have also examined the ability of its agonists/antagonists (i.e. alpha-bungarotoxin) to modulate Abeta/Abeta-AChE neurotoxicity. Our results suggest that Abeta/Abeta-AChE dependent neurotoxicity results in loss of function of Wnt signaling components and indicate that compounds that mimic this signaling cascade may be candidates for therapeutic intervention in Alzheimer's patients.

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TRANSCRIPTIONAL CONTROL OF THE CHOLINERGIC GENE LOCUS (CGL): A MOSAIC MODEL FOR REGULATION OF THE CHOLINERGIC PHENOTYPE

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While VACHT and ChAT transcription are coordinated, low levels of R-exon expression indicate independent transcription of each gene from the rodent CGL in vivo (Hsiao et al., J. Mol. Neurosci. 9, 223, 1997; Schuetz et al., Neuroscience 104, 633, 2001). Co-expression of VACHT and ChAT with the high-affinity choline transporter, transcribed from a separate gene (Okuda et al., Nat. Neurosci. 3, 120, 2000), is much less tightly coupled in certain cholinergic neurons. An 8.7 kb fragment of the human CGL extending ~2 kb past the VACHT gene is sufficient for VACHT expression in somatomotor neurons of transgenic mice (Schuetz et al., Neuroscience 96, 707, 2000). An additional 2.5 kb downstream fragment is required for diencephalic expression (Schuetz et al., this meeting), in contrast to expression from the murine CGL in transgenic mice (Naciff & Dudaen, J. Neurochem. 72, 17, 1999).

Co-transmitter expression in cholinergic neurons is also highly species-specific. Cholinergic neurons of the rat stellate ganglion co-express VACHT and TH before birth, and at sweat gland terminals soon after. TH expression is switched off in the adult rat, but persists in the adult mouse and primate. Human sweat gland terminals are also positive for VMAT2, consistent with human 'noradrenergic sweating'. Cyclohexylamine(COX), rate-limiting for prostaglandin biosynthesis, is co-expressed in cholinergic nucleus basalis neurons in primates, and irreversibly down-modulated in simian AIDS. Thus, prostaglandins may be involved in inflammation-associated cholinergic dysfunction.

These findings suggest a 'mosaic' model for CGL transcription that allows fine-tuning of cholinergic traits, and differential co-transmitter expression, in different types of cholinergic neurons.

NEURONAL NICOTINIC RECEPTORS, ALLOSTERIC POTENTIATING LIGANDS (APs), AND ENDOGENOUS METABOLITES: IMPLICATIONS FOR TREATMENT OF ALZHEIMER'S DISEASE (AD)

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The severity of symptoms in AD, a progressive neurodegenerative disorder that affects millions worldwide, is closely related to the extent of reduction of nicotinic cholinergic activity in the brain. Thus, augmenting nicotinic functions has emerged as a promising therapeutic approach for treatment of AD. Research from our laboratories has demonstrated that nicotinic receptor (nAChR) activity can be increased by substances referred to as APs. Galantamine, a weak anticholinesterase, is the prototypical AP. By increasing activity of presynaptic nAChRs, galantamine causes sustained facilitation of synaptic transmission in the brain. Methamidophos, Metrifonate, Tacrine and Donepezil, in contrast, are anticholinesterase agents devoid of AP activity, and they facilitate synaptic transmission only transiently. Recent studies also revealed that (i) kynurenic acid, a tryptophan metabolite whose levels are increased in the brain of AD patients, is a non-competitive antagonist at alpha7 nAChRs (IC50, 7 micromolar) and increases alpha4beta2 nAChR expression in the brain, and (ii) acetyl-L-carnitine, a metabolite whose levels are decreased in the brain of AD patients, potentiates nAChR activation by nicotinic agonists. These findings lay the groundwork for development of new therapeutic avenues for treatment of patients with AD.

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THE RATIONALE FOR USING GALANTAMINE TO TREAT DIFFERENT DEMENTIA TYPES

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Objective: Considerable evidence has established that impaired cholinergic neurotransmission is central to Alzheimer's disease (AD). Cholinergic deficits related to nicotinic acetylcholine receptors (nAChR) coupled with degeneration of neuronal circuits are now believed to contribute to cognitive impairment across a range of dementias, including vascular dementia (VaD). Here, we present a rationale for using drugs that affect nAChR activity to treat dementia. Methods: We examined preclinical and clinical evidence to define the nature and contribution of cholinergic impairment to symptoms of AD and other dementias to determine a rational treatment strategy. Results: Human and animal models indicate that, regardless of the underlying cause of dementia, cognitive deficits involve decreased nicotinic cholinergic neurotransmission and reduced numbers of nAChR. Cholinergic function is compromised in animal models of VaD. Studies have shown reduced binding of nAChR ligands in the cortex and hippocampus of spontaneously hypertensive stroke-prone rats. Decreased choline acetyltransferase activity has also been observed in the cortex, hippocampus and striatum of VaD patients compared with controls. Patients withBinswanger or multiple small infarct VaD subtypes have lower cerebral fluid concentrations of acetylcholine. Galantamine (Reminyl(R)), an acetylcholinesterase (AChE) inhibitor that allosterically modulates nAChR, may be particularly useful for treating various dementias. Recent research has shown that, as well as having broad, long-term efficacy in AD, galantamine is effective in patients with VaD and AD with concomitant cerebrovascular disease. Conclusion: Drugs like galantamine, with a nicotinic mode of action in addition to AChE inhibitory activity, provide an effective treatment option for various types of dementia.

MOLECULAR AND FUNCTIONAL DIVERSITY IN NICOTINIC ACETYLCHOLINE RECEPTOR GENE FAMILIES OF C. ELEGANS AND D. MELANOGASTER

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Genetics, genomics and electrophysiology are transforming our understanding of the largest known gene family of nicotinic acetylcholine receptor (nAChR)27 subunits in C. elegans (27 members) and the exclusively neuronal nAChR gene family in D. melanogaster (10 members). In C. elegans, several genetic screens have enabled identification of nAChR subunits, along with novel proteins that act upstream and downstream of functional nAChRs. The C. elegans genome project has identified many new candidate nAChR subunits and the calculated electrostatic potential energy profiles for the M2 channel-lining regions predict considerable functional diversity. The respective roles of subunits are under investigation using forward and reverse genetics. Electrophysiological and reporter gene studies have demonstrated roles for particular subunits in levamisole-sensitive muscle nAChRs and a role for nAChRs in pharyngeal pumping. Recombinant homomeric and heteromeric C. elegans nAChRs have been expressed in Xenopus laevis oocytes. In D. melanogaster, three new nAChR subunits have been cloned, one of which shows multiple variant transcripts arising from alternative splicing and A-to-I pre-mRNA editing. Thus, studies on the genetic model organisms C. elegans and D. melanogaster have revealed different routes to generating molecular and functional diversity in the nAChR gene family and are providing new insights into the in vivo functions of individual family members.

GENETIC DISSECTION OF AN ACETYLCHOLINE RECEPTOR INVOLVED IN NEURONAL DEGENERATION

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The C. elegans DEG-3 gene codes for a subunit of a nicotinic acetylcholine receptor that can mutate to cause neuronal degeneration. The degeneration causing mutation, deg-3(n662), is a gain of function mutation, affecting a residue in transmembrane domain II, that apparently interferes with receptor desensitization. In order to identify genes needed for DEG-3 activity we screened for mutations that suppress the behavioral defects associated with the deg-3(n662) mutation. This screen led to the identification of mutations in three genes: deg-3 itself, des-2 an acetylcholine receptor subunit, and ric-3 a novel gene specifically required for the maturation of acetylcholine receptors. Co-expression of these genes in Xenopus oocytes shows that DES-2 is essential for DEG-3 channel activity and that RIC-3 affects channel activity both quantitatively and qualitatively. The reconstruction of DEG-3 channel activity in oocytes also suggests that des-3(n662)-induced degenerations are a result of constitutive activity of the mutant channel in the presence of physiological choline concentrations; choline is an agonist of the DEG-3 channel. High calcium permeability of this channel may also contribute to the degeneration process. Thus genetic analysis has identified elements affecting acetylcholine receptor activity and is providing insights into the degeneration process.

A COMMON AGONIST AND POTENTIATOR FOR α7 NICOTINIC AND 5-HT, SEROTONIN RECEPTORS

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α7 nicotinic receptors (α7 nAChRs) as well as serotonin 5-HT receptors (5-HT(R)s) belong to the family of ligand-gated ion channels. Two receptors exhibit some cross-pharmacology, e.g., high concentrations of the α7 nAChR agonist nicotine inhibit 5-HT(R)-mediated responses, and high concentrations of the 5-HT(R) agonist 5-HT inhibit α7 nAChR-mediated responses.

Using voltage clamp and calcium imaging techniques, we are studying the pharmacological properties of 5-HT(R)s in murine NIE-115 neuroblastoma cells and of cloned human 5-HT(R)s and α7 nAChRs expressed in Xenopus oocytes. We have found that a recently described α7 nAChR agonist (Philips et al., Astra Aesc USA, patent WO99/03859) is activating human α7 nAChRs expressed in oocytes. However, the same compound also activates human 5-HT(R)s. The EC50, nH and Emax are 2.2 μM, 0.8 and 83% for α7 nAChRs, and 1.4 μM, 1.8 and 66% for 5-HT(R)s, respectively. In the presence of 30 μM D2-6E5, this agonist still induces robust inward currents and raises [Ca2+] in NIE-115 cells. These responses are blocked by the selective 5-HT(R) antagonist MDL 72222.

5-Hydroxytryptophol (5-HT) potentiated 5-HT(R)- as well as α7 nAChR-mediated responses. The threshold concentration of 5-HT to potentiate both types of receptor is 100 μM, and maximum potentiation is observed at both 1 mM 5-HT. On both types of receptors, 1 mM 5-HT shifts the agonist concentration-effect curve towards lower agonist concentrations and enhances the maximum effect of both ACh and 5-HT.

In this study, the general notion that 5-HT(R)s and α7 nAChRs share some pharmacological properties is extended by the finding that both types of receptor share a common agonist and a common potentiator.
BETA-AMYLOIDS, TAU HYPERPHOSPHORYLATION AND COGNITION ARE BENEFICIALLY AFFECTED BY M1 MUSCARINIC AGONISTS - PERSPECTIVES IN ALZHEIMER'S DISEASE TREATMENT

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M1 agonists from the AF series [AF102B (prescribed in USA for Sjögren's Syndrome), AF150(S) & AF1267B] - i) restored cognitive impairments in several animal models for Alzheimer's disease (AD) with an excellent safety margin, ii) elevated the non-amyloidogenic amyloid precursor protein (alpha-APPS) levels, iii) attenuated vicious cycles induced by beta-amyloid (Abeta), and inhibited Abeta- and oxidative-stress-induced apoptosis; and iv) decreased tau protein hyperphosphorylation in vitro and in vivo [review: Fisher Jpn J Pharmacol 84: 101, 2000]. Unlike M1 agonists, nicotinic agonists and cholinesterase inhibitors increased tau hyperphosphorylation [Hellsén-Lindahl et al J Neurochem 74, 777, 2000]. In aged mice, Aβ(40) (natural model for AD; Bons. Alz. Res 1, 83, 1995), prolonged treatment with AF150(S) restored cognitive and behavioral impairments and decreased tau hyperphosphorylation, paired helical filaments and astroglia [Bons. Maestro-Frances and Fisher, unpublished]. In rabbits, with Abeta sequence identical to the human Abeta. AF267B & AF150(S) decreased CSF Abeta1-42 & 1-40, while AF102B reduced Abeta1-40 [Beach et al Brain Res, 905, 226, 2001]. Finally, AF102B decreased CSF Abeta42 in AD patients [Nitsch et al Ann Neurol, 48, 913, 2000]. In summary, M1 agonists may represent a unique therapy in AD due to their combined beneficial effects on the three major hallmarks of AD - cholinergic hypofunction, Abeta and hyperphosphorylated tau.

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CROSS-TALK BETWEEN APOLIPOPROTEIN E, THE AMYLOID PRECURSOR PROTEIN AND BRAIN INFLAMMATION

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Transgenic mice expressing either human apoE3 or apoE4 on a null mouse apoE background were employed to investigate the role of isoform-specific interactions between apoE: the amyloid precursor protein (APP) and brain inflammatory processes in the phenotypic expression of the pathophysiological effects of the apoE genotype. Immunoblot measurements of the brain levels of APP and of its soluble fragment APPs, revealed that they were lowest in the apoE4 transgenic mice. Closed head injury increased the levels of brain APPs of all mice groups. This effect was markedly and significantly larger in the apoE3 transgenic mice than in the other mice groups.

LPS was injected to the mice i.e.v. and the resulting levels of activation of microglia and astrocytes in the brains of the different mice groups were monitored immunohistochemically. This revealed that LPS treatment of 12 months old mice triggered astrogliosis in the apoE4 deficient and in the apoE4 transgenic mice, but not in the control and apoE3 transgenic mice. In contrast, LPS dependent microglial activation was not significantly affected by either apoE deficiency or the human apoE genotypes.

In conclusion, the present findings show that APP metabolism and brain inflammatory processes are modulated in vivo in an isoform specific manner by apoE, and suggest that the pathological effects of apoE4 in Alzheimer's disease may be mediated by similar cross talk interactions.

CHOLINEnergic DEFICITS AND NON-CogNITivE BEHAVIOURAL CHANGES IN PATIENTS WITH DEMENTIA


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Non-cognitive behavioural changes such as depression, aggressive behaviour, psychosis and overactivity occur frequently in patients with dementia, in addition to cognitive impairment, and often determine the need for institutionalisation. The biochemical basis of such changes is poorly understood. Clinical trial data indicate that cholinomimetics improve non-cognitive behaviours. We wished to investigate the relationship between markers of the cholinergic neurotransmitter system and non-cognitive behavioural symptoms assessed during the course of declining illness.

Brains from 46 patients with dementia (36 with Alzheimer's disease, AD, and 10 with mixed or other dementias) were examined together with 32 normal controls.

The patients with dementia had been evaluated every 4 months, often over several years, for cognitive performance (Mini-Mental State Examination) and behaviour (Present Behavioural Examination). Choline acetyltransferase activity (ChAT) and density (Bmax) of muscarinic M1 and M2 receptors in frontal and temporal cortex were studied by radioligand binding protocols. None of the patients were receiving cholinomimetic drugs. ChAT activity and M2 density, but not other neurochemical markers, were reduced in AD compared with controls. Loss of ChAT activity correlated with cognitive impairment. Lowered ChAT activity also correlated with increasing overactivity in patients with dementia in both frontal and temporal cortex while M2 receptor density was increased in frontal cortex of AD patients with delusions and in temporal cortex of those with hallucination compared to patients without psychotic symptoms. Disturbance of the cholinergic system may underlie both cognitive and some non-cognitive behavioural changes in dementia, providing a basis for rational therapy.

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ACHETYCHOLINERSTERASE REGULATION IN SKELETAL MUSCLES

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Muscle acetylcholinesterase (AChE) is concentrated in the rat neuromuscular junctions (NMJ) as the asymmetric (A) molecular form, composed of the catalytic tetramers and the collagen Q (ColQ) subunit. In immature muscles, ColQ is expressed all along the fibers. It is segregated to the NMJs in mature fast muscles but not in the slow soleus muscle. Distribution of the A AChE forms behaves accordingly to the ColQ expression. Regulation of the expression of the extrajunctional A AChE forms in muscle fibers is a nerve dependent phenomenon. A forms of AChE in the NMJ are bound to the basal lamina largely by ionic interactions involving Ca2+, but about 1/3 is bound more firmly. Expression of the extrajunctional AChE in muscle fibers is higher in the fast than in the slow rat muscles, in accordance with about 3-4-fold difference in AChE mRNA level. This is probably due to different patterns of muscle activation of the two types of muscles because the level of AChE mRNA is high in phasically activated fibers and decreases after low-frequency tonic stimulation. The soleus is an antigravity muscle and differs from fast muscles also in regard to loadbearing, but muscle load is not decisive for differences in AChE expression. However, thyroid hormones have an enhancing effect on the level of muscle AChE mRNA. Hyperthyroidism increased AChE mRNA levels in the slow muscle (but not in the fast), whereas hypothyroidism decreased AChE expression in fast and slow muscles.
CHOLINERGIC CORTICAL TERMINATIONS ESTABLISH CLASSICAL SYNAPSES AND UNDERGO AGE-RELATED ATROPHY

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For many years it has been assumed that cortical cholinergic synapses communicate with target-neurons in a "non-synaptic" fashion (volume transmission). We have recently re-examined this problem at the ultrastructural level by applying a highly reliable marker of presynaptic cholinergic boutons: antibodies against the vesicular acetyl choline transporter (VACHT). Our observations demonstrate that - contrary to the above view - cholinergic neurons in the cerebral cortex establish classical synapses. Thus, the electron microscopical analysis of lamina V in the rat cerebral cortex revealed that the majority of cholinergic boutons (i.e. those immunoreactive to VACHT) established typical axo-dendritic synapses of the symmetric type, and are rarely seen in cell soma or dendritic spines. When they are seen in the latter cases, they were asymmetric synaptic contacts. Combining intracellular labelling of tissue slices with high-resolution immunocytochemistry, we observed a preferential relationship of cholinergic boutons with dendritic shafts of pyramidal neurons. We further investigated the fate of cholinergic synapses during aging. This revealed that the age-related loss of cholinergic boutons precedes that of the overall presynaptic population. In addition, cholinergic pre-synaptic boutons undergo a marked atrophy with aging. These manifestations of cortical cholinergic synaptic disconnection should be the major reason for cholinergic participation in age-dependent cognitive decline.

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DEVELOPMENT OF CHOLINERGIC PROJECTIONS TO CORTEX: POSSIBLE ROLE OF NEUROTROPHINS IN TARGET SELECTION

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The goal of this project is to understand how growing cholinergic axons identify and form synaptic contacts with their target cells; the septal cholinergic projection pattern to the dentate gyrus was used as a model system. In the first set of studies, distributions of cells expressing mRNA for selected neurotrophins were studied in developing rats and mice using either autoradiographic or immunocytochemical techniques. These studies demonstrated that NT-3 and BDNF expression develops in dentate gyrus granule cells in spatial and temporal patterns that parallel, and slightly precede, the patterns of ingrowth of septally derived cholinergic axons. Patterns of NGF mRNA expression were not similar to patterns of cholinergic ingrowth. In a second set of studies, organotypic slice cultures were prepared using slices of medial septum paired with a slice of hippocampus, including the dentate gyrus. Normal co-cultures made from tissue from rats or wild-type mice display organotypic patterns of septal cholinergic ingrowth into the dentate gyrus. In contrast, chimeric co-cultures made from septum from wild type mice and hippocampus from NT-3 null mice show profuse cholinergic axonal growth into the hippocampus, but the organotypic pattern of axons in the dentate gyrus is not formed. Instead, AChE positive axons in these cultures grow directly through the dentate gyrus without forming terminal branches in the molecular layer. These data suggest that expression of NT-3 by dentate gyrus granule cells serves to attract or retain septum-derived cholinergic projections to the inner molecular layer of the dentate gyrus.

ACTIVATION OF THE CHOLINERGIC SYSTEM DURING COGNITIVE PROCESSES

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The nucleus basalis-cortical-amygdalar, and septo-hippocampal cholinergic pathways play a critical role in information processing. However, the cognitive events in which they are activated are not yet fully identified. Therefore, the aim of our investigations was to study the relationship between ACh release from cerebral cortex, hippocampus and basolateral amygdala and behaviors in adult male rats, used to be handled, during exploration of a novel environment (arena), habituation, and fear conditioned response. ACh release was investigated by microdialysis coupled to HPLC detection and quantification.

Three 5 min exposures to an arena, with 60 min intervals, were accompanied by a rapid two fold increase in cortical ACh release followed by a prompt drop to basal levels when the rats returned to the home cage. If the exposures lasted 30 min, cortical and hippocampal ACh outputs were significantly smaller during the second exposure than during the first. Motor activity increased mainly during the first exploration of the arena and was significantly correlated to cortical but not hippocampal ACh release. These results indicate that the cortical and hippocampal cholinergic pathways are activated during exploration of a novel environment, probably subserving arousal and attention, but not when habituation takes place. In the amygdala, the facilitation of fear response consolidation by H3 histamine receptor agonists and the impairment by H3 receptor antagonist are associated with an increase or decrease, respectively, in ACh release, suggesting its role in fear memory formation.

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PRECLINICAL AND CLINICAL STUDIES ON THE ROLE OF MUSCARINIC RECEPTORS IN THE PHARMACOTHERAPY OF SCHIZOPHRENA

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Emerging preclinical and clinical evidence supports a role for cholinergic muscarinic receptors (MR) for novel pharmacotherapy of schizophrenia. Anatomical studies indicate that (MR) are uniquely distributed to modulate sensory information, dopaminergic and glutamatergic neurotransmission which may be dysregulated in psychosis. Hyperactivity of dopamine in limbic tracts and hyporeactivity in cortical regions is hypothesized to cause positive and negative symptoms of schizophrenia, respectively, and MR have been shown to modulate these dopamine tracts. For example, the muscarinic M1/M4 preferring agonist xanomeline increases cortical extracellular dopamine and Fos expression, similar to effects of atypical antipsychotics. In electrophysiological studies, xanomeline with acute and chronic administration decreased firing of the mesocorticobasal dopaminergic A10 tract, but not the motoric A9 tract. Behavioral investigations have shown that muscarinic agonists, like dopamine antagonists, inhibit dopamine-agonist-induced behaviors including hyperactivity, climbing and disruption of prepulse inhibition. models for positive symptoms. Knockout mice with ablated M4 receptors are hyperactive and hyperresponsive to dopamine D1 agonists, suggesting a dynamic balance between dopamine and M4 receptors. Mice with M1 receptors deleted have deficits in cognition and muscarinic agonist-induced in vivo phosphoinositide hydrolysis. Consistent with preclinical studies, preliminary clinical investigation indicates that xanomeline may be effective for pharmacotherapy of psychosis and promoting cognition in schizophrenia. Studies in knockout mice have shown that activation of the muscarinic M2 and M3 receptors produce the majority of parasympathomimetic side effects. Thus, we hypothesize that a combined M1 agonist to promote cognition and a M4 agonist for antipsychotic-like effects would treat the symptom domains of schizophrenia without causing parasympathomimetic side effects.
CENTRAL CHOLINERGIC NEURONS IN CULTURE:
REGULATION OF SURVIVAL AND FUNCTION

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Selective degeneration of basal forebrain cholinergic neurons has been
associated with cognitive impairment seen in Alzheimer's disease. We
examined the properties of cultured septal cholinergic neurons of the rat
brain, grown in presence of growth factors or hippocampal neurons, which
are innervated by cholinergic neurons in vivo. Individual cholinergic
neurons, visualized by selective staining with a fluorescent antibody to the
p-75 NGF-receptor and recorded with a patch pipette expressed spontaneous
activity consisting of both excitatory and inhibitory synaptic currents. The
presence of target hippocampal neurons, but not of NGF, enhanced
spontaneous network activity in septal cultures. Exposure to NGF caused an
increase in the size of cholinergic cell somata and primary dendrites, an
effect that was only partially shared by the presence of hippocampal
neurons. Exposure of cultures to glutamate agonists as well as to H2O2,
resulted in a higher proportion of cell death among cholinergic vs. non-
cholinergic neurons, indicating that the former neurons are more sensitive to
neurotoxic insults. The role of afferent hippocampal neurons in
neuroprotection is currently analyzed. These studies are expected to
contribute to the understanding of the unique role of cholinergic neurons in
cognitive aspects of the brain, and the unique properties of the septo-
hippocampal cholinergic system.

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HOW IS THE BRAIN SUPPLIED WITH CHOLINE, BUT
PROTECTED AGAINST EXCESS CHOLINE?

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In the brain, choline (Ch) is released from the cytoplasm of neuronal cells
by depolarization, is taken up from the blood at elevated plasma levels
(enhanced pharmacologically or nutritionally), and finally is catalytically
released from ACh and from phospholipids. An excessively high extracellular level of Ch leads to nicotinic receptor activation or
desensitization. However, the overall level of (Ch) is remarkably constant
at about 4-6 µM due to homeostatic mechanisms. Surplus Ch is removed
from the brain by a net-release into the venous blood. K+-induced
depolarization of synaptosomes caused the release of Ch due to the fact that
the transmembrane Ch gradient follows the Nernst equation. However, in
brain slices or in hippocampal tissue of adult rats (using the microdialysis
technique) K+ failed to enhance (Ch) apparently due to mechanisms
counteracting the K+-induced Ch release. In contrast, high K+ did elevate
(Ch) in brain slices of one week old rats, which have not yet fully
developed cellular uptake mechanisms. Brief electrical stimulation of
cholinergic nerves caused a biphasic change of the synaptic Ch level: a
transient increase (due to ACh hydrolysis) was followed by a long-lasting
hemicholinium-sensitive decline. Long-lasting activation of cholinergic
pathways by pharmaco-logical or behavioral means lead to a deficit of
synaptic Ch which became rate-limiting for ACh biosynthesis; the reduced
ACh release was reversed by Ch supplementation. On the basis of the above
analysis of Ch homeostasis, the significance of temporal and spatial
fluctuations of the synaptic Ch level is still a matter of speculation.

CONTROL OF ACETYLCHOLINE.Release UNDER
STIMULATORY CONDITIONS BY ITS BIOSYNTHETIC
PRECURSORS; GLUCOSE AND CHOLINE

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We tested the effects of glucose and choline, the biosynthetic precursors of
acetylcholine, on passive avoidance behaviour and hippocampal
acetylcholine release measured by microdialysis in awake mice. Glucose (10
and 30 mg/kg) or choline chloride (6-60 mg/kg), given by intraperitoneal
injection immediately after training, dose-dependently enhanced retention in
an inhibitory avoidance task. Combinations of low doses of glucose (10
mg/kg) and choline chloride (20 mg/kg) which alone were submaximally
effective significantly increased retention latencies in a synergistic manner.
an effect which was sensitive to atropine (0.5 mg/kg). This beneficial effect
vanished when higher doses of glucose or choline were combined. Basal
hippocampal acetylcholine release in mice habituated to their environment
was not affected by administration of glucose and choline. However, when
hippocampal acetylcholine release was stimulated either by infusion of
scopolamine (0.3 microM) or by transferring the mice into a novel
environment, the combination of glucose plus choline further increased
acetylcholine release to a significant extent. We conclude that low doses of
glucose and choline act synergistically to improve memory storage, an
effect which is due to facilitation of acetylcholine release.
This finding reinforces the view that central cholinergic functions are under
certain conditions influenced by dietary intake of precursors.

SIGNALLING PATHWAYS THAT REGULATE THE CHOLINERGIC
GENE LOCUS EXPRESSION

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The overall goal of our studies is to reveal how cholinergic neurons acquire
and maintain their neurotransmitter characteristics. The two genes that
determine the cholinergic phenotype are choline acetyltransferase (ChAT)
and the vesicular acetylcholine transporter (VACHT). They occupy one
genomic locus and are likely controlled by shared transcriptional
mechanisms. We employed model cholinergic cell lines and embryonal
spinal cord and septal neurons to investigate the regulation of ChAT and
VACHT expression by ciliary neurotrophic factor (CNTF) and bone
morphogenetic proteins (BMPs). By using pharmacological inhibitors of
MAPK kinase (MEK1), as well as expression of recombinant constitutively
activated MEK1, we demonstrated the negative effect of the MEK1 pathway
on CNTF-induced cholinergic promoter activity. We reported previously
that BMP-9 is a potent inducer of the cholinergic phenotype in vitro and in
vivo. Transient transfection of reporter constructs into primary neurons from
E14 mice revealed that BMP-9 activates transcription from the proximal -
but not from the distal - promoter region of the cholinergic locus. In search
of candidate immediate transcription factors involved in this BMP action,
we used DNA microarray technology (Affymetric GeneChip) to analyze the
changes in gene expression patterns caused by BMP-9. In rat and mouse
E14 neuronal cultures, BMP-9 activated the expression of multiple genes,
including growth factors, growth factor receptors, and transcription
regulators. It is likely that the products of some of these BMP-9-induced
genes are involved in the modulation of cholinergic gene locus expression.
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TREATMENT OF DEMENTIA WITH CHOLINESTERASE INHIBITORS
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Dementia is a heterogeneous group of disorders with several identifiable etiologies and mechanisms. It is becoming recognized that the border between Alzheimer's disease (AD) and the second most common cause of dementia, vascular dementia (VaD), is blurred and a rational approach should direct itself to cardiovascular factors, such as hypertension, which could prevent AD as well as VaD. The current treatment of patients with dementia includes primarily cholinesterase inhibitors. These agents (donepezil, rivastigmine, and galantamine) have a limited symptomatic effect, expressed on concentration, cognition and behavior, and affecting activities of daily living and independence. However, pathological and biochemical data also point to central cholinergic hypofunction in both disorders. Therefore, it is logical to examine the effect of ChEI's in VaD as well as in cases with mixed dementia.

Limited attempts to affect other parameters (non-cholinergic neurotransmitters and post-receptoral processes) have been limited and have largely failed.

ChEIs are of limited value for several reasons. They cause significant side effects, such as gastro-intestinal and cardio-vascular. Their potential value is also limited because they enhance the activity of the ever-declining acetylcholine. The use of muscarinic agonists in dementia has been limited. Reasons for failure of the previous studies will be outlined, and ways to overcome them will be suggested. In particular, the use of combined ChEI's with M1 agonists will be supported.

CHOLINESTERASE INHIBITORS STABILIZE COGNITIVE DECLINE IN ALZHEIMER'S DISEASE
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Cholinesterase inhibitors (ChEIs) are the only drugs approved in US and Europe for the indication of Alzheimer Disease (AD) treatment.

Long-lasting clinical efficacy of one year or longer produced by ChEI treatment in AD patient suggests that long-term effects of these drugs may not be solely symptomatic and related to an elevation of synaptic acetylcholine levels. Six most extensively clinically tested ChEIs (tacrine, eptastigmine, donepezil, rivastigmine, metrifonate and galantamine) produce a stabilization of cognitive and non cognitive function. Data collected from over 8,000 cases demonstrate that patients responders to treatment with ChEIs change little cognitively and behaviorally from the baseline at the beginning of the study for six-twelve months as compared to placebo treated patients. Long-term effects of one year or longer are seen using four different ChEIs (donepezil, metrifonate, rivastigmine and galantamine). Additional clinical observations suggest non-symptomatic effects. Effects on APP metabolism and release in vitro and in vivo and on beta-amyloid toxicity and aggregation may contribute to long term clinical efficacy seen in AD patients treated with ChEI (1).


GENDER DIFFERENCES IN THE ACTIONS OF CHOLINESTERASE INHIBITORS
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Introduction: Cholinesterase (ChE) inhibitors are currently the most effective treatment for Alzheimer's disease (AD). In normal elderly women and in those with AD, physostigmine produced a larger increase in plasma ACTH, and cortisol than in men, but blood levels did not differ. This study determined whether there was a sex difference in the effects of tacrine and rivastigmine and its relation to hormone levels. Method: The effects of tacrine and rivastigmine were compared on spatial memory impaired by scopolamine, body temperature and ChE activity in males and females.

Results: Both drugs caused significantly greater hyperthermia, antagonism of scopolamine-induced memory impairment and ChE inhibition in the brain, but not in the heart or skeletal muscle in females than in males. The sex difference resulted from higher amounts of drug reaching the female brain. Orchidectomy increased drug activity and brain levels in males to that in females, while ovariectomy had no effect. Adrenalecctomy increased brain levels of tacrine and activity in both sexes. Testosterone decreased brain levels and actions of ChE inhibitors both in castrated males and females.

Conclusions: The pharmacological effects of ChE inhibitors in the brain are greater in females than in males because testosterone reduces the amount of drugs reaching its target sites. It remains to be seen if women with AD are also more sensitive than men to the cognitive effects of these drugs.

AMYLOID PRECURSOR PROTEIN PROCESSING PROPERTIES OF THE NOVEL NEUROPROTECTIVE CHOLINESTERASE - MONOAMINE OXIDASE INHIBITOR. TV3326 AND ITS OPTICAL ITS-ISOMER. TV3279
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TV3326 (N-propargyl-3-(3R)-aminoindan-5-s-ethyl, methyl carbamate) was developed in order to combine the neuroprotective effects of the antiParkinson drug rasagiline, a selective inhibitor of monoamine oxidase (MAO)-B with the cholinesterase (ChE) inhibitory activity of rivastigmine as a potential treatment for Alzheimer's disease (AD). TV3326 is a cholinesterase and brain selective MAO A and B inhibitor, while its optical S-isomer, TV3279, has only cholinesterase inhibitory activity. Both drugs retain many of the cell culture and In Vivo neuroprotective-antianapoptotic properties of rasagiline and antagonize scopolamine-induced impairment in spatial memory. Reports indicate that cholinesterase inhibitors process amyloid precursor protein (APP) via α-secretase cleavage pathway. However their mechanism of action has not been fully established. Both TV3326 and TV3279 stimulated the release of the non-amyloidogenic α-secretase dependent soluble APP (sAPP) in PC12 and human neuroblastoma (SH-SY5Y) cells in a dose dependent (0.1-100μM) manner. This effect was blocked by hydroxymethylacetic acid-based metalloproteinase inhibitor Ro-31-7920, indicating the mediation via α-secretase cleavage. Using signal transduction inhibitors, we have shown that protein kinase C (PKC), mitogen-activated protein (MAP) kinase and tyrosine kinase dependent pathway may be involved in the enhancement release of sAPP by TV3326 and TV3279. In addition both drugs induced the phosphorylation of p44 and p42 MAP kinase and their effects were abolished by specific inhibitors of MAP kinase activation, PD98059 and U0126. Both drugs on oral chronic treatment (150umole/kg/day X 14days) decrease rat and mice hippocampal APP isolectroprotein. Since the generation of sAPP precludes the formation of amyloidogenic derivatives, the demonstration that TV3326 and TV3279 can stimulate the non-amyloidogenic α-secretase pathway and decrease rat hippocampal full length APP suggests that these neuroprotective drugs may prevent the basic APP pathogenic mechanism underlying AD.
NOVEL BIFUNCTIONAL COMPOUNDS ELICITING CHOLINERGIC AND ANTI-INFLAMMATORY ACTIVITY FOR THE TREATMENT OF CNS IMPAIRMENTS

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The development of new drugs for treatment of various CNS degenerative diseases such as dementia of Alzheimer's type (AD) is based mainly on the use of cholinergic compounds such as cholinesterase inhibitors (CHEIs). It was shown that anti-inflammatory drugs, free radical scavengers and antioxidants could also serve for the amelioration of the inflammatory processes occurring in various CNS diseases. The clinically used drugs, which presently demonstrate efficacy in mild to moderate AD are CHEIs (e.g., Aricept, Exelon and Reminyl). Certain quaternary CHEIs (such as pyridostigmine (Pyr) and polar non-steroidal anti-inflammatory NSAIDs (e.g. ibuprofen and diclofenac) could hardly cross the blood-brain barrier (BBB).

We have designed and synthesized a series of novel bifunctional compounds that contain covalently coupled cholinergic up-regulators (CURE) such as quaternary CHEIs, muscarinic and nicotinic agonists with various NSAIDs. These two moieties are coupled by a hydrophobic linker that renders permeability through the BBB.

This report is focused mainly on bifunctional NSAID-CURE compounds that are conjugates of NSAIDs with CHEIs. Most conjugates contain PYR as CHEI moiety coupled via a hydrocarbon octyl linker (PO) to the following NSAIDs: ibuprofen (IBU), diclofenac (DICLO), indomethacin (INDO), aspirin (ASP) and naproxen (NAP). IBU-PO, DICLO-PO, INDO-PO, ASP-PO and NAP-PO inhibit both ACHE and BChE with dissociation constants (K) and binucleotide kinetic rate constants (k) that range between 10^-10 M and 10^-10 M^2 min^-1, respectively. Some of these bifunctional compounds also inhibit cyclooxygenase (COX) I and COX II at micromolar level. Thus, the NSAID-CHEI conjugates exert both anti-ACHE and anti-COX activity at equimolar concentrations even in their intact non-hydrolyzed form. The acute toxicity of these compounds is 10-20 fold lower than that of PYR with LD50 values that range at 50-100 mg/kg ip in mice. Anti-inflammatory activity of IBU-PO was examined in carrageenan-induced peripheral and CNS inflammation in rats. Pretreatment with IBU-PO (5mg/kg, ip) decreased significantly the rat paw edema level and whole brain edema. IBU-PO (5mg/kg, ip) increased by 8-fold the survival time of mice that were exposed to hypoxic asphyxia as compared to control animals. IBU-PO (5mg/kg, ip) decreased significantly the brain edema and improved the neurological severity score in closed head injury model in mice. These findings suggest that the new NSAID-CHEI bifunctional chimeras could be useful for treatment of CNS impairments such as cerebral-vascular dementia and for reducing the neuronal damage caused by either acute cerebral ischemia or closed head injury.

STRUCTURE AND ACTIVATION OF MUSCARINIC ACETYLCHOLINE RECEPTORS

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Systematic scanning mutagenesis has been applied to the transmembrane domain of the M1 muscarinic acetylcholine receptor. In addition to the negatively-charged aspartic acid residue in transmembrane (TM) helix 3, the aromatic rings of tyrosine residues in TM 3, 6 and 7 play a key part in binding acetylcholine (ACh), and triggering the conformational change which activates the receptor, while hydrogen bonding residues in TM 3, 5, 6 and 7 help to anchor ACh in the ground state binding site. Residues in TM 4 may contribute to a peripheral ligand docking site. Amino acids which make inter-helical contacts which help to stabilize the ground state of the receptor have been identified. These include the Asn and Tyr residues in the highly-conserved Asn-Pro-X-X-Tyr sequence in TM 7. The contacts made by these residues are probably rearranged during receptor activation, to produce a G-protein binding site. The results are presented in the context of a 3-dimensional model of the M1 muscarinic receptor, based on the structure of rhodopsin.

ROLES OF EXTERNAL LOOPS OF MUSCARINIC RECEPTORS IN INTERACTIONS BETWEEN N-METHYLSCLOPAMINE AND ALLOSTERIC MODULATORS

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Allosteric modulators are known to modify the affinity of muscarinic receptors for orthosteric ligands but the sites of their attachment to receptors and the nature of the conformational change which they induce in receptors have been little clarified. It is assumed that they bind to a domain which is located more extracellularly than the orthosteric binding site. Neuroumecular blockers atropineum and gallamine have a high affinity for and strong allosteric effects on the M2 receptors. Their affinity for and allosteric efficacy on the M3 subtype is much weaker.

We performed mutations or exchange of extracellular loops in the M3 receptors, trying to make this subtype more similar to the M2 subtype and to discover sites important for allosteric interactions. Transferring the second outer loop from M2 to M3 receptors enhanced the affinity for both atropineum and gallamine, without changing their allosteric efficacy. On the other hand, M3 receptors in which the third outer loop had been modified so as to correspond to the same loop of the M2 receptors displayed not only higher affinities for both allosteric modulators, but also much stronger negative cooperativity between N-methylscopamine (NMS) and gallamine. Atropineum lost its negative effect on the binding of NMS (typical of M3 receptors) and acquired the ability to enhance the binding of NMS (positive cooperative action, typical of M2 receptors).

The third outer loop apparently plays important roles both in the high-affinity binding of gallamine and atropineum and in the mechanism of their negative and positive allosteric effects.

GENERATION AND ANALYSIS OF MUSCARINIC ACETYLCHOLINE RECEPTOR KNOCKOUT MICE

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Identification of the physiological and pathophysiological roles of the individual muscarinic acetylcholine receptor subtypes (mAChRs; M1-M5) has proven a challenging task, primarily due to the lack of ligands endowed with a high degree of receptor subtype selectivity and the fact that most tissues and organs express multiple mAChRs. To circumvent these difficulties, we generated mutant mouse lines ('KO' mice) in which specific mAChR genes (M1, M2, M3, or M4) had been inactivated by gene targeting techniques. The different mAChR mutant mice and their wild-type littermates were subjected to a battery of physiological, pharmacological behavioral, biochemical, and neurochemical tests. The M2 and M4 receptor mutant mice showed several striking phenotypes, as reported previously (Gionveza et al., PNAS, 96, 1692 and 10483, 1999). M3 receptor KO mice displayed a significant decrease in food intake, associated with reduced body weight and low serum leptin and insulin levels, probably due to disorption of a hypothalamic cholinergic pathway involved in the regulation of appetite. Pharmacological analysis of M2 receptor single and M2/M4 receptor double KO mice indicated that muscarinic agonist-induced analgesic responses are mediated by both M2 and M4 receptors. Neurochemical studies showed that autoinhibition of ACh release is mediated primarily by M2 receptors in hippocampus and cerebral cortex, but predominantly by M4 receptors in the striatum. These results provide a rational basis for the development of novel muscarinic drugs.

-18-
STRUCTURAL AND FUNCTIONAL CONSERVATION OF SNARE COMPLEXES

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The SNARE proteins syntaxin1, synaptobrevin2, and SNAP-25 play a key role during synaptic exocytosis. It was suggested that their assembly into stable membrane-bridging complexes gradually brings the membranes into close apposition. Thus, complex formation may provide the energy for initiating bilayer merger. The SNARE assembly pathway is only partly understood. Hence, we believe that detailed structural, kinetic, and thermodynamic investigations will bring a closer understanding of the function of SNARE proteins during membrane fusion. SNARE complex assembly can be described as a folding reaction, since it is accompanied by extensive structural rearrangements from less structured monomers to a tightly packed parallel four-helix bundle. Furthermore, a switch from a closed to an open conformation of syntaxin is thought to be a prerequisite for complex formation. We found that assembly and dissociation of the synaptic SNARE complex exhibits a marked hysteresis. It implies that assembly and dissociation do not follow the same path — probably due to a high energetic barrier between both native states.

Consequently, the ATPase NSF is necessary for disassembly. In addition, it suggests that SNARE assembly between two membranes is unidirectional — probably towards fusion. In an analogous study, we showed that an endosomal SNARE complex, composed of four SNARE proteins only distantly related to the synaptic SNAREs, has an almost identical structure to the synaptic complex. Moreover, the endosomal SNARE complex displayed a similar hysteresis. Taken together, our data suggest that SNARE function is conserved for all membrane fusion steps.

PRECLINICAL STUDIES OF GALANTAMINE USING A FORM OF ASSOCIATIVE LEARNING SEVERELY IMPAIRED IN ALZHEIMER'S DISEASE

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Classical eyeblink conditioning is a well-characterized model paradigm for the study of the neurobiology of learning, memory, and aging which also has application in the differential diagnosis of neurodegenerative diseases. Studies of eyeblink conditioning in neurological patients along with brain imaging studies of conditioning in normal adults document parallels in the neural substrates of this form of associative learning in humans and non-human mammals. Disruption of septohippocampal cholinergic neurotransmission impairs acquisition of conditioned eyeblink responses and slows the rate of learning. Alzheimer's disease (AD) profoundly disrupts the hippocampal cholinergic system, and patients with AD consistently perform poorly in eyeblink conditioning. Some nicotinic acetylcholine receptor (nAChR) subtypes are lost in AD making the use of allosteric modulation of nAChRs a promising therapeutic strategy. Galantamine modulates nAChRs to promote acetylcholine neurotransmission and also acts as an acetylcholinesterase (AChE) inhibitor. Galantamine was tested in preclinical experiments. Young and older rabbits received galantamine (3.0 mg/kg) for 15 days during conditioning, and the drug significantly improved learning, reduced AChE levels, and increased nAChR binding. In a second experiment, 53 retired breeder rabbits were tested over a 15-week period. A continuous dose of 3.0 mg/kg galantamine over 15 weeks ameliorated learning deficits significantly during acquisition and retention. nAChR binding was significantly increased in rabbits treated for 15 days with 3.0 mg/kg galantamine, and all galantamine-treated rabbits had lower levels of brain AChE. In a third experiment, galantamine reversed the effect of the nicotinic antagonist, mecamylamine. The efficacy of galantamine in a learning paradigm severely impaired in AD is consistent with outcomes in clinical studies.
XIIth INTERNATIONAL SYMPOSIUM ON CHOLINERGIC MECHANISMS-FUNCTION AND DYSFUNCTION & 2nd MISRAHI SYMPOSIUM ON NEUROBIOLOGY

Poster Abstracts
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ACETYLCOLINESTERASE mRNA EXPRESSION IN RAT SPINAL CORD

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The function of acetylcholinesterase (AChE) in the CNS is not as well defined as in the neuromuscular junction (NMJ). Besides terminating the cholinergic inputs to motor and other neurons, high concentrations of acetylcholinesterase (AChE) in the spinal cord might also reflect other cholinergic and non-cholinergic roles of this molecule. Transportation of the asymmetric A12 form along axons of motor neurons suggests that AChE synthesized in motor neurons also contributes to the pool of basal lamina-bound AChE in the NMJ. To elucidate the role of AChE in the spinal cord, we studied the expression of AChE in this part of the CNS. By combining in situ hybridization with fluorescent nuclear labeling and choline acetyltransferase staining, we found that glial cells are AChE mRNA negative and that motor neurons are the only cells expressing this mRNA in the ventral horns of the rat spinal cord. Further, we quantitated by competitive RT-PCR the expression of AChE mRNA in the spinal cord between E15 and E21 of rat embryonic development when the NMJs are formed. We found that expression of AChE mRNA follows the same temporal pattern of expression as mRNA encoding the neuronal isoform of agrin (AGR19), which is released from the motor nerve ending and is bound to the synaptic basal lamina in the NMJ. The same pattern of expression of AChE and AGR19 in the motor nerve may be a reflection of common processing and targeting of these two molecules to the same location and therefore suggests that, at least during the initial stages of NMJ formation, a part of AChE in NMJ may be contributed by a motor neuron.

FUNCTIONAL EXPRESSION AND STOICHIOMETRY OF THE NOVEL HUMAN α9α10 HETEROMERIC NICOTINIC ACETYLCHOLINE RECEPTOR


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We have cloned a novel human neuronal nicotinic acetylcholine receptor (nACHR) subunit, termed α9 (Genbank accession number AJ295237). The α10 nucleotide sequence maps to chromosome 11 (p15.5) and the predicted α10 polypeptide translation product is 66% similar to that of the human α9 subunit. α10 alone does not form a functional nACHR in oocytes but when it is co-expressed with α9 oocytes responded to bath applied ACh (30μM) with much larger currents (1.0±1.34μA; N=27) compared to oocytes expressing only α9 (0.7±0.6μA; N=23). Hence, co-expression of α10 with α9 does not dramatically change the EC50 value to ACh (10.3μM for α9α10 versus 14.4μM for α9 alone). To conclusively prove that α10 co-assembles with α9 to create a functional receptor we introduced a reporter mutation in α9 (L277T) and α10 (L276T) and determined EC50 values for various α9α10 combinations. Mutating α10 shifted the ACh concentration-response curve to the left (EC50=2.9μM; 95% confidence limits (CL) 2.5 to 3.3) while mutating α9 resulted in an intermediate sensitivity to ACh (EC50=4.6μM; 95% CL 4.2 to 5.1). nAChRs composed only of mutant α9 were the most sensitive to ACh (EC50=0.5μM; 95% CL 0.4 to 0.7). These observations confirm that the α10 subunit co-assembles with α9 to produce a functional heteromeric α9α10 nAChR. Furthermore, they suggest a pentameric receptor stoichiometry in which there are more α10 subunits than α9, most likely two α9 and three α10.
LARGE SCALE EXPRESSION OF THE EXTRACELLULAR AND CYTOPLASMIC DOMAINS OF THE DROSOPHILA ADHESION PROTEIN, GLIOTACTIN
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The transmembrane cell adhesion protein Gliotactin (Gli) from Drosophila (Auld et al, 1995), a member of the acetylcholinesterase (AChE) alpha/beta hydrolase superfamily, shares 24% sequence identity with Torpedo californica AChE (TcAChE), including two residues of the catalytic triad and two of three disulfide bonds. The extracellular portion is believed to share a common global fold with AChE, while the intracellular portion has no significant similarity to any protein in the sequence databases. This protein is therefore of interest for several reasons. Confirmation of a similar fold for Gli and AChE would have important implications for recently suggested non-classical functions of AChE. In addition, the intracellular portion of Gli is an excellent candidate for structural genomics. Therefore, the C-terminal, intracellular portion of Gli (Gli nt) and the N-terminal, cholinesterase-like portion of Gli (Gli ct), were cloned separately into pET21b. Gli ct was expressed in soluble form in E. coli (BLR) at 7 mg/L and purified to 95% homogeneity using Ni-NTA chromatography followed by size exclusion chromatography. Small crystals have been obtained, and optimization of crystallization conditions is currently underway. Gli nt was also expressed in soluble form in E. coli (Origami(DE3)pLysS) at 1 mg/L, purification is currently underway.

Reference:

STRUCTURAL REORGANIZATION OF THE ACETYLCHOLINE BINDING SITE OF TORPEDO NICOTINIC RECEPTOR REVEALED BY DYNAMIC PHOTOLUMINESCENCE LABELING
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We have explored by photoaffinity labeling the structural changes that occurred at the acetylcholine (ACh) binding site of the Torpedo nicotinic receptor (nAChR) during activation by a tritiated photoactivatable agonist [3H]DCTA (diazocyclohexadienonypropyl-trimethylammonium). The agonistic nature of DCTA was confirmed both at the muscular and neuronal nicotinic receptors reconstituted in Xenopus oocytes. In a previous study, the identification of labeled residues by [3H]DCTA allowed to characterize the amino acids involved in the binding of the ester moiety of ACh in the desensitized state [1]. In the present work, the agonist character of DCTA was used to label functional states. A stopped-flow apparatus was adapted for dynamic photolabeling. After selected mixing times between nAChR and [3H]DCTA in a delay loop, samples were irradiated by intense UV light preceding the collection for biochemical analysis. We quantified a saturable increase of specific photolabeling on the alpha and gamma subunits. We further analyzed this incorporation either after rapid mixing (500 ms) or after equilibration (50 min) of [3H]DCTA with nAChR, hence the probe explores transient state(s) and the stable desensitized state, respectively. Comparative analysis showed a differential photoincorporation of [3H]DCTA and suggests that in the course of agonist-induced desensitization of the receptor, the site labeling peptide loops from adjacent alpha and gamma subunits move closer together.


COEXPRESSION OF ALPHA10 AND ALPHA9 NICOTINIC ACETYLCHOLINE RECEPTORS IN RAT DORSAL ROOT GANGLIONEURONS
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Acetylcholine excites sensory neurons of dorsal root ganglia (DRG) via both muscarinic and nicotinic acetylcholine receptors (nAChR). Until recently, 9 different α and 4 different β subunits of nAChR that form pentamers to yield functionally active receptors have been identified. However, binding and immunoprecipitation studies revealed a population of nAChR in DRG that shall contain at least one additional, unknown subunit. This year, a new member of the nAChR family (α10) has been identified in cochlear hair cells. It forms heteromers with α9 that share a crucial feature with the unknown nAChR of DRG, i.e. sensitivity to α-bungarotoxin. Here, we asked whether these subtypes are also expressed in rat DRG, and addressed this issue by RT-PCR, in-situ hybridization, and using a newly raised antibody against the α10 subunit. All sensory neurons, but no glial cells, expressed both α9 and α10 mRNA, and exhibited α10-immunoreactivity. These in-situ hybridization and immunohistochemical findings were confirmed by RT-PCR. These data show that coexpression of α9 and α10 nAChR subunits is not restricted to hair cells but occurs also in DRG sensory neurons where it is likely to represent the previously unknown nAChR (supported by the DFG, SFB 547).

MAPPING THE ACETYLCHOLINE BINDING SITES OF TORPEDO NICOTINIC RECEPTOR USING PHOTOLUMINESCENCE LABELING: PAST, PRESENT AND FUTURE
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Topographical mapping of residues contributing to the ACh binding sites was achieved with site-directed antagonists or agonists labels. From these studies and mutagenesis three discontinuous domains of the alpha-subunit (loops A, B and C) with additional residues on the gamma and delta subunits (loops D, E and F) were identified (1).

The structural reorganization that occurs at the cholinergic binding domains on desensitization was studied at the molecular level first with [3H]JDF using flash photolysis coupled to a stopped flow apparatus and more recently with the agonist [3H]DCTA. [3H]JDFCA was able for the first time to explore, at the molecular level, transient states of nAChR upon agonist activation showing a modified labeling pattern for amino acids labeled in loop C and for gamma and delta fragment's peptides (see poster T. Gruetter et al.).

To investigate more precisely the gamma (or delta) peptides determinants involved in the structural rearrangement upon agonist activation, we used a formerly described photosensitive agonist [3H]IAC5 whose labeling pattern on Torpedo nAChR ACh binding area involved all four subunits (2). This pattern was modified upon receptor desensitization according to preliminary dynamic photolabeling experiments.

Identification of gamma and delta residues involved in subunits reorganization around the binding sites and modelisation of nAChR Torpedo at the alpha-gamma and alpha-delta interfaces based on the X-ray structure of AchBP will give new insights in allosteric transitions upon desensitization.

NFkB REGULATES THE ACTIVITY OF HUMAN ACETYLCHOLINESTERASE PROMOTER IN MUSCLE
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Precise expression of biological active acetylcholinesterase (AChE; EC 3.1.1.7) is required for development and maintenance of the neuromuscular junctions. Previous studies reveal that the expression of human AChE in muscle can be regulated by the classical cAMP-dependent pathway that involved cAMP-dependent protein kinase (PKA) and cAMP-responsive element binding protein (CREB), via the regulatory element called cAMP-responsive element (CRE) on promoter of the enzyme. In the present study, we investigated the role of another regulatory element namely NFkB, in regulating the AChE promoter activity. A 2.2 kb human AChE promoter was tagged with a luciferase reporter gene, and the construct was transfected in chick myotube cultures for analyzing the luciferase activity. The transcriptional activity of human AChE promoter was shown to be regulated by NFkB signaling. One of the possible nerve-derived factors that utilizes this mechanism is the adenosine 5-triphosphate (ATP) and its correspondent receptor called P2Y purinoceptor. All these data suggested that the AChE promoter could be regulated by multiple signaling pathways for precise transcriptional control at the post-synaptic muscle.

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THE INITIAL BINDING OF ACETYLCHOLINESTERASE AND PERLECAN OCCURS INSIDE THE CELL PRIOR TO EXTERNALIZATION
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The collagen-tailed form (A12) of acetylcholinesterase (AChE) is attached to the extracellular matrix on the surface of myotubes in culture and on the basal laminae at sites of nerve-muscle contact "in vivo". This basal lamina-associated AChE is tightly linked to the extracellular matrix through the heparan sulfate proteoglycan perlecan. Where and how this linkage occurs is not known. One possibility is that the initial interaction between AChE and perlecan occurs post-translationally during intracellular trafficking. To test this hypothesis, we incubated quail muscle culture extracts with protein-A Sepharose beads bound with anti-perlecan monoclonal antibody and assayed them for bound AChE. Anti-perlecan antibodies immunoprecipitated about 5-10% of total AChE activity. To determine whether this binding occurred intracellularly, muscle cultures were treated with DFP, an irreversible AChE inhibitor, followed by incubation in complete medium to allow de novo synthesis of AChE. This time course study showed that AChE binding to perlecan occurred approximately 2-3 hours after synthesis, coincident with the assembly of A12 AChE in the Golgi apparatus. At this time A12 AChE is approximately 15% of the total enzyme synthesized, and about the same percentage of the newly synthesized intracellular AChE pool binds to anti-perlecan beads. We concluded that the newly synthesized A12 AChE is assembled with perlecan inside the cell prior to externalization.

ROLE OF MDX NERVE AND MUSCLE IN REGULATING NEUROMUSCULAR JUNCTION PROPERTIES: A STUDY USING MUSCLE TRANSPLANTS
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Neuromuscular junctions (NMJs) of mdx (X-linked dystrophy) mutant mice have fewer junctional folds, altered shape, lower AChR density, and accelerated AChR degradation rate. The relative contributions of nerve and muscle to this organization are unknown. The stemmatonoid muscle of mdx and wild-type mice were surgically exchanged into host counterparts, and 7.5 mo. later examined for expression of dystrophin protein and NMJ fine-structure. The dystrophin staining patterns were maintained in the transplants, and when the wild-type was the host, the transplanted mdx muscle exhibited extensive recovery of junctional folds. To analyze the NMJ shape, we labeled AChRs in 4.5 mo. transplants with fluorescent alpha-bungarotoxin, Bgt. Labeled AChRs were redistributed at the NMJ, when the wild-type was the host, the shape of NMJ in the donor mdx muscle was more contiguous than when the mdx was the host. To determine if AChR number was affected by host phenotype, we labeled AChRs in 7.5 mo. transplants by injecting 125I-Bgt. Two days later, the donor muscle was assessed for AChR radioactivity using a gamma counter. Consistent with the change in junctional folds and NMJ shape, the mdx transplant into wild-type host showed 125I-Bgt binding similar to control wild-type muscles, while that of the wild-type transplant into mdx host was comparable to control mdx muscles. Together, these findings suggest that the donor phenotype of the muscle is maintained and that the host (and nerve) influences NMJ fold formation, shape, and total number of AChRs.

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CRYSTAL STRUCTURE OF THE TETRAMERIZATION DOMAIN OF ACETYLCHOLINESTERASE AT 2.3A RESOLUTION
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Tetramerization of acetylcholinesterase (AChE) is achieved by the interaction of 2 peptide motifs: a 40-residue 'tryptophan amphiphilic tetramerization' (WAT), at the C-terminus of the catalytic subunit, and a 17-residue 'proline-rich attachment domain' (PRAD), localized near the N-terminus of the CoQ collagenic tail polyepitope, with 4:1 WAT:PRAD stoichiometry. The two peptides were produced by chemical synthesis. Met21 of WAT replaced by selenomethionine, to facilitate a MAD diffraction experiment. The synthetic WAT and PRAD were mixed at a 4:1 ratio, and co-crystallized. The monoclinic crystals obtained diffracted to 2.3A resolution, and MAD data sets were collected at the synchrotron. The structure was solved with the program SOLVE which produced a traceable electron density map. The structure was refined to an R-factor of 24.6% with the 2 PRADs seen in full and the 8 WATs having disordered C-termini. The WAT chains assume an α-helical conformation, and are all parallel. The PRAD has a polyproline II conformation and threads its way anti-parallel to the WAT chains. Most of the 3 highly conserved Trp residues in each WAT chain are stacked against the 8 Pro residues or 3 Phe residues of the single PRAD. An AChE tetramer structure can be modeled based on the structure of the WAT/PRAD complex.
SCALING UP OF PRODUCTION, PURIFICATION AND REFOLDING OF A CHIMERIC THREE-FINGERED TOXIN WITH SPECIFICITY FOR ACETYLCHOLINESTERASE

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Chimera II (rcII) is an engineered hybrid toxin constructed from two members of the “Three-Finger-Toxins” family: fasciculin 2 from the green mamba, *Dendroaspis angusticeps*, a potent inhibitor of acetylcholinesterase (AChE), and toxin α from the black neck spitting cobra, *Naja nigricollis*, a potent inhibitor of the nicotinic acetylcholine receptor. The 3D structures of both toxins are known and reveal a striking resemblance in their cores with significant difference in their fingers. The structure of the fasciculin 2/AChE complex has also been solved.

The 3D structure of rcII consists of the globular core of toxin α (3 anti-parallel β-strands) together with 2 of the extended loops (fingers) connecting these strands from fasciculin 2. rcII inhibits *E. electricus* AChE with an affinity that is 15-fold less than native fasciculin 2. The chimera was expressed, purified, crystallized and its structure solved (Ricciardi et al. J. Biol. Chem. [2000] 275, 18302-18310; Le Du et al. J. Mol. Biol. [2000] 296, 1017-1026).

Scaling up of production of rcII was initiated to produce amounts adequate for determination of the structure of rcII/Torpedo californica AChE (TcAChE) complex and for more detailed kinetic and physicochemical studies. It included large-scale production of bacterial host cells (10 L.), to yield 150 mg of fusion protein per liter of bacterial culture. Additional HPLC steps were added in the purification of the unfolded chimeric polypeptide to yield 10 mg of unfolded polypeptide per liter of initial bacterial culture, which were then taken for the refolding step.

ROLE OF SPONTANEOUS MUTATIONS OF NEURONAL NICOTINIC RECEPTORS IN ADNFLE

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The association between a mutation in either the alpha4 or beta2 coding gene and a form of nocturnal encephalopathy (ADNFLE) suggests that activation of the nicotinic acetylcholine receptor can be at the origin of brain seizures. Thus, three spontaneous mutations in the alpha4 (S248F, S76ins3, S522L; Steinlein et al., 1997) and two different substitutions of valine 287 in the beta2 subunit (V287L, V287M; De Fusco et al., 2000; Phillips et al., 2001) have been identified. These mutations modify amino acids within or near the second transmembrane domain of the protein (TM2). To identify the effects caused by these five mutations on the functional nicotinic receptor properties, exogenous expression of these proteins were designed. Determination of a series of physiological and pharmacological characteristics revealed that the single common trait identified so far is an increased sensitivity of the receptors to their endogenous ligand acetylcholine. Based on our understanding of the receptor distribution in the different brain areas, their development and the neuronal network circuitry, we hypothesize that an increased of the acetylcholine sensitivity may cause an unbalance in the fine-tuning of the cortico-reticular thalamic and thalamo-cortical network in favour of the latter. The use of specific agonists or antagonists of the alpha4beta2 receptor should help to elucidate the last steps in these critical brain pathways and therefore shine a new light on the fundamental mechanism of epilepsy.

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EXPRESSION OF COLQ AT THE NEUROMUSCULAR JUNCTION

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ColQ, a specific collagen has been identified in Torpedo and mammals (Krejci et al., 1991, 1997). So far the main function attributed to this collagen is to anchor acetylcholinesterase (AChE) in the synaptic basal lamina at the neuromuscular junction (NMJ). Our data obtained by RT-PCR and in situ hybridization show that the ColQ gene is produced both by the motoneuron and the muscle cell. However, several variants of ColQ have been identified. On-going experiments are determining the structure of ColQ produced by the motoneuron. Is ColQ secreted in extrasympathetic domains where AChE is low? Studies from ColQ -/- muscle cells in culture transfected with a GFP-ColQ cDNA suggest that ColQ can be secreted in the absence of AChE. We also transfected these cells with mutants of the ColQ heparin binding sites (HBS) or mutants of ColQ C-terminus and show that in both cases no spontaneous clusters of AChE can be detected, suggesting a complex mode of AChE anchoring in the synaptic basal lamina.

INTERACTION OF RECOMBINANT SOLUBLE NEUROLIGINS-1 WITH NEUROXIN-BETA

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Neurolignis are postsynaptic transmembrane proteins of the alpha/beta-hydrolase fold family with high sequence identity to the cholinesterases. They form heterologous cell contacts with the neurexins. To examine the neurilogn-1 (NL1) interactions with neuroxin1-beta (NX1-beta), and to define the NL1 binding surface interacting with NX1-beta, we expressed soluble recombinant forms of NL1 by truncating the carboxyl-terminal end prior to the transmembrane span at positions D638 and D691. LCMS of peptide digests showed the pairing of the disulfide bonds and the carbohydrate composition of N- and O-linked glycosylation sites. NL1 truncated at D691 was crystallized by vapor diffusion and data up to 4.35 Å were collected on the ESRF beamline ID29. Attempts to solve the structure by molecular replacement were unsuccessful but efforts to solve a structure of a truncated NL1 devoid of O-glycans are underway. By surface plasmon resonance, we studied the association of purified NL1s with NX1-beta. Elimination of the NL1 transmembrane span yielded soluble protein that associates with NX1-beta with a KD in the nM range, while the homologous protein AChE does not bind NX1-beta. Purified soluble NL1 binds two splice forms of NX1-beta with slight differences in affinity. Incomplete post-translational processing of NL1 and enzymatic removal of certain oligosaccharides or sialic acids enhance activity, while enzymatic deglycosylation of neuroxin-beta does not alter binding, indicating that glycosylation of NL1 may provide a further level of control in the neurexin-neurolin association. Work supported by USPHS Grant GM-18360 to P.T.
CLONING AND CHARACTERIZATION OF ACETYLCHOLINESTERASE GENE IN CHICKEN

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Acetylcholinesterase (ACHE) is a highly polymorphic enzyme with multiple forms and its regulation is under strict regulation. To understand more about the species-specific regulation of ACHE gene in avian versus mammalian species, we have attempted to clone acetylcholinesterase gene in chicken by rapid amplification of genomic end (RAGE). Antisense primers complementary to the 5-prime coding region of the chicken ACHE cDNA were used as nested PCR primers, which resulted in the isolation of the largest genomic fragment of about 2.1 kb. This genomic DNA contained a 269 bp sequence at the 3-prime end corresponding to 19-380 bp of the published chicken ACHE cDNA with minor sequence discrepancy. The cDNA at position 906GATTGCCC98 was found to be CCAGATTGCCC, resulting in a change of deduced amino sequences from ArgPhcAla to ProAspPhcAla. TATAA box was found upstream the start codon and a BLAST search of the sequence immediately upstream to the start codon, that may confer the functional promoter activity, revealed little homology to the published ACHE genes. The putative promoter fragment was subcloned into pGL3 basic vector for the luciferase promoter-reporter assay. Deletion analysis with a truncated 366 bp fragment upstream to ATG resulted in a ten-fold increase of the activity. PMA, forskolin and cAMP can induce the luciferase activity increase 60% to 150% in NG108-15 cells. Similar response to the drugs was also found in the chick myotubes. Further characterization of chick ACHE promoter is under investigation.

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MUSCARINIC RECEPTOR REGULATION OF EVOKED ACETYLCHOLINE RELEASE IS AFFECTED BY ACETYLCHOLINESTERASE INACTIVATION AT THE MOUSE NEUROMUSCULAR JUNCTION

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In the present work we used antibodies and Western blotting of lysates obtained from innervated and non-innervated regions of the mouse diaphragm to demonstrate that mouse neuromuscular junctions (NMJ) express mAChR subtypes M1 through M4 and that localization of all subtypes is restricted to the innervated part of the muscle. To elucidate the roles of the mAChR subtypes in regulating ACh release evoked by the presynaptic action potential, we evaluated the quantal content of endplate potentials by the method of failures on isolated phrenic-hemidiaphragm preparations removed from normal and collagen Q-deficient mice.

Muscarine reduced evoked ACh release in normal NMJs, but it enhanced ACh release both in collagen Q-deficient NMJs lacking acetylcholinesterase (ACHE) and in normal junctions when ACHE was inhibited by fasciculin-2. The muscarine-electicted depression of ACh release in normal NMJs was completely abolished by pre-treatment with PTX or methoctramine, a selective M2 antagonist, but was not affected by M4 selective muscarinic toxin MT-3. These results indicate the involvement of the M2 mAChR in decreasing ACh release. The muscarine-induced increase of ACh release in ACHE-deficient NMJs was not affected by PTX, but was completely blocked by a specific M1 antagonist MT-7. Our results show that the M1 and M2 mAChRs have opposite presynaptic functions in modulating ACh release, and that regulation of transmitter release by the two receptor subtypes depends on the functional state of ACHE at the NMJ.

CANDOXIN A NEW SNAKE TOXIN SPECIFIC FOR THE ALPHA7 nACHR


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Toxins isolated from animals provide important pharmacological tools for studies of ligand-gated ion channels. Snake toxins like alpha-bungarotoxin are known to inhibit almost irreversibly the muscle nicotinic acetylcholine receptor (nAChRs). In addition, alpha-bungarotoxin was shown to powerfully block the alpha7 neuronal nAChRs. Snake toxins are divided in short and long-chain according to the number of amino acids they contain. A further classification is often made in function of their number of disulfide bridges. The different snake toxins share the same structural form, constituted by three loops (I, II and III) protruding from a globular core whereas the loop II contains the principal amino acids participating to the nAChRs recognition. Candoxin is a new toxin that was isolated from the Malayan krait Bungarus candidus. Constituted by 66 residues and displaying the typical three fingers motif with five disulfide bridges, candoxin shares 30% homologies with the alpha-bungarotoxin. Candoxin is a member of the short chain toxins and is closely related to neurotoxins (98% homologies) that were isolated from Bungarus multicinctus. When tested on heterologously expressed receptors, candoxin was found to strongly antagonize either the muscle or the alpha7 rat nAChRs. The main difference with other known toxin is, however, the almost irreversible blockade of candoxin at the neuronal alpha7 receptors, while the muscle receptors readily recovered from blockade within minutes. Therefore, candoxin provides a new tool for the studies of the neuronal nAChRs in the central nervous system.

E.COLI EXPRESSED EXTRACELLULAR DOMAIN OF RAT ALPHA 7 NICOTINIC ACETYLCHOLINE RECEPTOR: PHYSICOCHEMICAL AND BINDING PROPERTIES


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The title domain (amino acid residues 1-208) was expressed as a fusion protein with glutathione S-transferase (GST). Refolding of GST-(1-208) protein in the presence of 0.1% SDS and the C116S mutation considerably decreased the formation of high-molecular weight aggregates. Gel-permeation HPLC was used to isolate the monomeric forms of the GST-(1-208) and its mutant almost devoid of SDS. CD spectra revealed that the mutation considerably increased the content of beta-structure and made it more stable at different conditions. The proteins obtained bound iodinated alpha-bungarotoxin with KD about 100 nM, the C116S mutation slightly increasing the affinity for the toxin. The proteins retained such selectivity features of the intact alpha7 as discrimination of long-chain alpha-neurotoxins and alpha-conotoxin Mim against short-chain neurotoxins and conotoxin G1. The alpha7 domain devoid of the fusion moiety was found to bind biotinylated alpha-cobratoxin which could be displaced by alpha-cobratoxin itself, d-tubocurarine or arahasine. Therefore, the obtained proteins seem promising for further structural analysis and for screening purposes as demonstrated recently (Utkin et al. 2001) by unreacting the alpha7 binding capacity in the so-called weak toxin.
NICOTINIC ACETYLCHOLINE RECEPTOR AND MUSK ARE CLUSTERED IN C2C12 CELLS VIA LIPIDIC RAFTS
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Clustering of nicotinic acetylcholine receptors (nAChRs) and signaling molecules such as the Muscle-specific Receptor Tyrosine Kinase MuSK in the postsynaptic membrane is the landmark of synaptic differentiation at the neuromuscular junction. Cholesterol-sphingolipid-enriched microdomains or rafts are part of the cell machinery ensuring correct intracellular trafficking of selected proteins and lipids, and have been shown to participate in the formation and maintenance of alpha-7 nAChR clusters in somatic sipes of ciliary neurons (Brandt et al. 2001). In addition, cells defective in sphingolipids biosynthesis were shown to express low amounts of muscle AChR, suggesting that sphingolipid metabolism may influence trafficking of the protein to the surface membrane (Roccato et al., 1999). In this work, we have asked whether rafts are engaged in AChR clustering in muscle cells. Using filamin, a fluorescent marker of cholesterol, and beta-cyodexin, a drug that depletes cells from cholesterol, we showed that cholesterol is present within agrin-induced AChR clusters in C2C12 cells and is necessary to maintain these clusters. Non-ionic detergent extraction on ice and flotation gradient centrifugation of COS7 cells transfected with AChR subunits cDNAs demonstrate that AChRs are contained in low density (5-30% interface) fractions enriched in two rafts markers, filamin and cavinin. Yet, in C2C12 myotubes, AChRs and MuSK are recovered in the 30% sucrose fraction following solubilization with Lubrol. We conclude that AChR as well as important signaling molecules such as MuSK are accumulated at postsynaptic sites via their association with large raft-like microdomains.

ACETYLCHEINESTERASE IS REQUIRED FOR NEURONAL AND MUSCULAR DEVELOPMENT IN ZEBRAFISH
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The neurotransmitter acetylcholine (ACh) has a crucial role in central and neuromuscular synapses of the zebrafish. After release into the synaptic cleft, ACh is rapidly degraded by acetylcholinesterase (AChE). We have identified a mutation in the ache gene of the zebrafish, which abolishes ACh hydrolysis in homozygous animals completely. Embryos are initially motile but subsequently develop paralysis. Mutant embryos show defects in muscle fiber formation and innervation, and primary sensory neurons die prematurely. The neuromuscular phenotype in ache mutants is suppressed by a homozygous loss-of-function allele of the alpha-subunit of the nicotinic acetylcholine receptor (nAChR), indicating that the impairment of neuromuscular development is mediated by activation of nAChR in the mutant. Here we provide genetic evidence for non-classical functions of AChE in vertebrate development.

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IDENTIFICATION OF SPECIES DIFFERENCES IN THE PHARMACOLOGY OF THE ALPHA-7 NICOTINIC RECEPTOR USING THE ANTAGONIST RADIOLIGAND [3H]- METHYLLYCOCONITINE
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It has been suggested that the impairment in cholinergic transmission observed early in Alzheimer's disease (AD) may partly underlie the cognitive decline associated with the condition. Recent reports that the alpha-7 nicotinic receptor (α7nAChR) co-localises with amyloid plaques and serves as a high affinity binding site for the beta-amyloid peptide, AB1-42, suggests this receptor may play a role in AD pathophysiology (Wang et al., 2000). Membranes were prepared from a cell line expressing human α7nAChRs (NIH-FP1-ha7, Peng et al., 1999), or rat brain tissue, and binding assays performed using the selective α7nAChR receptor antagonist [3H]-methyllycococitine ([3H]-MLA: Davis et al., 2000). A range of compounds were used to examine the pharmacology of [3H]-MLA binding sites, and for rat forebrain the rank order of potency was: MLA > ([JAR-17779 = 2020+ taurine = (+) JAR-17779 = JAR(R)/2403 = mecamylamine. When membranes expressing the ha7 receptor were used, antagonist affinities and that of the acetylcholinesterase inhibitors were similar to rat. In contrast, the affinities of the agonists for ha7nAChRs were approximately 10-fold higher than compared to rat tissue. Furthermore, under the conditions used in the present study, beta-amyloid had no effect on [3H]-MLA binding to human or rat α7nAChRs. Preliminary studies indicate [3H]-MLA binding to mouse membranes is consistent with rat data. In conclusion, we have demonstrated species differences in the pharmacology of [3H]-MLA binding sites; however, the lack of effect of beta-amyloid at both human and rat binding sites requires further clarification. Wang et al. (2000) J.Biol. Chem. 275(8) p5626. Peng et al. (1999) Brain Res. 825 p172. Davis et al. (2000) Eur.J.Neurosci. 12: p374.

THE 14.3.3 GAMMA PROTEIN IS PART OF THE MUSK SIGNALING COMPLEX AT THE COMPLEX AT THE NEUROMUSCULAR JUNCTION
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The Muscle-Specific receptor tyrosine Kinase, MuSK, forms a receptor complex activated by nerve-derived agrin that orchestrates the differentiation of the neuromuscular junction (NMJ). To identify partners and/or effectors of MuSK, co-cloning and immunopurification experiments have been performed in purified postsynaptic membranes from Torpedo electrocytes. In a first series of experiments, this approach has lead to the identification of a PDZ-containing protein MAGI-1 (Strohle et al. J.Cell Biol., 153: 1127-1132, 2001). In this work, a second polypeptide corresponding to the 14.3.3 gamma protein was identified by MALDI-TOF mass spectrometry (MS) analysis of the MuSK cross-link products. Moreover, MuSK and 14.3.3 gamma protein colocalized at the NMJ and co-immunoprecipitated following transient transfection in COS7 cells, suggesting a direct interaction between these two proteins. Transfections experiments in C2C12 myotubes show that the 14.3.3 gamma protein downregulates the transcription of the epsilon subunit of the AChR activated by ARIA/neuregulin1. Members of the 14.3.3 family of proteins play a central role in the regulation of various signalling pathways. Our data indicate that in addition to its role in AChR aggregation, the MuSK complex may be involved in the regulation of AChR transcription at the NMJ, possibly via the regulation of the MAP Kinase pathway by the 14.3.3 gamma protein.
CONSTRUCTION AND CHARACTERISATION OF A CHIMERIC HUMAN ALPHA 7 NICOTINIC ACETYLCHOLINE / MOUSE SHT3 RECEPTOR

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The neuronal nicotinic acetylcholine receptor alpha 7 subunit can assemble as a homo-pentamer to form a functional ligand-gated ion channel. Each subunit comprises four membrane-spanning regions linked by alternate cytoplasmic and extracellular loops, together with extracellular N- and C-terminal domains. Chimeric alpha 7 / SHT3 constructs have been described previously as tools for investigating the role of the various domains of the receptor. We report on the construction and characterisation of a novel human alpha 7 / mouse SHT3 chimera. A chimera was constructed comprising the N-terminal region of the human alpha 7 nicotinic acetylcholine receptor linked at valine 202 with the transmembrane / C-terminal regions of the mouse SHT3 receptor. Expression in Xenopus oocytes or HEK-293 cells resulted in functional channels that were sensitive to ligands of nicotinic acetylcholine, but not SHT3 receptors. Currents obtained from oocytes injected with cDNA for the chimera desensitised more slowly than those obtained by injection of wild-type alpha 7. The response of both wild-type and chimeric receptors was potentiated by SOH-indol. Expression in mammalian cells was initially demonstrated by surface alpha-bungarotoxin binding and single-cell calcium imaging in transiently transfected HEK-293 cells. Subsequently, stable clones were produced and functional clones selected by assessing agonist induced calcium increase in a HPLR. The best responding clones were used to characterise the chimera pharmacologically using standard nACHR ligands.

SEgregation of Phosphatidic Acid-Rich Domains in Reconstituted Acetylcholine Receptor Membranes


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Purified Acetylcholine Receptor (AcChR) from Torpedo has been reconstituted at high (3509:1) and low (560:1) phospholipid to protein molar ratios into vesicles containing egg phosphatidylcholine, cholesterol and different diacylglycerols (dimeristoyl phosphatidylethanolamine, phosphatidylglycerol and phosphatidic acid) as probes to explore the effects of the protein on phospholipid organization by differential scanning calorimetry, infrared and fluorescence spectroscopy. All the experimental results indicate that presence of the AcChR protein, even at the higher phospholipid to protein molar ratio, directs lateral phase separation of the monoaminergic phospholipid form of the phosphatidic acid probe, causing the formation of specific phosphatidic acid-rich lipid domains that become segregated from the bulk lipids and whose extent (phosphatidic acid quenched into the domain, out of the total population in the vesicle) is protein dependent. Furthermore, fluorescence energy transfer using the protein tryptophan residues as energy donors and the fluorescence probes trans-erythroic acid or diphosphonate, as acceptors, establishes that the AcChR is included in the domain. The other dimeristoyl phospholipid probes, under identical conditions, could not mimic the protein-induced domain formation observed with the phosphatidic acid probe and produce ideal mixing behaviour. Since phosphatidic acid along with cholesterol have been implicated in functional modulation of the reconstituted AcChR, it is suggested that such a specific modulatory role could be mediated by domain segregation of the relevant lipid classes. Supported by grant PM98-0098 from the DGICYT of Spain.

ALPHA-CONOTOXINS PnIA AND A10L-PnIA STABILISE DIFFERENT STATES OF THE CHICK NEURONAL ALPHA 7 ACETYLCHOLINE RECEPTOR

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The alpha-conotoxins PnIA and the single amino acid substitution [A10L]PnIA are inhibitors of neuronal nicotinic acetylcholine receptors (nACHRs) in dissociated neurons (1) and recombinant alpha7 receptors expressed in Xenopus oocytes. The effects of PnIA, [A10L]PnIA and alanine scan mutants of [A10L]PnIA were investigated in homomeric chick alpha7 nACHRs and alpha7 receptors using the L247T mutant (2). PnIA inhibited the ACh-evoked current in the wild type alpha7 receptor and the alpha7 L247T mutant in a similar manner with IC50 values of 349 and 194 nM, respectively. All [A10L]PnIA alanine mutants investigated inhibited the ACh-activated current in wild type alpha7 nACHRs. In contrast, when co-applied with ACh, [A10L]PnIA and the alanine scan mutants all potentiated alpha7 L247T responses and in addition [A10L]PnIA was able to activate a current in the absence of ACh. Because it is assumed that the L247T mutation renders conductive one of the desensitised states, these data indicate that PnIA toxin inhibits the alpha7 by stabilising a desensitised non-conducting state. Activation of the L247T mutant with [A10L]PnIA further demonstrates that this mutation stabilises another state of the receptor.

Rapsyn escorts the nicotinic acetylcholine receptor along the exocytic pathway via the association with lipid rafts

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The 43kDa receptor-associated protein rapsyn is a myristoylated peripheral protein that plays a central role in nicotinic acetylcholine receptor (AcChR) clustering at the neuromuscular junction. In a previous work, we demonstrated that rapsyn interacts with the exocytic pathway and is specifically cotransported with AcChR via post-Golgi vesicles targeted to the innersurface of the Torpedo electrosome (Marchand et al., 2000). In this work, to further elucidate the mechanisms for sorting and assembly of postsynaptic proteins, we analyzed the dynamics of the intracellular trafficking of fluorescently labeled rapsyn in the transient expression COS-7 cell system. Our approach was based on fluorescence, time-lapse imaging and immunoelectron microscopies, as well as biochemical analyses. We report that the acetylcholine receptor is transported with the rapsyn-associated peripheral proteins to the cell surface via a microtubule-dependent transport. Using cotransfection experiments of rapsyn and acetylcholine receptors, we report that these two molecules codepend on the exocytic and are transported in a raft-membrane. Triton extraction on ice and flotation gradient centrifugation demonstrated that rapsyn and AcChR are recovered in low density fractions enriched in two rafts markers, cavin-1 and flotillin-1. These data show that sorting and targeting of these two companion molecules is mediated by association with cholesterol-sphingolipid-enriched rafts microdomains. This raises the interesting hypothesis of the participation of the raft machinery in the targeting of signaling molecules at synaptic sites.
IDENTIFICATION AND CHARACTERIZATION OF DIVERSE FAMILY OF NEUROTOXIN-LIKE PEPTIDES FROM THE SOUTH AMERICAN CORAL SNAKE
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Snake venoms are unique mixtures with reference to their biochemical and pharmacological properties. A bite by Elapidae snake often causes serious neurogenic symptoms, such as convulsions and paralysis. The mixture of toxins may target the various synaptic proteins including nicotinic and muscarinic acetylcholine receptors (AChRs), potassium and calcium channels. To elucidate the molecular and physiological basis of the mode of toxin actions, we isolated cDNAs for neurotoxin-like peptides from the venom gland cDNA library of South American coral snake (Micruroides euryxanthus). We have identified ten new neurotoxin-like peptides so far, named CTx1 to CTx10. They are 56-67 amino acid residues long in mature form, and having eight cysteine residues in the corresponding position; the cysteine framework is critical for folding into a three-finger structure of neurotoxins. The CTx2 has two additional cysteine residues, which may possibly make the fifth disulfide bridge within the first loop. Seven peptides are more or less similar to the representative neurotoxins: short and long neurotoxins, α-Agrin, pre-synaptic toxin, muscarinic toxin, cardioxin. However, the intercysteine interval sequences of the peptides CTx3, CTx4 and CTx8 are fairly diverged from the known neurotoxins. Each recombinant toxin (rCTx) was prepared in E.coli, yeast and/or Xenopus oocytes and characterized. The peptides interact with acetylcholine binding protein (AChBP) in variable kinetics and blocked ACh-induced muscle- and neuronal-type nicotinic AChR responses expressed in Xenopus oocytes.

ORIGIN OF ACETYLCOLINESTERASE IN THE DEVELOPING NEUROMUSCULAR JUNCTION
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Most of the acetylcholinesterase (AChE) in the neuromuscular junction (NMJ) is bound to the basal lamina located between the nerve ending and postsynaptic membrane of the muscle fiber. Potentially, this AChE can originate in both, muscle and/or nerve. Present evidence strongly supports muscular origin of synaptic AChE, however neural origin has never been disproved and is supported by another line of evidence. In this work we studied the origin of synaptic AChE at the early stages of the NMJ formation. We employed an in vitro model in which motor neurons originating from the embryonic rat spinal cord explants form well differentiated NMJs with human myotubes. By immunocytochemical staining with species specific antibodies against human and rat AChE, we were able to distinguish human AChE (muscle origin) from the rat AChE (neural origin) in the NMJ. Phase-contrast microscopy and fluorescent identification of acetylcholine receptors were used for the visualization of the NMJs. A strong signal was observed after staining with anti-human AChE antibodies indicating relatively strong muscular AChE contribution. faint signal was observed at the NMJ also after the staining with rat anti-AChE antibodies suggesting neural origin of a part of synaptic AChE. At present we can conclude that at the early stages of NMJ development (3-4 weeks of co-culture) majority of AChE is of muscular origin but that a small part might also be contributed by the motor neuron. Further experiments will be necessary to assess quantitatively the neuronal origin of AChE in developing NMJ.

MODELS OF THE EXTRACELLULAR DOMAIN OF THE NICOTINIC RECEPTORS AND OF AN AGONIST AND CA++ BINDING SITES
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We constructed a three-dimensional model of the amino-terminal extracellular domain of three major types of nicotinic acetylcholine receptor, (alpha7)5, (alpha4)2(beta2)3 and (alpha1)2(beta1)3 gammmadelta, on the basis of the recent X-ray structure determination of the molluscan Acetylcholine Binding Protein. Comparative analysis of the three models reveals that agonist binding pocket is much more conserved than the overall structure. Differences however exist in the side chains of several residues. In particular, a phenylalanine residue, present in beta2 but not in alpha7, is proposed to contribute to the high affinity for agonists in receptors containing the beta2 subunit. The semi-automatic docking of agonists in the ligand binding pocket of (alpha7)5 led to positions consistent with labeling and mutagenesis experiments. Accordingly, the quaternary ammonium head group of nicotine makes a pi-cation interaction with W148 (alpha7 numbering), while the pyridine ring is close both to the cysteine pair 189-190 and to the complementary component of the binding site. The intrinsic affinities inferred from docking give a rank order epibatidine > nicotine > acetylcholine, in agreement with the experimental values. Finally, our models offer a structural basis for the potentiation by external Ca++.

PROBING THE BINDING INTERFACE BETWEEN THE NICOTINIC ACETYLCHOLINE RECEPTOR AND A SHORT ALPHA-NEUROTOXIN THROUGH RECEPTOR-BIOTINYLTOXIN-STREPTAVIDIN TERNARY COMPLEXES
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We investigated the interacting surface between a short curarimimetic toxin and a muscular-type nicotinic acetylcholine receptor, looking for the ability of various biotinylated Naja nigricollis alpha-neurotoxin analogues to bind simultaneously the receptor and streptavidin. All these derivatives, modified at positions 10 (loop I), 27, 30, 33, 35 (loop II), 46, 47 (loop III) or N-terminal still shared high affinity for the receptor, and in the absence of receptor, they all bound soluble streptavidin. However, the proportion of the toxin/receptor complex, that bound onto streptavidin-coated beads, varied both with the location of the modification and with the length of the linker between biotin and the toxin. In the receptor/toxin complex, the concave side of loops II and III was not accessible to streptavidin, contrary to the N-terminus of the toxin and, to a certain extent, to loop I. On the convex face, loop III was the most accessible, while the tip of loop II, especially Arg30 seemed to be closer to the receptor. The present data demonstrate that short toxins neither lay parallel nor strictly perpendicular to the receptor extracellular wall, nor do they penetrate deeply into a crevice. These results fit nicely with 3D-models of interaction between long neurotoxins and their receptors and support the idea that short and long curarimimetic toxins bind in a nearly similar way.
THE AGONIST BIPHASIC DOSE-RESPONSE CURVE OF THE HUMAN ALPHABETA2 RECEPTOR BECOMES MONOPHASIC IN THE PRESENCE OF PKC INHIBITORS OR IN LOW LEVELS OF EXTRACELLULAR CALCIUM IONS

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In the last year it has become widely accepted that the agonist dose-response curve of human alpha4beta2 (h4b2) nicotinic acetylcholine (nACh) is biphasic. However, the molecular basis for the presence of multiple components in the agonist concentration-response curve, and their pharmacological nature, have not yet been fully determined, although their modulation by phosphorylation and chronic exposure to ligands has received some attention (Mileo, 1995). We have used the potent nicotinic agonist epibatidine to construct a characteristic two-component dose-response curve for h4b2 nACh receptors expressed in oocytes. The two components were modulated by chronic exposure to both competitive antagonists and agonists, and by Ca++, in a concentration dependent manner. Nicotine upregulated the high affinity component, whereas the high affinity component was abolished by antagonists. To investigate the mechanism whereby chronic ligand exposure modulated the two components, we used activators and inhibitors of Protein Kinase C to modify phosphorylation of the h4b2 nACh receptor. Inhibition of Protein Kinase C caused a significant reduction in the size of the high affinity component only. This study shows that the two components of the epibatidine dose-response curve for the h4b2 nACh receptor can be differentially modulated.

MOLECULAR CLONING OF NICOTINIC ACETYLCHOLINE RECEPTOR SUBUNIT GENES FROM THE PEACH-POTATO APHD, MYZUS PERSICAE

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Nicotinic acetylcholine receptors (nAChRs) play a major role in excitatory synaptic transmission in insects and are the primary target site for the recently introduced chloronicotinyl insecticides such as imidacloprid. These compounds display a high selectivity towards insect nAChRs and are particularly active against 'sucking pests' such as aphids and whiteflies. Despite its commercial importance as a target for insecticides and the recent cloning of nAChR subunit genes from a range of insect species, the insect nAChR is not well characterized at the molecular level and the subunit combinations required to reconstitute functional insect nAChRs are yet to be determined. We are investigating the molecular diversity of nAChR subunit genes in an important agricultural pest, the peach-potato aphid, Myzus persicae. Five genes have already been cloned as full-length cDNAs (Mpα1-4, Mpβ1) and others are currently being isolated based on their homology to nAChR subunit-like genes identified in the full genome sequence of Drosophila melanogaster. Heterologous expression studies of the Myzus subunit genes in Drosophila S2 cells have revealed some evidence for co-assembly of certain alpha/beta subunit combinations, although functional reconstitution of receptor complexes as judged by radioligand binding (epibatidine and imidacloprid) have so far been entirely dependant on co-expression of the Myzus alpha subunits with rat beta2 rather than Mpβ1. This suggests that one or more key Myzus nAChR subunits required for functional reconstitution of the native receptor have yet to be cloned, and ongoing efforts to identify and express these subunits will be presented.

NFkB REGULATES THE ACTIVITY OF HUMAN ACETYLCOLINESTERASE PROMOTER IN MUSCLE

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Precise expression of biological active acetylcholinesterase (AChE; EC 3.1.1.7) is required for development and maintenance of the neuromuscular junctions. Previous studies reveal that the expression of human AChE in muscle can be regulated by the classical cAMP-dependent pathway that involved cAMP-dependent protein kinase (PKA) and cAMP-responsive element binding protein (CREB), via the regulatory element called cAMP-responsive element (CRE) on promoter of the enzyme. In the present study, we investigated the role of another regulatory element namely NFkB, in regulating the AChE promoter activity. A about 2.2 kb human AChE promoter was tagged with a luciferase reporter gene, and the construct was transfected in chick myotube cultures for analyzing the luciferase activity. The transcriptional activity of human AChE promoter was shown to be regulated by NFkB signaling. One of the possible nerve-derived factors that utilizes this mechanism is the adenine 5-prime-triphosphate (ATP) and its correspondent receptor called P2Y purinoceptor. All these data suggested that the AChE promoter could be regulated by multiple signaling pathways for precise transcriptional control at the post-synaptic muscle. Acknowledgments: supported by grants from the Research Grants Council of Hong Kong (HKUST 60999/98M, 611200/M & 2999C).

NEW ESSENTIAL RESIDUES IN CHOLINESTERASE ACYL POCKET

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The statistical analysis of changes of the bimolecular rate constant (k2) of inhibition of cholinesterases (ChEs)—acetylChE from human, mouse and flies Musca domestica and Calliphora vicina, and horse butyrylChE—by dialkylphosphates (OPI, 67 compounds of 13 series) with the general formula (AlkO)2PO(O)N at elongation of alkyl radicals and change of their branching in comparison with three physical-chemical characteristics (hydrophobicity (hyd), polarity (pol), and volume of the side chain (vol)) of 6 amino acid residues in acyl and alkoxyl pockets variable in the studied ChEs (No 282, 287, 288, 290, 330, 335 in Torpedo ray acetylChE sequence) is performed. It has been shown that depending on structure of alkyl radicals, the rate of ChE interaction with OPI is determined by steric hindrances to sorption (residues 282, 287, 290, 335), hydrophobic interactions (288) or polarity of microenvironment (287). In particular, gradients of the inhibition constants at gradual elongation of alkyls from methyl to hexyl ones are correlated to pol(282), hyd(288), vol(335), vol(282) and vol(287) accordingly. Revealed dependences are statistically significant in most cases; however, rather low values of the correlation coefficient indicate influence of structure of the OPI leaving part. The decrease of the statistical significance with elongation of alkyl radicals seems to be due to an increase of the number of possible conformational states of the OPI molecule. Thus the list of essential residues in ChE acyl pocket has been enlarged.
VARIABILITY OF SUBSTRATE SPECIFICITY IN CHOLINESTERASES OF VERTEBRATES AND INVERTEBRATES

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The data on substrate specificity of cholinesterases from vertebrates and invertebrates, obtained during about 40 years work of Soviet school of cholinesterase researchers and from available foreign journals, were summarized. The relative rate of hydrolysis (ratios of \( V / V_{\text{Km}} \) or \( k / k_{\text{cat}} \)) and relative "affinity" of substrate to active center (ratios of \( V / K_{\text{M}} \) or \( k / k_{\text{cat}} \)) and relative "affinity" of substrate to peripheral "anionic" site (ratios of \( k_{\text{inhib}} / k_{\text{cat}} \)) of the choline (acetyl-, propionyl-, butyrylcholine, acetyl-\( \beta \)-methylcholine) and/or of corresponding thiocholine substrates by 59 cholinesterases from 49 different animals (chordates, insects, mollusks, nematodes) were analyzed. The characteristic features of the enzymes from different groups of animals were revealed. The absence of regular trends of parameters of cholinesterase substrate specificity during evolutionary development was shown. It is supposed, that the evolution of the cholinesterase active center occurs by neutral mutations and results both in divergence, and in convergence of their catalytic properties. This review, named as "Substrate inhibition — one of the aspects of substrate specificity of cholinesterases from vertebrates and invertebrates", is accepted in *J. Evol. Biochem. Physiol.*, 2001, vol. 37, no. 3 (English version of the Journal is published by Kluwer Academic/Plenum Press).

2-AMINOPERIMIDINE IS AN EFFECTOR OF CHOLINESTERASES

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Few hydrophobic quaternary nitrogen compounds were reported in the literature as activators of cholinesterases. We have studied the effect of 2-aminoperimidine (AP), which consists of a guanidine group fused to a naphthalene moiety, on the activity of cholinesterases. AP was found to inhibit the reaction of acetylcholinesterase (AChE) and butyrylcholinesterase (BChE) with their specific choline esters with Ki in the micromolar range. With a series of p-nitrophenyl (pNP) esters as substrates of BChE the reaction rate was increased in the presence of AP. The maximum rate enhancement, 9.5 fold, was obtained with pNP butyrate in the presence of 10 \( \mu \)M AP. On the other hand, AP was an efficient inhibitor of the hydrolysis of \( \alpha \)-nitrophenyl (\( \alpha \)NP) esters. The opposite effect of AP on the hydrolysis of pNP and \( \alpha \)NP esters is probably due to a diverse mode of interaction of the esters with the enzyme's active site in the enzyme-substrate-AP ternary complexes. A spectacular effect of AP was on the covalent inhibition of BChE by diphenyl carbamoyl chloride (DPC), which inhibits the enzyme with \( K_{i}=80,000 \) 1/(M.min). Addition of AP to the inhibition mixture led to a dramatic enhancement of the inhibition rate up to 300 fold, yielding inhibition rate constant of 2.4 \( \times \)10\(^{7}\) 1/(M.min), which is similar to that obtained by the most powerful inhibitors of cholinesterases. It seems that AP induces conformational changes in the cholinesterase molecule which can lead to striking activation effects.

PERIPHERAL BINDING OF ETHOPROPAZINE TO HORSE SERUM BUTYRYLCHOLINESTERASE

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The inhibition of purified horse serum butyrylcholinesterase (BChE) with ethopropazine (0.25-20 \( \mu \)M) was studied in order to evaluate the binding sites on the enzyme. Activities were measured spectrophotometrically with acetylthiocholine (ATCh; 0.05-80 mM) as substrate at 37°C in 1 M phosphate buffer pH=7.4. The pH-curve for the ATCh hydrolysis fitted well the Weibull equation: \( K_s = 0.25 \) and 2.0 mM respectively, \( \beta=3.2 \). This equation assumes two binding sites for the substrate on the enzyme, catalytic and peripheral, and the beta-value above unity indicates apparent substrate activation. Inhibition of BChE with ethopropazine was non-competitive at substrate concentrations up to 1.0 mM. The enzyme-inhibitor dissociation constant was 0.81 \( \mu \)M. Competition between ATCh and ethopropazine occurred at substrate concentrations above the \( K_s \) value for ATCh. Such inhibition pattern indicates binding of ethopropazine to the peripheral, non-productive, site on the enzyme.

SOME CONSIDERATIONS TO MOLECULAR MECHANISM OF CHOLINESTERASE CATALYSIS

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In due time we have offered the scheme of cholinesterase catalysis assuming formation of octagonal cyclic enzyme-substrate complex by the substrate carboxyl group and the enzyme imidazole (Brestkin and Rozengart. *Nature*, 1965, V.205, N 4969, P. 388-389). This scheme was concurred by many researchers and cited in several monographs on cholinesterases. However the modern concept of esteratic hydrolysis, based, in particular, on the data of X-ray crystallographic analysis, comprehends participation of catalytic triad Ser-His-Glu and oxynion hole, which nitrogens form hydrogen bonds with the substrate carboxylic oxygen. With the purpose to solve finally the problem on possibility of formation of cyclic enzyme-substrate complex, we have carried out researches by the method of molecular mechanics using programs ZMM and MVM (author D.B.Tikhonov, unpublished). It is shown, that the cyclic complex closure is impossible sterically in both investigated variants of interaction of acetylcholine with the triad (with Ser or with Glu). It confirms the scheme, accepted now, of reciprocation of the serine hydroxyl hydrogens during both acylation and deacylations. At the same time it is necessary to note, that the discussed problem on the one or two-proton mechanism of the catalysis is devoid of sense, as the protons in hydrogen bonds, both Ser-His and His-Glu, are delocalized perenniually, in free enzyme, in transition state and in acylated enzyme, and it should talk only about shift of the equilibrium. This delocalization of protons in catalytic triad is one of the causes of extremely high rate of the esteratic hydrolysis.
PHOSPHORYLATION OF ACETYLCHELINESTERASE AND THE PROPENSITY FOR REACTIVATION ANALYZED BY CHIRALITY AND MUTAGENESIS

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Mice acetylcholinesterase (ACHE) and its mutants were inhibited with Sp- and Rp-cyclopropyl-(CHMP), isopropyl-(PrMP), and 3,3-dimethylbutyl (DMMBP) methylphosphonothioate thiocholine esters. Double and triple mutants of ACHE had high level inhibition rates for both inhibitors except for Sp-CHMP which inhibited the mutant at a similar rate as wild type ACHE. On the other hand, modification of aromatic residues in the active centre of ACHE into aliphatic residues found in butyrylcholinesterase, F259S, F297I, and Y337A, enhance inhibition of Rp isomers thus approaching inhibition rates of butyrylcholinesterase. Upon the F297I/Y337A mutation, Rp esters of CHMP and PrMP became more reactive than Sp esters while reaction with the Sp esters was slightly reduced, displaying inverted stereo-specificity. Similar multiple mutations at these positions have been analyzed for reactivation by the oximes, 2-PAM and HI-4. Similar to the reactivation rates, the Sp formed conjugates show the more rapid reactivation rates. Certain multiple mutations yield substantial enhancements of reactivation rates. Binary combinations of oximes and these mutant enzymes may form effective scavenging agents.

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THE FIRST TWO NATURALLY OCCURRING ACTIVATORS/REACTIVATORS OF ACETYLCHELINESTERASE

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There is no satisfactory way to provide protection against toxic effects of acetylcholinesterases (antiCHE) in cases of environmental contamination by pesticides, chemical warfare, accidental overdoses of anaesthetics or accidental ingestion of antiCHE present in adulterated or contaminated food or fodder. Although some synthetic reactivators of acetylcholinesterase (ACHE) are known, practically occurring activator or reactivator has been reported thus far. We report isolation; physico-chemical properties and biochemical activities of two novel activator of ACHE (EC 3.1.1.7) found in wheat leaves. The compounds have been named as wheecheac (wheat ChE activator) and trititchac (Triticum ChE activator). Wheecheac is a novel diacylglycerol (DAG). It is an activator of ACHE at low concentrations (50 percent activation at 7 nM and 30 nM) but inhibitor at higher concentrations (50 percent inhibition at 10 microM). It is the first report of direct regulation by DAG of any enzyme other than protein kinase C. Wheecheac partially reactivates the neostigmine- and succinylcholin- inhibited electric eel ACHE (EC 3.1.1.7). Trititchac is a novel organophosphate (OP). It is the first organophosphate reactivator of ACHE in a scenario where organophosphates are almost synonymous with antiCHEs. Trititchac has 50 percent activation at 30 microM for electric eel ACHE. It also partially reactivates ACHE inhibited by neostigmine, succinylcholine and Phosphohemide. Discovery of activation/reactivation of ACHE by naturally occurring chemical compounds has tremendous implications in physiology, therapeutic- particularly for cholinergic dysfunction, and study of plant-animal co-evolution.

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KINETICS OF INTERACTION OF ETHOPROPAZINE ENANTIOMERS WITH BUTYRYLCHOLINESTERASE AND ACETYLCHELINESTERASE

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The association and dissociation rates of (+)-ethopropazine and (-)-ethopropazine with wild-type mouse and horse butyrylcholinesterases (BChE), and mutant mouse acetylcholinesterase (ACHE) were studied in order to analyze elements of stereo selectivity in two similar but distinct enzyme templates. Reaction traces at 23 oC in 0.1 M phosphate buffer pH 7.0 were recorded in millisecond time frame using Applied Photophysics stopped-flow apparatus equipped with fluorescence detection. Dissociation rate constants evaluated for BChEs were three to four-fold faster for the enantiomer (130 (1/min) vs. 39 (1/min) for horse BChE), while association rate constants for enantiomers were similar (1.4 (1/nM*min) vs. 1.0 (1/nM*min) for horse BChE), resulting in lower equilibrium dissociation constant and better binding of (+)-ethopropazine with BChE. In the ACHE template the preferential binding of (+)-ethopropazine was observed with Tyrs373Aia mutant, but with an order of magnitude greater stereo specificity. Binding preference for Tyr124Gln ACHE mutant, however, was reversed and (+)-ethopropazine (Kd of about 1.8 uM) bound several fold better than (-)-enantiomer. Inhibition of enzyme acetyltihiocholine hydrolysis by ethopropazine enantiomers yielded equilibrium inhibition constants similar to equilibrium dissociation constants derived from stopped-flow rate measurements. In conclusion, active cleft gorge of ACHE, lined with larger number of aromatic residues than the gorge of BChE, provides narrower and more stereo selective environment for binding of ethopropazine.

POLYURETHANE IMMOBILIZED ENZYMES: OF SENSING AND DECONCATINATING MATRIXES


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During combat, personnel have been exposed to organophosphates (OPs). Other exposures to chemicals contain insects or terrorist acts in subways or sports events. For successful survival of exposed persons with minimal adverse effects, it is important to have rapid and simple detection of OPs and also uncomplicated deconcatination and detoxification procedures. To accomplish this, we are developing enzyme-immobilized polyurethanes configured as (1) biosensors for OPs or (2) as sponges to soak up and inactivate the OPs. As a biosensor for OPs, the polyurethane matrix is composed of cholinesterase or other OP hydrolizing enzymes to both indicate the presence of the OP agents, and to differentially indicate the type of OP present in the field. One of the advantages this immobilization technology affords the enzymes is that they are resistant to denaturing events, and are now suitable for sampling OPs in diverse environments such as soil, large bodies of water, as well as conventional airborne contamination. In the second configuration, polyurethane sponges are synthesized with enzymes and agents for external treatment of contaminated skin and other sensitive and exposed surfaces. To detoxify OPs, the cholinesterase is combined with oximes so the catalytic activity of OP-inhibited enzyme is continuously restored. Additional post-synthesis components include compounds to improve the extraction of OPs from guinea pig skin. Resulting sponges provided protective ratios of about 15 and 30-fold for soman and VX, respectively, when tested in a guinea pig model. These immobilized enzyme biosensors and sponges, by virtue of their high capacity for enzymes, stability, specificity, sensitivity, and resistance to harsh environmental conditions, can be used under diverse conditions encountered by troops and civilian first responders in the field.
The inhibition of acetylincholinesterase (AChE) by organophosphorus (OP) nerve agents (sarin, soman, GF and VX) and pesticides (paraxon and its analogs) poses a continuous threat due to the possible use in battlefield, terrorist attack or in agriculture. Antidotes containing oxime compounds to reactivate the inhibited enzyme are highly valued for treatment against OP poisoning. One of these reactivators, HI-6, is significantly more effective in alleviating toxicity due to soman and GF compared with the old generation of oximes such as 2-PAM, TM64 and LDH6. However, HI-6 shows is less effective as an antidote for the treatment of OP pesticides poisoning compared to other oximes. The mechanism responsible for this observed selectivity of HI-6 is unknown.

In the present study, the mechanism of HI-6-induced reactivation of AChE-OP conjugates was investigated by using mutant mouse AChEs and different OPs such as paraoxon, 7-(methylthio)phosphonyl-oxyl-1- methylquinolinium iodide (MEPQ), soman, sarin, and Sp and Rp isopropyl methylphosphonic thiocolines. The effect of mutations on HI-6 induced reactivation was examined and results indicate that HI-6 can assume two possible orientations in the active site gorge of AChE: the reactive center against a different types of AChE-OP conjugates. For all enzyme conjugates formed with the Sp anamorphs of organophosphonates, Tyr 124 at the peripheral site is an important element in facilitating reactivation by HI-6 since its mutation to Gln drastically reduced reactivation by HI-6. On the other hand, HI-6 is less effective in reactivating enzyme conjugates formed with the Rp enantiomers and paraoxon, and mutations at the peripheral site do not have a significant effect on reactivation by HI-6. The structure-function correlation suggested by the study with mutant enzymes was further corroborated by reaction studies with an analog of HI-6, which is devoid of the other oxygen. These results suggest that the reactivation potency of HI-6 depends on its orientation in the active site gorge, which is dictated by the structure of the OP.

**ATTEMPTS TO ENGINEER AN ENZYME-MIMIC OF BUTYRYLCHOLINESTERASE BY SUBSTITUTION OF THE SIX DIVERGENT AMINO ACIDS IN THE ACTIVE CENTER OF ACETYLCHOLINESTERASE**

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The active center gorge of human acetylcholinesterase (HuAChE) is lined by 14 aromatic residues, whereas in the closely related human butyrylcholinesterase (HuBChE) 3 of the aromatic active center residues (Phε295, Phe297, Tyr337) as well as 3 of the residues at the gorge entrance (Tyr72, Tyr124, Trp286) are replaced by aliphatic amino acids. For all the prototypical noncovalent active center and peripheral site ligands tested, the hexa-mutant of HuAChE (Y72D/Y124Q/W286A/F295L/F297V/Y337A) displayed a reactivity phenotype closely resembling that of HuBChE. These results support the accepted view that the active center architectures of AChE and BChE differ mainly by the presence of a larger void space in BChE. Nevertheless, reactivity of the hexa-mutant HuAChE toward the substrates acetylthiocholine and butyrylthiocholine and the transition state analog m-(N,N,N-trimethylammonio)tri-fluoroacetophenone (TMFTA) is about 45-170-fold lower than that of HuBChE. Most of this reduction in reactivity is manifested by the triple active center mutant F295L/F297V/Y337A. We propose that the hexa-mutant HuAChE, unlike BChE, is impaired in its capacity to accommodate certain tetrahedral species in the active center. This impairment is probably related to the enhanced mobility of the catalytic histidine His447, as observed in molecular dynamics simulations of the hexa-mutant or the F295L/F297V/Y337A HuAChE enzyme but not in the wild type HuAChE.

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**MALDI-TOF/MS ANALYSIS OF ACETYLCHOLINESTERASE-LIGAND CONJUGATES: A TOOL FOR RESOLUTION OF MECHANISTIC PATHWAYS**

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Understanding reaction pathways of phosphorylation and reactivation of AChE and "aging" of the corresponding AChE adducts is both a biochemical and a pharmacological challenge. Here we describe experiments which allowed to resolve some of the less well understood reaction pathways of phosphorylation, and "aging" of acetylcholinesterase (AChE) involving phosphoroamidates (P-N agents) such as tabun or the widely used pesticide methamidophos. Tryptic digests of phosphorylated AChEs (from human and Torpedo californica), ZipTip peptide fractionation and Matrix Assisted Laser Desorption Ionization Mass Spectrometry (MALDI-TOF/MS) enabled reproducible signal enrichment of the isotopically resolved peaks of organophosphoroamide-conjugates of the AChE active site Ser peptides. For tabun and its hexa-deuterio-analog we find, as expected, that the two phosphoramidate adducts of the active site peptide differ by 6.05 mass unit but following aging we find that the two corresponding phospho-peptides have identical molecular weights. We further show that the aging product of paraxoxon-AChE adduct is identical to the aging product of the tabun-AChE conjugate. These results unequivocally demonstrate that the pathway of aging of tabun adducts of the human or the Torpedo californica AChEs proceeds through P-N bond scission. For methamidophos we show that phosphorylation of AChE involves elimination of the thiomethyl moiety and that the spontaneous reactivation of the resulting organophosphate adduct generates the phosphorous free AChE active site Ser-peptide.

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X-RAY STRUCTURE OF TORPEDO ACHE COMPLEXED WITH BIFUNCTIONAL LIGANDS RELATED TO HUPA: NOVEL DRUGS FOR TREATMENT OF ALZHEIMER'S DISEASE

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Reversible AChe inhibitors of AChe slow the cognitive decline experienced by Alzheimer patients. (+)-Huperzine A (Hupa), an alkaloid isolated from the shrub Huperzia serrata, is one such inhibitor, but the search for more potent and selective drugs continues.

Based on the bivalent strategy and on structural studies on TaAChe, dimerization of a pharmacologically inactive fragment of Hupa (monomer E) produced a drug with twice the potency of the natural product. We soaked two alkylene-linked bis-Hupa-like AChe inhibitors, (S,S)(+-)-E12E and (S,S)(+-)-E10E, into trigonal TaAChe crystals and solved the X-ray structures using the difference Fourier technique, both to 2.15 angstroms resolution. The structures revealed one E unit bound to the 'anionic' subsite of the active site, at the bottom of the active-site gorge. The E and the second E unit adjacent to Trp279 in the 'peripheral' anionic site at the top of the gorge, with both bifunctional molecules thus spanning the active-site gorge. The results confirm that the increased affinity of the dimeric Hupa analogs for AChe is conferred by binding to the two 'anionic' sites of the enzyme.

CRYSTALLIZATION AND DETERMINATION OF THE X-RAY STRUCTURE OF HUMAN ACHE

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For certain AChe inhibitors, crystallographic studies of their complexes with AChe from a particular species may fail to explain differences in binding affinity observed for AChe from different species. Thus, it would be preferable to study the binding affinity in solution and the X-ray structure of the complex for AChe from the same species. To date, the only structural reports on complexes of AChe with potential drugs for the management of Alzheimer's disease (AD) are of AChe from Torpedo californica (Tc), whereas studies on complexes with human AChe (hAChe) would obviously be preferable. We have recently crystallized a recombinant catalytic domain (55E residues) of hAChe, and solved its X-ray structure at 3.2 A resolution by molecular replacement. The data revealed a P6(1) space group with unit cell dimensions: 210.9 210.9 115.3 angstroms and angles of 90, 90, 120 degrees. The asymmetric unit contains a dimer with a non crystallographic 2-fold symmetry similar to the crystallographic 2-fold symmetry previously seen for the native dimer. X-ray data have been collected from native hAChe crystals soaked with the anti-Alzheimer drugs: (+)-huperzine A, tacrine, huprine X and ENA-713. The structure determination of these complexes is currently being pursued.
3D STRUCTURE OF TORPEDO CALIFORNICA ACETYLCOLINESTERASE COMPLEXED WITH HUPRINE X

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Huprine X is a novel AChE inhibitor, with one of the highest affinities reported for a reversible inhibitor (Camps et al Mol. [2000] Pharmacol. 57, 409-417). It is a synthetic hybrid containing the 4-aminquinolone substructure of one anti-Alzheimer drug, tacrine, and a carbobicyclic moiety resembling that of another, (+)-huperzine A. The crystal structure of its complex with Torpedo californica AChE (TcAChE), determined to 2.1 angstroms, shows that huprine X binds to the anionic site, and also binds to the esteratic site. Its aromatic portion occupies the same binding site as tacrine, whereas the carbobicyclic unit occupies the same binding pocket as (+)-huperzine A. Inhibition data show that huprine X binds human AChE (hAChE) and Torpedo AChE 28-fold and 34-fold, respectively, more tightly than tacrine. This difference stems from the fact that the aminoquinoline moiety of huprine X makes interactions similar to those made by tacrine, but additional bonds to the enzyme are made by the huperzine-like substructure and the chlorine atom of huprine X. Furthermore, both tacrine and huprine X bind more tightly to TcAChE than to hAChE, suggesting that their quinoline substructures interact better with Phc330 in TcAChE than with Tyr337 in hAChE.

LIGAND INDUCED CONFORMATIONAL CHANGES IN THE OMEGA LOOP OF ACETYLCOLINESTERASE REVEALED BY FLUORESCENCE SPECTROSCOPY

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We have used a combination of cysteine substitution mutagenesis and site-specific fluorescence labeling to characterize the structural dynamics of mouse AChE. Cysteines were substituted at positions 262 (distal disulfide loop), 124 and 287 (gorg entry and rim), 76, 81 and 84 (outer rim of the omega loop). In turn, these Cys residues have been specifically labeled with fluorophores, and fluorescence spectra and decay of anisotropy of the fluorescent side-chains analyzed in relation to the kinetic parameters of ligand binding. Residue 262 shows rapid decay of anisotropy and no evidence for a global conformational change upon ligand binding. Ligands which occlude the gorge opening, such as fasudil or extended bisquaternary ligands, cause blue shifts in the spectra of acrylodan when conjugated to residues 124 and 287; this reflects an increase in local solvent exclusion at these positions by the bound ligand. Unexpectedly, the binding of active center ligands and fasudil induce a large red shift in emission of acrylodan conjugated at the three outer residues on the omega loop, 76, 81 and 84, indicating an enhanced solvent exposure upon ligand binding. Concomitantly, time resolved fluorescence anisotropy reveals a more rapid rate of anisotropy decay at positions 81 and 84 upon active site ligand binding. Thus, ligand binding at the active center induces a local change in conformation of the omega loop not revealed in crystal structures.

bis-ACTING GALANTHAMINE DERIVATIVES AS IMPROVED DRUGS IN THE SYMPTOMATIC TREATMENT OF ALZHEIMER'S DISEASE

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The alkaloid galanthamine (GAL), isolated from the Amaryllidaceae family of plants, shows strong, reversible anticholinesterase activity. As such it has been tested as a possible alternative to current anticholinesterases such as Aricept1, used in the palliative treatment of Alzheimer's Disease. GAL is already in use in Austria and has been approved for use in the UK and the USA, under the trade name Reminyl. It interacts with several residues in the active site of acetylcholinesterase (AChE) at the bottom of the "gorge," including Trp84, which binds the quaternary ammonium group of acetylcholine. In an effort to improve the efficacy of this drug, derivatives have been synthesized with the aim of interacting with both the active site and the second cation-binding site at the top of the gorge of AChE, viz. the peripheral binding site. The crystal structures of complexes of two such compounds with Torpedo californica AChE have been solved and refined and will be presented.

COMPARISON OF TWO REACTION SCHEMES FOR THE HYDROLYSIS OF ACETYLCHELCHOLINE BY BUTRYRYCOLINESTERASE

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Catalytic parameters calculated from the schemes derived by Webb (Model A) and by Stojan (Model B) were compared for the hydrolysis of acetylthiocholine (ATCh) by purified horse serum butryrycholinesterase (BChe) (measured by conventional and stopped-flow techniques). In Model A the acetylated enzyme is omitted from the scheme, while in Model B the Michaelis complex is omitted. The enzyme-substrate dissociation constants Ks and Kss in Model A were 0.25 and 2.0 mM (conv. tech.) and 0.17 and 6.3 mM (stopped-flow tech.). These were attributed to binding of ATCh to the catalytic and peripheral site of BChe. The constants K1 and K2 in Model B (stopped-flow tech.) were 0.223 and >1000 mM. These are attributed to binding of ATCh to an unidentified site in the free enzyme and to the peripheral site in the acetylated enzyme. As the Ks and K1 values are almost the same, both constants are likely to refer to the same enzyme-substrate complex. The constants Kss and K2 are very different; they are both attributed to peripheral binding of ATCh, but they obviously refer to different complexes. Both models postulate that peripheral binding of a substrate affects rate constants of substrate hydrolysis; this also follows from the evaluated constants for ATCh hydrolysis.
QUANTAL ACETYLCHOLINE RELEASE THROUGH MEDIATOPHORE PROTEOLIPID OVER-EXPRESSED IN NEUROBLASTIC CELLS

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This review summarizes a series of experiments demonstrating that reconstitution of mediatophore in transfected cells support quantal acetylcholine (ACh) release. Mediatophore is a proteolipid initially identified in presynaptic membrane of Torpedo cholinergic nerve terminals. It is formed by homo-oligomeric assembly of 15-16 kDa subunits showing a very high homology with either the 'c' subunit of the V-ATPase transmembrane sector (Vo) and the subunit of a recently identified proteolipid fusion pore in yeast vacuoles. Transfection of the Torpedo 15-16 kDa subunit in non cholinergic neuroblastoma cells (N18TG-2) enable them to perform rapid, Ca2+-dependent and quantal release of ACh from a cytosolic pool, in response to electrical stimulation. Transfection of choline acetyltransferase (ChAT) supports ACh release only if mediatophore is transfected as well. Furthermore, evoked release by mediatophore transfected cells is modulated by co-expression of ChAT but also vesicular ACh transporter (VACHT). Together these data indicate that mediatophore proteolipid plays a central role in the process of quantal ACh release. Mediatophore may either work as an transmitter permeable channel using cytosolic transmitter, or form a fusion pore which can allow kis and run off or full exocytosis of synaptic vesicles. One question remains: can these different modes of release co-exist at the same active zone, depending on the presynaptic activity and vesicle life cycle, or are they differentially expressed in specific types of synapses according to their architecture and function? 

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CAPILLARY ZONE ELECTROPHORESIS DETECTS UNWANTED CHOLINESTERASE-BOUND HIDDEN LIGANDS THAT MODULATE ENZYME CONFORMATIONAL STABILITY

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Proteins of pharmacological interest have to exhibit their nakedness to become therapeutic drugs. Thus, detection of compounds present in biopharmaceuticals is of central concern. Among them, cholinesterases (ChEs) are enzymes of major importance for detoxification of poisonous esters. ChEs are characterized by an asymmetrical distribution of charged residues, thought to increase enzyme efficiency, and a remarkable conformational plasticity, to allow adjustment of the active site and the allosteric control of activity. Likewise, ChEs display high catalytic efficiency of an active site position at the bottom of a deep gorge. The gorge can be partially or fully occupied by ligands (substrates or inhibitors). Accordingly, a suitable method allowing to detect unwanted ligands and their influence on the functional conformation and stability of these enzymes was essential. We have developed a capillary zone electrophoresis (CZE) approach for that purpose [1]. The factors causing discrepancies between data for thermal unfolding of ChE by electrophoretic and by calorimetric methods were investigated. The presence of unwanted hidden ligands bound to purified cholinesterase was first demonstrated. The role of hidden bound ligand in stabilization of AChE samples was emphasized. Our results raised several questions concerning the real conformation of the native state of enzymes such as ChEs. Finally, CZE was proven to be a pertinent tool to validate the conformity of purified enzymes to a status of biopharmaceutical. [1] Rochu D. et al. 2002, Electrophoresis, in press. This work was supported in part by DSP/STTC CO No 99-C0029 to P.M.

CRYSTAL STRUCTURE OF TORPEDO CALIFORNICA ACETYLCHOLINESTERASE WITH A NOVEL GALANTHAMINE DERIVATIVE: IMPLICATIONS FOR THE DESIGN OF NEW ANTI-ALZHEIMER DRUGS

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Galanthamine, a tertiary alkaloid extracted from several species of has received recent attention as a centrally acting, selective, competitive, and reversibly showing to produce significant improvement of cognitive performances in Alzheimer's disease patients. This compound is less potent than tacrine and E2020 but has excellent pharmacological and pharmacokinetic profiles and exhibits very low hepatotoxicity and fewer side effects. Galanthamine, together with physostigmine, is one of very few drugs showing a dual activity, acting not only as an inhibitor of AChE but also as an allosteric potentiator of the nicotinic response induced by ACh and competitive agonists. The crystal structure of Torpedo californica AChE with galanthamine has been determined [1,2] and computerized molecular docking studies successfully predicted its bound conformation [3]. Structure-based drug design is an important tool in the development of second generation candidate drugs based on a lead compound. The crystal structure of Torpedo californica AChE with a novel galanthamine derivative was solved to 2.3 Angstroms resolution. The structure revealed that the galanthamine moiety binds at the base of the active site gorge, interacting with both the aeyl-binding pocket and the choline binding site. The charged nitrogen of the piperidine moiety, tethered to galanthamine via a functional linker, makes a cation-pi interaction with the phenyl ring of Tyr334. [1] H. M. Greenblatt et al. (1999) FEBS Lett. 463, 321-326. [2] C. Bartolucci et al. (2001) Proteins 42, 182-191. [3] C. Pilger et al. (2001) J. Mol. Graph. and Modell. 19, 288-296. This work was supported by SanofiAventis Pharmazeutika AG, Vienna, Austria.
X-RAY STRUCTURE OF SOMAN-AGED HUMAN BUTYRYLCHOLINESTERASE

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Acetylcholinesterase (AChE) and butyrylcholinesterase (BChE) phosphorylated by branched organophosphates (OP) undergo a dealkylation reaction ('aging'), leading to OP-enzyme conjugates that cannot be reactivated by oximes. Previous studies from our laboratory revealed that aged-BChE conjugates are more stable to high temperature and pressure than the native enzyme. The increase in stability was shown to be related to changes in the water structure network in the active site gorge and to the formation of a salt bridge between PO- and protonated His338. Such a bridge was observed in the crystal structure of soman-, sarin- and DFP-aged AChE. These x-ray structures also revealed that the reactivation of aged conjugates was prevented by the stabilizing effect of H-bonding between a P bound oxygen and the oxazolone hole, and by the 'drying' effect of the acyl-binding pocket, preventing the nucleophilic attack. To complete our understanding of the molecular mechanism of the aging reaction of phosphorylated BChE, the X-ray structure of a engineered form of human BChE has been determined. The structure of the soman-BChE aged conjugates was solved to 2.4 Ångstrom resolution and compared to the aged conjugates of Torpedo californica AChE. Because the aged conjugate has no activity, we used this feature to determine the binding location of butyrylthiocholine in the active site of the phosphorylated enzyme. The structure of the ternary complex aged BChE/butyrylthiocholine was solved to 2.3 Ångstrom resolution. This later structure was aimed to provide new information on the peripheral site of BChE and the related phenomenon of substrate activation.

STUDIES ON DYNAMICAL TRANSITIONS IN CHOLINESTERASES

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Acetylcholinesterase (AChE) hydrolyses the neurotransmitter, acetylcholine, very rapidly, as required for termination of impulse transmission at cholinergic synapses. Its 3D structure reveals a deeply buried active site accessed by a narrow gorge [1]. This unanticipated structural characteristic raises cogent questions concerning traffic of substrates and products to and from the active site. It is obvious that substantial 'breathing' motions of the protein are required for penetration of substrates to the active site via the gorge and for clearance of products via routes as yet undefined. It is, therefore, important to characterise the dynamics of the enzyme on various time scales. We have chosen incoherent elastic neutron scattering (IENS) [2] to investigate global atomic dynamics of both hydralazine hemochromate AChE (DmAChE) and of DmAChE dried by lyophilization as a function of temperature, and compared them to corresponding samples of human butyrylcholinesterase, an enzyme structurally very similar to AChE, whose biological function is still unknown. The atomic mean square displacements (MSD) obtained by this technique reveal a dynamical transition for both hydrated samples. Such a transition has been observed previously for myoglobin [3], as well as for bacterialidendeprotein [4], and has been associated with the onset of biological function. Surprisingly, MSDs of the dry samples exceed those of their hydrated counterparts in the intermediate temperature range. Future objectives include theoretical interpretation of these findings, as well as investigation by IENS of the influence of covalently and reversibly bound inhibitors on the dynamics of these enzymes. [1] Suarez JL, Harel M, Eronoff F, Oefner C, Goldman A, Toker L & Simm J (1991). Atomic structure of acetylcholinesterase from Torpedo californica: A prototypic acetylcholine-binding protein. Science 233, 872-879. [2] Zacc1, G. (2001). How Soft is a Protein? A Protein Dynamics Force Constant Measured by Neutron Scattering. Science 284, 1603-1607. [3] Toker L, Cusack S & Petry W (1989). Dynamical transition of myoglobin revealed by inelastic neutron scattering. Nature 337, 754-756. [4] Ferrand M, Dourmashkin A, Petry W & Zacc1 G (1993). Thermal motions and function of bacteriendependin in purple membranes: Effects of temperature and hydration studied by neutron scattering. Proc. Natl. Acad. Sci. USA 90, 9664-9672.

TETANIC FADE IS REVEALED BY BLOCKING PRESYNAPTIC NICOTINIC RECEPTORS CONTAINING ALFA4BETA2 AND ALFA3BETA2 SUBUNITS AFTER REDUCING THE SAFETY FACTOR OF NEURONAL TRANSMISSION

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The effects of subtype specific nicotine receptor (nAChR) antagonists were studied on nerve-evoked tetanic (50Hz, 5sec) contractions and [3H]acetylcholine release from rat isolated hemidiaphragms. nAChR antagonists reduced tetanic peak tension with a rank potency order of alfa-bungarotoxin(BTX)-d-tubocurarine(TC) >> mecamylamine(Mecaa) > hexamethonium(Hex). Depression of tetanic tension by dihydro-beta-erythroidine(DHBE, 0.03-10 microM, an alfa4beta2 and alfa3beta2 antagonist), methyllycaconitine(MLA, 0.003-3 microM, an alfa7 antagonist) and alfa-conotoxin MII(CTX MII, 1-300 nM, a selective alfa3beta2 antagonist), did not exceed 30% TC (0.1-0.7 microM), Mecaa (0.1-300 microM) and Hex (0.03-3 nM) induced both tetanic fade and peak tension depression. With DHBE (0.03-10 microM) and CTX MII (1-300 nM), tetanic fade was only evident after decreasing the safety factor of neuromuscular transmission (McGiz, 6-7 nM). Neither BTX (3-100 nM) nor MLA (0.003-3 microM) produced tetanic fading. The antagonist rank potency order to reduce (50%) evoked [3H]-ACh release from motor nerve terminals, was CTX MII(100 nM) ≡ TC(1 microM), DHBE(1 microM) > Hex(1 nM); BTX (300 nM) failed to affect ACh release. The results suggest that neuromuscular block is linked to the activation of BTX-sensitive nAChRs containing alfa1-subs units, whereas preferential blockade of neuronal alfa4beta2- and alfa3beta2-containing receptors cause tetanic fade by reducing nicotinic autofacilitation.

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MOLECULAR CHARACTERISATION OF ACETYLCHOLINESTERASE FROM THE PEACH-POTATO APHID MYZUS PERSICAE(SULZ)

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The peach-potato aphid, Myzus persicae, is an extremely important pest that occurs worldwide causing direct feeding damage to many field and glasshouse crops. It is, more importantly, a major vector of many plant viruses including virus yellows in sugar beet. A modified acetylcholinesterase (AChE) that confers specific insensitivity to prinema carb and tricazam has been previously reported in Myzus persicae. In order to determine the genetic basis of this insensitivity, the AChE gene was amplified from both sensitive and insensitive forms using RT-PCR. However, no mutations were identified which could account for this insensitivity. The presence of a second gene was thus proposed. Alternative degenerate primers were designed based on published ace sequences from both vertebrate and invertebrate organisms. This led to the amplification of a partial sequence that differed from Myzus AChel sequence. Gene specific primers have been designed based on this sequence to characterise any mutations found in an insensitive form of the enzyme. Additionally, AChE has also been purified using affinity chromatography. The results of direct sequencing of this AChl protein is compared with amplified sequence from PCR to verify the second Ace sequence.
PECULIARITIES OF KINETIC BEHAVIOUR OF FISH'S ABRAMS BALLERUS BLOOD SERUM CHOLINESTERASE

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The cholinesterase (ChE) of fish's blood serum is the typical acetylcholinesterase (AChE) with the same substrate specificity as AChE from human erythrocytes. On the other hand, the blood serum of some fish's blood plasma contains mainly butyrylcholinesterase (BuChE). The blood plasma ChE (FChE) of fishes (blue bream - Abramis ballerus, roach - Rutilus rutilus) was purified for study of kinetic behaviour and sensitivity to anticholinesterase compounds. The kinetics of choline and thiocyanate ethers hydrolysis has been studied at the presence of new purified (FChE). The sensitivity of enzyme to organophosphates, carbamates and reversible ChE inhibitors has been determined. The results of this and the following experiments indicate that the new purified FChE can be classified as BuChE, so the velocity of butyrylcholine and butyrylthiocholine hydrolysis is more than other substrates. At the same time, this type of new enzyme differs from other typical BuChE, so the hydrolysis rate of butyrylcholine by FChE is in 10 - 13 times more rapid as compared to hydrolysis of acetylcholine. The sensitivity of FChE to organophosphates is in 100-1200 times higher than the sensitivity of all types of commercial ChE. It is very unexpectedly and unordinary that FChE has a very small sensitivity to active carbamates and reversible ChE inhibitors. The results obtained in this investigation suggested that there might be an essential difference between the active sites of FChE and another types of cholinesterases.

LETHAL EFFECTS OF HEAD-TO-TAIL 3-ALKYLPIRIDINIUM POLYMERS ISOLATED FROM THE MARINE SPONGE RANIERA SARA: ACHÉ INHIBITION OR UNSPECIFIC BINDING TO SERUM PROTEINS?

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The toxic aqueous extract polymeric 3-alkylpyridinium salts were isolated from the marine sponge Raniera sarai. Both in vitro and in vivo experiments were performed to evaluate the role of acetylcholinesterase (AChE) inhibition in the toxin lethality. Binding of the polymer to the Drosophila AChE was estimated by the fluorescence of the protein polymer complex (excitation at 290 nm). On male Wistar rats ECG, blood pressure and breathing pattern were monitored. In vitro the polymers show a quick initial reversible binding, followed by slow binding and finally by irreversible inhibition of AChE. In vivo doses lower than 1 mg/kg caused bradycardia and prolongation of expiration. At doses above 2.7 mg/kg all treated animals died, but signs were not typical of acetylcholinesterase inhibition. Binding of the R. sarai inhibitor seems to occur at the active site gorge. However, unusual inhibition kinetics suggests that there are several affinity binding sites. At lower doses, the toxin caused changes of the measured parameters typical of AChE inhibition. At lethal doses the possible AChE inhibitory effects were probably covered by the other, more pronounced lethal effects of the toxin. At both concentrations only small fraction of the toxin could act in neuromuscular junction since the huge molecular conglomerates (20 nm) of the toxin molecules develop in vitro. Those aggregates could hardly cross the capillary wall and enter the neuromuscular junction in vivo. The influence of the toxin on platelets aggregation, and later fibrin segregation could be the primary and most important in the toxin lethality.

STRUCTURAL INSIGHTS INTO THE INTERACTIONS AT THE ACETYLCHOLINESTERASE PERIPHERAL ANIONIC SITE

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The AChE peripheral anionic site, located at the entrance of the enzyme active site gorge, consists of a matrix of overlapping loci for the binding of the diverse ligands that serve as allosteric activators or inhibitors; yet, the molecular mechanisms coupling this site to the remote active center to modulate catalysis remain unclear. The peripheral site has also been proposed to be involved in cellular adhesion through formation of heterologous protein interactions that occur during synaptogenesis or the onset of degenerative neuropathologies. Based on a new crystal form of mouse AChE and combined spectrophotometric crystal analyses, unique structures of the enzyme with a free peripheral anionic site, and as three complexes with pentantridinium and pyrrolyl peripheral site inhibitors, were solved in the 2.20-2.35 A resolution range. A comparative analysis with earlier structures of AChE in complexes with the peptide fasciculin or with organic bifunctional inhibitors unveils new structural determinants contributing to ligand interactions at the AChE peripheral anionic site, and permits a detailed topographic delineation of its locus. Hence, these structures provide templates for ligand docking that may be useful for designing compounds directed to the enzyme surface and able to modulate specifically surface interactions involved in non-catalytic AChE function. Supported by USPHS grant GM418360 and DAMD grant 17-1-8014 to PT, the NSF-CNRS collaborative project 39026 to PM and PT, the AFM to PM and PE Bougis (CNRS-UMR6560).

PIPERONYL BUTOXIDE: A SPECIFIC INHIBITOR OF INSECTICIDE RESISTANT ACETYLCHOLINESTERASE

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The methyldioxyphenyl compound piperonyl butoxide (PBO) has a long history as an insecticide synergist, yet it is a compound whose mode of action is still poorly understood. PBO was long assumed to be solely a cytochrome P450, however, it is known to be capable other functions, such as esterase inhibition. The objective of this study was to explore to effects of PBO on insect acetylcholineserase (AChE). In vitro, PBO inhibited organophosphate or carbamate insensitive AChE in Helicooverpa armigera, Helicoverpa punctigera, B-orange Isetimia tabaci and Aphis gossypii. Inhibition was particularly strong in H. armigera. In vivo studies also supported these findings. PBO appeared to enhance the activity of insecticide sensitive AChE (H. armigera, H. punctigera, B. tabaci, A. gossypii however, this is likely consequence of PBO interaction with the substrate acetylthiocholine iodide. Inhibition of insensitive AChE provides very exciting possibilities for the control of an intractable resistance mechanism. While PBO is unlikely to be acting as a conventional inhibitor of insensitive acetylcholinesterase, observations suggest that PBO is entering a physicochemical reaction with insensitive AChE, making it unavailable to the substrate. While it is unknown, as to how structure differs between sensitive and insensitive AChE, it is commonly the case that the mutation conferring insensitivity decreases the affinity of the protein for its natural substrate. PBO may have a three dimensional structure able to penetrate the AChE active site.
SIGNIFICANCE OF PARAMETERS BETWEEN VARIOUS KINETIC SCHEMES FOR CHOLINESTERASES

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Cholinesterases show homotropic pseudo-cooperative phenomena. At intermediate substrate concentrations the activities of these enzymes exceed the expected values according to Michaelis-Menten mechanism and at very high substrate concentrations they are inhibited. A kinetic model [Stojan et al. FEBS Lett. 440 (1998) 85-88] was put forward which can describe the data of entire substrate concentration interval and also in the presence of the inhibitors. However, the criticism has been raised, since some steps in the reaction mechanism were roled together, thus not reflecting subsequent events during the catalytic process. Although some evaluated constants can be explained in terms of classical kinetic parameters, we performed an extensive mathematical characterisation to evaluate the significance of determined kinetic parameters. We have compared their values and derived the appropriate connections with Webb model and with the traditional reaction scheme for vertebrate acetylcholinesterase.

STUDIES OF ACETYLCHOLINESTERASE FROM THE PEACH-POTATO APHID, MYZUS PERSICAE(SULZ.)

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Acetylcholinesterase (AChE) is the target site for two important classes of insecticides, organophosphates (OPs) and carbamates. Widespread use of these compounds has led to the development of resistance in many insect species, often due to an insensitive form of the enzyme. To identify the structural changes that confer protection and triazine insensitivity to acetylcholinesterase (AChE) in the peach-potato aphid, Myzus persicae, the ace gene (600 amino acids) from susceptible and resistant clones were sequenced using RT-PCR and RACE. However, no mutational changes were identified between the sensitive and insensitive enzymes. Possible explanations for this observation include the presence of post-translational modification, or the possibility that the sequence obtained is not that of the synaptic AChE. Since primers were derived from conserved regions of dipteran sequences, novel reversible inhibitors synthesised by Syngenta were employed to elucidate possible differences between sternorrhyncha and non-sternorrhyncha species. Multidimensional scaling and hierarchical cluster analysis of inhibition data utilising these reversible inhibitors against a variety of insect species revealed compelling evidence that the sequence obtained was not that of synaptic AChE.

EXPLORING THE ACHE GORGE WITH GALANTHANINE

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Acetylcholinesterase (AChE) plays a key role in the development of Alzheimer's disease as this enzyme is responsible for cleavage of the neurotransmitter acetylcholine (ACh), and, according to recent investigations, also promotes aggregation of beta-amyloid peptides, which causes plaque formation in synaptic areas. While the catalytic triad responsible for ACh-cleavage is located 20 Angstrom deep in the enzyme at the end of narrow gorge, the beta-amyloids seem to attach to the peripheral anionic site (PAS) centered around amino acid Trp279 at the mouth of the gorge. We have investigated the potential of galanthamine, and some derivatives of this alkaloid with varying N-substituents, to interfere with both these functions in the AChE-gorge. Our studies result in a correct prediction of the orientation and conformation of galanthamine in the active site of AChE from Torpedo californica (TeAChE) with a RMS deviation of about 0.5 Angstrom with respect to the crystal structure of the TeAChE-galanthamine complex. The docking studies, furthermore, revealed a second binding site for galanthamine which is located at the PAS. Based on these findings, we have performed a molecular modeling study to investigate bis-galanthamine derivatives that have two galanthamine moieties connected by a methylene spacer of varying length as dually acting AChE-ligands. Our results propose that such ligands indeed can simultaneously interfere with both biological functions of the enzyme, the neurotransmitter cleavage and the plaque formation and should therefore serve as the basis for a further development of bis-functional Alzheimer drugs.

HYSTESIS IN BUTYRYLCHELINESTERASE CATALYSIS: EVIDENCE FOR SUBSTRATE-INDUCED CONVERSION OF THE ENZYME FROM LATENT TO OPERATIVE FORM

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Human butyrylcholinesterase (BuChE, EC. 3.1.1.8) was found to reach slowly steady-state velocity with the neutral ester N-methyllycoxyl acetate as substrate. The lag rate decreased with increasing substrate concentration. The kinetic mechanism of this hysteresis process was interpreted in terms of slow conformational change of the enzyme from an inactive to an active form that binds and hydrolyses substrate [1]. Hysteresis by peripheral site mutants (D70G, Y332A, D70GY332A) was similar to that of wild-type enzyme. In addition, hysteresis was shown for hydrolysis of positively charged substrates: benzoylcholine with wild-type; butyrylhicoline with the D70H mutant and different active site mutants, e.g., A328C and G117H/E197Q. This clearly indicated that hysteresis is a general property of BuChE which does not involve the peripheral site. Resolution of the X-ray structure of BuChE provided evidence that the native BuChE is a 'dormant' enzyme [2], and Nicolet et al., this symposium). Therefore, lags in progress curves for substrate hydrolysis can be interpreted as the time needed to convert the whole population of dormant enzyme forms to an operative one, performing catalytic turnover at steady-state velocity. A new mechanistic model for catalysis of BuChE is proposed. [1] P. Masson, M.T. Froment, S. Fort, F. Ribes, N. Bec, C. Balny and L.M. Schoepfer. Butyrylcholinesterase-catalyzed hydrolysis of N-methyllycoxyl acetate: analysis of volume changes upon reaction and hysteresis behavior, Biochim. Biophys. Acta (2002) in press. [2] F. Nachon, Y. Nicolet, N. Viguier, P. Masson, J.C. Fontecilla-Camps and O. Lockridge. Engineering of a monomeric and low-glycosylated form of human butyrylcholinesterase: expression, purification, characterization and crystallization, Eur. J. Biochem. (2002) 269, 630-637.
OXACHEIN, A NOVEL POTENT INHIBITOR OF ACETYLCHOLINESTERASE FROM A PLANT—OXALIS CORNICULATA L.

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Anticholinesterases (antiChE) have been used as medicines, pesticides and agents of chemical warfare. A variety of natural and synthetic antiChE are known. However, the search is on for new and more effective antiChE. We report a novel and potent antiChE compound from roots of a plant, Oxalis corniculata L. and name it as oxachein (Oxalis ChE inhibitor). Oxachein has dull white needle shaped crystals having melting point 145-145.2°C. Based on data obtained by employing NMR, MS, X-ray diffraction and other analytical techniques, oxachein is 2-(3,5-dimethoxyphenyl)-5-(3,5-dihydroxyphenyl)-6-(4-hydroxyphenyl)-5,6-dihydrofuro-(3,2-f)benzofuran (MW = 496). To the best of our knowledge, this compound has neither been synthesized nor reported from any natural source. Oxachein is a very potent inhibitor of AChE (EC 3.1.1.7) from electric eel (50 percent inhibition at 0.8 microM, 100 percent inhibition at 5 microM). However, it is a weak inhibitor of CH from roots of a plant Cicor aristatum L. (chickpea) causing only 23 percent inhibition at a concentration of 10 microM. Lineeweaver-Burk plot study shows non-competitive mode of inhibition of CH AChE. Unlike some other antiChE from plants, oxachein is neither an alkaloid nor a terpenoid. It may serve as a prototype for novel antiChE in drug development programmes.

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A CALCIUM-PROTON ANTIPORT IN CHOLINERGIC AND GLUTAMATERIC SYNAPTIC VESICLES

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Neurotransmission in rapid synapses implies reactions where 'time is gained at the expense of sensitivity' (Katz, 1988). Indeed, bursting of nicotinic or glutamate AMPA receptors typically occurs in response to a brief exposure to relatively high transmitter concentrations; desensitization occurs if exposure to the ligand is prolonged. Similarly, quantal transmitter release, supported by a plasmalemma protein lipid (mediatorphore), is activated by a brief high local increase of Ca2+ concentration; and also desensitises if Ca2+ is not rapidly removed. Calcium has long been known to be transported into synaptic vesicles and reticulum by a Ca2+-ATPase, a high affinity system working rather slowly. In addition, a H+/Ca2+ antiport has recently been demonstrated to operate in synaptic vesicles of CNS cortex. We are investigating the physiological role of the H+/Ca2+ antiport in the mammalian brain and Torpedo electric organ by using either intact synapses, or synaptosomes or isolated vesicles. Our and other data support the following sequence of events in presynaptic nerve terminals submitted to intense stimulation 1) Ca2+ concentration abruptly rises in limited microdomains under the presynaptic membrane, activating mediatorphores and triggering release; 2) The local Ca2+ concentration drops rapidly in a first phase (H+/Ca2+ antiport) and then more slowly (Na+/Ca2+ antiport, Ca2+ ATPase, and other buffering processes); 3) Ca2+ accumulation in synaptic vesicles causes neurotransmitter leakage from the organelle, fuelling the local pool available for release; 4) Accumulated Ca2+ is expelled to synaptic cleft by exocytosis.

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MECHANISM OF ACETYLCHOLINESTERASE INHIBITION BY FASCICULIN: A 5 NANOSECOND MOLECULAR DYNAMICS SIMULATION

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In our previous molecular dynamics simulation of mouse acetylcholinesterase (EC 3.1.1.7), the enzyme revealed complex gorge fluctuation. Now we report another 5 ns simulation, with acetylcholinesterase complexed with fasciculin 2. Fasciculin 2 binds to the gorge entrance of acetylcholinesterase with excellent complementarity and many polar interactions. In this simulation the protein-protein complex, we continue to observe a two-peaked probability distribution of the gorge width, though fasciculin 2 appears to block access of ligands to the gorge. The gorge width distribution is altered when fasciculin is present: the gorge is more likely to be narrow. Though the gorge is sterically blocked by fasciculin 2, there are large increases in the opening of alternative passages, namely the side door and the back door. The catalytic triad arrangement in the acetylcholinesterase active site is disrupted with fasciculin bound. These data suggest that, in addition to the steric obstruction, fasciculin may inhibit acetylcholinesterase by allosteric and dynamical means. Additional data from these simulations can be found at http://mccammon.ucsd.edu/.

PROF. RENE COUTEAUX AND HIS PRESYNAPTIC "ACTIVE ZONE"

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Couteaux had a notion of the active zone around 1961. He wrote his idea only as a personal communication to J. Eccles in 1961. Eccles in "The Physiology of Synapses" (1964) cited the term of Couteaux erroneously from one of his publications in 1961. To my sticky question for the historical background, Couteaux wrote his answer on a piece of paper in 1997 (Fig. 1). Indeed, the active zone was used in public for the first time in an article of Couteaux R. and Pécor-Dechavanne M. (CR Acad Sci Paris 1970, 271, 2346). We read there "What we call hypothetically the active zones". Couteaux had not claimed paternity of the term. He knew that similar ideas existed already, though electron microscopic proof was not available. Couteaux needed absolutely an image of the exocytosis of synaptic vesicles from a specialized dense part of the presynaptic membrane of the motor nerve terminal. For this purpose, he spent almost ten years.
MITOGEN-ACTIVATED PROTEIN KINASE KINASE INHIBITS CILIARY NEUROTROPIC FACTOR-ACTIVATED CHOLINE ACETYLTRANSFERASE GENE EXPRESSION

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The expression of choline acetyltransferase (ChAT), which synthesizes acetylcholine (ACh) is upregulated by ciliary neurotrophic factor (CNTF). We investigated the role of the mitogen-activated protein kinase (MAPK) pathway in regulating CNTF-stimulated ChAT expression. Using the murine septal cell line SN56T1, we found that PD98059 and U0126, two structurally distinct inhibitors of MAPK kinase (MEK1), increased both basal and CNTF-stimulated ACh production. Transient transfections with ChAT promoter-luciferase reporter construct demonstrated synergy between PD98059 and CNTF at the transcriptional level. Moreover, in co-transfection studies, overexpression of constitutively active MEK1 completely inhibited the CNTF-mediated induction of the cholinergic promoter. To elucidate the mechanism of this inhibition, we examined the signaling pathways evoked by CNTF in SN56T1 cells. Tyrophostin AG-490, a Jak2 inhibitor, abrogated both the activation of the transcription factor Stat3 (measured as Tyr705 phosphorylation) and the increase in ACh production stimulated by CNTF, indicating that Stat3 is a mediator of the CNTF effect. PD98059 did not significantly altered CNTF-induced Tyr705 phosphorylation of Stat3, but it inhibited Ser277 phosphorylation, demonstrating that the latter is MEK1-dependent. The results indicate that MEK1 inhibits the CNTF-mediated stimulation of ChAT expression, possibly as a part of a feedback mechanism.

PLANT CHOLINESTERASE ACTIVITY AS A BIOSENSOR FOR TOXINS IN THE ENVIRONMENT

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Plant cholinesterases are sensitive to derivatives of carboxylic acid, neostigmine and physostigmine and organophosphorus compounds, disopropyl phosphofluoridate, dichlorvos, phosphon and quaternary ammonium compounds as many other artificial and natural toxins (Roschchina, 2001). Cholinesterase activity of pollen Humulus lupulus, Tulipa sp., Salix caprea, Allium cepa and Hippopastrum hybridum was studied with Ellman reagent and its new reagent azoanil as a biotest on the pesticides - dichlorvos, rigor and some synthesized organophosphates GD-42 and GD-7, carbamates neostigmine and its derivatives and plant toxins physostigmine, allicin, atropine, solanine (Roschchina, 2001). There are differences between the pollen sensitivity to physostigmine and neostigmine for certain species, for instance, more lower to first inhibitor for Hippopastrum and Salix pollen. If to compare 1μ for carbamate inhibitors between cholinesterases from plant and animal tissues, pollen cholinesterases seem to be most sensitive among the plant enzymes. Organophosphates demonstrate lower specificity to cholinesterases than carboxylic inhibitors. Cholinesterase of author of Hippopastrum is sensitive to only organophosphate GD-42, but not to GD-7 or isop-OMPA. Natural toxins, except atropine, inhibited the plant cholinesterase with the same degree as carbamates. Pollen cholinesterase can be used as biosensor on the toxins of the environment.


EFFECTS OF METHOMYL ON SPLEENS AND APOPTOSIS

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Methomyl, a carbamate insecticide widely used for crop protection, is known as an acetylcholinesterase inhibitor. Its reported toxic effects are mostly cholinergic effects. However, some toxic effects reported from methomyl may not be related to the anticholinesterase action. Our studies reported that 24 hour single dose exposure of 8 mg/kg methomyl po. caused a significant decrease in rat splenocyte viability, induced oxidative stress, and showed the conformational change of proteins at amide-I and II regions in rat spleen cells. However, effect of methomyl on protein conformation in spleens could not be blocked by anticholinergic agent, atropine. Therefore, toxicity of methomyl could also be generated by mechanisms other than acting through acetylcholinesterase and acetylcholine. Effects of methomyl on leukocyte cells have been studied. The results showed that 6 hour exposure of methomyl could reduce mitochondrial transmembrane potential and induce caspase-dependent apoptosis in MM6, THP-1, and Jurkat cell lines at final concentrations of 18, 18, and 6 mM, respectively. Its action on IL-6 signaling pathway was found to play roles. Methomyl did not induce apoptosis in Raji cells, but caused a cell cycle arrest in the G0/G1 phase. The ability of methomyl to induce protein conformational change in spleens and apoptosis in the leukocytic cell lines is probably generated from its action as a sulfhydryl binding agent, which can interfere with intracellular signals. The study of effects of methomyl on signal transduction alterations will provide more information of mechanisms of methomyl toxicity.

EFFECTS OF CARBAMATE INSECTICIDES ON RAT NEURONAL α4β4 NICOTINIC RECEPTORS AND RAT BRAIN ACETYLCHOLINESTERASE

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Carbamate insecticides are generally known to inhibit the enzyme acetylcholinesterase (AChE). In addition, some carbamates, e.g., physostigmine, modulate neuronal nicotinic ACh receptor (nAChR) function by potentiating the ACh-induced ion current. The aim of this study was to investigate the effects of carbamate insecticides on nAChRs and to compare these effects with those on AChE. Rat α4β4 nAChRs, expressed in Xenopus laevis oocytes, were exposed to different concentrations of insecticides. ACh-evoked currents were measured using a two-microelectrode voltage clamp. Unlike physostigmine, all 6 carbamates tested inhibited 1 mM or 1 μM ACh-induced ion currents. Potencies ranged from 5 μM for fenoxycarb to 1.2 mM for aldicarb. Inhibition of rat brain AChE activity was determined spectrophotometrically for all compounds. Fenoxycarb and EPTC caused no inhibition, whereas bendiocarb and propoxur proved to be the more potent inhibitors of rat brain AChE with potencies of 0.75 μM and 2.4 μM, respectively. The results demonstrate that the potentiating activity of physostigmine does not solely depend on the presence of a carbamate moiety. There is no correlation observed between inhibition of the rat α4β4 nAChR and inhibition of rat brain AChE. Some carbamates are more potent inhibitors of the rat α4β4 neuronal nAChR than of rat brain AChE, suggesting that neuronal nAChRs are an additional target for carbamate insecticide neurotoxicity.
SITE-SPECIFIC ANALYSIS OF GLYCAN STRUCTURES ON PLASMA-DERIVED HUMAN (Hu) AND HORSE (Eq) BUTYRYLCHOLINESTERASES (BChE)

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The successful use of plasma-derived cholinesterases as pretreatment drugs for organophosphate toxicity, stems from their long mean retention times. The relatively high stability of these enzymes in circulation may be attributed to the number and structures of their carbohydrate residues. Therefore, the site-specific carbohydrate (CHO) structures of several soluble BChEs were determined. Purified proteins were fragmented by enzymatic or chemical means. Glycopeptides were purified by reverse-phase HPLC and identified by N-terminal sequencing. Asparagine-linked glycans were released by treatment with PNGase F. The free reducing ends were labeled with N-aminophenyl-1,3,6-triazinol (Glyko, Inc.), and subjected to fluorosphere-assisted carbohydrate electrophoresis analysis following sequential digestion with various glycosidases. Site-specific CHO structures were determined for 8/9 Hu and 3/8 Eq BChEs. For Hu BChE, the major structure consisted of an asialo, galactosylated biantennary oligosaccharide without core fucosylation. Hu BChE site 8 (aaf481) and 9 (aaf486) were inseparable with the cleavage strategies employed, which precluded an absolute assignment of CHO for these sites. Glycans of Eq BChE were similar except that the sites displayed less microheterogeneity and the majority of structures were sialated. While all the sites for Hu BChE displayed some degree of microheterogeneity, some sites were more heterogeneous; for example site 4 (aaf 241) had 25% oligo-mannose structures, while sites 5 (aaf 256) and 7 (aaf 455) consisted of 11% and 12% tri-galactose, tri-antennary structures, respectively. The notable lack of siaylation of Hu BChE CHO's was probably due to the prolonged storage of the glycopeptides. CHO structures for fewer Eq BChE sites were determined, but a direct comparison (Hu site 3 and Eq site 2) indicated a high degree of homology between them.

MECHANISM AND STRUCTURAL REQUIREMENTS OF XANOMELINE WASH-RESISTANT BINDING TO M1 MUSCARINIC RECEPTORS

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Xanomeline is a novel agonist functionally selective for muscarinic M1 receptors. It is unusual in that it binds to receptors both in a reversible manner (with micromolar affinity) and in a wash-resistant, quasi-irreversible manner (with micromolar affinity). We investigated the two modes of its binding on membranes from CHO cells expressing M1 muscarinic receptors, with the following conclusions:

(1) Reversible binding of xanomeline occurs at the classical binding site and is competitive with that of classical muscarinic ligands. (2) Stable binding (half-life of >24 h) is to an "exosite", distinct from the classical site. (3) Stable binding to the exosite is preceded by initial high-affinity binding, but an interaction of xanomeline with the classical binding site is not a prerequisite for its binding to the exosite. (4) Comparison of xanomeline with its analogues indicates that the wash-resistant binding of xanomeline depends on the presence of the O-hexyl arm in its molecule and that an at least five-carbon long O-alkyl chain is required for wash-resistant binding.

The length of the O-alkyl chain required and thermodynamic data suggest that the wash-resistant binding depends on the penetration of the O-alkyl chain of xanomeline between individual alpha-helices of the binding pocket and its direct interaction with lipids surrounding the receptor.

RECOVERY FROM DESENSITIZATION OF A NEURONAL NICOTINIC RECEPTOR

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We have shown (J. Pharm. Exp. Ther. 289:656,1999) that skeletal muscle nicotinic acetylcholine receptors (nACHR) recover from desensitization depending on concentration and duration of exposure to agonist and its identity. Recovery after ACh exposure was significantly slower than that induced by nicotine (Nic) for all concentrations. Further, the muscle type nACHR displays a 'molecular memory' probably resulting from the conformational state that the receptor undergoes in response to agonist-specific desensitization.

Using whole cell patch voltage-clamping (EPC9) combined with rapid agonist delivery (Warner Instruments, Fast-Step), we examined responses to ACh and Nic in SHSY-5Y cells expressing the ganglionic, neuronal nACHR (nAChR) (α3 β4; α5 β2). Our data indicate that recovery from desensitization induced with ACH or Nic proceeds in a double exponential fashion except for the shortest durations of exposure we used (1 s or less) in which recovery kinetics were best fit with a single exponential. In all cases, recovery from desensitization proceeded considerably faster when ACH induced desensitization. Further, when ACH was used to desensitize and Nic to measure recovery, the kinetics of recovery were similar to those found when Nic alone was used to measure responses. Similarly, when Nic was used to desensitize and ACH to monitor recovery, kinetics were similar to those whose cells were exposed to ACH alone.

We conclude that the neuronal receptor examined here behaves differently from the muscle type with respect to desensitization and recovery and that this difference is consistent with the known addictive effects of nicotine on neuronal cholinergic receptors.

IS THE G-PROTEIN-COUPLLED M2 MUSCARINIC RECEPTOR A VOLTAGE SENSOR?

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G-protein coupled receptors (GPCRs) are not considered to exhibit voltage sensitivity. Here, using Xenopus oocytes, we examined whether a classical GPCR, the muscarinic M2-receptor (m2R), is by itself a voltage sensor. Oocytes expressing m2R were voltage-clamped at -60 mV or at +40 mV, and the relationship between the G-protein-gated inwardly rectifying K+ channels (GI-RK) response and acetylcholine (ACH), (a positively charged agonist) or oxotremorine (OXO), (an uncharged agonist), concentration was established at these two holding potentials. We found, for both agonists, that at +40 mV there is a shift to the right of the dose response curve. Direct binding experiments of [3H]-ACH to individual oocytes expressing m2R showed that the specific binding of [3H]-ACH was reduced by depolarization. These results suggest that the m2R senses changes in membrane potential.
SOME BASIC RULES GOVERNING OLGOSACCHARIDE-DEPENDENT CIRCULATORY RESIDENCE OF GLYCOPROTEINS ARE REVEALED BY MALDI-TOF MAPPING OF THE MULTIPLE N-GLYCANS ASSOCIATED WITH RECOMBINANT BOVINE ACETYLCHOLINESTERASE

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Pharmacokinetic studies of recombinant bovine acetylcholinesterase (rBoAChE), revealed that this enzyme is cleared more rapidly than the native serum-derived FBS-AChE. Extensive MALDI-TOF analysis of sialylated and desialylated glycans purified from rBoAChE, revealed that these are comprised of a complex array of diverse structures differing in branching, monosaccharide substitutions and relative abundances. The exact structures of the different glycans were confirmed by a series of exoglycosidase treatments followed by MALDI-TOF analysis. The most prevalent structure was the biantennary fucosylated form, (Man3-GlcNAcβ2-3Galβ1-4Galβ1-4Glcβ1-2), which constitutes approximately 40-50 percent of the total glycans. 20-30% of the glycans were of the triantennary form, while tetraantennary glycans were present at very low levels. Most importantly, the glycans of rBoAChE were found to be heavily undersialylated, containing -4.5 terminally exposed β-gal per enzyme subunit.

To allow efficient sialylation, rBoAChE was produced in an engineered HEK-293 cell line clone which expresses high levels of recombinant sialyltransferase. MALDI-TOF analysis of the glycans of rBoAChE produced in these cells demonstrated that the vast majority of these glycan forms were now highly sialylated. Pharmacokinetic studies of highly-sialylated rBoAChE established that this enzyme was retained in the circulation for extended periods of time, as compared to undersialylated rBoAChE. These studies, emphasize the pivotal role of glycan sialylation in determining the circulatory fate of cholinesterases, and provide the basis for detailed determination of their glycan structures.

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EFFECT OF POST-TRANSLATION MODIFICATIONS OF HUMAN ACETYLCHOLINESTERASE ON ITS CIRCULATORY RESIDENCE

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Homogenous preparations of HuAChE differing in their oligomerization status were generated: (a) monomers - represented by the oligomerization-impaired CS80A-HuAChE mutant (b) dimeric wild-type (WT) and (c) tetramers of WT-HuAChE generated in vitro by complexation with a synthetic CalQ-derived Proline Rich Attachment Domain (PRAD) peptide. Three different series of each of these three oligomer preparations were produced: 1) partially sialylated - derived from HEK-293 cells; 2) fully sialylated – derived from HEK-293 engineered cells expressing high levels of sialyltransferase; 3) desialylated – following treatment with sialidase to quantitatively remove sialic acid termini. The oligosaccharides associated with each of the various preparations were extensively analysed by MALDI-TOF. With the enzyme preparations comprising the fully sialylated series, a clear linear relationship between oligomerization and circulatory mean residence time (MRT) was observed. Thus, monomers, dimers and tetramers exhibited MRTs of 110, 195, and 740 min, respectively. As the level of sialylation decreased, this differential behavior became less pronounced, and eventually following desialylation all oligomers had the same MRT (5 min). These observations suggest that multiple removal systems contribute to the elimination of acetylcholinesterase from the circulation. The studies presented here demonstrate also that by combined modulation of sialylation and tetramerization it is possible to generate a HuAChE displaying a circulatory residence exceeding that of all other known forms of native or recombinant human AChE.

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CHANGES IN NEURONAL CHOLINERGIC RECEPTOR BINDING SITES AT DIFFERENT AGES IN TRANSGENIC MICE OVEREXPRESSING HUMAN ACETYLCHOLINESTERASE

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Deficits in the cholinergic circuits of the human brain are observed in neurodegenerative disorders like Alzheimer’s and Parkinson’s disease. An overexpression of neuronal human acetylcholinesterase (AChE) in mice causes alteration in the cholinergic transmission. The objective of this study was to investigate how overexpression of AChE activity influences the plasticity of cholinergic neurons, particularly the nicotinic and muscarinic receptor subtypes. AChE transgenic (Tg+) mice at different ages, from 3 days old up to 1 year, were compared to age-matched non-transgenic (Tg-) mice. The nicotinic receptor binding sites were quantified, in the cortex and the striatum using [3H]Acetylcholine (alpha4beta2) and in the cortex and the hippocampus using [3H]alpha-bungarotoxin (alpha7). In addition, muscarinic receptor binding sites were quantified, in the cortex and the striatum using [3H]Pirenzepine (M1). A significantly increased [3H]Acetylcholine binding was found in the cortex and the striatum in AChE Tg+ mice in comparison to Tg- mice in various age groups. No major alteration in [3H]alpha-bungarotoxin binding sites were observed in Tg+ compared to Tg- mice. However, an up-regulation of [3H]alpha-bungarotoxin binding sites was observed in Tg+ compared to Tg- mice. In the hippocampus, the binding sites were detected at any age. The increase in alpha4beta2 and M2 receptor binding sites observed in AChE Tg+ mice was found at all ages and thus not influenced by aging processes.

TRANSGENIC OVEREXPRESSION OF READTHROUGH ACETYLCHOLINESTERASE (ACHE-R): DISTRIBUTION OF ACHE-R AND CFOS IN BRAIN IN RELATION TO BEHAVIOR

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Acute stress induces a cascade of events, which involves increased release of acetylcholine and feedback activation of acetylcholinesterase gene expression, shifting into the "readthrough" acetylcholinesterase variant ACHE-R, which contributes to downregulating stress-induced cholinergic excitability. To explore whether chronic elevation of ACHE-R affects behavioral and physiological functions, we employ FVB/N transgenic (Tg) mice over-expressing human ACHE-R. The present report correlates behavioral and physiological functions with brain distribution of ACHE-R and of cFOS, an immediate early gene that is responsive to stress. Tg AChE-R mice appeared healthy but their body weight was lower compared to FVB/N mice. Low body weight correlated with appearance of ACHE-R-filled neurons in the lateral hypothalamic area and with increase in c-FOS positive cells in lateral and ventromedial hypothalamus. Compared to FVB/N mice, Tg-AChE-R mice were significantly impaired in learning in a serial choice maze. This was correlated with appearance of ACHE-R-filled neurons and c-FOS positive cells in the hippocampal dentate gyrus. Additional mice, were implanted with transmitters and their motor activity recorded in the home-cage. Increased motor activity in Tg-AChE-R mice correlated with appearance of ACHE-R-filled neurons and c-FOS positive cells in striatum, parietal and retrosplenial cortex. We propose that chronic elevation of ACHE-R alters regulation of neuronal excitability in several brain regions that, in turn, may contribute to the alterations in behavioral and physiological functions under conditions of chronic stress.
STUDY ON THE MECHANISM OF BLOCKADE OF ACETYLCOLINE RELEASE BY SNAKE PRESYNAPTIC PLA2 NEUROTOXINS ON NERVE TERMINALS

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Several animal venoms contain toxins with phospholipase A2 (PLA2) activity. These enzymes hydrolyze the sn-2 ester bond of 1,2-diacyl-sn-phosphoglycerides producing fatty acids and lysophospholipids. Some snake venoms contain presynaptic PLA2 neurotoxins that cause a persistent blockade of neurotransmitter release from nerve terminals. Three subsequent phases can be distinguished at the neuromuscular junction (NMJ) poisoned by PLA2 neurotoxins: a short initial phase with either decreased or unchanged ACh release, is followed by a longer phase of facilitation of ACh release, which then fades into the third phase of complete and irreversible inhibition of neurotransmission. Electron microscopy studies of poisoned NMJ revealed appearance of many clathrin-coated W-shaped plasma membrane invaginations, indicating a blockage of endocytosis. Therefore PLA2 neurotoxins both promote fusion of small synaptic vesicles (SSVs) with the presynaptic membrane and inhibit their retrieval, thus causing release of ACh, depletion of vesicles and enlargement of nerve terminals. On the basis of these data we are investigating whether PLA2 neurotoxins block nerve terminals entering the lumen of synaptic vesicles and hydrolyse phospholipids of the inner leaflet of the membrane. The suggested entry of PLA2 neurotoxins inside SSVs would account for the finding that electrical activity of the nerve terminals actively promotes intoxication. In order to test our hypothesis we are studying the intracellular localization of two snake PLA2 neurotoxins (beta-bungarotoxin and taipoxin) on mouse phrenic nerve-hemidiaphragm NMJ intoxicated with these toxins. If this working hypothesis proves true, the presynaptic PLA2 neurotoxins could be employed as tools to investigate specific aspects of acetylcholine release and of vesicles fusion and recycling.

THE ROLE OF READTHROUGH ACETYLCOLINESTERASE IN THE PATHOPHYSIOLOGY OF MYASTHENIA GRAVIS

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Alternative splicing induces, under cholinergic imbalances, overproduction of the rare "readthrough" acetylcholinesterase variant, ACHE-R. We explored the pathophysiological relevance of this phenomenon in patients with myasthenia gravis (MG) and rats with experimental autoimmune MG (EAMG), both neuromuscular junction diseases with depleted acetylcholinesterase receptors. In MG and EAMG, we detected serum ACHE-R accumulation. In EAMG, we alleviated electromyographic abnormalities by nanomolar doses of EN101, an antisense oligonucleotide that selectively lowers ACHE-R in blood and muscle, yet leaves unaffected the synaptic variant, ACHE-S. While animals treated with placebo or conventional anticholinesterases continued to deteriorate, 4 weeks daily oral EN101 administration improved survival, neuromuscular strength and clinical status in moribund EAMG rats. The efficacy of targeting only one ACHE splicing variant highlights potential advantages of mRNA-targeted therapeutics for chronic cholinergic imbalances.

"READTHROUGH" ACETYLCOLINESTERASE FORMS NEURONAL COMPLEXES WITH PKC BETA II AND ITS WD CARRIER RACK1

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Expression of neuronal "readthrough" acetylcholinesterase (ACHE-R) is robustly increased under psychological stress, when it serves to hydrolyse the stress-elevated acetylcholine. However, the ACHE-R protein also accumulates intracellularly.

Therefore, we used a yeast two hybrid screen to search for intracellular protein partners of ARPI (ACHE readthrough peptide 1), the C-terminal ACHE-R domain that does not participate in acetylcholine hydrolysis. Here, we report that ACHE-R, through ARPI, interacts intracellularly with RACK1, the WD domain PKC beta II carrier protein. Ex vivo, triple ARPI/RACK1/PKC beta II complexes were found in transfected COS cells and ACHE-R/RACK1/PKC beta II complexes in native pheochromocytoma cells. In vivo, physiological stress induced parallel increases in ACHE-R, RACK1 and PKC beta II in parietal cortex and hippocampal CA1 neurons. Moreover, the stress-protected ACHE-R overexpressing transgenic mice display elevated levels of RACK1 and PKC beta II, accompanied by RACK1 translocation to the perikaryal periphery and the appearance of dense co-labeled ACHE-R/PKC beta II punctated neuronal clusters in several stress-responding brain regions. These findings present a non-catalytic intracellular capacity for ACHE-R which may trigger PKC beta II-dependent signalling processes that augment physiological responses to diverse external stimuli.

CHRONIC ACETYLCOLINESTERASE OVEREXPRESSION INDUCES MULTILEVELED ABBERRATIONS IN NEUROMUSCULAR PHYSIOLOGY

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Overexpression of acetylcholinesterase (ACHE) is a notable consequence of exposure to anticholinesterase drugs and/or poisons. However, the physiological consequences of such chronic overexpression with regards to neuromuscular function were not yet carefully analyzed. Here, we report detailed dissection of the different components of neuromuscular transmission in transgenic mice overexpressing the neuronal "synaptic" variant of ACHE, which were previously reported to develop altered muscle morphology (Andres et al., 1997). Transgenic diaphragms presented pronounced fatigue, which reflected both neurotransmission fading and muscle mechanical aberrations. Under tetanic stimulation protocol, transgenic (TG) muscles fatigued to a larger extent than wild type (WT) muscles, either when stimulated directly (40% vs. 25% decay from initial force, within 10 contractions of 10 s intertrain interval, respectively) or via the phrenic nerve (68% vs. 49%, respectively), probably due to an impaired recovery process. ACHE overexpression further affected synaptic transmission with significantly higher quantal content (0.095 in TG vs. 0.068 in WT, P < 0.05, at 0.2 mM Ca2+ and 2.4 mM Mg2+). Furthermore, adjusted quantal size was exposed by treatment with physostigmine (10 μM) revealing higher amplitude (2.08 mV in TG vs. 1.54 mV in WT, P < 0.01) and half decay time (2.87 ms in TG vs. 2.29 ms in WT, P < 0.02) in the TG neuromuscular junction. Our observations predict multileveled neuromuscular malfunctioning under disrupted cholinergic homeostasis, which is relevant for chronic anticholinesterase exposure.
EXPRESSION OF THE CHOLINERGIC GENE LOCUS IN THE TRACHEAL EPITHELIUM OF THE RAT

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Cholineacyltransferase (ChAT) catalyzes the biosynthesis of acetylcholine (ACH) from acetyl-CoA and choline in the axoplasm. The vesicular acetylcholine transporter (VACHT) is responsible for the translocation of ACh into the interior of synaptic vesicles. The gene coding for ChAT contains the entire intronless sequence for VACHT within its first intron. Northern blot analyses have demonstrated mRNAs of different sizes in the central and peripheral nervous system of the rat. A ChAT mRNA (cChAT) of a single size (about 4 kb) was detected in brain and spinal cord. In peripheral tissues mRNAs of different sizes and a peripheral ChAT (pChAT) which seems to be a splice variant of cChAT could be detected. The occurrence of ChAT in non-neuronal tissue is known, but still unknown is the molecular identity of ChAT in the respiratory epithelium. Thus, we amplified the whole coding region of the rat tracheal epithelial ChAT mRNA. By RT-PCR a 2071 bp fragment could be detected, which shares a 99% sequence identity with cChAT. The expression of VACHT could be demonstrated for the first time in rat tracheal epithelium by RT-PCR. Amplification of the whole coding region revealed a sequence identity of about 99% to the VACHT known from cholinergic neurons. In the present study we demonstrate the simultaneous expression of ChAT and VACHT in rat tracheal epithelium. The obtained sequencing data strongly suggest that the ChAT and VACHT proteins of rat tracheal epithelium are identical to that known from central cholinergic neurons. (SFB 547, project C2)

MUSCARINIC RECEPTORS AND TRP-CHANNELS IN PRIMARY SENSORY NEURONS OF THE RAT

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In sensory neurons acetylcholine interacts with G-protein coupled muscarinic receptors (MR), thereby activating a variety of second messenger systems (IP3, DAG, PLC, PKC). TRP-channels are a family of Ca2+-permeable cation channels that are activated by depletion of intracellular Ca2+-stores or subsequent to the stimulation of PLC isomers. In in vitro expression systems, MR are coupled via Gq/11 proteins to transient receptor potential (TRP) channels. Stimulation is followed by an increase in [Ca2+]. Similarly, MR activation leads to an increase in [Ca2+] in sensory dorsal root ganglion (DRG) neurons. Therefore, we investigated the presence and localisation of TRP-channels in DRG neurons at transcriptional (RT-PCR), translational (immunohistochemistry) and functional (Ca2+ imaging) level. Total RNA of lumbar DRG contained mRNA for five out of seven channel subtypes. Proteins for the TRP-1, 3- and 6-channel could be demonstrated in subpopulations of sensory neurons. Perikarya that expressed the marker of presumably nociceptive neurons, the vanilloid receptor-1, were also immunoreactive for the TRP-1, 3- and 6-channels. Activation of Gq/11-coupled MR using muscarine was followed by an increase in [Ca2+]. The rise in [Ca2+] was abolished in presence of Nickel/Cadmium or after depletion of intracellular Ca2+-stores by application of thapsigargin. This further suggests the involvement of TRP-channels in the MR-mediated Ca2+-signalling. (SFB 547, Teilprojekt C2, SFB 535, Teilprojekt A10)

DIVERSE MOLECULAR MECHANISMS UNDERLYING CONGENITAL MYASTHENIC SYNDROMES

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Mutations of muscle acetylcholine receptors cause defective neuromuscular transmission through a variety of molecular mechanisms. AChR deficiency is the most common of the congenital myasthenic syndromes (CMS). However, it is not always clear how mutations within the AChR epsilon-subunit coding regions affect mRNA and protein levels at the endplate. We found different homozygous mutations, located in the M3-M4 cytoplasmic loop, in four patients with typical AChR deficiency. Surface expression in HEK293 cells showed that each is a null mutation. Surprisingly, in situ hybridisation in biopsies showed normal expression of epsilon-subunit mRNA. Thus the mutant epsilon-subunit mRNA transcripts are neither up-regulated nor preferentially degraded. Moreover, there was no compensatory increase in gamma-subunit mRNA suggesting that the normal low level of the gamma-subunit in human muscle is sufficient for survival of patients with epsilon-subunit null alleles. Other CMS are due to abnormal ion channel function. In one case, abnormal fetal development had resulted in multiple joint contractures (arthrogryposis) of the fingers at birth, as well as CMS presenting in infancy. Mutational screening revealed heteroallelic mutation within the AChR delta-subunit, d535ins2 and d539K. d536ins2 is a null mutation, but both adult and fetal AChRs containing d539K show abnormally short burst lengths, predicting a “fast channel” phenotype. Thus d539K causes dysfunction of fetal as well as adult AChRs explaining the presence of joint contractures which result from reduced fetal movement. This is the first description of the association of AChR gene mutations and arthrogryposis, but any mutation that disrupts fetal AChR function could underlie additional cases.

ACETYLCHOLINESTERASE KNOCKOUT MICE HAVE INCREASED SENSITIVITY TO SCOPOLAMINE AND ATROPINE

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It is generally accepted that continued stimulation of cells with agonists results in a state of desensitization or downregulation. The purpose of this work was to test the hypothesis that AChE-/- mice have reduced levels of functional muscarinic receptors. The toxicity of the antimuscarinic drugs, scopolamine and atropine was tested in acetylcholinesterase wild-type (AChE+/-), heterozygous (AChE+/-) and nullizygous mice (AChE-/-). Mice were injected i.p. with scopolamine or atropine dissolved in saline. In the case of a lethal dose, mice died within 2–15 minutes. The LD50 for scopolamine was 100 mg/kg in AChE+/-, 180 mg/kg in AChE+/-, and 5 mg/kg in AChE-/- mice. The LD50 for atropine was 250 mg/kg in AChE+/-, 215 mg/kg in AChE+/-, and 96 mg/kg in AChE-/- mice. The higher sensitivity of AChE-/- mice to muscarinic antagonists is consistent with the interpretation that these mice have fewer muscarinic receptors. These results correlate with data obtained in our laboratory (Bin Li et al.) demonstrating that AChE-/- mice are less sensitive to the muscarinic agonists, oxotremorine and pilocarpine. Taken together, these results support the conclusion that AChE-/- mice have adapted to excess acetylcholine by downregulating muscarinic receptors.

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DOWNREGULATION OF MUSCARINIC RECEPTORS IN MICE DEFICIENT IN ACETYLCOLINERESTASE

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This study examined how acetylcholinesterase (ACHE) knockout mice have adapted to the absence of ACHE. ACHE hydrolyzes acetylcholine to terminate cholinergic neurotransmission. Overstimulation of cholinergic receptors by excess acetylcholine is lethal. However, ACHE-/- mice live to adulthood. The hypothesis was tested that adaptation occurred through downregulation of cholinergic receptors. Muscarinic receptors were investigated by treating mice with muscarinic receptor agonists, pilocarpine and oxotremorine (OXO). In response to 200 mg/kg pilocarpine i.p., 10/10 ACHE+/+, 5/10 ACHE+/–, and 0/6 ACHE-/- mice had seizures. Results indicated that ACHE-/- mice have reduced numbers of functional muscarinic receptors. A second group of mice was treated with 1 mg/kg OXO s.c. OXO specifically stimulates muscarinic receptors inducing tremor, hypothermia, and salivation. 6/6 ACHE+/+, 6/6 ACHE+/–, and 0/6 ACHE-/- mice had severe tremor, a drop in surface body temperature of 12

TARGETING OF THE HUMAN VESICULAR ACETYLCOLINERESTASE TRANSPORTER TO CHOLINERGIC SUBDIVISIONS IN TRANSGENIC MICE

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The vesicular acetylcholine transporter (VACHT) and choline acetyltransferase (ChAT) are encoded within a single regulatory unit, called the cholinergic gene locus (CGL). To identify regions in the CGL that are important for the cell type-specific expression of VACHT in vivo, we tested fragments of the human CGL for their ability to confer correct expression of human VACHT to mouse cholinergic neurons. In our previous work (Neuroscience 96 (2000):707-22) we identified an 8.7 kb fragment from the human CGL that restricted expression of the embedded VACHT to somatodentritic neurons in transgenic mice. In the present study, we report the generation and analysis of two additional human CGL transgenes. The addition of 2.5 kb of downstream sequence to the existing transgene resulted in an extension of the expression of human VACHT to the cholinergic neurons of the medial habenular nucleus. The removal of 4.5 kb from the 5’-end of this construct completely abolished human VACHT expression in mice. Our data provide strong evidence for a mosaic model for CGL regulation in separate subdivisions of the mammalian cholinergic nervous system.

THE DIURNAL ACTIVITY OF ACETYLCOLINERESTASE INHIBITORS

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Objective: Acetylcholinesterase inhibitors (ACHEi), particularly donepezil, can induce sleep disorders in Alzheimer's disease patients. It is believed that acetylcholinesterase (ACHE) displays diurnal variation, and that this circadian phenomenon may be linked to ACHEi-induced sleep disturbances. We examined the potential effects of ACHEi on the diurnal activity of ACHE. Methods: A literature search was conducted to identify studies investigating the sleep-activity cycle of patients taking ACHEi. Results: Published evidence suggests that ACHE activity follows a circadian pattern. Drugs such as donepezil, which exhibit highly potent ACHEi activity, seem to interrupt this natural diurnal activity. As donepezil has a long half-life (average, 70 hours), it remains in the body for long periods of time. Thus, even if given in the morning, donepezil concentrations can still remain high enough to create daytime acetylcholine levels at night, interrupting sleep. Donepezil may override the circadian rhythm of the cholinergic system, disturbing the sleep-activity cycle and resulting in high incidences of insomnia and other sleep-related events, and raising hypnic medicative use. Galantamine (Reminyl(R)) has a half-life of 6-8 hours, is a less potent ACHE than donepezil, and allosterically modulates nicotinic acetylcholine receptors. Unlike donepezil, galantamine appears not to perturb the diurnal activity of ACHE. This could explain the lower incidence of sleep disturbances and concomitant hypnic use with galantamine. Conclusion: While donepezil may override the natural diurnal activity of ACHE resulting in increased sleep disturbance, galantamine appears not to perturb this activity, giving a lower incidence of insomnia and sleep-related disorders.

THE MUSCARINIC M1 RECEPTOR AS A THERAPEUTIC TARGET FOR COGNITIVE DEFICITS: PRECLINICAL PHARMACOLOGY AND KNOCKOUT MOUSE STUDIES

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Muscarinic M1 receptors are localized to brain regions involved in learning and memory and have been implicated in the regulation of cognitive processes at the behavioral and molecular level. However, the specific muscarinic receptor subtypes involved in cognitive processes have not been definitively identified. With the availability of muscarinic receptor knockout mice, the effects of various receptor subtypes on learning and memory can be examined. In the present studies, spatial learning and memory as well as open field activity were assessed in M1 knockout (KO) mice and age-matched wild type (WT) controls. Significant deficits were found for M1-KO mice compared to controls in task completion time, reference memory and total errors. M1-KO mice had a significantly greater rate of ambulation on Days 1, 3 and 14. Results indicate that M1-KO mice exhibit impaired learning and memory in a spatial discrimination task and that this effect was independent of differences in activity level which were short-lived with extended monitoring. Within the hippocampus, the M1 receptor, but not the functionally similar M3 receptor, stimulated G protein activation and calcium mobilizing signaling pathways. Broader studies with mice bearing a genetic deletion of each of the 5 muscarinic receptor subtypes suggests that muscarinic receptor-mediated activities, such as tremor, salivation, hypothermia, heart rate, and smooth muscle function are regulated predominantly through the M2 and M3 receptors. Taken together, these studies indicate that the M1 receptor is a likely target for the development of therapeutics for the treatment of cognitive deficits.

-47-
Acetylcholinesterase (AChE, EC 3.1.1.7) functions in nerve impulse transmission, and possibly as a cell adhesion factor during neurite outgrowth. These functions predicted that a mouse with zero AChE activity would be unable to live. It was a surprise to find that AChE-/- mice were born alive and survived an average of 14 days. The emaciated appearance of AChE-/- mice suggested an inability to obtain sufficient nutrition and experiments were undertaken to increase caloric intake. Pregnant and lactating dams (+/-) were fed 11% high fat chow supplemented with liquid Ensure. AChE-/- pups were weaned early, on day 15, and fed liquid Ensure. Although nullizygous animals showed slow but steady weight gain with survival over 1 year (average 100 days), they remained small at all ages compared to littermates. They demonstrated delays in temperature regulation (day 22 vs 15), eye opening (day 13 vs 12), righting reflex (day 18 vs 12), descent of testes (week 7 vs 4), and estrus (week 9 vs 6-7). Significant physical findings in adult AChE-/- mice included body tremors, abnormal gait and posture, absent grip strength, inability to eat solid food, pinpoint pupils, decreased pain response, vocalization, and early death caused by seizures or gastrointestinal tract ileus. Behavioral deficits included urination and defecation in the nest, lack of aggression, reduced pain perception, and sexual dysfunction. These findings support the classical role for AChE in nerve impulse conduction and further suggest that AChE is essential for timely physical development and higher brain function.

**ROLE OF MUSCARINIC RECEPTORS IN THE ACTIVATION OF THE SUBICULO-ACUMBENS PROJECTION**
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The nucleus accumens receives limbic inputs from a number of brain regions, including the ventral subiculum. Activation of this projection, following microinjection of N-ethyl-D-aspartate (NMDA) or carbachol increases locomotor activity. Using in vivo microdialysis, ventral subiculum application of NMDA increases levels of dopamine in the nucleus accumens. Experiments were conducted to ascertain, in the nucleus accumens, the neurochemical consequences of carbachol administration using microdialysis, and to explore the cholinergic receptor subtype(s) involved in any evoked response. In anaesthetized rats, ventral subiculum administration of carbachol increased dopamine levels in the nucleus accumens. Administration of nicotine or the alpha-7 nicotinic acetylcholine receptor agonist. AR-R17779 failed to evoke a response. An involvement of muscarinic receptors was suggested from the significant reduction in response to carbachol following co-administration with atropine. Sensitivity of the subiculo-accumens projection to muscarinic cholinergic receptor stimulation was confirmed by a significant increase in nucleus accumens dopamine following ventral subiculum administration of the broad-spectrum muscarinic agonist oxotremorine M. In further studies utilizing subtype selective agonists, xanomeline (M1 and M4 preferring agonist) failed to increase dopamine. However, (5R-[4R]-6-[6-butyrylthio]-1,2,5-thiadiazol-3-yl]-1-aracizyclo[7.2.1]dectoc (BuTAC - M2 and M4 partial agonist, and M1, M3 and M5 antagonist) evoked a significant response. These data show that the subiculo-accumens projection is sensitive to muscarinic receptor stimulation, and suggests an involvement of the M2 receptor subtype.

**BRAIN PENETRATION AND BEHAVIOURAL PROPERTIES OF A POTENT ALPHA 7 NICOTINIC ACETYLCHOLINE RECEPTOR AGONIST IN THE RAT**
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The homomorphic alpha-7 nicotinic receptor is the second most abundant nicotinic receptor in the brain and has been implicated in a number of psychiatric and neurological disorders. We have evaluated PSAB-OFP (R)-(-)+3'-phenylsphin[1-azabicyclo[2.2.2]octane-3,3(3H)-furo[2,3-b]pyridine], Phillips et al., Astra Arcus USA. potency W099/02859 a potent alpha-7 agonist in a battery of behavioural assays in the rat. Initial studies confirmed that after systemic administration (5 mg/kg s.c.) the Cmax in rat brain was 5.8 microM with a T1/2 of 1.3 hr. We then went on to evaluate the effects in the rat of PSAB-OFP in locomotor activity, pre-pulse inhibition and on performance in the radial and Morris mazes. In addition the compound was examined against hyperalgesia induced by formalin and carrageenan and in 6-hydroxydopamine lesioned rats. Results indicated that PSAB-OFP decreased spontaneous locomotor activity 20-40 min after injection (20 % decrease at 10 mg/kg), but failed to alter stimulant-induced activity, pre-pulse inhibition or cognitive performance in either the Morris water maze or 8-arm radial maze. The compound was also inactive in hyperalgesia models and had no functional or neuroprotective actions in the 6-OHDA model. PSAB-OFP was thus inactive in a wide range of behavioural assays. It is not clear whether this reflects a relatively unimportant role for the alpha-7 receptor, insufficient receptor exposure to the compound and/or rapid receptor desensitisation. The 5-HT3 agonist cross-reactivity of this molecule also makes it difficult to make concrete conclusions on the role of alpha-7 receptors in these in vivo tests.

**FINE-TUNING MODULATION OF NEURONAL MUSCARINIC M1 (FACILITATORY) AND M2 (INHIBITORY) RECEPTORS ACTIVATION BY ADENOSINE AT THE RAT NEUROMUSCULAR JUNCTION**
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The crossstalk between adenosine and muscarinic autoreceptors regulating evoked [3H]-acetylcholine ([3H]-ACH) release was investigated on rat phrenic nerve-hemidiaphragm. Motor nerve terminals possess facilitatory M1 and inhibitory M2 autoreceptors. Dicyclomine (3 nM-10 microM) caused a biphasic (inhibitory/facilitatory) effect, indicating that M1-facilitation prevails during 5 Hz-trains. Co-activation of M2 receptors was partially attenuated, since pirenzepine (1 nM, an M1-antagonist) significantly enhanced inhibition by oxotremorine (10 microM), CGS 21680C (2 nM), an A2A-adenosine receptor agonist, (1) potentiated oxotremorine (10 microM) inhibition, and (2) shifted Me-Ca-A-343 (3 microM)facilitation into a small inhibitory effect. Conversely, the A1-receptor agonist, R-PIA (100 nM), reduced the inhibitory effect of oxotremorine (10 microM), without changing facilitation by Me-Ca-A-343 (3 microM). Synergism between A2A- and M2-receptors is regulated by a reciprocal interaction with M1 receptors that can be prevented by pirenzepine (1 nM). During 50 Hz-bursts, facilitation (M1) of [3H]-ACH release by Me-Ca-A-343 (3 microM) disappeared, while M2 inhibition became predominant. This muscarinic shift results from the interplay with A2A-receptors, because it was prevented by the A2A antagonist, ZM 241385 (10 nM). Thus, when muscarinic M1 facilitation is fully operative, inhibition of ACh release is mediated by adenosine A1 receptors. During high frequency bursts, tonic activation of A2A receptors promotes M2-autoinhibition by braking M1 receptor counteraction.

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IDENTIFICATION OF SIGNALING PROTEINS DOWNSTREAM OF THE TYROSINE KINASE MUSK IN CLUSTERING OF ACETYLCHOLINE RECEPTORS

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During development of the neuromuscular junction, nerve-released agrin causes clustering of acetylcholine receptor complex (ACHRs) in the muscle membrane. The muscle specific kinase (MuSK) is part of the agrin receptor complex and undergoes agrin-induced dimerization and autophosphorylation, thereby starting a signaling pathway that eventually drives AChR phosphorylation and clustering. We observed previously that treatment of C2C12 myotubes with the kinase inhibitor staurosporine prevents aggregation of AChRs but leaves MuSK active, implying the activity of other kinases downstream of MuSK. To analyse possible interaction partners and/or substrates of MuSK, we immunoprecipitated MuSK and identified associated phosphorylated proteins by phosphotyrosine immunoblotting. One major band running at about 60 kDa was specifically coprecipitated by MuSK and phosphorylated on tyrosine in response to agrin treatment. As Src-family kinases bind to the AChR and are activated by agrin, we investigated the possibility that these kinases associate with MuSK and are phosphorylated due to agrin. In myotubes derived from mice lacking both Src and Fyn, no change in the phosphorylation extent of the MuSK-associated protein was observed. Staurosporine and the specific Src-family kinase inhibitors PP1 and C28PT7615 did not affect the phosphorylation degree of the protein. Instead, it was reduced by treatment with herbimycin, in parallel with the reduction of MuSK phosphorylation. Finally, in myotubes lacking rapsyn, phosphorylation of the MuSK-associated protein was still observed. Thus, so far, the identification of the MuSK-associated protein as a member of the Src-family could not be confirmed, and we presently consider other muscle proteins of 60 kDa.

CHEMICAL MODIFICATION OF RECOMBINANT HUMAN ACETYLCHOLINESTERASE BY POLYETHYLENE GLYCOL GENERATES AN ENZYME WITH EXCEPTIONAL CIRCULATORY LONGEVITY

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One of the major obstacles to the fulfillment of the therapeutic potential of recombinant human acetylcholinesterase (rhHuAChE) as a bioscavenger of organophosphates is its short circulatory residence. Post-translation-related factors such as sialylation level and subunit assembly were recently shown to determine the circulatory fate of AChE and demonstrated the ability to generate recombinant AChE with improved pharmacokinetic traits. Here we show that the pharmacokinetic performance of rhHuAChE can be increased significantly by the controlled attachment of polyethylene glycol (PEG) sidechains to lysine residues. The increase in mean residence time (MRT) of the PEG-modified monomeric enzyme is linearly dependent, in the tested range, on the number of attached PEG molecules as well as on their size. It appears that even low level PEG-conjugation can overcome the deleterious effect of sub-optimal post-translation modifications such as under-sialylation. Attachment of as many as four PEG molecule to monomeric rhHuAChE had minimal effect if any on either the catalytic activity or on the reactivity of the modified enzymes towards active center inhibitors such as edrophonium and diisopropylfluorophosphate (DFP) or to peripheral site ligands such as propidium. BW284C51 and even towards the bulky snake-venom toxin - fasciculin-II. At the highest tested ratio of attached PEG-20,000 to rhHuAChE (4:1), an MRT of over 2100 minutes was attained (compared to MRT of 42 minutes for the non-modified enzyme), a value unmatched by any other known form of recombinant or native plasma derived AChE reported to date. This provides an important step toward the generation of a pharmaceutically efficient recombinant AChE-based bioscavenger for prophylactic treatment of organophosphate poisoning.

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DRAMATIC DEPLETION OF CELL SURFACE ACETYLCHOLINE MUSCARINIC RECEPTORS m2R DUE TO LIMITED DELIVERY FROM INTRACYTOPLASMIC STORES IN NEURONS OF ACETYLCHOLINESTERASE (ACHE) DEFICIENT MICE

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The abundance of G protein-coupled receptors (GPCR) located at the neuronal membrane depend on complex intraneuronal trafficking involving delivery of GPCRs from the cytoplasm to the membrane. We have studied here the effect of the chronic AChE deficiency in neurons of AChE gene KO mice on the subcellular distribution of the m2R, using immunohistochemistry at light and electron microscopic levels. 1) In AChE"/" mice in vivo, m2R is abundant at the plasma membrane in striatum, hippocampus and cortex. b) In AChE"/" mice, m2R is almost absent at the membrane but is abundant in endoplasmic reticulum and Golgi complex. 2) Dynamic studies show that the balance between membrane and cytoplasmic m2R depends on the cholinoergic influence: a) In AChE"/" mice, the blockade of muscarinic receptors restores m2R at the membrane. b) In AChE"/" mice in vitro(organotypic culture), when acetylcholine is produced by interneurons (striatum), m2R is located in the cytoplasm as in vivo. The supplementation of AChE"/" neurons with AChE provokes a translocation of m2R from the cytoplasm to the membrane. c) In vitro, when AChE"/" hippocampus is disconnected from its cholinoergic afference, m2R is located at the membrane. When AChE"/" hippocampus is co-cultured with AChE"/"septum, its cholinoergic afference, m2R keeps the cytoplasmic distribution seen in hippocampus in vivo. Our data suggest that the neurochemical environment may contribute to the control of abundance and availability of cell membrane GPCR, and consequently to the control of neuronal sensitivity to neurotransmitters, by regulating their delivery from intracytoplasmic stores to the membrane.

HUPERZINE A AND DONEPEZIL ATTENUATE STAUROSPORINE-INDUCED APOTOPSIS IN RAT CORTICAL NEURONS VIA BCL-2 AND BAX REGULATION AND INHIBITION ON CASPASE-3

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Stauroporine treatment results in apoptotic cell death ad DNA fragmentation. bcl-2, bax and Caspase-3 are known to regulate the apoptotic cell death. This study sought to examine effects of huperzine A and donepezil on stauroporine-induced neuronal apoptosis and potential mechanisms in primary cultured rat cortical neurons. Treated with 0.5 micromolar stauroporine for 24 hours results in significant decrease in cell viability, alteration of neuronal morphology and DNA fragmentation. Pretreatment of the cells with huperzine A and donepezil for 2 hours prior to stauroporine exposure markedly elevated the cell survival at 0.1-10 micromolar concentrations and reduced stauroporine-induced nuclei fragmentation at 1 micromolar concentration. 1 micromolar huperzine A and donepezil pretreatment also reduced the upregulation of pro-apoptotic gene bax, the downregulation of anti-apoptotic gene bcl-2 as well as activation of Caspase-3. Thus our results provide the first evidence that huperzine A and donepezil protect neurons against stauroporine-induced apoptosis via the upregulation of bcl-2 and downregulation of bax and inhibition on Caspase-3 activity.
INSECT GROWTH REGULATORS INHIBIT ACETYLCHOLINESTERASE ACTIVITY IN B-BIOTYPE BEMISIA TABACI IN AUSTRALIA
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B-biotype Bemisia tabaci (Stennorrhyncha: Aleyrodidae) was first detected in Australia in October 1994 and poses a threat to many plant industries. This insect is a severe pest worldwide and is resistant to most conventional insecticides. Insect growth regulators such as buprofezin and novaluron, are being investigated for use against B-biotype B. tabaci in Australia, although resistance has already been detected. In addition to disrupting the nymphal molting process, we have found that buprofezin and novaluron are inhibitors of acetylcholinesterase in insect growth regulator susceptible Australian B-biotype B. tabaci. This is a new mode of action for insect growth regulators. This inhibition did not occur in the buprofezin resistant strain and our results indicate that insect growth regulator resistant B-biotype B. tabaci have evolved an insensitive form of acetylcholinesterase as a resistance mechanism.

VESICULAR ACETYLCHOLINE TRANSPORTER TRAFFICKING AND ADAPTOR PROTEINS
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In neuronal cells, the neurotransmitter acetylcholine is transported from the cytoplasm into synaptic vesicles by the vesicular acetylcholine transporter (VAcT). During biogenesis VAcT is trafficked into the synaptic vesicle. Adaptor proteins have been shown to play a major role in synaptic vesicle recycling. It has been proposed that the cytoplasmic tail of VAcT is involved in its trafficking to the small synaptic vesicle.

In this study we evaluated whether VAcT is sorted into synaptic vesicles through clathrin associated protein complexes. We constructed a GST-fusion VAcT cytoplasmic carboxyl tail and this was incubated with a rat brain extract. Complexes were captured on glutathione beads and analyzed by immunoblotting with Adaptin gamma to detect Adapter Protein-1 (AP1 -1) complexes, Adaptin alpha to detect AP-2 containing complexes, and AP-180 to detect AP-180 containing complexes.

Surprisingly, AP-1, AP-2, and AP-180 proteins were all detected indicating that they all bind to the VAcT cytoplasmic tail. We also tested mutants at the phosphorylation site and at the dileucine motif of VAcT to map the site of interaction with adaptor proteins. Mutant Leu485/Glu486A showed no binding to AP-1, however binding to the AP-2 and AP-180 was retained. Mutants Ser480Ala and Ser480Glu retained binding to the APs but showed different binding intensities indicating that the relative amounts of these complexes changed.

On the basis of these results, we propose that VAcT trafficking involves association with clathrin associated protein complexes, and the cytoplasmic carboxyl tail of VAcT contains multiple trafficking signals.

BIOCHEMICAL CHARACTERISATION OF MICE TRANSGENIC FOR A MUTATION IN AMYLOID PRECURSOR PROTEIN (APP) KNOWN TO CAUSE FAMILIAL ALZHEIMER'S DISEASE
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Alzheimer's disease (AD) is characterised by a deficit in markers of the cholinergic system (Francis et al., 1999). Tg2576 mice, which overexpress a transgene of human APP carrying the Swedish mutation (K670N, M671L), demonstrate age-dependent amyloid deposition, senile plaques and cognitive impairment (Hsiao et al., 1996). To investigate whether these mice also show cholinergic dysfunction, regional biochemical markers of this system were determined.

Brains from Tg2576 mice and non-transgenic (non-Tg) littermates of 3 ages (4, 8 and 13 months) were dissected to yield cortex, hippocampus and striatum and frozen (-70 C) until assayed. Membranes were prepared as previously described and binding to receptors determined using a single ligand concentration (Minger et al., 2000).

Cortical muscarinic M1 and M2 binding was significantly reduced (by 70% and 60% respectively) in Tg2576 mice compared to non-Tg littermates at 13 months of age. There were no significant reductions in younger age groups. although a 25-30% decrease was seen in these receptors in 8 month old animals. No significant changes in binding to nicotinic receptor subtypes (alpha 4 beta 2 or alpha7) were seen.

In conclusion, although the loss of muscarinic receptors in the oldest Tg2576 mice exceeds that seen in AD, it does coincide with cognitive impairment and beta deposition in these animals (Hsiao et al., 1996).

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SYMPATHETIC SUPERIOR CERVICAL GANGLIA (S.C.G.) OF CAT CHOLINERGIC RELAY OF HYPOTHALAMIC-STIMULATED ORGAN-SPECIFIC VASCULAR CHANGES-RELEVANCY TO NORMAL AND TO CLINICAL DYSAUTONOMIC FUNCTION
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Electrical stimulation of prefrontal lateral hypothalamus (LH) induces a moderate systemic BP rise. We examined this in cat model whether it is result of a general systemic or a localized vascular bed effect. Muscarinic and nicotinic mediation of sympathetic ganglia neurotransmission has been shown by use of ganglionic blockers ATN and Hexamethonium. We obtained potentiation of the LH-induced BP rise by systemic ATN or applied in vivo in situ to the desheathed s.c. ganglia thus showing the BP rise cholinergic muscarinic mediation at the ganglia, proposing this a blocking of a ganglionic attenuating mechanism. This cholinergic muscarinic mediation of sympathetic ganglia relay appeared to inherently allow specificity in target activation. Our study has confirmed that this BP rise is a result of discrete localized blood flow changes in just one or two individual organs: it is abolished by bilateral cutting of either one of s.c.g. exiting nerves the medial or the lateral branch. In compliance with the concept of segmental sympathetic innervation the respective s.c.g exiting nerves were traced and identified in vivo, the medial to pharyngeal and to lower respiratory organs, the lateral branch to neck vasculature. Organ-specific blood flow neuro-regulation via segmental autonomic output from s.c.g. was thus demonstrated- as differentially innervating, neck vs pharyngeal and lower respiratory organs. While normal significance of this finding is implicit, we suggest possible bearing on dysautonomic swelling into the trachea due to anomalous fuction of the innervation.
THE EFFECT OF DIET RESTRICTION, SEPARATION STRESS AND TYROSINE ADMINISTRATION ON THE CHOLINERGIC SYSTEM IN MICE

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Background: We have previously shown that weight loss leads to alterations in autonomic tone which may affect cognitive function. We have investigated the effect of tyrosine on hippocampal cholinergic activity in two animal weight loss models - Diet restriction (DR) and self-induced weight loss caused by Separation stress. Methods: Female Sabra mice were fed either 100% (control), or DR to 60% and 40% of daily nutritional requirements. Mice in the Separation group were housed in a cage fitted with six individual Plexiglas partitions of size 11×10×12 cm. They could smell each other and see their neighbours without physical contact except when transferred to regular cages for the 2-h feeding schedule. Control groups were housed 6 to the same cage without partitions. M1 receptors were evaluated by Pirenzepine binding while choline uptake by IHC-3, choline acetyltransferase according to (Lau et al, 1987) acetyl cholinesterase by (Eldman et al, 1961 and Fattranska et al, 1987). M1 mRNA and protein by Northern and Western blotting respectively. Results: DR to 40% significantly decreased choline uptake (p<0.05) and M1 receptor number (Bmax) (p<0.05), without changes in affinity (Kd), choline acetyltransferase (ChAT) or acetyl cholinesterase (AChE) activity. Tyrosine administration significantly increased choline uptake (p<0.05) and M1 density in the 40% DR (p<0.01) without changes in affinity. ChAT activity was decreased after tyrosine-significantly after 40% DR (p<0.05) while AChE was not affected. M1 mRNA and protein were not influenced by DR or tyrosine. Separation stress increased M1 receptor density (p<0.05) and its mRNA signal (p<0.001), without changes in choline uptake, ChAT and AChE. Tyrosine further increased M1 receptor density of stressed mice (p<0.05).

Conclusion: Tyrosine might be a potential therapy for the management of problems associated with stress induced weight loss.

EFFECTS OF LITHIUM CHLORIDE ON MEMORY PERFORMANCES OF MICE IN ELEVATED PLUS-MAZE TEST

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It has been frequently reported that cognitive functions of patients use lithium are impaired although there is an inconsistency in this situation. Therefore the effects of lithium chloride (LiCl) on memory performances of mice were investigated in elevated plus-maze test. In this test transfer latency (TL) the time it took for each mouse to move from the open arm to either of the enclosed arms where feels itself safe is recorded twice in 24 hours. On second trial day, learning leads a shortening in TL values and its prolongation indicates memory impairment. Neither acute nor chronic LiCl administrations have altered memory performances of mice. The results suggest that lithium may not always affect memory as negative. Besides scopolamine, an anticholinergic agent induced prolongation of TL was reversed by LiCl. So it can be considered that lithium may ameliorate some kinds of memory impairments and the cholinergic system may be involved in its actions on memory.
EFFECTS OF 7-NITROINDAZOLE ON MEMORY PERFORMANCES OF RATS TRAINED FOR THREE-PANEL RUNWAY TASK: HIPPOCAMPAL CHOLINERGIC ENZYME ACTIVITIES

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Nitric oxide synthase (NOS) inhibition leads memory deficits in some animal experiments and does not impair memory in some others. Roles of cholinergic and nitricergic systems in working and reference memory performances of rats were assessed regarding the relation between them. Animals trained for working and reference memory performances in three-panel runway were injected intraperitoneally with selective neuronal NOS inhibitor, 7-nitroindazole (7-NI) and observed 45 minutes later from this administration in the runway recording error (wrong door choosing) and latency (time to reach the food pellet in goal box). The activities of choline acetyltransferase (CAT) and acetylcholinesterase (ACHE) were measured in hippocampi of rats taken immediately after the test. The results have showed that systemic administration of 7-NI did not affect the two kinds of memory performances and there was not any alteration in hippocampal CAT and ACHE activities in rats administered NOS inhibitor comparing control group. In conclusion, nitric oxide does not seem to contribute to mechanisms underlying memory performances of rats under these experimental conditions and the results suggest that the cholinergic system is not involved in the possible impairing effects of NOS inhibition on memory.

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HYDROCORTISONE AFFECTS THE DENSITIES OF CARDIAC MUSCARINIC AND ADRENERGIC RECEPTORS

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Gluocorticoid hormones affect the expression of a number of proteins but information on their effects on neurotransmitter receptors in the heart is incomplete and controversial. We investigated the effects of repeated administrations (1-12 days) of high doses of hydrocortisone (50 mg/kg) to adult rats on the densities of cardiac muscarinic and alpha-1, beta-1, beta-2 and putative beta-4 adrenergic receptors, on the coupling of muscarinic receptors with G proteins and on the control of adenyl cyclase.

Hydrocortisone enhanced the densities of muscarinic receptors in the atria and both ventricles. The density of beta-1 adrenoceptors became enhanced in the atria, and that of beta-2 adrenoceptors was raised in the atria and ventricles. The density of putative beta-4 adrenoceptors first increased and then diminished in the atria and did not change in the ventricles. It varied independently of the density of beta-1 adrenoceptors, although the putative beta-4 adrenoceptors are now proposed to represent an atypical state of beta-1 adrenoceptors. The alpha-1 adrenoceptors underwent a decrease followed by a transient increase in the atria and no change in the ventricles. According to the effects on GTP on the binding of carbachol, the coupling of muscarinic receptors with G proteins was not affected. The activity of adenyl cyclase (determined by HPLC) and its stimulation by isoprenaline and inhibition by carbachol were not significantly altered.

The findings raise the question of possible roles of glucocorticoids in the control of the expression of neurotransmitter receptors in healthy and diseased hearts.

ACETYLCHOLINE AND NO-MEDIATED CGMP SYNTHESIS IN THE RAT BRAIN

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About 30 years ago acetylcholine was linked to cGMP in the brain for the first time. Only recently it was shown that NO-mediated cGMP synthesis takes place in cholinergic fibers in the forebrain of the rats (De Vente et al. Exp. Brain Res. 136, 480-491 2001). NO - cGMP signal transduction is found throughout the central nervous system. The NO stimulated intracellular increase in cGMP will be short lived due to the presence of phosphodiesterases (PDE). PDE activity in the basal ganglia is generally very high. PDE's present a very complex group of enzymes, containing a large number of isozymes classified into 11 subfamilies. Each family has its own structural characteristics, substrate preference and inhibitor profile. It is not known which PDE isozymes are present and in control of cGMP levels in cholinergic neurons. We incubated brains slices in vitro in the presence of different PDE inhibitors and different NO donor compounds. cGMP synthesizing structures were visualized and characterized using immunoocytochemistry. cGMP synthesis was found to be present virtually all of cholinergic fibers in the cerebral cortex and in almost all cholinergic fibers of the basal ganglia further details will be presented on the presence of PDE isozymes in these cholinergic fibers.

EEG EVALUATION OF HUPERZINE A, A REVERSIBLE CHOLINESTERASE INHIBITOR

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Huperzine A (HupA), an alkaloid isolated from the Chinese club moss. Huperzia serrata, is characterized as a potent, reversible cholinesterase inhibitor that crosses the blood brain barrier. This compound has previously been shown to provide protection against seizures, neuropathology and mortality induced in laboratory animals by subsequent challenge with organophosphate nerve agents (i.e. soman). Because of HupA's potential as a neuroprotective drug, identification of potentially serious CNS side effects is a necessary part of its evaluation. In this study the EEG and behavioral effects of a single i.v. injection of HupA (0.5-2.0 mg/kg, n=5/dose) were evaluated over a 24 hour period. Male Sprague-Dawley rats were implanted with cortical EEG electrodes and jugular vein catheters several days prior to administration of HupA. Immediately following treatment, dose dependent symptoms of muscle fasciculation, intense sedation and a significant delay in the onset to normal EEG slow-wave sleep (maximum =167 ± 10 min/h=10) were apparent. There was no evidence of seizure activity or marked cortical slowing. However, at the three highest doses of HupA evaluated (1.5, 1.75, 2.0 mg/kg), computer-aided spectral analysis revealed significant shifts in the EEG frequency pattern relative to baseline controls. EEG power increased in the 4-8 Hz frequency band over dose-dependent consolidation of the EEG waveform centered at a peak frequency of 5.5 Hz. Light microscopic evaluation of HupA treated brains removed 24 h post-injection failed to reveal histological evidence of lesions. In rats, even near-lethal doses of the cholinesterase inhibitor HupA were apparently without significant functional or pathological central nervous system intoxication.

-52-
HUPERZINE A AND CHOLINESTERASE INHIBITORS: GLUTAMATE AND BENZODIAZEPINE RECEPTOR INTERACTIONS
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Huperzine A (HupA) is a natural alkaloid that exhibits a unique dual pharmacology: it is a potent and specific acetylcholinesterase inhibitor over butyrylcholinesterase, but also a neuroprotective agent. We demonstrated that HupA protected primary neuronal cells against excitatory amino acid induced neurototoxicity. Part of its pharmacological action resides in its ability to non-competitively inhibit the passage of calcium ions through NMDA ion channels. Thus, we established that HupA non-competitively inhibited 3H-MK801 and 3H-TCP binding to synapticosomal plasma membranes, (pseudo Ki of 5-10 uM). MK801 and TCP are specific probes for the NMDA ion-channel PCP site. HupA did not interact at the NMDA glycine, polyamine, or agonist sites as determined by ligand binding studies. In addition, similar results were observed for the (+)stereoisomer of HupA, which exhibits markedly reduced anticholinesterase activity. Additional evidence that the neuroprotective effects of HupA were separate from cholinesterase activity is that cholinesterase inhibitors such as tacrine, physostigmine, or E2020 showed markedly less effect in displacing 3H-MK801. Unlike MK801, HupA, in the presence of an irreversible cholinesterase inhibitor, dissociated from rat brain NMDA receptors in synapticosomal plasma membranes with both a fast and slow component. This may be another advantage of HupA over high affinity ion-channel blockers that exhibit side effects such as PCP. HupA also exhibited antagonism of central benzodiazepine receptors measured with 3H-RY-80, although it was less strong than observed for 3H-MK801. These data suggest that neuroprotection by HupA against excitatory-induced cell death is at least partially based on the blockage of calcium ions through NMDA ion channels, and that the effect is independent of its efficacy as a cholinesterase inhibitor.

CHOLINERGIC MODULATION OF CHEMOTAXIS IN HUMAN MELANOMA CELLS
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Neural crest cells transiently express muscarinic acetylcholine receptors during migration from the neural tube to the definitive location in the epidermis. Differentiated melanocytes of normal human skin do not express muscarinic receptors. After malignant transformation receptors re-appear in primary and metastatic human melanomas. We assume that an embryonic trait which is involved in morphogenesis, is reactivated in a pre-malignant or malignant state and mediates cellular movements during invasive growth. For further characterization, we used the human melanoma cell line SK-Mel 28. Pharmacological characterization via dose–response curves of calcium mobilization after stimulation by acetylcholine indicated expression of the M3 subtype with a Kd of approximately 2x10^-7M. Molecular weight of muscarinic receptors of SK-Mel 28 cells determined by western blot with M3 subtype specific antibodies was 66kDa which is in accordance to published data for the M3 subtype. In time lapse videomicroscopy, stimulation of muscarinic receptors with acetylcholine and carbachol induced contractions and a change in cell shape. In the present study, an effect on chemotactic activity by cholinergic treatment is demonstrated. By chemotaxis assays with modified Boyden chambers, an increase of movement towards the chemotactic factor fibronectin was found after addition of carbachol in the upper compartment. This indicates a modulatory effect on cell movement of the muscarinic cholinergic system in non neural cells.

CALCIUM MOBILISATION AND CELLULAR CONTRACTION OF EMBRYONIC LENS VESICLE AND NEURAL TUBE ON MUSCARINIC CHOLINERGIC STIMULATION
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Embryonic epithelial structures like the neural tube and the lens vesicle exhibit cholinesterase activity during phases of morphogenetic movements. Cholinesterase activity coincides with the expression of muscarinic receptors. Here we show that the isolated lens vesicle and the intact neural tube in the chick embryo react on muscarinic cholinergic stimulation by calcium mobilisation and concomitant cellular contraction. The lens vesicle was stained with Fura2-AM and studied by digital time lapse epifluorescence video. On stimulation with carbachol we observed a peak of intracellular calcium release followed by a plateau phase of extracellular influx, which was reversed by addition of atropine. The peak reaction was accompanied by contraction of the lens. Addition of atropine led to relaxation to the initial diameter. Experiments with the neural tube were performed by perfusion of the neural canal in the intact embryo. After perfusion with atropine a characteristic, but reversible deformation of the neural tube occurred. The experiments demonstrate, that the embryonic muscarinic system is involved in cellular movements of non-neural cells during morphogenesis.

COMBINED ANDROGEN-DONEPEZIL TREATMENT IN POST-STROKE REHABILITATION
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There is a theoretical basis for thinking that a combination of androgenic steroid DHEA and the anti-cholinesterase, donepezil may have an enhanced (greater than additive) effect in the post-stroke syndrome. Steroids increase potassium conductance in central neurons and in particular, hippocampal neurons (eg. Colino and Halliwell. Nature 328:73, 1987). Under certain circumstances, this effect leads to hyperpolarization and a modulatory effect on neurotransmitter function (Beck et al., Neuropharmacol 14:27, 1996). More specifically, androgens have been shown to facilitate the responses of hippocampal neurons to cholinomimetics (including ACh's). In fact, Karczmar and Dun showed that androgens enabled non-cholinergic neurons to respond to the direct actions of cholinergic agents. (Karczmar and Dun, unpublished observations) Androgens produce trophic in several systems, including neuronal systems (Jones, NY Acad Sci 743:141, 1994) They improve muscle strength, mood and memory, particularly in elderly, hypo-gonadal subjects. (Tenover, J Clin Endocrinol Met 75:1092, 1992) Taken together, these considerations suggest that DHEA will enhance the beneficial effects of donepezil in this population and hence speed recovery from stroke. We will describe a study which aims to determine if the hypothesized synergy between androgens and donepezil promotes enhanced recovery from stroke. Patients with recent strokes receive donepezil alone, DHEA alone, or a combination of the two agents. They are followed over a six-month period with monitoring of their neurological, neuropsychological, and psychological functioning. In addition, patients will be followed with repeated MRI and EEG studies. Preliminary data from this ongoing study will be presented.

We acknowledge the support of the Illinois AMVETS and the Chicago Association for Research and Education in this work.
LOCALISATION OF THE HIGH-AFFINITY CHOLINE TRANSPORTER-I IN RAT SKELETAL MUSCLE AND SPINAL CORD

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Acetylcholine (Ach) is synthesised by choline acetyltransferase in the cytoplasm of cholinergic neurons. The vesicular acetylcholine transporter (VACHT) imports Ach into synaptic vesicles. After exocytotic release it is split into acetate and choline. Choline is taken up into the synaptic terminal via a high-affinity choline transporter (CHT) for resynthesis of Ach. The first high-affinity CHT (CHT1) was recently cloned, and in situ hybridization showed its expression in spinal motorneurons. We generated polyclonal antisera against a synthetic peptide corresponding to aa residues 29-40 of the rat CHT1 sequence. These antisera were used in immunofluorescence to analyse the distribution of the CHT1 protein either singly or in combination with antisera against VACHT to label cholinergic terminals. Alexa-488 conjugated a-bungarotoxin to label motor end plates. Perikarya of spinal motorneurons were moderately CHT1- and VACHT-immunoreactive, while the recurrent cholinergic synapses were intensely CHT1- and VACHT-immunolabelled. In skeletal muscles, motor end plates showed an intense CHT1- and VACHT-immunoreactivity, while preterminal axons showed a distinct CHT1- but only weak VACHT-immunolabelling. The results show a preferential localisation of the CHT1 protein at the neuron-neuronal and neuro-muscular synapse, well in line with its anticipated function in the synaptic transmitter recycling. (supported by the DFG, SFB 547).

CHOLINERGIC-GLUTAMATERGIC INTERACTIONS IN HIPPOCAMAL NEURONS: POSSIBLE ROLE IN THE NORMAL AND DISEASED HIPPOCAMPUS

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In the cortex, released Acetylcholine (ACh) binds to both pre- and post-synaptic receptors and assumed to serve as a modulatory neurotransmitter and thus set the response of the nerve cell to an incoming stimulus. The electrophysiological details of such modulation are only partly understood. We have tested the role of Ach on excitatory synaptic transmission in the CA1 area of the hippocampus, using extracellular and intracellular recordings. In whole cell voltage clamp experiments, the frequency, but not amplitude, of spontaneously occurring excitatory synaptic current (EPSCs), was increased following either local application of Ach or the addition of cholinesterase inhibitors (ChEIs) to the bathing solution. In contrast, the addition of the muscarinic antagonist, atropine, reduced EPSCs frequency, suggesting a role for basal Ach release in modulating glutamate transmission. The effect of ChEIs on the local neuronal network was tested using extracellular recordings of evoked responses. In control animals following ChEIs exposure a mild (+2 fold) increase in the amplitude of population spikes was observed. In contrast, weeks following stress, or injections of low concentrations of the AChEI, di-isopropyl-fluoro-phosphonate (DFP), a marked sensitivity of the evoked synaptic response to either AChEIs or Atropine was noted. These results suggest that Ach-regulated synaptic transmission might alter in response to environmental conditions, and be associated with marked alterations in cortical function.

-54-

HUPERZINE A, A PROMISING ANTI-ALZHEIMER'S AGENT, REDUCES STAuroSPORINE-INDUCED APOPTOSIS IN NG108-15 CELLS

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The present study investigates the effects of huperzine A (HupA) and taurine (Tac), two anti-Alzheimer's agents, on staurosporine-induced apoptosis and potential mechanisms in neuroblastoma hybrid NG108-15 cell lines. Preincubation with HupA and Tac significantly attenuated 0.1 μM staurosporine-induced chromatin condensation, nuclei fragmentation and DNA laddering, and inhibited or delayed expression of the pro-apoptotic gene product Bax, and increased Bcl-2 levels. Acetylcholinesterase (AChE) enzymatic and Western blot assay demonstrated that staurosporine treatment induced an abnormal AChE activity that was resistant to HupA and Tac. These results suggest that HupA and Tac might exert significant protection against staurosporine-induced apoptosis via suppressing the abnormal AChE and modulating expression of apoptosis-related proteins in apoptotic cells, which might be beneficial for its therapeutic usage.

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DOPAMINE RELEASE FROM RAT STRIATAL SLICES IN VITRO AND FUNCTIONAL EFFECTS IN 6-OHDA TREATED RATS IN VIVO ARE MEDIATED BY BETAZ CONTAINING NICOTINIC ACETYLCHOLINE RECEPTORS

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Several studies have reported that nicotinic acetylcholine receptors (nAChRs) are located on dopaminergic cell bodies and terminals and that activation of neuronal nAChRs plays a role in the reinforcing properties of nicotine and cigarette smoking. It is also clear that there is a negative correlation between smoking and the incidence of Parkinson's Disease and that nicotine can increase dopamine release in vitro and in vivo. We have utilised several new ligands for nicotinic receptor sub-types to investigate the nAChRs involved dopamine release in vitro and the functional effects of dopamine in vivo. We have evaluated the effects of SIB-1508Y and TC-2559 (beta2 preferring) nicotine (a broad spectrum agonist), SIB-1553A (beta4 preferring) and Astra II (alpha7 selective) on dopamine release from rat striatal slices and on rotational behaviour in rats with unilateral 6-hydroxydopamine lesions of the substantia nigra. Nicotine, TC-2559 and SIB-1508Y produced a dose-dependent, mecamylamine sensitive increase in dopamine release in vitro. In contrast, SIB-1553A and Astra II did not stimulate dopamine release. In vivo, SIB-1508Y (10 mg/kg s.c.), nicotine (0.5 mg/kg s.c.) and TC-2559 (5 mg/kg) produced a significant increase in ipsiversive rotations (p < 0.05), while SIB-1553A (20 mg/kg s.c.) and Astra II (10 mg/kg s.c.) had no effect. The SIB-1508Y-induced increases in rotational behaviour were blocked by mecamylamine. These results indicate that dopamine release in vitro and in vivo is mediated by beta2 containing nicotinic receptor sub-types and suggest that beta2 nAChR agonists may provide a useful symptomatic treatment for Parkinson's Disease.
THE EFFECTS OF GALANTAMINE IN PATIENTS WITH REFRACTORY SCHIZOPHRENIA RECEIVING RISPERIDONE

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Objective: The prevalence of smoking in patients with schizophrenia is higher than in the general population. Some schizophrenia patients may have abnormal nicotinic neurophysiology, since sensory gating deficiencies can be corrected by nicotine. Galantamine (Reminyl(R); GAL) is a reversible acetylcholinesterase inhibitor (AChEI) and an allosteric modulator of nicotinic acetylcholine (ACh) receptors (nAChR), for treating mild-to-moderate Alzheimer's disease. This ongoing 4-week, dose-finding study aims to investigate the effect of GAL on behaviour in risperidone (RIS)-treated patients with schizophrenia. Methods: All patients are being treated with fixed-dose RIS (1-6 mg daily). Patients (n = 24) are divided into 4 groups. Groups 1-3 have 2 patients on placebo (PLA) and 6 patients on GAL. (Group 1: 8 mg BID; Group 2: 12 mg BID; Group 3: 16 mg BID; standard dose-escalation is undertaken for GAL. All patients in Group 4 receive PLA. Psychopathology is assessed using the Brief Psychiatric Rating Scale and Clinical Global Impression at baseline and at days 7, 14, 21 and 28. Cognitive psychomotor performance, neuropsychological functioning and smoking behaviour are also assessed at varied time-points. Results: In two patients with refractory schizophrenia, preliminary findings with GAL have shown improvement in episodes of agitation in one patient (8 mg BID) and improved social and hygiene manners in another patient (12 mg BID). Conclusion: The results from this ongoing study and subsequent studies may aid in establishing a therapeutic use for GAL in patients with refractory schizophrenia.

CEREBRAL METABOLIC ACTIVATION WITH CHOLINESTERASE INHIBITOR THERAPY IN ALZHEIMER'S DISEASE

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Objective: The clinical response of patients with Alzheimer's disease (AD) to cholinergic treatment is similar with different cholinesterase inhibitors, suggesting a common neuronal system affected by general cholinergic augmentation. We performed a study to compare functional patterns of metabolic response and identify brain regions affected by increased cholinergic tone due to treatment with various cholinergic AD treatments. Methods: Demographically similar mild AD patients (n = 12) treated either with metrifonate, donepezil or galantamine (Reminyl(R)) were studied from a pool of 30 patients. Pre- and post-treatment [18F]-fluorodeoxyglucose positron emission tomography (FDG-PET) was performed, and patients were followed using the Mini-Mental State Examination and the Neuropsychiatric Inventory. Patients' normalized PET studies, registered to the AD probabilistic anatomic atlas, were subjected to a voxel-by-voxel subtraction of the post-treatment minus pre-treatment studies. Sub-volume thresholding was used to correct random lobar noise, allowing the production of 3-D functional significance maps for total brain voxel comparison. Results: Significant post-treatment anterior cingulate, dorsolateral frontal, and supramarginal activation was observed with cholinergic treatment in pooled groups. Different mechanisms of action of the AD treatments studied could underlie differential activation patterns among the patient groups. More results will be presented that may establish this further. Conclusion: FDG-PET has potential as a useful technique to enhance our understanding of possible differences in the effects a number of cholinergic treatments in the brains of patients with AD. For instance, attentional networks, with greatest activation centred on the anterior cingulate, appear to be a key neurophysiological target activated by cholinergic AD treatments.

THE ROLE OF LIPID PEROXIDATION IN THE MECHANISM OF NEUROTOXICITY OF ORGANOPHOSPHATES

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The present study was undertaken to elucidate the relations between lipid peroxidation, organophosphates (OPs) toxicity and delayed, long lasting, non-cholinergic changes. In the experiments on rats we studied the influence of OPs intoxication by paraaxon, sarin, malathion, soman on lipid peroxidation in rat cerebral hemispheres. The level of lipid peroxidation was measured as the amount of common phospholipids, peroxidate lipids and malondialdehyde (MDA) in reaction with thioarbituric acid. Results were compared to those with pre-treatment with atropine and reversible cholinesterase inhibitor - galanthamine alone or together with different antioxidants (tocopherol, oxybutynacyl, insole). The rate of reaction of conditioned reflex of active avoidance was measured. OPs caused a rapid, dose-dependent increase of peroxidate lipids and MDA 15-30 days after intoxication. With paraaxon and sarin pre-treatment with atropine and galanthamine totally prevents the all symptoms of intoxication and changes in lipid peroxidation. Comparatively such type of prophylaxis in malathion and soman poisoned rats didn't normalize the biochemical and physiological parameters. The protective effect of antioxidants against soman and malathion - induced lipid peroxidation was shown. Therefore, malathion and soman - associated lipid peroxidation is likely to arise mainly as a primary change which may, however, play a significant role in delayed neurotoxicity and conditioned reflex activity.

RESCUE OF THE NEURODEGENERATIVE PHENOTYPE IN AD11 ANTI-NGF MICE

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We have obtained transgenic mice expressing a neutralizing anti-NFG recombinant antibody, in which the levels of antibodies are three order of magnitude higher in adults than in newborns (Ruberti et al., J. Neurosci. 20, 2000). The analysis of brains of these mice revealed that they display a progressive neurodegeneration characterized by neuronal loss, cholinergic deficit, tau hyperphosphorylation, extracellular deposits of amyloid precursor protein (APP) and behavioral deficits (Capsoni et al., Proc. Natl. Acad. Sci. USA 97, 2000). Beta-amyloid cerebralvascular deposition and beta-amyloid plaques are observed as well (Capsoni et al., submitted). In this study, we performed pharmacological treatments of AD11 anti-NGF mice with acetylcholinesterase inhibitors and agents that increase NGF levels and/or activity. Different time windows were examined. Amelioration of the neurodegenerative phenotype was achieved both with acetylcholinesterase inhibitors and NGF, thus showing the dependency of the phenotype from NGF deprivation. This study was partially supported by Telethon (grant D.122) and SIRS s.r.l. (now Lay Line Gen.-mice S.p.A.)
ACUTE CHOLINERGIC RESCUE OF SYNAPTIC PLASTICITY IN THE NEURODEGENERATING CORTEX OF ANTI-NERVE GROWTH FACTOR MICE

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Deficits in cholinergic systems innervating cerebral cortex are associated to cognitive impairment during senescence and in age-related neurodegenerative pathologies. However, little is known about the role of cholinergic pathways in modulating cortical plasticity. Basal forebrain cholinergic neurones are a major target for Nerve-Growth Factor (NGF). In order to investigate the relationship between cholinergic innervation and cortical synaptic plasticity, we exploited a transgenic mouse model in which the activity of NGF in the adult nervous system is neutralized by the expression of blocking antibodies to NGF itself (anti-NGF mice) [Roberti, F., et al. (2000). J. Neuroscience. 20, 2589-2601]. In six months old anti-NGF mice, we show that the reduction in cholinergic innervation of the cortex is associated to different forms of synaptic plasticity impairment. A local, acute increase in the availability of acetylcholine rescues these synaptic plasticity deficits, thus indicating that cholinergic system mediates the impairment of cortical plasticity at this early stage of the neurodegenerative process triggered by NGF neutralization. Our results represent an important step to unveil the pivotal role of cholinergic transmission in modulating adult cortical plasticity. This study was partially supported by Telethon (grant D.122) and SIRS s.r.l. (now Lay Line Genomics S.p.A.).

THE EFFECT OF NICOTINE ON EXPRESSION OF NICOTINIC RECEPTORS IN THE BRAIN OF PATIENTS WITH ALZHEIMER'S DISEASE

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Epidemiological studies have suggested some association between smoking and Alzheimer’s disease (AD). Nicotine treatment may compensate for some of the cholinergic deficits e.g. decreased number of nicotinic acetylcholine receptors (nAChRs) seen in AD. In the present study the effect of smoking on nAChRs in AD was investigated in autopsy brains using receptor binding and Western blotting techniques. The nAChR subtypes were measured using radioactive ligands 3H-cytisine (alpha 4), 3H-epibatidine (alpha 3, 4) and 125I-aBTX (alpha 7). The binding of 3H-cytisine (2.0 nM) and 3H-epibatidine (0.1 nM) was significantly increased in the hippocampus, cerebellum, frontal and temporal cortices and 125I-aBTX (3.5 nM) in the temporal cortex of smoking controls (SC) (69.5±3.2 years, n=4-11) compared to non-smoking controls (NSC) (78.9±3.5 years, n=9-10). A significantly increased binding of 3H-epibatidine and 3H-cytisine in the temporal cortex and 3H-cytisine in the cerebellum was observed in smoking AD (SAD) (73.3±3.5 years, n=8-13) compared to non-smoking AD (NSAD) (80.2±3.2 years, n=8-13). Binding levels for both ligands in SAD was similar to those in NSC. Up-regulation of 3H-cytisine, 3H-epibatidine and 125I-aBTX binding in the temporal cortex of SC compared to NSC was positively correlated with an increased level of alpha 3, 4 and 7 protein expression. This study showed an increased number of nAChRs, especially alpha 4 in SAD compared to NSAD, which may be relevant to a neuroprotective effect of nicotine.

A PEPTIDE FROM THE C-TERMINAL OLIGOMERISATION DOMAIN OF HUMAN SYNAPTIC T-FORM ACETYLCHOLINESTERASE FORMS CLASSICAL AMYLOID FIBRILS

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Acetylcholinesterase (AChE) is abnormally localised in amyloid plaques, which are a key neuropathological feature of Alzheimer’s disease. It has recently been shown to interact in vitro with the beta-amyloid peptide, which is the principal constituent of the Alzheimer plaques, and to increase the rate of its assembly into amyloid fibrils (Alvarez, 1997 & 1998; Inestrosa, 1998). We report that a synthetic peptide corresponding to residues 586 to 599 of the synaptic or tailed (T) splice variant of human AChE can itself form typical amyloid fibrils similar to those formed by beta-amyloid. Under physiological buffering conditions, the peptide rapidly aggregates into fibrillar structures that have all the classical features of amyloid. Electron microscopic analysis of negatively stained aggregates reveals that they are composed of long fibrils which are 6-7 nm in diameter. The fibrils bind both Congo Red and thioflavin-T and cause the expected metachromatic shift in the absorbance spectrum of Congo red and in the fluorescence excitation spectrum of thioflavin-T. Far-UV circular dichroism spectroscopy reveals that fibril formation by the peptide is accompanied by a change in its secondary structure from random coil to beta-sheet. Amyloid fibrils have a generic ‘cross-beta’ structure consisting of repeating beta-sheets running perpendicular to the fibril axis. Furthermore, the peptide is cytotoxic in vitro, as determined by its effect upon the reduction of MTT by PC-12 pheochromocytoma tissue culture cells. This fibrillogenic region of the C-terminal oligomerisation domain of T-form AChE might be responsible for an interaction of the enzyme with beta-amyloid.

NICOTINE INDUCES GLUTAMATE RELEASE FROM HIPPOCAMPAL MOSSY FIBRES SYNAPTOSES

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Nicotine is known to modulate the release of several neurotransmitters from mammalian CNS synaptosomes: Dopamine, ACh, Noradrenaline and GABA. In addition, electrophysiological evidence suggests that nicotine also promotes the release of glutamate. In the present work glutamate release was elicited from rat hippocampal mossy fibre synaptosomes by using nicotinic agonists, and measured by a luminescence assay. Glutamate release was elicited by micromolar concentrations of nicotine. The maximal effect was observed with 25 mM nicotine, and blocked at 90% of control by curare 100 mM. The effects of other antagonists is presently under investigation. In parallel experiments, carbocyanide fluorescence was used to measure the membrane potential. When promoting glutamate release, nicotine did not induce significant depolarisation. Nonetheless, the amount of transmitter released by maximal doses of nicotine corresponded to that induced by approximately 25 mM KCl. We are currently investigating the pharmacological profile of mossy fibres presynaptic receptors.

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NICOTINIC CHOLINERGIC ACTIVATION OF MAGNOCYTOCELLULAR ENDOCRINE NEURONS OF THE HYPOTHALAMUS

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Using IR DIC videomicroscopy, we performed whole-cell recordings in hypothalamic slices containing the supraoptic (SO) and paraventricular (Pa) nuclei. Acetylcholine (ACH), locally applied in the presence of atropine, evoked a rapidly rising inward current in SO and Pa magnocellular neurons, which persisted in the presence of blockers of synaptic transmission. It could be reversibly suppressed by methyllycaconitine, a selective antagonist of alpha7-containing nicotinic receptors, but was insensitive to dihydro-beta-erythroidine, an antagonist preferentially on non-alpha7 nACHRs.

The effect of ACh could be mimicked by (-)-2PABH, a recently synthesized nicotinic agonist specific for alpha7 nACHRs. ACh also desensitized nicotinic receptors. Desensitization was pronounced and recovery from desensitization was rapid, consistent with the properties of alpha7-containing nACHRs. Nicotinic currents could not be evoked in Pa parvocellular neurons, suggesting that these neurons are devoid of functional nicotinic receptors. Light microscopic autoradiography showed that [125I]alpha-bungarotoxin binding sites are present in the SO and in all magnocellular divisions of the Pa, but are undetectable in other areas of the Pa. Immunohistochemistry, performed using antibodies directed against vasopressin and oxytocin, indicated that responsiveness to nicotinic agonists was a property of vasopressin as well as of oxytocin magnocellular neurons, in both the SO and Pa nuclei. In conclusion, by directly increasing the excitability of magnocellular endocrine neurons, nicotinic agonists can influence the release of vasopressin and oxytocin from the neurohypophysis. By contrast, they appear to have no direct effects on paraventricular parvocellular neurons.

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UNDERSTANDING THE DUAL MODE OF ACTION OF REMINYL(R) USING A VIRTUAL SYNAPTIC CLEFT

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Objective: Galantamine (Reminyl(R)) GAL) is both an acetylcholinesterase (AChE) inhibitor and an allosteric modulator at nicotinic acetylcholine (ACh) receptors (nAChR). As these two cholinergic effects are interdependent, the overall effect of GAL on transmission is a complex function of the synaptic concentration of GAL. We have addressed this problem using a computer simulation based on mathematical descriptions of known physicochemical interactions. Methods: The virtual synaptic cleft is a computer model incorporating the latest neuroanatomical and neurophysiological data on the cholinergic synapse. It includes a full description of the kinetic transitions associated with both the alpha4beta2 and the alpha7 nAChR, and describes the potentiating interaction between GAL and the alpha4beta2 nAChR. Results: Overtly potent AChE inhibition can drive nicotinic receptors into desensitization, especially under phasic activation of the synapse. As a consequence, the beneficial effects of increasing ACh are reduced. By introducing neuropathology data from AD tissue, the model allows assessment of cholinergic transmission deficits, and evaluation of the effect of pure AChE inhibition and of GAL on synapses from mild to severe AD. The virtual synaptic cleft also allows for the interaction between cholinergic and dopaminergic neurotransmission (modelled after the neurophysiology of the striatum); simulations show a unique effect of GAL on dopamine levels. Conclusion: This computer simulation allows investigation of the molecular interactions of complex neurological systems, where different subsystems are active simultaneously. The web-based nature of this application ensures maximal flexibility for worldwide use and increases its usefulness as a knowledge reference database.

ALTERED ACTIVITY OF CHOLINERGIC ENZYMES IN MUSCLES AND BRAIN OF THE OBESE-DIABETIC (OBD) MOUSE

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The activity of acetylcholinesterase molecular forms was determined in the diaphragm, extensor digitorum longus (EDL) and soleus muscles, and the brain of obese diabetic (ob/ob) mice of different ages and their lean littermates. Three peaks of activity, representing the G1, G4 and A12 molecular forms were separated in the muscles and two peaks, representing the G1 and G4 forms in the brain. In the diaphragm, at 10-12 weeks of age, the activity of all molecular forms of the enzymes was significantly higher in the diabetic mouse than in controls. In the EDL the activity of only the G4 form was significantly higher than in the controls. In the soleus there were no significant differences between the two groups of mice. In the brain tissue the activity of both the G1 and G4 molecular forms was significantly lower in the obese mice than in the lean mice. Neuropathy is often present in non-insulin dependent (type 2) diabetes and defects in glutamatergic transmission have been reported. However this is the first report of changes in the cholinergic system. The importance of these findings in diabetes is as yet unclear but they may be relevant to the presence of cognitive defects in the condition.

EFFECT OF PYRIDOSTIGMINE ADMINISTRATION ON ACETYLCHOLINESTERASE AND CHOLINEACETYLTRANSFERASE ACTIVITY IN THE GUINEA-PIG STRIATUM AND CEREBELLUM

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Pyridostigmine bromide or saline (controls) was administered continuously for 6 days in guinea pigs, via osmotic pumps. The activities of G4 functional acetylcholinesterase (AChE) and cholineacetyltransferase (ChAT) were measured in striatum and cerebellum, (regions of high and low levels of acetylcholine respectively). Measurements were made on day 6, and at 1, 7, or 13 days after the treatment. In the pyridostigmine-treated animals at 6 days the activity of G4 AChE in the striatum was slightly higher than in controls, but the ChAT activity was approximately twice that in controls. At one day after the treatment ChAT activity and AChE activity were both lower than in controls. Thereafter both enzymes gradually increased up to 13 days after the treatment. A different pattern was seen in the cerebellum. In the pyridostigmine-treated animals at 6 days AChE activity was lower than in controls, but on day 7 it was higher. There was no significant change in ChAT activity at any time point examined. In the case of AChE the activity was measured at a time when the inhibitory action of the drug would long have worn off. Therefore the changes in activity of both enzymes were probably due to altered expression of the enzymes. Pyridostigmine does not readily cross the blood-brain barrier. Therefore the changes in the enzyme activities in the brain tissues could be secondary to a peripheral action of the drug. Alternatively the drug or the anaesthetic used could have permeabilised the barrier.

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NICOTINIC BETA4 RECEPTOR MEDIATED ACETYLCHOLINE RELEASE FROM RAT INTERPEDUNCULAR NUCLEUS

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Previous studies (Grady et al., 2001) have shown that acetylcholine (ACh) release from mouse interpeduncular nucleus (IPN) is predominantly mediated by nicotinic receptors containing the beta4 subunit. We studied the effects of different nicotinic agonists on ACh release from rat IPN. In addition, we used immunolocalisation techniques to determine the distribution of the beta4 subunit within the rat brain. ACh release was measured from superfused synaptosomes that had been pre-labelled with [3H]-choline. Nicotine and cystine stimulated ACh release in a dose-dependent manner, which could be blocked by both mecamylaniline (10 microM) and dihydro-beta-erythroidine (100 microM), but not by alpha-bungarotoxin (40nM). In the immunolocalisation studies a rabbit polyclonal antibody raised against a peptide mapping the intracellular loop of the rat beta4 receptor was evaluated, and its monospecificity confirmed using a range of techniques. Beta4 immunoreactivity was demonstrated in a number of brain regions, but was particularly prominent in the IPN and medial habenula. The data indicates that the beta4 subunit is present in the IPN of the rat and that nicotinic receptors containing this subunit are modulating ACh release in this area of the brain.


CALCIUM CONDUCTANCE AND CHOLINE SENSITIVITY OF SLOW CHANNEL SYNDROME ACETYLCHOLINE RECEPTOR MUTANTS

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The slow channel myasthenic syndrome is caused by mutations in genes for the acetylcholine receptor (AChR), which lead to prolonged AChR openings. It is thought that the myasthenic weakness is due to an "endplate myopathy", resulting from excess calcium entry during the prolonged ACh-induced bursts; and also from choline-induced openings that occur with much greater frequency in mutant channels than in wild type. However, the calcium conductance and choline-induced calcium permeability of mutant and wild-type nACHRs have not been compared.

Single channel conductance and calcium permeability of epsilon L221F and wild type nACHRs were studied using outside-out patches from transiently transfected HEK 293 cells. As reported for other mutants, the mutant channels were 20 fold more sensitive to cholinergic agonists. However, the voltage/current relationships at different calcium concentrations were not different between mutant and wildtype, indicating similar calcium permeability. Using ratemetric recordings of FurA-2 AM-loaded cells, internal calcium changes with 5 millimolar choline (plus 1 millimolar atropine) were larger than those with 25 micromolar nicotine, but this difference was not greater in mutant than wildtype. This result does not confirm the marked difference in choline sensitivity between wildtype and mutant AChR channels reported by Zhou et al (PNAS 1996).

BEYOND THE USUAL SUSPECTS. A CHOLINERGIC ROUTE FOR PANIC ATTACKS

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For unknown reasons and through poorly understood mechanisms, people at risk for panic attacks are hypersensitive to suffocative stimuli and experience hyperventilation and anxiety after exposure to heightened concentrations of carbon dioxide. Similarly to the physiological reflex response to hypercapnia in animal and man the anxious response to carbon dioxide in people with panic disorder is at least partially controlled by the central muscarinic receptors (Battaglia et al., 2001). A falsifiable hypothesis (Battaglia, 2002) is offered here that some modifications of the cholinergic functions could underlie human individual differences in carbon dioxide sensitivity and proneness to experience panic attacks. The hypothesis is based upon experimental evidence (Kaufert et al., 1998) that stressful and potentially harmful stimuli prime relatively long lasting changes in cholinergic genes expression and cholinergic receptors' regulation. The adaptive sequel of these modifications include protection of the brain from overstimulation, and, at the level of the corticolimbic circuits, promotion of passive avoidance and learning after stress. The extension of the same modifications to the cholinergic receptors involved in chemoception, however, could lower the threshold for reaction to suffocative stimuli, including carbon dioxide. The exaggerated sensitivity to carbon dioxide observed in humans suffering from panic attacks could then be thought of as an evolutionary cost of the involvement of the cholinergic system in shaping otherwise adaptive responses to stress and threatening stimuli.

SELECTIVE HISTOCHEMICAL STAINING OF PERINEURONAL ACETYLCHOLINESTERASE (AChE) IN THE LIVING ENTERIC NERVOUS SYSTEM (ENS) OF RAT AND GUINEA-PIG


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AChE activity was localized only in perineuronal space of the ENS, when they were stained alive. Absence of staining of intracellular AChE was due to impermeability of the living membrane to the histochemical medium. The rat ENS was stained by means of intracellular perfusion of modified Karnovsky's histochemical medium. After fixation and cleavage of the intestinal layers, initial weak histochemical reaction was intensified by dianinobenzidine. In case of the guinea-pig ENS (protected with a blood-brain barrier-like structure), fresh longitudinal muscle-myenteric plexus preparations were immersed in the histochemical medium, showing intense staining without diaminobenzidine intensification. The cleaved face of preparations of the rat and guinea-pig was adequate for SEM observation. A high contrast image of histochemical precipitates (cupric ferricyanide) was obtained by the backscattered electron imaging (BSEI). The Cu and Fe of the precipitates were detected by energy dispersive X-ray analysis (EDXA), and elementary X-ray mapping of Cu and Fe on the whole specimens provided images superposable to those of the BSEI, or even finer than those of the BSEI. TEM observation confirmed fine perineuronal localization of the AChE activity. It is plausible that the perineuronal AChE plays a role for hydrolysis of ambient ACh and regulates extra-synaptic communication among neurons.
NICOTINIC ACETYLCHOLINE RECEPTOR α5 SUBUNITS MODULATE OXOTREMORINE-INDUCED SALIVATION AND TREMOR

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Objective: To determine function role of neuronal nicotinic acetylcholine receptor (nAChR) α5 subunits in modulating the effects of oxotremorine (OXO) on autonomic functions and tremor.

Background: In 12 distinct nAChR subunits (α2-α10 and β2-β4), α5 subunits are expressed throughout central and autonomic nervous systems and have unique properties in modulation of ACh transmission.

Design/Methods: OXO-induced autonomic functions and tremor were measured in mice lacking α5 subunits (α5−/−) and wild-type (WT) control mice. OXO was injected subcutaneously in gradually increasing doses (0.01, 0.03, 0.1, 0.3 and 0.5 mg/kg in 40 min intervals). The effects of OXO were tested on awake mice (hypothermia, tremor and salivation) and on mice under anesthesia (bradycardia, defecation and salivation).

Results: Injection of OXO produced dose-dependent whole body tremor, salivation, and hypothermia with maximal responses obtained at a dose of 0.3 mg/kg, in both α5−/− (n=4) and WT control (n=7) awake mice. However, α5−/− mice showed significantly greater intensities of salivation and tremor responses to low OXO doses. For example at 0.03 mg/kg α5−/− mice reached near maximal responses (about 90%), while WT mice reached only 40% and 50% of maximal salivary (F1,6=17.2) and tremor (F1,11=28) responses, respectively (p<0.01, one way ANOVA, Dunnett multiple comparison). The hypothermia, bradycardia and defecating effects induced by OXO were of similar magnitudes in the two mouse strains.

Conclusion: The increased OXO effects in α5−/− mice might be due to elimination of α5 inhibitory effects, and support the participation of α5 subunits in cholinergic transmission in autonomic ganglia.

Supported by the Sieratzki Chair of Neurology, Tel Aviv University and the Miriam Turjanski de Gold and Dr. Roberto Gold Fund for Neurological Research.

DEFICIENCY OF β4 NICOTINIC ACETYLCHOLINE RECEPTOR SUBUNITS CAUSES AUTONOMIC CARDIAC AND INTESTINAL DYSFUNCTIONS

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Objective: To investigate the physiological and pharmacological functions of β4 neuronal nicotinic acetylcholine receptors (nAChRs) subunits in autonomic nervous system (ANS).

Background: nAChR β4 subunits are expressed in ANS and influence many properties of physiology and pharmacology.

Design/Methods: Autonomic functions were measured in knockout mice lacking nAChR subunit β4 (β4−/−) and wild-type (WT) mice.

Results: β4−/− mice grew to normal size without showing any obvious physical, neurological or autonomic deficits. There was no difference between β4−/− and WT mice on the rectal temperature changes during exposure to cold stress (6 °C) and following 30 mg/kg morphine, as well as on pupillary size changes following morphine. Heart rate at rest, stressed by cage shaking, during exposure to cold stress or anesthetized was not significantly different between the β4−/− and WT mice. During high frequency of vagal stimulation, the WT mice, but none of the β4−/− mice developed cardiac arrest. Deficiency of β4 subunits strikingly increased the sensitivity to a low dose of C5, (3 mg/kg). A greatly reduced ileal contractile responses to nicotinic agonists choline, dimethylphosphoriperezaminium iodide and nicotine (10 mg/kg each), and epinephrine (0.1 mg/kg) were seen in β4−/− mice.

Conclusion: β4 subunits are important components in the ANS. Deficiency of β4 subunits altered ion channel properties, conductance and sensitivity and affinity of receptors to agonists and antagonists leading to reduction of ganglionic transmission to end-organ.

Supported by the Sieratzki Chair of Neurology, Tel Aviv University and the Miriam Turjanski de Gold and Dr. Roberto Gold Fund for Neurological Research.

AUTONOMIC FUNCTION OF NEURONAL NICOTINIC ACETYLCHOLINE RECEPTORS α5 SUBUNITS

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Objective: To determine the function role of neuronal nicotinic acetylcholine receptor (nAChR) α5 subunits in autonomic ganglia.

Background: In 12 distinct nAChR subunits (α2-α10, β2-β4), α5 subunits have unique properties in their sequences and their combinations with other subunits. The functional role of α5 subunits in autonomic ganglia are not well-known.

Design/Methods: Autonomic functions were measured in mice lacking α5 subunits (α5−/−) and wild-type (WT) control mice to characterize the properties of α5 subunits under physiological conditions and following pharmacological interventions.

Results: All α5−/− mice grew to normal size showing no obvious physical, neurological or autonomic deficit. Similitudes between α5−/− and WT mice: The rectal temperatures in ambient temperatures of 21 °C and cold exposure (6 °C); Pupil size at rest and following morphine (100mg/kg); Heart rate at rest, under anesthesia and stressed by cage shaking and cold exposure. Deficiency of α5 subunits strikingly increased the sensitivity to a low dose of hexamethonium (C5) leading to a nearly complete blockade of bradycardia in response to vagal stimulation as well as elimination of rebound post vagal-stimulation tachycardia. Such a dose of C5 only slightly depressed the effects of vagal stimulation in control mice. An impairment of cardiac parasympathetic ganglionar transmission was observed during high frequency cervical vagal stimulation. Another strikingly difference was that deficiency of α5 subunits significantly increased ileal contractile responses to choline and epinephrine (but not to dimethylphosphoriperezaminium iodide and nicotine).

Conclusion: α5 nAChR subunits are normally present in ANS ganglia, probably modulating postgangnic nAChR channels responses to endogenous ACh and regulating responses to ganglion drugs in receptor complexes. Such effects of α5 subunits may lower the safety factor in transmission systems.

α7 ACETYLCHOLINE RECEPTOR IN SCHIZOPHRENIA: DECREASED mRNA LEVELS IN PERIPHERAL BLOOD LYMPHOCYTES

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Central cholinergic systems are known to control basic functions of the brain. Recent studies have suggested that α7 nicotinic acetylcholine receptor (α7 AChR) may be associated with some aspects of schizophrenia. Reduced amounts of brain α7 AChR in schizophrenics as compared with healthy controls has been reported. In search for peripheral biological markers for schizophrenia that may enable measurable and rapid diagnosis of this disorder we have investigated α7 mRNA levels in peripheral blood lymphocytes (PBLs) of schizophrenic patients and healthy controls. Blood samples were collected from 34 medicated and unmedicated (drug naive) schizophrenic patients, and from 21 healthy smokers and nonsmokers. RNA was prepared from isolated lymphocytes and its amount and quality determined. PCR products, specific for human α7 AChR, were quantified by densitometry using Scion image analysis software. A significant decrease (20-95%) of α7 mRNA levels in PBLs of schizophrenic patients has been observed, compared with controls. The decrease in α7 mRNA levels was not a result of medication, since unmedicated patients showed the same range of reduction as hospitalised schizophrenic patients. In addition, we have ruled out the possibility that the observed decrease in α7 mRNA levels resulted from nicotine consumption by smoking, as healthy smokers exhibited the same levels of α7 mRNA as nonsmokers. We propose that α7 AChR mRNA may serve as a peripheral marker for schizophrenia. Moreover, if the observed changes in α7 mRNA levels in PBLs indeed reflect the state of this receptor in the brain, our findings support the assumption that a deficit in α7 AChR is involved in the pathophysiology of schizophrenia.
USE OF THE MORPHING GRAPHICS TECHNIQUE TO VISUALIZE CONFORMATIONAL DIFFERENCES BETWEEN ACHES FROM DIFFERENT SPECIES AND INHIBITOR-INDUCED CONFORMATIONAL CHANGES

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There are currently more than 25 AChE structures deposited in the Protein Data Bank, from four different species and/or complexed or conjugated with a repertoire of ligands. A method for sorting and characterizing differences between these structures is presented.

Pairs of AChE structures were aligned using LSQMAN, and rmsd values were calculated. Intermediate models between the two structures were produced by LSQMAN, in Cartesian space, by taking the initial and final coordinates, and interpolating the predicted intermediate coordinates. The intermediate models were then collated into a single QuickTime movie file easily viewable on most computers.

This morphing approach highlighted a conformational difference in loop 319-324 (hAChE numbering) between hAChE and TcAChE earlier reported by Kryger et al. (Acta Cryst. 2000) D56:1385). A similar conformational difference in the same loop between DmAChE and TcAChE was pinpointed utilizing the novel procedure.

A series of movies was compiled, comparing native TcAChE with its complexes and conjugates with a number of inhibitors. These reveal significant inhibitor-induced conformational changes at the top of the active-site gorge. A major conformational change was visualized for the conjugate of TcAChE with diisopropylphosphorofluoridate (DFP).

The simple morphing technique developed thus provides a valuable tool for locating and assessing conformational differences between closely related protein structures.
INDEX
<table>
<thead>
<tr>
<th>Name</th>
<th>Program Page</th>
<th>Abstract Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adams, D.J.</td>
<td>22</td>
<td>29</td>
</tr>
<tr>
<td>Adams, M.E.</td>
<td>12</td>
<td>4</td>
</tr>
<tr>
<td>Adani, R.</td>
<td>18</td>
<td>19</td>
</tr>
<tr>
<td>Ahdut, R.</td>
<td>13</td>
<td>6</td>
</tr>
<tr>
<td>Albuquerque, E.X.</td>
<td>16</td>
<td>13</td>
</tr>
<tr>
<td>Alewood, P.F.</td>
<td>22</td>
<td>29</td>
</tr>
<tr>
<td>Alkondon, M.</td>
<td>16</td>
<td>13</td>
</tr>
<tr>
<td>Allen, T.B.</td>
<td>31</td>
<td>55</td>
</tr>
<tr>
<td>Alvarez, A.</td>
<td>15</td>
<td>13</td>
</tr>
<tr>
<td>Amit, T.</td>
<td>18</td>
<td>18</td>
</tr>
<tr>
<td>Amitai, G.</td>
<td>18</td>
<td>19</td>
</tr>
<tr>
<td>Andrews, J.</td>
<td>21</td>
<td>23</td>
</tr>
<tr>
<td>Andrews, M.C.</td>
<td>25</td>
<td>38</td>
</tr>
<tr>
<td>Anglister, J.</td>
<td>11</td>
<td>3</td>
</tr>
<tr>
<td>Anglister, L.</td>
<td>12,21</td>
<td>5,25</td>
</tr>
<tr>
<td>Angus, L.M.</td>
<td>12</td>
<td>5</td>
</tr>
<tr>
<td>Antil-Delbeke, S.</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>Antollini, S.</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>Antonin, W.</td>
<td>13</td>
<td>20</td>
</tr>
<tr>
<td>Ariel, N.</td>
<td>14,24</td>
<td>8,9,34</td>
</tr>
<tr>
<td>Arpagus, M.</td>
<td>12</td>
<td>6</td>
</tr>
<tr>
<td>Ashani, Y.</td>
<td>14</td>
<td>10</td>
</tr>
<tr>
<td>Auld, V.J.</td>
<td>21</td>
<td>24</td>
</tr>
<tr>
<td>Aviv, E.</td>
<td>17</td>
<td>11</td>
</tr>
<tr>
<td>Avraham, Y.</td>
<td>30</td>
<td>51</td>
</tr>
<tr>
<td>Azeeva, E.A.</td>
<td>22</td>
<td>27</td>
</tr>
<tr>
<td>B</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Badet, B.</td>
<td>25</td>
<td>36</td>
</tr>
<tr>
<td>Baier, J.</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>Baker, S.R.</td>
<td>28</td>
<td>48</td>
</tr>
<tr>
<td>Balass, M.</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>Bancila, V.</td>
<td>26,31</td>
<td>41,56</td>
</tr>
<tr>
<td>Baptista, G.</td>
<td>22</td>
<td>30</td>
</tr>
<tr>
<td>Barak, D.</td>
<td>14</td>
<td>8,9,34</td>
</tr>
<tr>
<td>Barak, R.</td>
<td>24</td>
<td>34</td>
</tr>
<tr>
<td>Bar-Am, O.</td>
<td>18</td>
<td>18</td>
</tr>
<tr>
<td>Bar-Her, N.</td>
<td>16</td>
<td>15</td>
</tr>
<tr>
<td>Barnard, E.A.</td>
<td>12</td>
<td>4</td>
</tr>
<tr>
<td>Bar-On, P.</td>
<td>14,24</td>
<td>8,34</td>
</tr>
<tr>
<td>Barrantes, F.J.</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>Barril, X.</td>
<td>24</td>
<td>36</td>
</tr>
<tr>
<td>Bartolucci, C.</td>
<td>25</td>
<td>37</td>
</tr>
<tr>
<td>Bass, C.G.</td>
<td>25</td>
<td>38</td>
</tr>
<tr>
<td>Battaglia, M.</td>
<td>31</td>
<td>58</td>
</tr>
<tr>
<td>Bauer, B.</td>
<td>17</td>
<td>10</td>
</tr>
<tr>
<td>Beattie, R.E.</td>
<td>22</td>
<td>29</td>
</tr>
<tr>
<td>Bednar, I.</td>
<td>27</td>
<td>44</td>
</tr>
<tr>
<td>Bee, M.S.</td>
<td>18</td>
<td>19</td>
</tr>
<tr>
<td>Beeson, D.</td>
<td>28,31</td>
<td>46,58</td>
</tr>
<tr>
<td>Behra, M.</td>
<td>22</td>
<td>28</td>
</tr>
<tr>
<td>Bekpinar, S.</td>
<td>30</td>
<td>52</td>
</tr>
<tr>
<td>Belanger, G.</td>
<td>12</td>
<td>5</td>
</tr>
<tr>
<td>Bell, E.</td>
<td>22</td>
<td>29</td>
</tr>
<tr>
<td>Ben Chaim, Y.</td>
<td>27</td>
<td>43</td>
</tr>
<tr>
<td>Name</td>
<td>Program Page</td>
<td>Abstract Page</td>
</tr>
<tr>
<td>-----------------------</td>
<td>--------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Ben Shushan, D.</td>
<td>30</td>
<td>51</td>
</tr>
<tr>
<td>Beni, S.M.</td>
<td>16</td>
<td>15</td>
</tr>
<tr>
<td>Berkovic, S.F.</td>
<td>21</td>
<td>26</td>
</tr>
<tr>
<td>Berman, H.A.</td>
<td>24,25</td>
<td>33,39</td>
</tr>
<tr>
<td>Bermudez, I.</td>
<td>23</td>
<td>31</td>
</tr>
<tr>
<td>Bernard, V.</td>
<td>28</td>
<td>49</td>
</tr>
<tr>
<td>Berry, E.M.</td>
<td>30</td>
<td>51</td>
</tr>
<tr>
<td>Berse, B.</td>
<td>13, 27</td>
<td>17,42</td>
</tr>
<tr>
<td>Bertrand, C.</td>
<td>22</td>
<td>28</td>
</tr>
<tr>
<td>Bertrand, D.C.</td>
<td>11, 21, 22, 31</td>
<td>3, 24, 26, 27, 29, 57</td>
</tr>
<tr>
<td>Bertrand, S.</td>
<td>21, 22</td>
<td>24, 26, 29</td>
</tr>
<tr>
<td>Bezakova, G.</td>
<td>12</td>
<td>3</td>
</tr>
<tr>
<td>Birikh, K.R.</td>
<td>27</td>
<td>45</td>
</tr>
<tr>
<td>Bixel, G.M.</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>Blank, P.</td>
<td>13</td>
<td>6</td>
</tr>
<tr>
<td>Blanton, M.</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>Bledi, Y.</td>
<td>12</td>
<td>7</td>
</tr>
<tr>
<td>Bloc, A.</td>
<td>25, 26, 31</td>
<td>37, 41, 56</td>
</tr>
<tr>
<td>Bloch, B.</td>
<td>28</td>
<td>49</td>
</tr>
<tr>
<td>Blum, B.</td>
<td>30</td>
<td>50</td>
</tr>
<tr>
<td>Blusztajn, J.K.</td>
<td>13, 27</td>
<td>17, 42</td>
</tr>
<tr>
<td>Bogoch, Y.</td>
<td>12</td>
<td>7</td>
</tr>
<tr>
<td>Bon, C.</td>
<td>25</td>
<td>37</td>
</tr>
<tr>
<td>Bon, S.</td>
<td>21</td>
<td>25</td>
</tr>
<tr>
<td>Boneva, N.</td>
<td>13</td>
<td>45</td>
</tr>
<tr>
<td>Bonini, I.</td>
<td>11</td>
<td>2</td>
</tr>
<tr>
<td>Bons, N.</td>
<td>16</td>
<td>15</td>
</tr>
<tr>
<td>Boot, J.R.</td>
<td>28</td>
<td>48</td>
</tr>
<tr>
<td>Bose, S.</td>
<td>22, 31</td>
<td>29, 58</td>
</tr>
<tr>
<td>Boss, A.</td>
<td>30</td>
<td>53</td>
</tr>
<tr>
<td>Botti, S.</td>
<td>21</td>
<td>24</td>
</tr>
<tr>
<td>Bourne, Y.</td>
<td>22, 25, 26</td>
<td>26, 39, 41</td>
</tr>
<tr>
<td>Boyd, A.</td>
<td>25</td>
<td>36</td>
</tr>
<tr>
<td>Brana, C.</td>
<td>28</td>
<td>49</td>
</tr>
<tr>
<td>Brandeis, R.</td>
<td>16</td>
<td>15</td>
</tr>
<tr>
<td>Brejc, K.</td>
<td>11</td>
<td>1</td>
</tr>
<tr>
<td>Brenner, T.</td>
<td>27</td>
<td>45</td>
</tr>
<tr>
<td>Brewer, C.J.</td>
<td>31</td>
<td>57</td>
</tr>
<tr>
<td>Brimijoin, S.</td>
<td>15</td>
<td>13</td>
</tr>
<tr>
<td>Broad, L.M.</td>
<td>16, 22</td>
<td>14, 29</td>
</tr>
<tr>
<td>Brochier, L.</td>
<td>25</td>
<td>38</td>
</tr>
<tr>
<td>Bronfman, M.</td>
<td>15</td>
<td>13</td>
</tr>
<tr>
<td>Broomfield, C.A.</td>
<td>27</td>
<td>43</td>
</tr>
<tr>
<td>Brovtsyna, N.B.</td>
<td>24</td>
<td>32</td>
</tr>
<tr>
<td>Brownlow, S.</td>
<td>28</td>
<td>46</td>
</tr>
<tr>
<td>Brydson, M.</td>
<td>28</td>
<td>46</td>
</tr>
<tr>
<td>Bueters, T.J.H.</td>
<td>14, 27</td>
<td>10, 42</td>
</tr>
<tr>
<td>Bunc, M.</td>
<td>25</td>
<td>39</td>
</tr>
<tr>
<td>Bymaster, F.P.</td>
<td>17, 18, 28</td>
<td>16, 19, 47</td>
</tr>
<tr>
<td>C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Caccin, P.</td>
<td>27</td>
<td>45</td>
</tr>
<tr>
<td>Cambi, F.</td>
<td>21</td>
<td>23</td>
</tr>
<tr>
<td>Camp, S.</td>
<td>13</td>
<td>6</td>
</tr>
<tr>
<td>Camps, P.</td>
<td>24</td>
<td>36</td>
</tr>
<tr>
<td>Cangioili, J.</td>
<td>17</td>
<td>16</td>
</tr>
<tr>
<td>Capsoni, S.</td>
<td>31</td>
<td>55, 56</td>
</tr>
<tr>
<td>Name</td>
<td>Program Page</td>
<td>Abstract Page</td>
</tr>
<tr>
<td>---------------------</td>
<td>--------------</td>
<td>---------------</td>
</tr>
<tr>
<td>Carlier, P.R.</td>
<td>24</td>
<td>35</td>
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