Neuronal nicotinic acetylcholine receptors (nAChRs) expressed in the brain are known to be important for cognition, learning and memory, and their deficiencies are shown to play a crucial role in Alzheimer’s disease (AD) pathogenesis. Neuronal nAChRs consist of various combinations of α2-α10 and β2-β4 subunits. The most abundant subtypes of nAChRs in the central nervous system are α4β2 and α7, whereas α3β4 predominates in the periphery. Administration of nAChR agonists with high affinity to the α4β2 nAChR has been proposed as one of the approaches for the treatment of AD. Further, the activation of α7 nAChRs has been recently shown to exhibit a neuroprotective action. The natural alkaloid epibatidine is known to possess a high affinity but lack of selectivity towards central neuronal nAChRs. Three novel analogs of epibatidine with -Cl, -F or -NH2 substitutions at the 3’ position of the pyridine ring, that have been recently developed and found to possess high binding affinity to brain nAChRs, were proposed to be tested: a) in vitro for their functional activity and nAChR subtype selectivity; and b) in vivo for a memory enhancement effect. The in vitro patch-clamp experiments demonstrated that, compared to the two other tested analogs, 3’-fluoro substitution in the epibatidine pyridine ring results in an analog with the most effectively increased efficacy and improved selectivity for α4β2 versus α3β4 nAChRs, while retaining an agonist effect on α7 nAChRs. These findings suggest that 3’-fluoro analog of epibatidine may serve as a novel candidate for a treatment of AD due to its potential memory enhancement, neuroprotection and minimized peripheral side effects. The in vivo studies were still in progress due to a temporary failure of positive nootropic control compounds to decrease the number of errors in the radial arm maze test. Continued work aims to replicate the nootropic effects of donepezil and rimonabant in the radial arm maze test, in order to proceed with evaluation of the novel 3’-fluoroepibatidine analog for memory enhancement. (Dr. Abdrakhmanova can be reached at 804/828-1797)

Shenandoah Mary A. Corcoran, Ph.D., OTR (Div. of Occupational Therapy, School of Health University Professions) “Caregiving Styles of Adult Children Who Provide Dementia Care” Thirty one individuals who provide care for a parent or similarly related person with dementia participated in this qualitative study of caregiving styles. Each participant was interviewed on three occasions (for an average of 55 minutes per occasion) and completed a questionnaire to gather information about sociodemographic characteristics and well-being. With regard to the elements of caregiving style (beliefs, meanings, and actions), filial caregivers reported a consistent set of beliefs about the nature, causes, and progression of dementia and the definition of an ideal caregiver (although most would not claim to embody that definition). Meanings associated with caring for a parent included priorities for care (trying to avoid future regrets, paying respects to an honored parent, and fulfilling commitments), costs, conflicts, self-image, and change. Actions included interacting with the parent (i.e., communication, managing medical routines, being vigilant), managing the system and environment (i.e., interacting with the staff at an assisted living facility or keeping things organized), and managing self and non-parental responsibilities (i.e., work duties and children). Turning to overall style, it was found that the context of care is an important factor in determining style, with the presence of other involved family members and living arrangement shaping patterns in thinking and action. Three caregiving styles have emerged 1) Informing – collecting and dispensing information about the parent and from the literature to influence the care decisions of others; 2) Arranging – juggling multiple roles and schedules including caregiving; and, 3) Monitoring and Managing – being vigilant about the health of the parent and acting on his/her behalf with formal care providers. (Dr. Corcoran can be reached at 540/665-5563)
Alzheimer’s disease has long been known to involve formation of fibrillar structures from a protein fragment termed amyloid-β. More recently, the interactions between this protein fragment and cell membranes have been implicated as critical aspects to the neuronal damage in Alzheimer’s patients. This research demonstrated that a peptide mimic of the amyloid-β peptide can exhibit many of the critical features of Aβ behavior, including self-association, binding to membranes, and acceleration of self-association by membranes. Particularly important, the mimic is also toxic to neurons. Further, like Aβ it shows the trend that intermediate concentrations of the peptide are most toxic. This suggests that at least some aspects of the disease may be valuably studied using such peptide mimics. Finally, the investigators have also studied the effect of some recently discovered molecules that manipulate the aggregation of amyloid-β. They have been able to distinguish the effects of these molecules on peptide association vs. membrane binding. The results may have implications for the design of new therapeutic molecules that can prevent the toxic interactions of amyloid-β with membranes. (Dr. Fernandez can be reached at 434 924-1351)

Alzheimer’s disease is related, in part, to a deficiency in the neurotransmitter acetylcholine in relevant brain areas. Acetylcholine activates several types of brain receptors, and one current treatment modality is to prevent the degradation of acetylcholine by agents that block its metabolism (i.e., cholinesterase inhibitors). This “shotgun” approach can lead to undesirable side effects. Another approach would be to activate selected acetylcholine receptors using a novel agent. There are growing implications for the involvement of the nicotinic acetylcholine (nACh) receptor type. Unfortunately, there are multiple subtypes of these receptors making it difficult to specifically target the particular receptor subtype of interest. A natural product, desformylflustrabromine (dFBr), isolated in small quantities from a marine organism, was found to potentiate the effects of ACh. But, it does so through a unique mechanism that does not involve direct receptor activation (i.e., it is a positive allosteric modulator). Being the first member of a novel mechanistic type of agent that selectively activates the actions of ACh at the target nACh receptor subtype of interest (i.e., α4β2 nACh receptors), it offered a new target for exploitation. The purpose of this work was to a) synthesize a sufficient quantity of dFBr as a water-soluble salt for pharmacological study, and b) identify which structural features are important for activity. The first goal was achieved, and structural features important for the potentiating action were identified. NIH funding is now being sought in order to utilize the information obtained so that activity might be optimized. (Dr. Glennon can be reached at 804/828-8487)

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