Dementia

Gary W. Small, M.D. Parlow-Solomon Professor of Aging UCLA School of Medicine Professor of Psychiatry & Biobehavioral Sciences Director, UCLA Center on Aging

James M. Ellison, M.D., MPH

Clinical Director, Geriatric Psychiatry Program McLean Hospital Associate Professor of Psychiatry, Harvard Medical School

<u>Self-Assessment Question 1</u> <u>Which of the following</u>

are required for a diagnosis of dementia?

- A. Cognitive decline is ACQUIRED
- B. MEMORY is affected
- C. In addition to memory, ANOTHER cognitive function is affected.
- D. Symptoms are not attributable to delirium or another psychiatric disorder.
- E. All of the above

Self-Assessment Question 2 Which of the following cognitive or behavioral domains is/are affected in dementia?

- A. Memory
- B. Executive function
- C. Behavior
- D. Activities of daily living
- E. All of the above

Self-Assessment Question 3 Which of the following statements is correct?

- A. Alzheimer's Disease affects greater than 30% of adults older than 85 years of age.
- B. Alzheimer's Disease is infrequent among adults less than 60 years of age.
- C. Alzheimer's Disease is the most common cause of dementia.
- D. All of the above
- E. None of the above

Self-Assessment Question 4 Treatment of AD with cholinesterase inhibitors is based on which of these rationales?

- A. Noradrenergic neurotransmission in the locus ceruleus is reduced in late AD
- B. Pathological stimulation of NMDA receptors is associated with excitotoxic death of neurons.
- C. The number of cholinergic neurons in the basal forebrain is reduced in late Alzheimer's disease.
- D. All of the above
- E. None of the above

Self-Assessment Question 5 Treatment of AD with memantine is based on which of these rationales?

- A. Noradrenergic neurotransmission in the locus ceruleus is reduced in late AD
- B. Pathological stimulation of NMDA receptors is associated with excitotoxic death of neurons.
- C. The number of cholinergic neurons in the basal forebrain is reduced in late Alzheimer's disease.
- D. All of the above
- E. None of the above

Major Points

- Dementia is underrecognized and undertreated in primary care and in mental health settings
- Dementia can be recognized and treated beneficially in primary care and mental health settings
- Both pharmacological and nonpharmacological interventions may benefit overall brain health and dementia course

Major Points (cont.)

- Neuroimaging with PET can show a pattern of regional glucose metabolism that improves early detection of Alzheimer's disease with greater specificity
- Novel approaches to <u>in vivo</u> plaque and tangle imaging will be useful in monitoring potential disease-modifying agents

Definition of Dementia

Acquired syndrome of decline in memory and at least 1 other cognitive function (e.g., language) sufficient to affect daily life, not explainable by delirium or other mental disorder.

Causes of Dementia Symptoms

Alzheimer's disease Vascular disease Lewy Body Disease Parkinson's Huntington's Frontotemporal dementias Head Injury Metabolic/Nutritional ♦B₁₂/Folate *****Thiamine Thyroid

Hepatic/Renal

Medications Alcohol/Toxins Infectious *HIV *Syphilis Meningitis Depression *NPH Neoplasms Autoimmune disorders

Diagnostic Criteria for Alzheimer's Disease (1)

Multiple cognitive deficits manifested by both of:

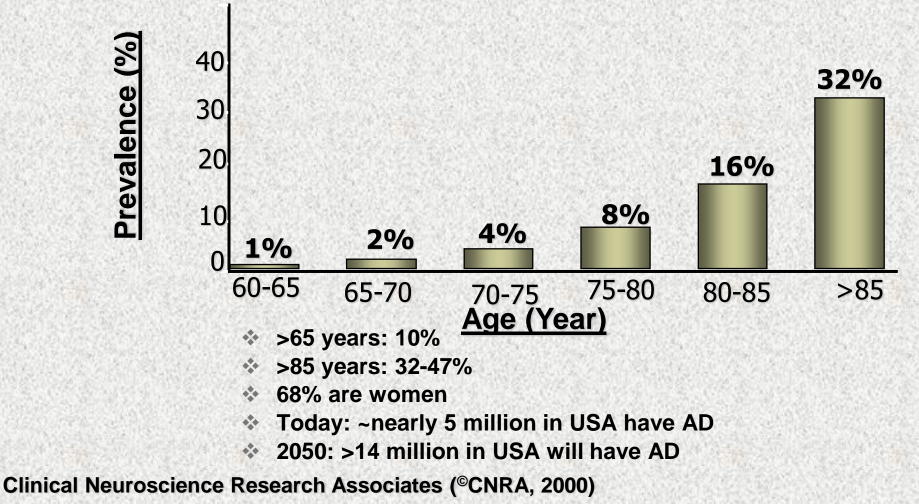
- Memory impairment
- One (or more) of the following cognitive disturbances: aphasia, apraxia, agnosia, disturbance in executive functioning
- Significant impairment in social or occupational functioning representing a significant decline from a previous level of functioning
 - Gradual onset and progressive cognitive decline

Diagnostic Criteria for Alzheimer's Disease (2)

Cognitive deficits are NOT due to any of the following:

- Other central nervous system conditions that cause progressive deficits in memory and cognition
- Systemic conditions known to cause dementia
- Substance-induced conditions
- Deficits do not occur exclusively during delirium
- Disturbance is not better accounted for by another Axis I disorder

Prevalence of Alzheimer's Disease in the U.S.



www.therubins.com.

Diagnosing AD: physical examination

- * Life-threatening conditions, e.g. mass lesions, vascular lesions and infections
- * Blood pressure and pulse
- * Vision and hearing assessments
- * Cardiac and respiratory function
- * Mobility and balance
- * Sensory and motor system examination (tone, reflexes, gait and coordination) and depressive symptoms (sleep and weight)

Physical examination

Diagnosing AD: laboratory tests

All patients

- * Complete blood count
- * Thyroid function
- * B12, folate, fasting homocysteine
- * BUN and creatinine
- * Calcium
- * Glucose
- * Electrolytes
- * Urinalysis
- * Liver function tests
- * Fasting lipid profile
- * ESR

Most patients

* ECG

Many patients

* Neuropsychological testing* Neuroimaging

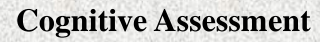
Some patients

*Specialized medical labs *RPR *CXR *LP

*ApoE genotype

Diagnosing AD: cognitive assessment with MMSE

	Score	Score
	Maximum	Actual
Cognitive area		
Mini Mental State Examination: test outline and scoring		
Orientation	k. Alexandra dia.	
*What is the (date, day, month, year, season)?	5	
* Where are you (clinic, town, country)?	5	
Memory		
*Name three objects. Ask the patient to repeat them	3	
Attention		
*Serial sevens. Alternatively ask the patient to spell world	5	
backwards (dlrow)		



Folstein et al 19756

Diagnosing AD: cognitive assessment with MMSE (2)

	Score	Score
	Maximum	Actual
Cognitive area		
Mini Mental State Examination: test outline and scoring		
Recall in the second se		
*Ask for the three objects mentioned above to be repeated	3	
Language		
*Name a pencil and watch	2	
*Repeat, 'No ifs, ands or buts'	1	
*A three stage command	3	
*Read and obey CLOSE YOUR EYES	1	
*Write a sentence	1	
*Copy a double pentagon	1	
	Total 30	
Cognitive Assessment	Folstein	et al 1975

Diagnosing AD: Neuropsychological Assessment¹

- No current "gold standard" single test identified
- Battery of tests improves sensitivity and specificity
- Typical neuropsychological battery:
 - RAVLT or CAVLT
 - WAIS-R
 - WMS-R
 - Rey-Osterrieth Complex Figure
 - Clock drawing / Trails
 - Mattis Dementia Rating Scale
- Predictive value of objective informant report is high
- 1. Petersen et al. 2001 and other sources

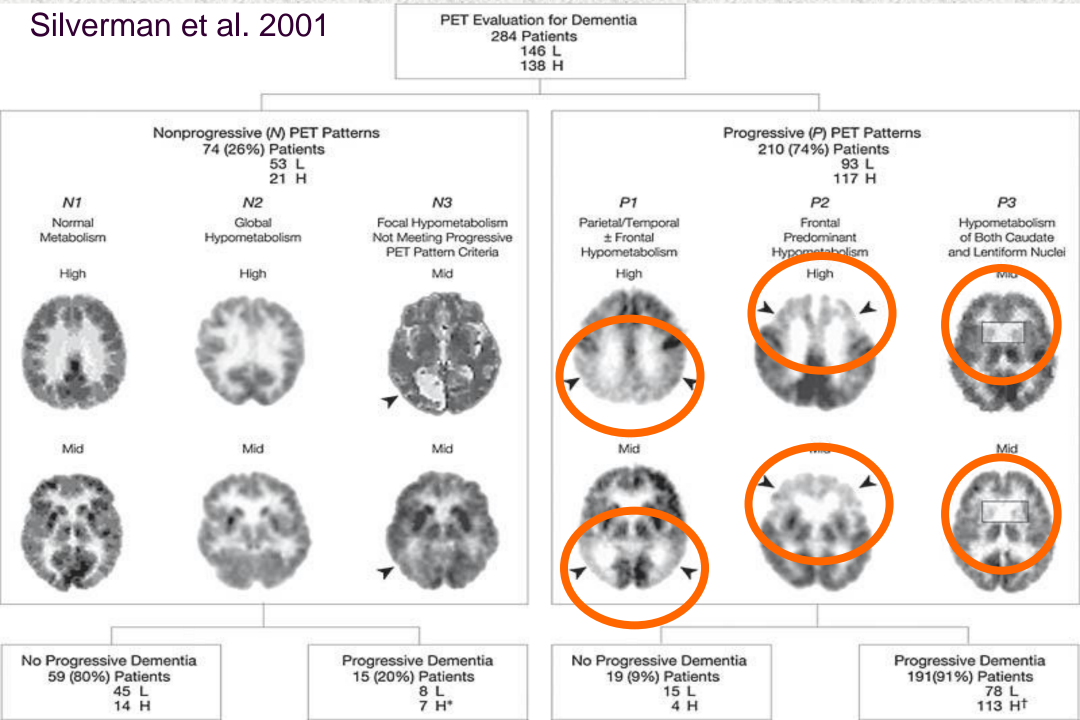
Diagnosing AD: Neuroimaging (Structural/Functional)



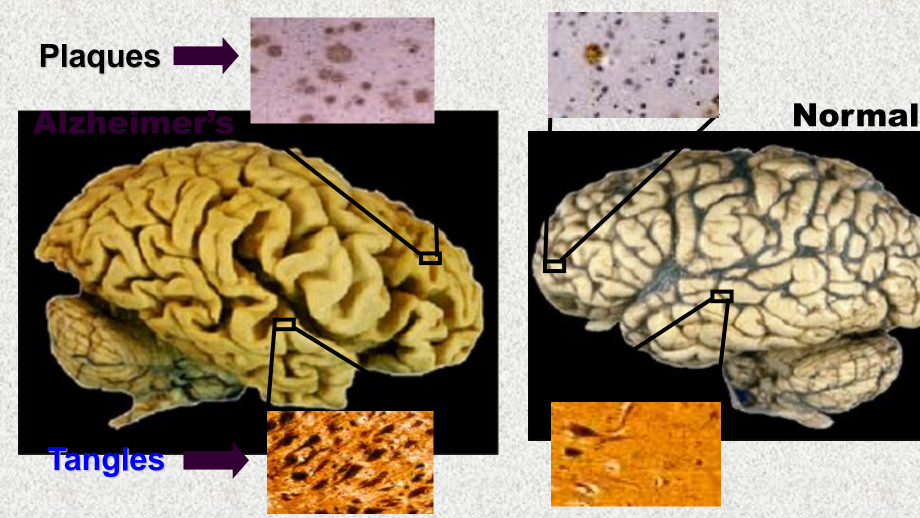




Reproduced from Doraiswamy PM, 1998



Amyloid Plaques and Neurofibrillary Tangles in Alzheimer's Disease and Normal Aging



Courtesy of Harry Vinters, M.D.

AAMI vs. MCI

- AAMI: defined vs younger controls; mild impairment, 1%/yr progress to dementia
- MCI: memory complaint, objective memory impairment on neuropsychological testing, often corroborated by observer. Called "amnestic" when only memory affected, "other" when memory not affected, subclassified as "single domain" vs "multiple domain". Estimated 10-15%/yr progress to dementia.

Risk/Protective Factors for Brain

Aging

Definite risks

- * Age
- Family history
- ApoEε4 genotype
- Other specific genotypes

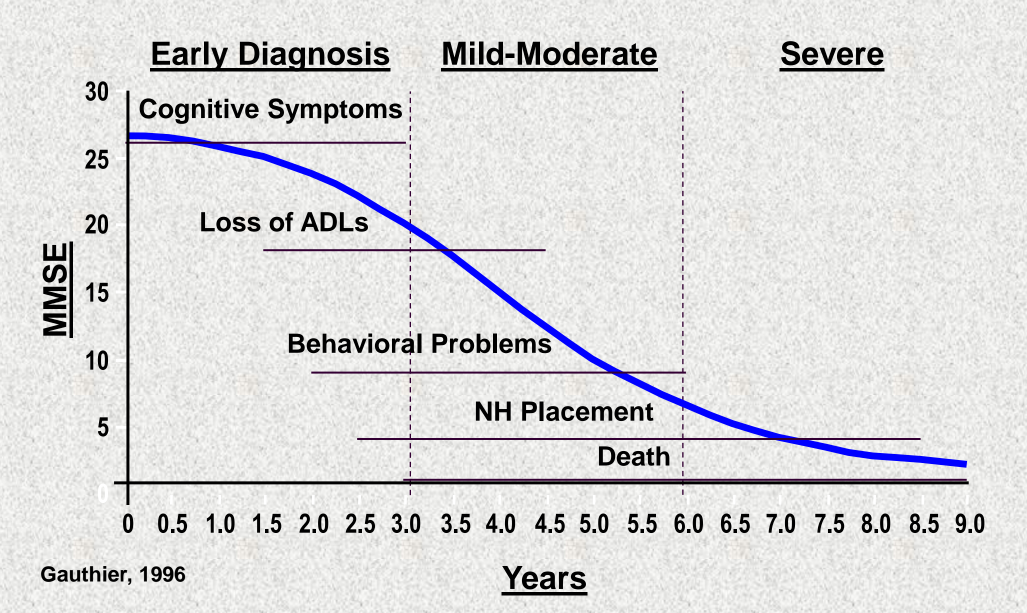
Probable/Possible risks

- Head trauma
- * Diabetes
- Hypertension
- Lower educational achievement
- Chronic stress

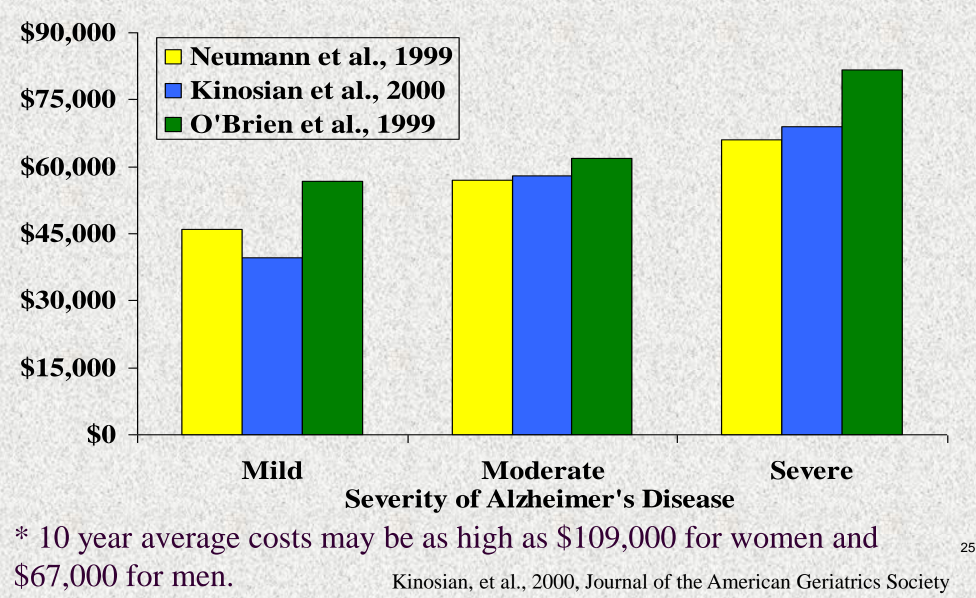
Possible protections

- Low-fat diet
- Aerobic exercise
- Cognitive stimulation
- * Wine
- Antioxidants
- Anti-inflammatory drugs

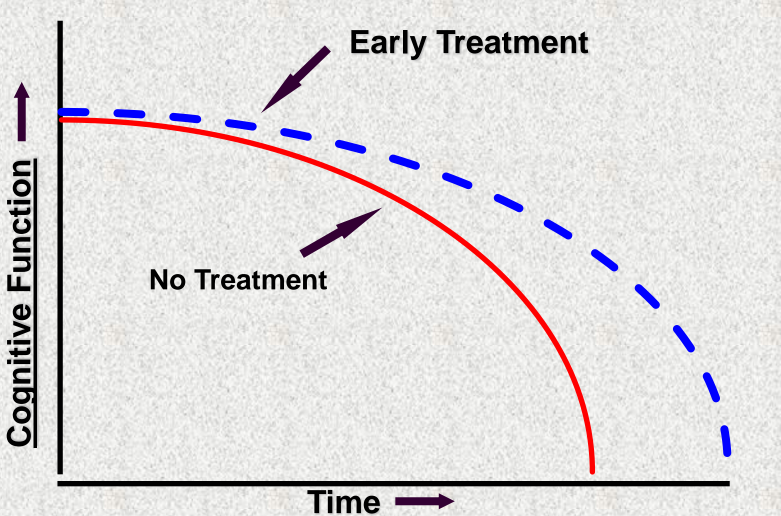
The Progress of Alzheimer's Disease



5 Year Cost of Care Models for Mild, Moderate, and Severe Alzheimer's Disease (as of 2000)



Delaying Onset Reduces Prevalence/Costs



Current Challenges in Dementia Diagnosis

Primary care physicians (PCPs)

- PCPs care for most dementia patients (64%)
- Sarrett et al, 1997 found that only 40% of PCPs knew Alzheimer's was most common cause of late-life memory loss (vs. 97% of experts)
- * PCPs usually do not use standardized dementia diagnostic criteria

Missed diagnosis*

- >75% of patients with moderate dementia
- 97% of patients with mild dementia

Underrecognition of dementia leads to increased:

- Motor vehicle accidents
- ER visits
- Hospitalization rates
- Medication errors
- * Mortality

*Callahan et al, 1995

When to Refer

Geriatric Psychiatrist

*Unclear dx, early onset, severe behavior/mood problem, non-responsive to tx, unable to tolerate drugs, caregiver stress

Neurologist

Parkinson sx, early onset, focal neurological signs, rapid progression, atypical presentation

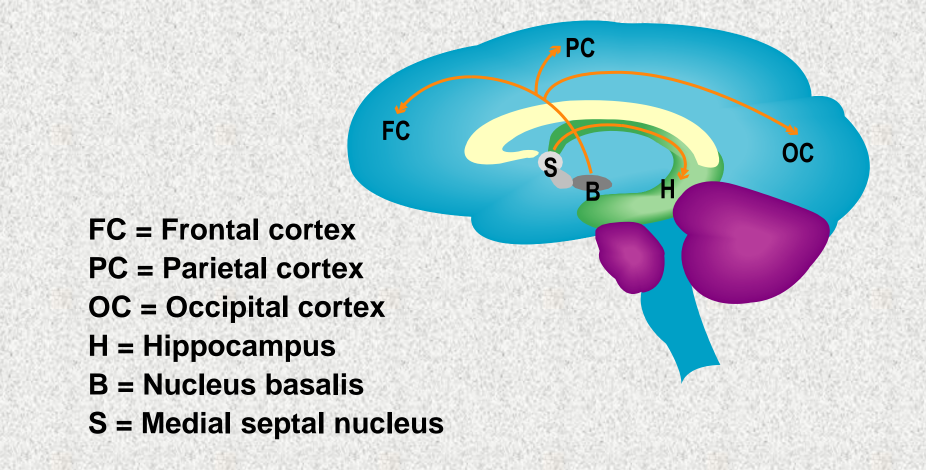
Geriatrician

Complex medical problems, functional assessment

When to Hospitalize

- Imminent danger to self or others
- **Severe mood problems, agitation, psychosis**
- Refusal to eat, severe sleep disturbance
- *Psychiatric illness complicated by alcohol/drug addiction
- Need for drugs or tests requiring hospitalization
 Need for IV or frequent IM injections

Cholinergic System Innervates Areas Associated with Memory and Learning



Adapted from: Coyle JT et al. Science. 1983;219:1184-1190

Rationale for Cholinergic Treatments of AD

- Cholinergic function including choline acetyltransferase (CAT) activity is reduced with aging¹
- Number of cholinergic neurons (particularly in basal forebrain) is reduced in late AD²
- In AD, nicotinic receptors in hippocampus and cortex are reduced^{1,3}

¹Bartus RT et al. Science. 1982;217:408-414; ²Whitehouse PJ et al. Science. 1982;215:1237-1239; ³Guan ZZ et al. J Neurochem. 2000;74:237-243 (from Small G: Dementia. ACNP Curriculum)

Cholinesterase Inhibitor Properties

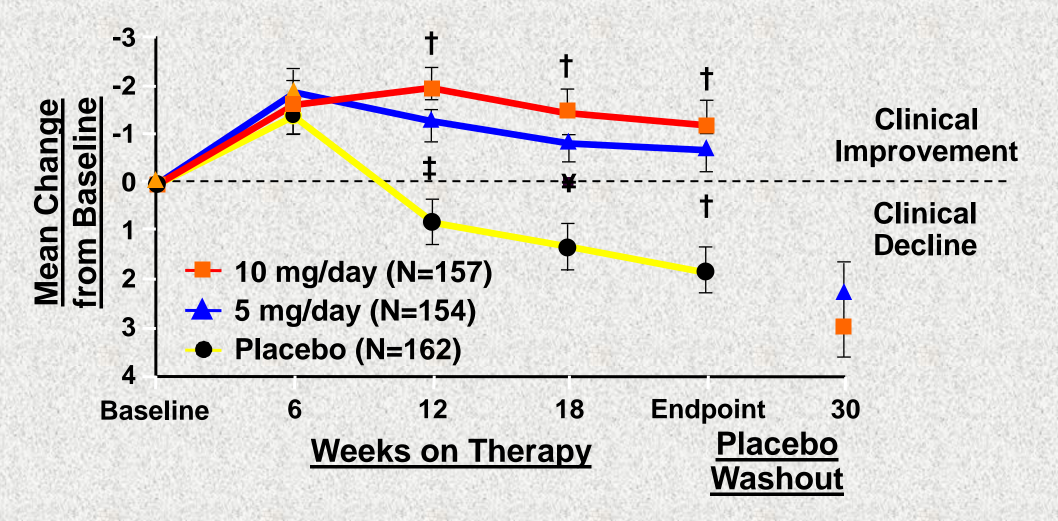
	Selectivity	Max serum conc	Absorbtion delay by food?	Serum Half-Life (hr)	Protein Binding %	Target Dose (mg/day)	Daily Dosing
Tacrine (Cognex)	AChE &BuChE	1-2 hr	Yes	1.3-2	75	80-160	qid
Donepezil (Aricept)	AChE	3-5 hr	No	70-80	96	5-10	qd
Rivastigmine (Exelon)	AChE &BuChE	0.5-2 hr	Yes	2	40	6-12	bid
Rivastigmine Transdermal (Exelon Patch)	AChE &BuChE		No	2	40	4.6 or 9.5 mg patch once daily	
Galantamine (Razadyne or Razadyne ER or generic galantamine)	AChE & Nic Mod	30-60 min	Yes	5-7 ER is longer	10-20	16-24	bid or qd

<u>Most Frequent Adverse Effects of</u> <u>Cholinesterase Inhibitors*</u>

Nausea	13% to 35%			
Anorexia	5% to 14%			
Dizziness	1% to 10%			
Diarrhea	0% to 11%			
Cost	>\$150/mo for branded drugs			

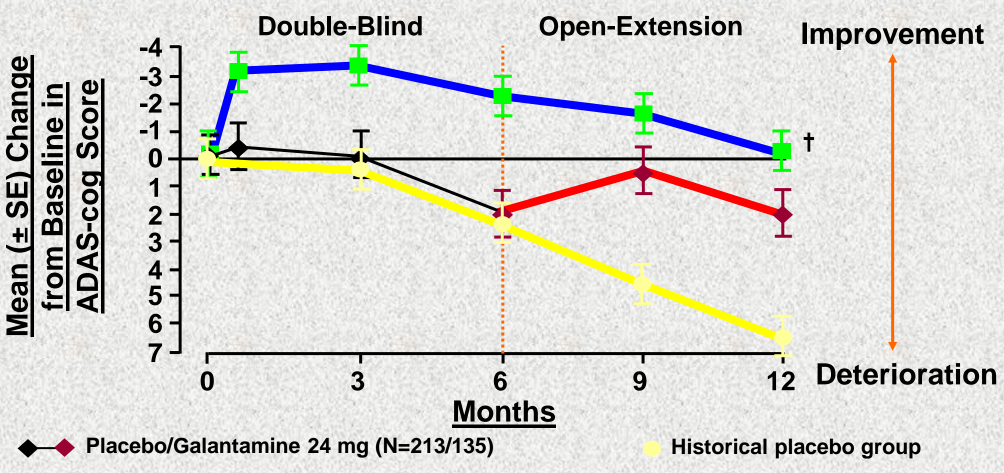
*These numbers are taken from package inserts for Chl's.

Effect of Donepezil on Cognition: ADAS-Cog*



*Alzheimer's Disease Assessment Scale-Cognitive Subscale. [†]p<0.0001; [‡]p<0.0007; [¥]p<0.0012 Rogers SL et al. Neurology. 1998;50:136-145

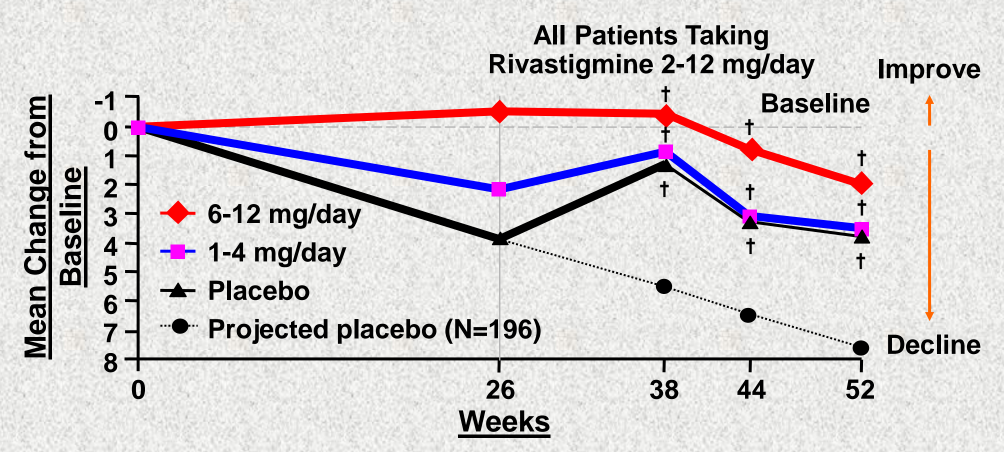
Effect of Galantamine on Cognition: ADAS-Cog*



Galantamine 24 mg/Galantamine 24 mg (N=212/116)

*ADAS-Cog = Alzheimer's Disease Assessment Scale-Cognitive Subscale; [†]p<0.05 vs. placebo/ Galantamine and not statistically different from baseline; Raskind MA et al. Neurology. 2000;54:2261-2268

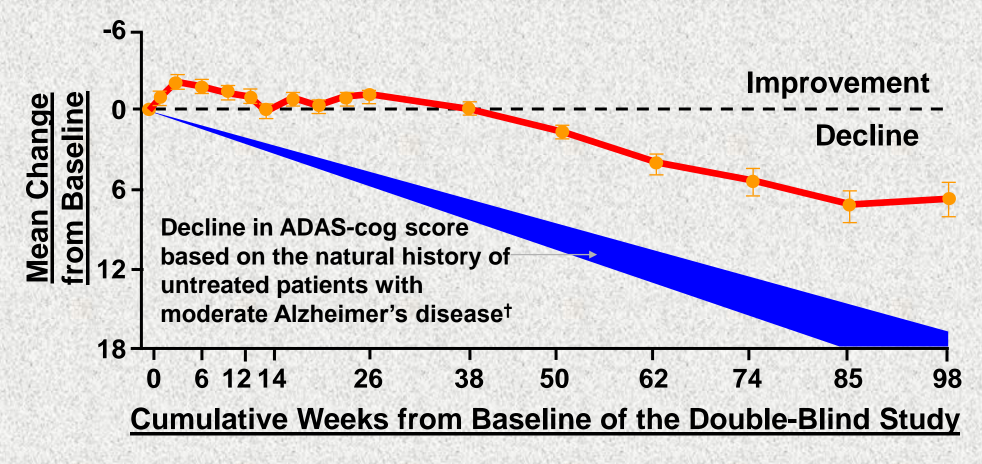
Efficacy of Rivastigmine on Cognition Through 52 Weeks: ADAS-Cog*



*Alzheimer's Disease Assessment Scale–Cognitive Subscale; [†]p<0.05 vs. projected placebo; Corey-Bloom J et al. Int J Geriatr Psychopharmacol. 1998;1:55-65; Adapted from: Messina J et al.

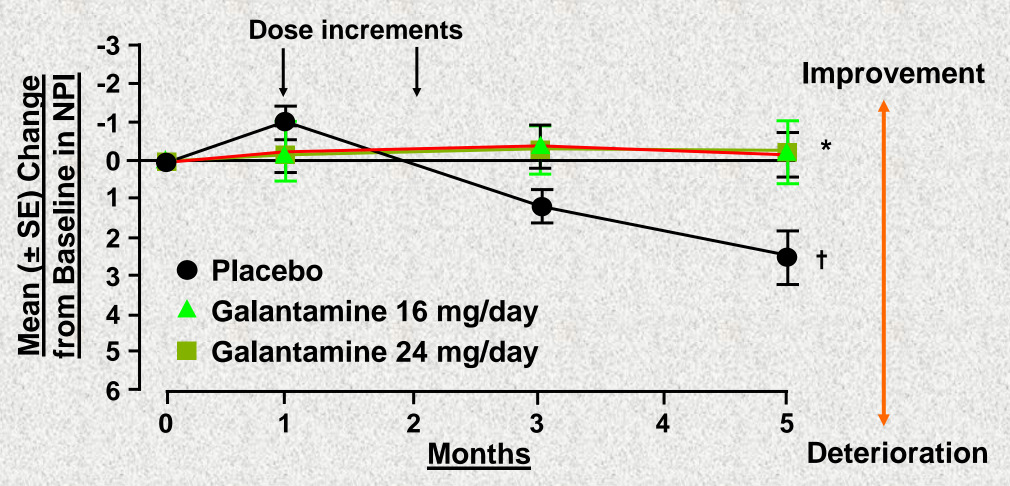
The 3rd Int'l Meeting for the College of Psychiatric and Neurologic Pharmacists. April 6-9, 2000. Washington, DC; Novartis Pharmaceuticals Corporation (Data on file)

Long-Term Effects of Donepezil on Cognition: ADAS-Cog*



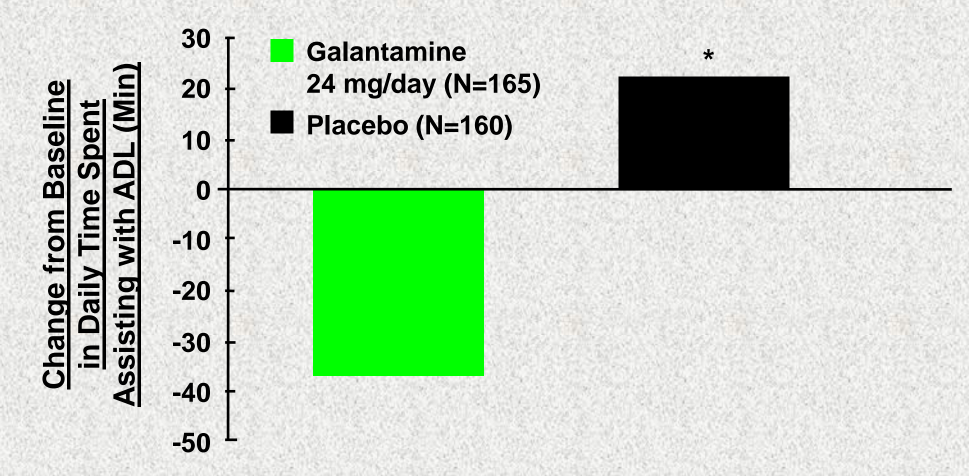
*Alzheimer's Disease Assessment Scale-Cognitive Subscale; Rogers SL, Friedhoff LT. Eur Neuropsychopharmacol. 1998;8:67-75; [†]Stern RG et al. Am J Psychiatry. 1994;151:390-396

Effect of Galantamine on Behavioral Symptoms: NPI



*p<0.05 vs. placebo (galantamine 16 mg and 24 mg); [†]p<0.05 vs. baseline; Adapted from: Tariot PN et al. Neurology. 2000;54:2269-2276

Change in Daily Time Spent by Caregiver Assisting with ADL



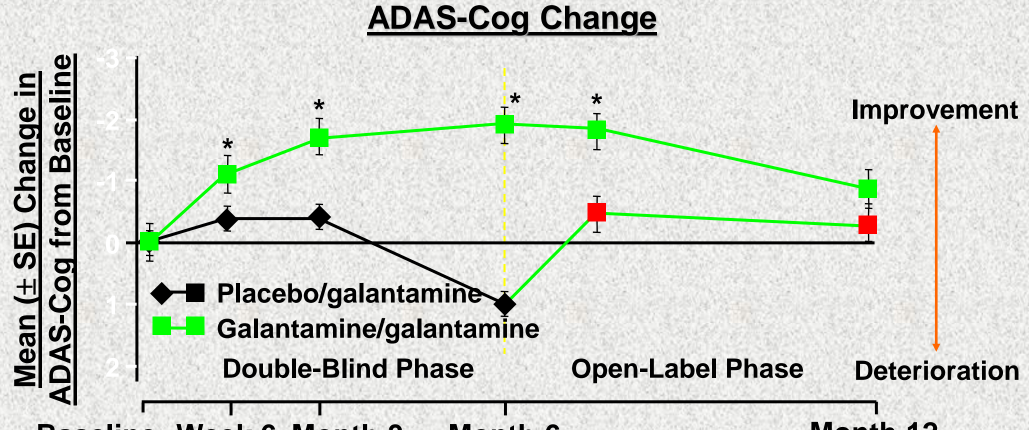
*p<0.05 vs. baseline; Lilienfeld S, Parys W. Dement Geriatric Cog Disord. 2000;11(suppl 1):19-27; Wilcock G et al. World Alzheimer Congress, 2000

Donepezil and Concomitant Treatments

Percentage of Caregivers Reporting Patient's Drug Use, by Category		
Drug Category	Donepezil (%) (N=108)	Nondonepezil (%) (N=268)
Antidepressant	25	43*
Antipsychotic	19	34*
Antianxiety	13	22†
Estrogen	10	* 8
Antiparkinsonian	5	7
Sedative-hypnotic	2	6†

*Pearson chi-square test, p<0.05; [†]Pearson chi-square test, p<0.10; Small GW et al. Clin Ther. 1998;20:838-850

<u>Galantamine in the Study of Alzheimer's</u> Disease, Vascular Dementia or Mixed Dementia



Baseline Week 6 Month 3 Month 6

Month 12

*p<0.001 vs. placebo; Erkinjuntti T et al. Lancet. 2002;359:1283-1290; Janssen Pharmaceutica Products, L.P. (Data on file)

AD2000 Study (UK)

Randomized Clinical Trial of Donepezil / Placebo in 566 AD patients followed for up to 4 years

Cognition (MMSE) and ADLs improved by donepezil over the first 2 years

No significant benefits in risk of institutionalization or progression of disability

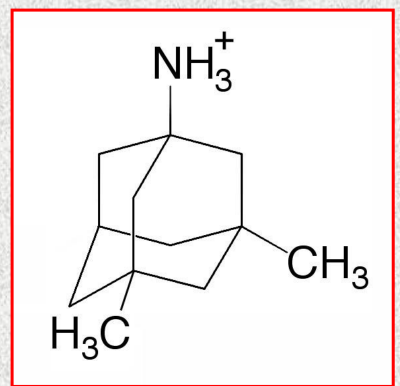
Conclusion: "Donepezil is not cost effective, with benefits below minimally relevant thresholds."

AD2000 Study - Limitations

- •Underpowered: 566 / 3,000 subjects enrolled
- •High attrition: 48% after 1 year, >80% after 2 years
- Lack of rigorous diagnostic criteria
- Repeated drug wash-out periods

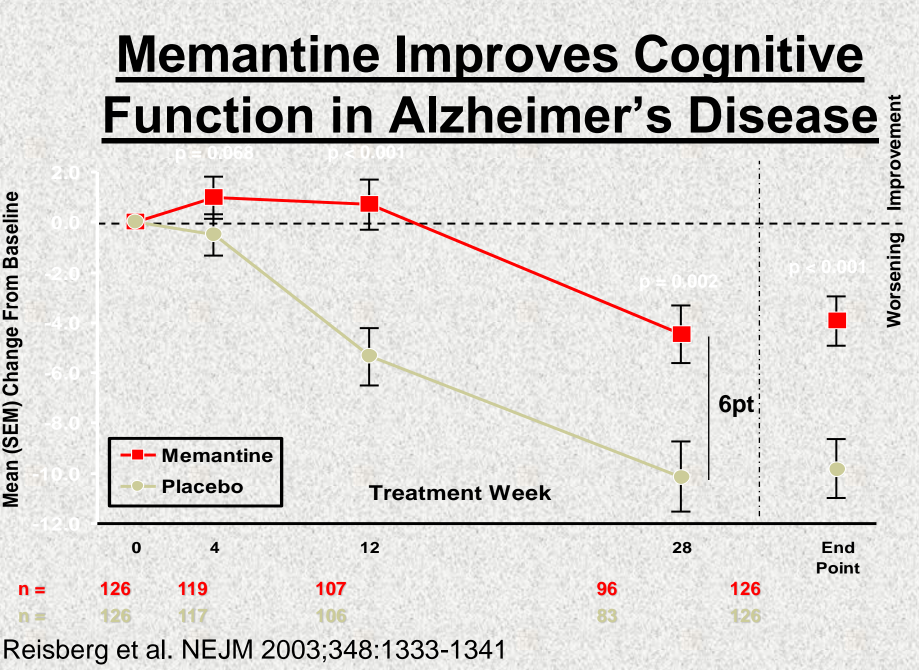
Pharmacology of Memantine

- Aminoadamantane derivative (1-amino-3,5-dimethyladamantane)
- NMDA receptor uncompetitive (open channel) antagonist low/moderate affinity
- 5-HT3 receptor allosteric antagonist of low/moderate affinity
- Binds with lower affinity to human nicotinic acetylcholine receptors – but may not be clinically relevant (does not alter AChE activity in the presence or absence of AChEls)



Memantine: Clinical Pharmacokinetics

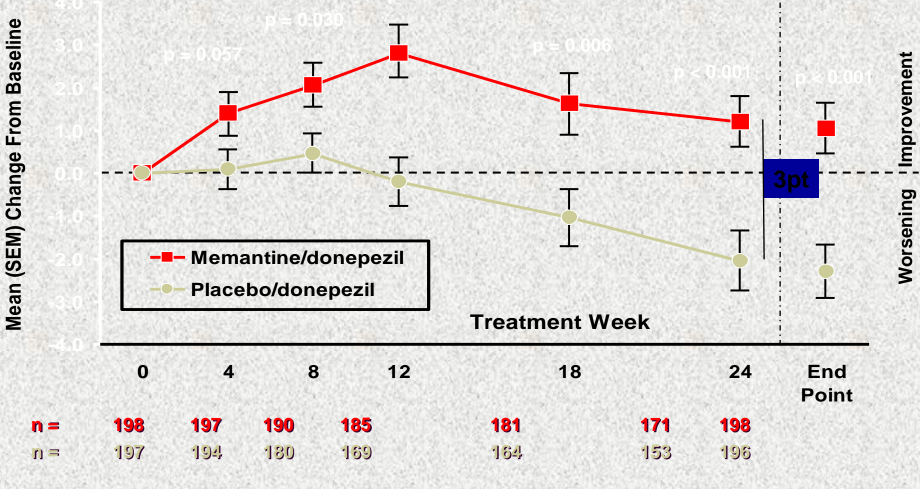
- 100% oral bioavailability
- Crosses blood-brain barrier rapidly
- No effects of food, age and gender
- Linear, dose-proportional kinetics (range 5- 40 mg)
- Elimination half-life of 60 80 hours
- Basis for dosing recommendation:
 - BID dosing better tolerated than QD dosing in early trials
 - Up-titration improves tolerability
- Metabolism:
 - Eliminated mostly in urine as parent drug and inactive metabolites
 - Lower dose recommended in moderate renal disease.
 - No or minimal effects on P450 isoenzymes
- No PK/PD interactions with donepezil



Mean (SEM) Change From Baseline

46

Memantine Added to Donepezil in AD Confers Additional Cognitive Benefits



Tariot et al. JAMA 2004;291:317-24

Adverse Events*

(Reported by ≥ 5% of Patients in Either Treatment Group)

Double-Blind, Placebo-Controlled Dementia Trials

	Placebo (N = 922) n (%)	Memantine (N = 940) n (%)
Any adverse event	624 (67.7)	662 (70.4)
Dizziness	49 (5.3)	64 (6.8)
Agitation	98 (10.6)	63 (6.7)
Confusion	42 (4.6)	58 (6.2)
Headache	31 (3.4)	54 (5.7)
Constipation	28 (3.0)	50 (5.3)
Fall 🕺	50 (5.4)	48 (5.1)
Inflicted Injury	64 (6.9)	44 (4.7)
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No significant effects on vital signs, tested lab parameters, or ECG

^{*} Adverse events were considered treatment emergent if not present at baseline or if present at baseline and increased in severity during the treatment period.

Noncognitive Behavioral Symptoms in Dementia: Critical Targets for Nonpharmacologic and Pharmacologic Therapy

* Mood

- Depression/Dysphoria
- Elation/Euphoria
- Anxiety
- Irritability
- Thought
 - Delusions
 - Hallucinations

* Activity

Apathy

- Aberrant motor behavior
- Vocalization
- Disinhibition
- Agitation/Aggression
- Sexual inappropriate behavior
- Disordered sleep
- Disordered Appetite/Eating

Behavioral Interventions: The First Line Treatment for NCBS

- Assure safety / Adequate supervision
- Behavioral analysis: Identify precipitants and response
- Behavioral interventions can include:
 - Caregiver education
 - Prosthetic (habilitative) environment
 - Activity/exercise
 - Individualized music therapy
 - Aromatherapy / massage
- Treatment should not exceed patient's capacity to learn/remember

Pharmacological Treatment of Psychosis/Agitation in Dementia

- No medication is FDA-indicated for treatment of psychosis or agitation in dementia; none has
- Intersection of the section of th
- Support for medications is based on short-term trials.
- Evidence base beyond antipsychotic medications is limited.
- Evidence-based off-label use of medications may be appropriate, is not illegal, and is common in practice.

Pharmacological Approaches to the Treatment of Behavioral and Psychological Symptoms of Dementia

Antipsychotics

- 2nd generation (atypical)
- 1st generation (typical)
- Mood stabilizing anticonvulsants
- Antidepressants
- Cholinesterase Inhibitors
- Miscellaneous agents

Class-Associated Severe AE and Mortality Concerns

- FDA Boxed Warning (April 11, 2005) notes "increased risk of death compared with placebo"
 - 17 PCTs, 5377 elderly pts with NCBS(3611 drug, 1766 placebo)
 - Deaths in drug treated patients: 4.5%
 - Deaths in placebo patients: 2.6%
 - **♦**OR = 1.6
 - Causes of death heart related or infectious
- Six drugs involved in trials: aripiprazole (3), olanzapine (5), risperidone (7), quetiapine (2), ziprasidone (1), haloperidol (2); Warning shared by clozapine and Symbyax (olanzapine/fluoxetine)

Current Standard of Care

- No clear consensus or treatment algorithm for agitation or psychosis in dementia
- Failure to show greater efficacy of antipsychotics for dementia-related agitation may be due in part to clinical trials designs
- Use of antipsychotics should be:
 - Short-term
 - With education of caregivers/family
 - Caution by clinicians
 - Careful documentation
- Further trials, new agents, FDA-approval needed.

Promising AD Therapies in Current Development (1)

Agent	Mechanism: AChEI	Status
Huperzine A	AcChE inhibitor and other effects	II

Agent	Mechanism: Immunotherapy	Status
ACC 001	Aβ fragment/carrier to induce Ab against Aβ Anti-Aβ monoclonal antibodies	I
Bapineuzumab Solanezumab	Anti-Aβ antibodies Anti-Aβ antibodies	

Promising AD Therapies in Current Development (2)

Agent	Mechanism: Amyloid-Lowering Agents	Status
LY 450139 MK 0752 Posiphen	Inhibits β amyloid synthesis/ γ secretase activity γ -secretase inhibitor β amyloid synthesis and β secretase activity inhibitor	II III I

For Reference: Additional Alzheimer's Disease Treatments Under Investigation (1)

のためある	Agent	Mechanism
	Dimebolin DHA EGb 761	Anti-tau agent Omega 3 FFA Improves blood flow, multiple effects
「たいない」のない	FK 962 GABA Aagonists	5HT agonist GABA A receptor agonists
あることである	GTS 21 Immune globulin Lecozotan	Synthetic nicotinic agonist Immune globulin Selective 5HT1A antagonist
100 Mar 100		

For Reference: Additional Alzheimer's Disease Treatments Under Investigation (2)

Agent	Mechanism
ABT 089 AC 1202 AL108 Alfatradiol Atorvastatin C 7617 C 9136 CERE 110	nAChR modulator Lowers oxidative stress Neuroprotective peptide (intranasal) 17-alpha-Estradiol Statin Neuroprotectant Neuroprotectant NGF agonist
Conjugated estrogens	Estrogens

For Reference: Additional Alzheimer's Disease Treatments Under Investigation (3)

Agent	Mechanism
Leuprorelin Levacecarnine LY 451395 MEM 1003 MEM 1414 Mifepristone MK 0249	GnRH analogue (acetyl-L-carnitine) Neuroprotective, unclear mechanism AMPA receptor agonist Ca channel modulator PDE4 inhibitor Glucocorticoid antagonist ?

Source (in part): http://newmeds.phrma.org, 2007

For Reference: Additional Alzheimer's Disease Treatments Under Investigation (4)

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	Agent	Mechanism
	MK 0952	PDE4 inhibitor
2	Modafinil	Mechanism unclear
27	Neramexane	NMDA receptor antagonist
22	NGX 267	Muscarinic M1 agonist
	Nicergoline	Alpha adrenergic vasodilator
2	PAZ 417	?
10000	PF 3084014	?
	Phenserine PRX	Inhibits AChE and β amyloid synthesis
17	03140	5HT4 agonist
	PRX 07034	Selective 5-HT6 receptor antagonist

For Reference: Additional Alzheimer's Disease Treatments Under Investigation (5)

Agent	Mechanism
R 1500 R-phenserine Rasagiline RN 1219 Rosiglitazone	? ? Selective MAO B inhibitor Humanized monoclonal antibody to Aβ PPARγ agonist
Selegiline transdermal SGS 742 SIB 1553A T817MA	Selective irrev. MAO B inhibitor GABA B antagonist nAChR agonist Neurotrophic agent

Source (in part): http://newmeds.phrma.org, 2007

For Reference: Additional Alzheimer's Disease Treatments Under Investigation (6)

Agent	Mechanism
Triacetyluridine TTP 488 Xaliproden	Uridine precursor (pro-drug) Decreases formation, deposition and accumulation of amyloid plaque 5HT 1a agonist and NGF agonist

Source (in part): http://newmeds.phrma.org, 2007

Acute Nonpharmacologic Approaches to Treating Noncognitive Behavioral Symptoms in Demented Patients

- Assure safety/adequate supervision
- Attend to environmental factors
- Educate support system
- Redirect and arrange pleasant experiences
- Do not rely on learning/memory
- Physical restraint is rarely necessary

Psychosocial Treatments for NCBS

- Psychoeducation for caregivers
- Individual behavioral plan for pt/caregiver
- Cognitive stimulation
- Music therapy
- Snoezelen
- Sensory stimulation
- Reminiscence therapy
- Validation therapy
- Reality orientation therapy
- Montessori activities
- Physical exercise
- Enforced social interaction

Evidence for benefit

Evidence for acute benefit only

Minimal evidence for benefit

Current Controversies over Pharmacotherapy of Behavioral and Psychological Symptoms of Dementia

- Use of Vitamin E: Still recommended?
- Use of antipsychotics for behavioral symptoms:
 - Limited efficacy / Unwanted metabolic effects / CVAE / Increased total mortality
 - Black box warnings
 - Shortcomings of data
- * Anticonvulsants
- Antidepressants
- Cholinesterase inhibitors/ Memantine

Additional Material on Early Detection of Alzheimer's Disease

Neuropathological Studies of Nondemented Elderly

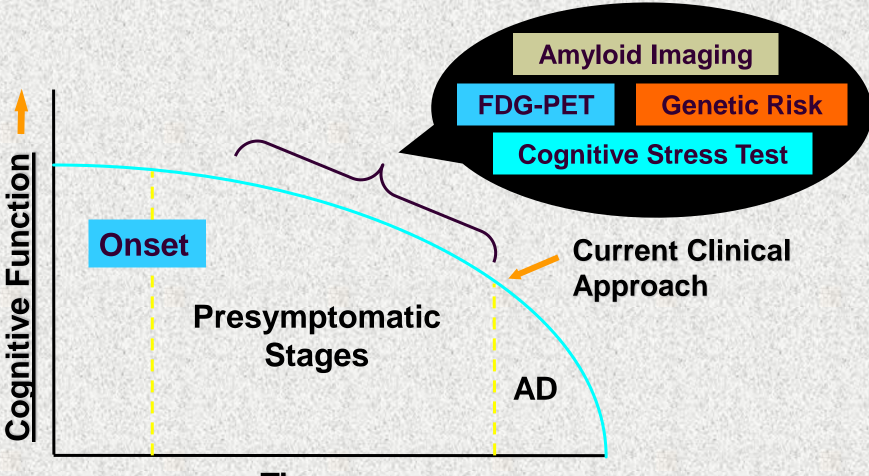
Morris et al¹

- Cerebral amyloid deposition in 21 elderly men, followed longitudinally
- *78% with high cortical plaque density had MCI
- Few or no plaques = cognitively intact

Braak & Braak²

 Neurofibrillary tangles appear to accumulate and extend from the entorhinal cortex as early as the third decade

Developing Tools for Early Detection of Alzheimer's Disease



<u>Time</u> —

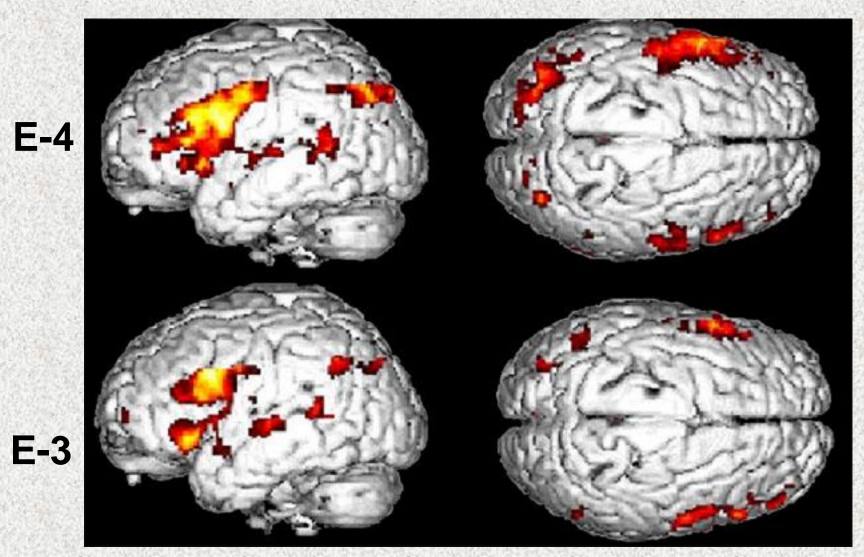
Genetic Risk

- Apolipoprotein E (APOE)—gene on chromosome 19
- **APOE-4** in 20% of population
- ***** APOE-4 increases risk, lowers onset age for AD
- APOE-4 may have modest effect in predicting cognitive decline in older persons, but APOE alone not considered a useful predictive test

Cognitive Stress Test with fMRI

- * Functional brain measures during memory performance may uncover subtle brain dysfunction not observed during mental rest (cf. treadmill ECG for cardiac disease)
- Combine neuroimaging and APOE-4 measures of genetic risk in order to identify abnormalities that may predict future cognitive decline

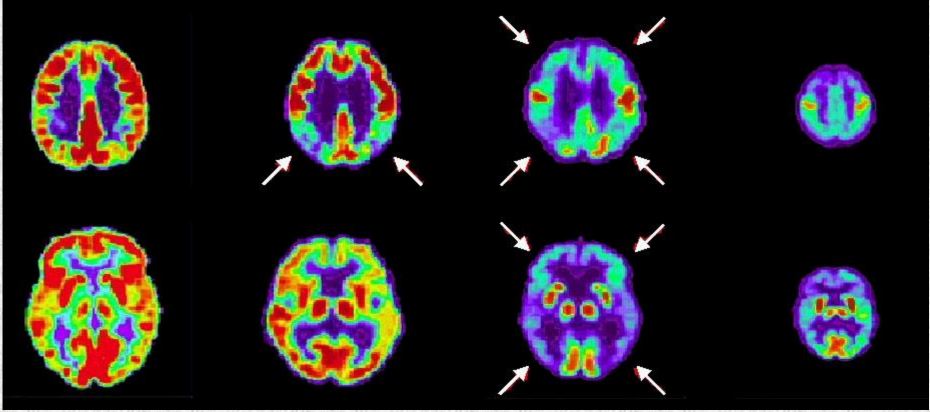
Increase Brain Activity During Memory Tasks



Bookheimer SY, Strojwas MH, Cohen MS, et al. N Engl J Med. 2000

Positron Emission Tomography (PET)

Cerebral Metabolism in Alzheimer's Disease Progression and in Normal Brains



Courtesy of Gary W. Small, M.D., UCLA

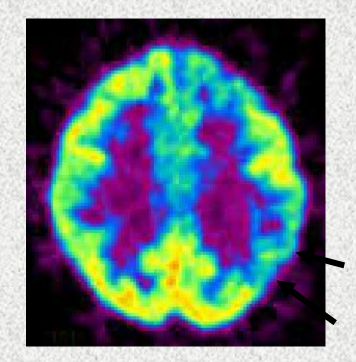
Accuracy of Early Diagnostic Assessment: Standard Clinical vs. FDG-PET

- Multiple clinical assessments over years in 134 patients
- Diagnostic accuracy¹
 - Sensitivity: 83-85%
 - Specificity: 50-55%
- Single baseline PET scan in 284 patients (138 autopsy diagnosis)
- Diagnostic accuracy²
 - Sensitivity: 93-95%
 - Specificity: 73-78%

¹Lim et al. J Am Geriatric Soc. 1999;47:564-569; ²Silverman et al. JAMA. 2001;286:2120-2127

Case Example

- 65-year-old woman diagnosed with depression and attention deficit after
 2 1/2 years of multiple neuropsychiatric evaluations including serial MRI scans
- PET showed typical Alzheimer's pattern
- Symptoms improved with cholinergic treatment



FDG-PET shows parietal deficit

Practical Consequences of Improved Diagnostic Accuracy

- Accurate diagnostic information and education reduces family/caregiver burden
- Decreased likelihood of repeated diagnostic assessments and testing
- *"Alzheimer's disease label" improves caregiver attitudes¹

Information about the disease improves quality of life for family/patient and delays nursing home placement²

¹Wadley et al. J Gerontol. 2001;56:244-252; ²Mittelman et al. JAMA. 1996;276:1725-1731

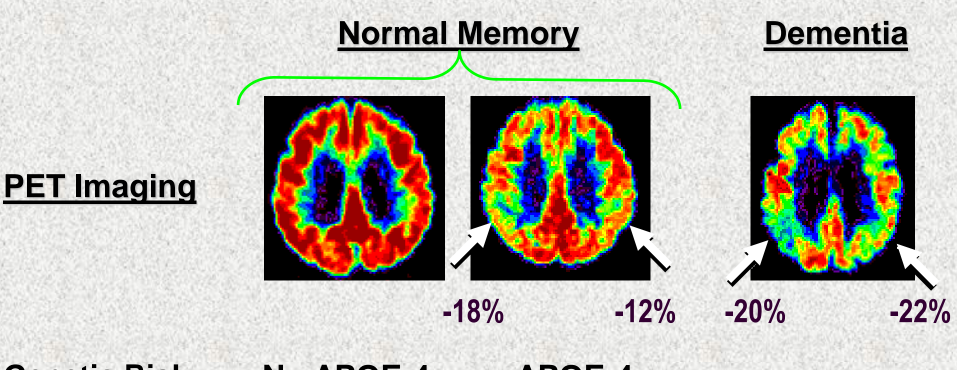
Practical Consequences of Improved Diagnostic Accuracy (Cont.)

Early, accurate diagnosis and treatment ...

- Maintains patients at higher levels of functioning leading to fewer MD/hospital visits¹
- Reduces caregiver burden²
- *Delays nursing home placement³
- Reduces use of other psychotropic drugs⁴

¹Small et al. J Am Geriatr Soc. 2002;50:321-327; ²Shikier et al. J Am Geriatr Soc. 2000;48:268-274; ³Knopman et al. Neurology. 1996;47:166-177; ⁴Small et al. Clin Therapeutics. 1998;20:838-850

Pet and Genetic Risk for Alzheimer Disease

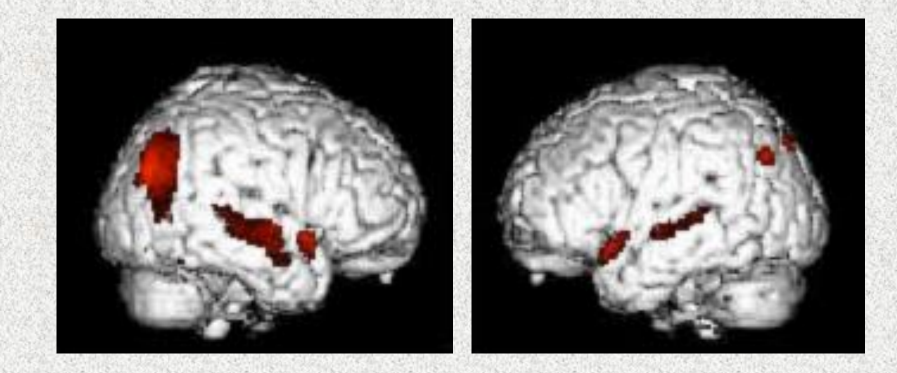


Genetic Risk No APOE-4 APOE-4

Lower inferior parietal metabolism in nondemented persons with a single copy of APOE-4

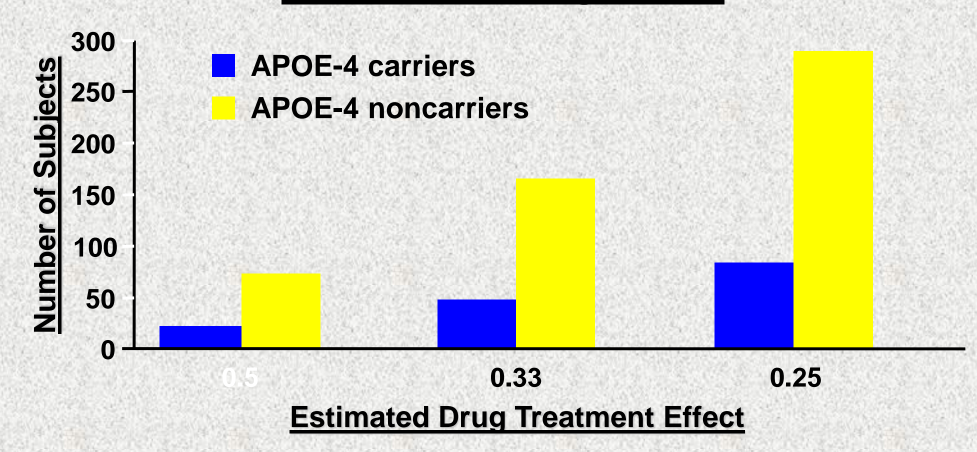
Small et al. PNAS. 2000;97:6037-6042

PET Scans Show Brain Function Decline in People at Genetic Risk for Alzheimer's Disease



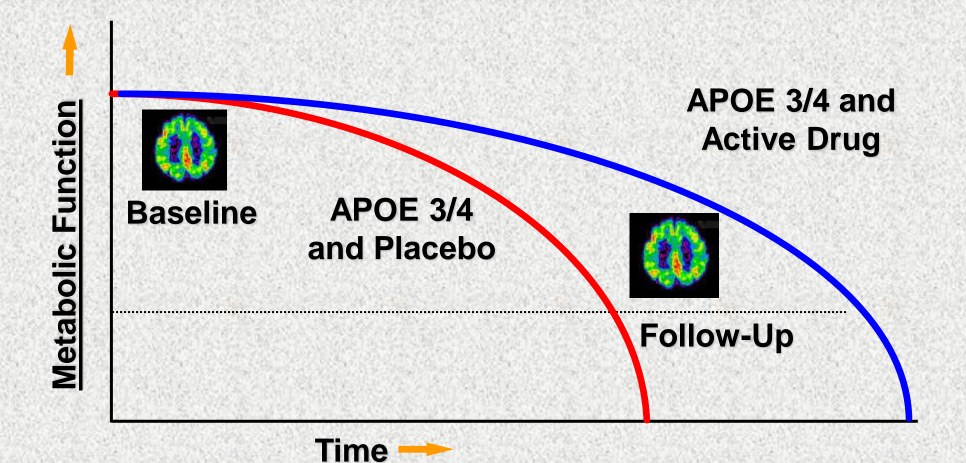
Small et al. PNAS. 2000;97:6037-6042

Number of Subjects Per Treatment Group Needed to Detect a Drug Effect in 2 Years Using PET*



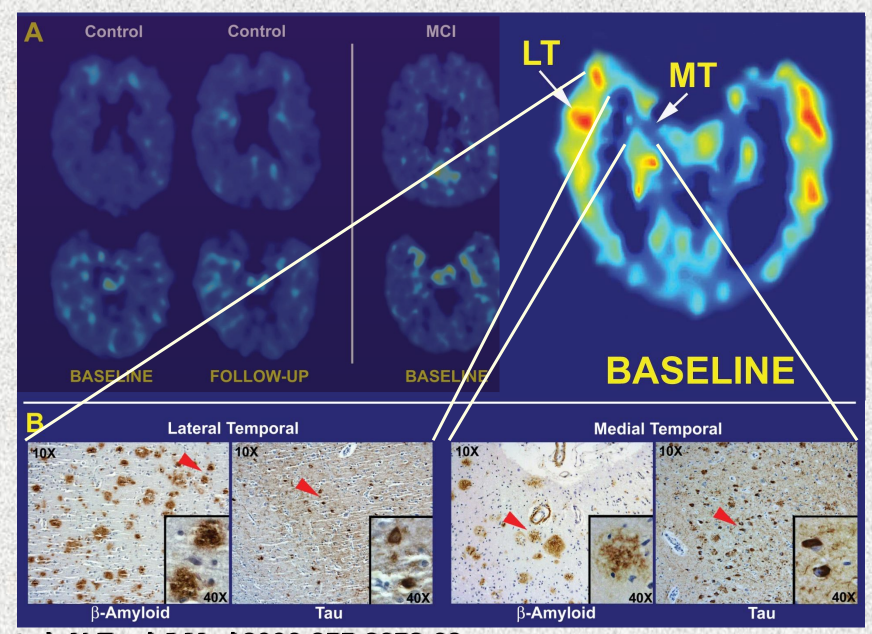
*Posterior cingulate metabolism; Based on data from: Reiman et al. PNAS. 2001;98:3334-3339

Possible Outcomes Using PET as a Surrogate Marker in AAMI Clinical Trials



AAMI = age-associated memory impairment

FDDNP-PET Plaque & Tangle Imaging



Small et al. N Engl J Med 2006;355:2652-63. LT = lateral temporal; MT = medial temporal.

Slowing Down Brain Aging

- Minimize stress
- Mental aerobics
- Physical exercise
- Healthy brain diet
- Lifestyle choices
- Medicines
- Memory training skills

Major Points

- Dementia is underrecognized and undertreated in primary care and in mental health settings
- Dementia can be recognized and treated beneficially in primary care and mental health settings
- Neuroimaging with PET can show a pattern of regional glucose metabolism that improves early detection of Alzheimer's disease with greater specificity

Major Points (cont.)

- Both pharmacological and nonpharmacological interventions may benefit overall brain health and dementia course
- Novel approaches to <u>in vivo</u> plaque and tangle imaging will be useful in monitoring potential disease-modifying agents

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Major Points (cont.)

- Neuroimaging with PET can show a pattern of regional glucose metabolism that improves early detection of Alzheimer's disease with greater specificity
- Novel approaches to <u>in vivo</u> plaque and tangle imaging will be useful in monitoring potential disease-modifying agents

Suggested Readings

- Cummings JL. Alzheimer's disease: from molecular biology to neuropsychiatry. Seminars in Clinical Neuropsychiatry. 8:31-6, 2003
- Cummings JL. Cole G. Alzheimer disease. JAMA. 287:2335-8, 2002
- Small GW. Rabins PV. Barry PP. et al. Diagnosis and treatment of Alzheimer disease and related disorders. Consensus statement of the American Association for Geriatric Psychiatry, the Alzheimer's Association, and the American Geriatrics Society. JAMA. 278:1363-71, 1997
- Small GW. What we need to know about age related memory loss. Br Med J 2002;324:1502-5.
- Small GW, et al. PET of brain amyloid and tau in mild cognitive impairment. N Engl J Med 2006;355;2652-63.

<u>Self-Assessment Question 1</u> <u>Which of the following</u>

are required for a diagnosis of dementia?

- A. Cognitive decline is ACQUIRED
- B. MEMORY is affected
- C. In addition to memory, ANOTHER cognitive function is affected.
- D. Symptoms are not attributable to delirium or another psychiatric disorder.
- E. All of the above

Self-Assessment Question 2 Which of the following cognitive or behavioral domains is affected in dementia?

- A. Memory
- B. Executive function
- C. Behavior
- D. Activities of daily living
- E. All of the above

Self-Assessment Question 3 Which of the following statements is correct?

- A. Alzheimer's Disease affects greater than 30% of adults older than 85 years of age.
- B. Alzheimer's Disease is infrequent among adults less than 60 years of age.
- C. Alzheimer's Disease is the most common cause of dementia.
- D. All of the above
- E. None of the above

Self-Assessment Question 4 Treatment of AD with cholinesterase inhibitors is based on which of these rationales?

- A. Noradrenergic neurotransmission in the locus ceruleus is reduced in late AD
- B. Pathological stimulation of NMDA receptors is associated with excitotoxic death of neurons.
- C. The number of cholinergic neurons in the basal forebrain is reduced in late Alzheimer's disease.
- D. All of the above
- E. None of the above

Self-Assessment Question 5 Treatment of AD with memantine is based on which of these rationales?

- A. Noradrenergic neurotransmission in the locus ceruleus is reduced in late AD
- B. Pathological stimulation of NMDA receptors is associated with excitotoxic death of neurons.
- C. The number of cholinergic neurons in the basal forebrain is reduced in late Alzheimer's disease.
- D. All of the above
- E. None of the above

Self-Assessment Question Answers

