# Energy Hypometabolism in Posterior Cingulate Cortex of Alzheimer's Patients: Superficial Laminar Cytochrome Oxidase Associated with Disease Duration

J. Valla,1,2 Jason D. Berndt,1 and F. Gonzalez-Lima1

<sup>1</sup>Institute for Neuroscience and Department of Psychology, University of Texas at Austin, Austin, Texas 78712, and <sup>2</sup>Harrington Alzheimer's Disease Research Laboratory, Arizona Alzheimer's Research Center, Phoenix, Arizona 85006

Among brain regions affected in Alzheimer's disease (AD), the posterior cingulate shows the earliest and largest decrement in energy metabolism. Positron emission tomography (PET) studies have shown that these decrements appear before the onset of memory deficits or other symptoms in persons at genetic risk for AD. This study compares in vivo imaging results and in situ postmortem analyses by examining the posterior cingulate (area 23) in 15 AD patients and 13 age-matched nondemented controls using quantitative cytochrome oxidase histochemistry as an intracellular measure of oxidative energy metabolic capacity. Each of the six layers of the posterior cingulate demonstrated a decline in cytochrome oxidase activity in AD relative to controls, whereas adjacent motor cortex showed no significant differences. This decrement did not appear to be mainly secondary to nonspecific decrement in mitochondrial enzymes, oxidative stress, cell loss, or histopathology. The cytochrome oxidase decrement was most severe in the superficial layer I (-39%), which demonstrated a correlation to disease duration. Covariance analyses suggest that superficial laminas undergo a functional uncoupling from the deeper layers of posterior cingulate cortex in AD, whereas no such effects are found in motor cortex or controls. These findings expand on previous results from PET studies by illuminating the layer-specific cytochrome oxidase contributions to energy hypometabolism. The findings suggest a decrement of cytochrome oxidase in posterior cingulate cortex, with progressive reduction within the superficial laminas linked to disease duration. Such decrement could contribute to some of the behavioral symptoms displayed by AD patients. This decrement appeared greater in women.

Key words: cytochrome oxidase; energy metabolism; posterior cingulate cortex; Alzheimer's disease; brain mapping; gender

Decrements in energy metabolism are one of the earliest detectable abnormalities in Alzheimer's disease (AD). Positron emission tomography (PET) studies on subjects homozygotic for the  $\epsilon$ 4 allele of the apolipoprotein E (APOE) gene, with no symptoms of AD, demonstrate metabolic reductions in posterior cingulate, parietal, temporal, and prefrontal cortices (Reiman et al., 1996; Small et al., 2000). Their largest decrement was in the posterior cingulate cortex (PCC) (Reiman et al., 1996), as in the case of AD (Minoshima et al., 1994). Another PET study (Minoshima et al., 1997) confirmed that although normal subjects show intense PCC metabolic activity, early-stage AD patients suffered a PCC decrement (21–22%) that was significantly greater than that seen in other cortical regions. This greater hypometabolism determined by PET is not merely an artifact of generalized atrophy (Ibanez et al., 1998) or AD histopathology that is lower in PCC than the other regions (Braak and Braak, 1998), although

degeneration of PCC occurs in AD (Brun and Gustafson, 1976). Severity of AD symptoms is correlated with hypoactivity in PCC but not temporal regions, as measured with PET and magnetic resonance imaging (Ishii et al., 1997; Hirono et al., 1998; Alsop et al., 2000). PCC hypoperfusion was also shown by subjects with only questionable dementia who later converted to AD in longitudinal SPECT studies (Johnson et al., 1998; Kogure et al., 2000). Cingulectomy has been linked to neglect and amnesia (Watson et al., 1973), and PCC is critically involved in memory retrieval as shown by PET studies (Nyberg et al., 1996; Cabeza et al., 1997). Together these studies suggest that regionally selective hypometabolism in PCC may be an important early event in the progression of AD.

In AD certain regions and cells are more vulnerable than others are in terms of energy metabolism but not histopathology (Gonzalez-Lima et al., 1998ab). For example, inferior colliculus neurons in AD patients versus age-matched controls revealed a selective reduction in cytochrome oxidase (CO) activity in dendritic neuropil around the largest and most metabolically active cell bodies in the central nucleus (Gonzalez-Lima et al., 1997). CO is the mitochondrial enzyme responsible for the activation of oxygen for aerobic energy metabolism and provides an intracellular measure of oxidative metabolic capacity because it is critically tied to ATP production inside mitochondria (Wong-Riley, 1989). CO is primarily localized in mitochondria near excitatory synapses in dendrites of neurons (Wong-Riley et al., 1998). Given the selective CO decrements in dendritic neuropil of AD neurons, it was of interest to investigate whether CO decrements may be preferentially localized to dendritic-rich cortical layers. The

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Correspondence should be addressed to F. Gonzalez-Lima, Professor and Head, Behavioral Neuroscience, University of Texas at Austin, Mezes Hall 330, Austin, TX 78712. E-mail: gonzalezlima@psy.utexas.edu.

J. D. Berndt's present address: Neuroscience Program, University of Wisconsin, Madison, WI 53706.

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Table 1. Summary of subject information for AD patients and aged controls

Subject	Age	Gender	PMI (hr)	Brain wt. (gm)	APOE	Clinical diagnosis/cause of death	Neuropathology diagnosis	
Alzheimer's cases								
3974	73	F	5	1020		Prob AD/unknown	CERAD definite AD	
9726	77	F	3.5	900	3/4	Poss AD/intestinal obstruction, hypovolemia	CERAD definite AD	
4353	77	F	6.1	1060		Prob AD/unknown	CERAD definite AD	
9701	78	F	5	1270	3/3	Prob AD/end stages of AD	CERAD definite AD	
9721	81	M	3	1150	3/4	Prob AD/pneumonia	CERAD definite AD	
9810	82	M	3.25	1145	3/4	Prob AD, COPD/CHF	CERAD definite AD	
9812	84	M	5.33	1050	4/4	Prob AD/CHF, inanition	CERAD definite AD	
4179	84	M	4.5	1200		Prob AD/unknown	CERAD definite AD	
9807	85	M	3.5	1220	3/3	Poss AD, CHF/RF	CERAD NP prob AD	
9817	86	F	4.5	780	3/3	Prob AD, depression/cardiac arrhythmia, ASHD	CERAD NP prob AD	
9830	86	F	3.5	1055	3/4	Prob AD, ASHD/cardiomyopathy	CERAD definite AD	
9727	88	M	5	1120	2/3	Prob AD/end stages of AD	CERAD definite AD	
9828	88	M	3	1170	3/3	Prob AD/CVA	CERAD definite AD	
9806	91	F	5	745	3/4	Prob AD/end stages of AD	CERAD definite AD	
9818	91	F	3.16	1010	3/3	Poss AD, COPD/CHF	CERAD NP prob AD	
Nondem	ented c	ontrols						
4143	71	M	11	1330		Control/unknown	CERAD NP poss AD	
9540	74	M	3	1230	3/3	Control/cardiac arrest	Normal brain, small infarcts	
9530	75	M	3	1560	3/3	Control/cardiopulmonary arrest	Pathologic aging, CERAD plaque score c, NP poss AD	
9537	77	M	3.5	1455	3/3	Coronary heart disease, hypertension/CHF	Pathologic aging, CAA	
9703	77	F	4.25	1235	3/4	Control/cardiopulmonary arrest	Pathologic aging, CERAD NP poss AD	
9904	77	M	3.5	1190	2/3	Head and neck cancer/RF	Astrocytoma, CERAD plaque score a, NP not AD	
9527	78	M	3	1240	3/3	COPD, ASHD, pulmonary fibrosis/RF	Normal brain, CERAD plaque score 0, NP not AD	
9521	79	M	3	1260	3/3	CVA/intracerebral hemorrhage	Hemorrhagic infarction	
9538	81	F	3.75	1225	2/3	Control/pneumonia	Pathologic aging, CAA, asymptomatic Lewy bodies	
9833	82	F	3	1185	3/3	Lung cancer	Subacute infarctions, CERAD plaque score a, NP not AD	
9751	84	M	4	1390	3/3	Control/cardiopulmonary arrest	CERAD NP poss AD	
9509	92	F	2	925	3/3	Normal pressure hydrocephalus with shunt	Normal pressure hydrocephalus with	
						1 , 1	shunt	
9536	92	F	2	985	3/3	Degenerative joint disease/CHF Pathologic aging, CERAD b, NP poss AD		
4338	95	F	7.05	1350		Control/unknown Not available		

ASHD, Arteriosclerotic heart disease; CAA, cerebral amyloid angiopathy; CERAD, Consortium to Establish a Registry for Alzheimer's Disease; CHF, congestive heart failure; COPD, chronic obstructive pulmonary disease; CVA, cerebral vascular accident; NP, CERAD Neuropathology protocol; RF, respiratory failure.

ability to do high-resolution laminar and cellular analyses is one large advantage histochemistry has compared with *in vivo* imaging, and as such, these techniques can complement one another. CO has been used to examine densitometrically the hippocampus of AD patients (Simonian and Hyman, 1993).

We assessed PCC laminar CO activity in AD patients and age-matched nondemented controls. The objectives were: (1) to confirm that CO activity shows more vulnerability in PCC than adjacent motor cortex, as suggested by the predominant *in vivo* hypometabolism of PCC in AD; (2) to investigate whether preferential CO decrement in AD may be localized to superficial cortical layers, as suggested by the predominance of CO in dendritic neuropil; and (3) to determine whether CO effects are correlated with disease duration, as expected of events relevant to AD progression.

# **MATERIALS AND METHODS**

Subjects and samples. Frozen tissue samples from 28 subjects were either dissected by us without intermediate thawing at the Sun Health Research Institute (Sun City, AZ) (23 cases) or received frozen from the Harvard Brain Tissue Resource Center (Belmont, MA) (5 cases). The acceptable postmortem interval (PMI) for inclusion was restricted to <12 hr to reduce the variability of the enzyme activities. Age was restricted to  $\geq$ 70 years in an effort to focus on late-onset, sporadic forms of AD. Table 1 summarizes the age, gender, PMI, brain weight, APOE status, clinical diagnosis, cause of death, and neuropathology findings available for each case

In the majority of cases, both area samples were dissected from the same section, corresponding to the medial aspect of a coronal section cut at the plane of the paracentral lobule dorsally and the posterior end of the splenium ventrally. The dorsal sample corresponded to Brodmann's area 4 in the paracentral lobule [primary motor cortex (PMC)], and the ventral sample corresponded to Brodmann's area 23 in the cingulate gyrus (PCC) (Fig. 1A). The investigated PCC is area 23 at the level of the

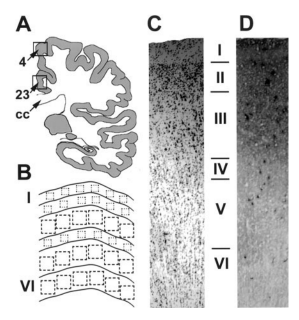


Figure 1. A, Schematic diagram indicating the sampling location for Brodmann's area 4 (PMC) and area 23 (PCC). CC, Corpus callosum. B, Schematic representation of the sampling from each layer of the cortex. C, Sample of Nissl-stained cortex (area 4). D, Sample of CO-stained cortex taken from a section immediately adjacent to the tissue in C.

splenium, not the retrosplenial cortex located posterior to the splenium. Occasionally a single area was available in a subset of the subjects, as reflected in the sample sizes reported below. This primary motor cortex region adjacent to PCC was chosen as a comparison region because it has high baseline metabolic activity similar to PCC, but it shows no significant hypometabolism in AD (Minoshima et al., 1997).

Tissue processing. Initially, several frozen sections were cut in various planes and Nissl-stained to determine a sectioning plane perpendicular to the horizontal cortical layers. Subsequently, the rest of the sample was sectioned into 40-µm-thick sections in this perpendicular plane and picked up on clean slides in a Frigocut 2800 cryostat at -15°C. Several adjacent series were created for analysis. One series was Nissl-stained with cresyl violet to aid with the determination of the laminar structure, and adjacent series were used for the histochemical stains. Experimenters were blind to the subjects' disease condition in all analyses below.

Cytochrome oxidase histochemistry. A series of sections was processed for CO quantitative histochemistry using previously described procedures (Gonzalez-Lima and Jones, 1994; Gonzalez-Lima and Cada, 1994, 1998; Gonzalez-Lima et al., 1997). Fresh-frozen tissue was used, and preincubation fixation was restricted to 5 min because of its detrimental effects on CO activity (Chalmers and Edgerton, 1989; Gonzalez-Lima and Cada, 1998). Briefly, slides were first treated in 10% sucrose phosphate buffer (0.1 M, pH 7.6) containing 0.5% glutaraldehyde (Grade II) for 5 min. Three changes at 5 min each of 10% phosphate buffer were followed by a preincubation for 10 min in Tris buffer (0.05 M, pH 7.6) containing 275 mg/l cobalt chloride, 10% sucrose, and 0.5% dimethylsulfoxide. The slides were then rinsed for 5 min in phosphate buffer and incubated at 37°C for 90 min in 700 ml of an oxygen-saturated reaction solution containing 350 mg of diaminobenzidine tetrahydrochloride, 52.5 mg of cytochrome c, 35 gm of sucrose, 14 mg of catalase, and 1.75 ml of dimethylsulfoxide in phosphate buffer. To stop the reaction and fix the tissue, a 30 min immersion in 10% sucrose phosphate buffer with 4% formalin (v/v) was used before dehydrating, clearing (xylene), and coverslipping with Permount. Sections from AD and control subjects from the PCC were stained in a single batch, and sections from PMC in AD and controls were stained in another to remove the possibility of interbatch variability as a confound in the comparison between subject groups. In addition, there were no interbatch differences across the two regions, as verified by the optical density (OD) of CO activity standards included in each batch. These same standards were used to convert tissue optical density measures to CO activity units via a regression equation based on their optical density and spectrophotometrically determined activity (Cada et al., 1995; Gonzalez-Lima and Cada, 1998).

Figure 1*B* shows the schematic sampling procedure for image analysis. First, the slides were coded so that the operator was blind to subjects' identity and group membership. Second, to rule out sampling bias because of better staining in the CO-stained regions, the areas for measurement were selected in the adjacent Nissl-stained sections. Selection was based on two morphological criteria: (1) the region must be free from histological artifacts, and (2) the region must show six well defined layers. Therefore, we sampled the most histologically intact area within each sample, which may reduce the differences between groups if the differences were caused by degenerative changes. The same region selected for cell counting in the Nissl-stained section was scanned in the adjacent CO-stained section by overlapping a template with the morphological delineations and landmarks.

Laminar delineation was accomplished with a low magnification camera lucida system using a six-layer definition of the cell populations in PCC and PMC. The laminas were defined by cell typing and population density by a single rater using the cresyl violet-stained sections. Examples of Nissl- and CO-stained cortex are shown in Figure 1, *C* and *D*, respectively.

The optical density of the CO-stained sections was analyzed using an image processing system consisting of a CCD camera, DC-powered light box, image capture board (Targa), and JAVA (Jandel Scientific, Corte Madera, CA) image analysis software. The system was calibrated using an optical density step tablet (Eastman Kodak, Rochester, NY), and the images were corrected for optical distortions using a subtraction of the background illumination. Six tissue optical density readings were taken from each layer from a single section for each subject with the aid of the overlaid camera lucida sketches. Each reading consisted of the average of the pixel values in a square sampling window of variable size. The size of the window was adjusted to encompass the width of the laminas and ranged in size from  $5 \times 5$  pixels ( $\sim 0.12 \times 0.12$  mm) to  $19 \times 19$  pixels  $(\sim 0.36 \times 0.36 \text{ mm})$ . The measurements were taken approximately equidistant across the visible sample, and any tissue artifacts were avoided (Fig. 1B). Each subject received seven final scores: a mean of the readings from each of the six laminas and a grand mean averaging across all six laminas. Statistical values are reported as mean  $\pm$  SEM.

Comparisons between the genders and agonal states used the grand mean scores of CO activity. Between-group multiple comparisons of the CO activity in PCC and PMC were analyzed with two-tailed t tests corrected for multiple comparisons with Hochberg's (1988) sharper Bonferroni procedure, as was the comparison between APOE  $\epsilon 4$  carriers and noncarriers. Within-group comparisons of laminar activity within each region and group were done with the two-tailed Mann-Whitney U test. All correlations presented are Pearson product moment correlations and their associated probability of being different from zero. To assess interlaminar correlations and differences between correlations, each subject's laminar data were normalized to an arbitrary overall average value of 100 to avoid artificial inflation of the correlations caused by subject variation. The correlation values found to be different from zero at p <0.05 were then compared for group differences using an r-to-z transformation corrected for multiple comparisons with Hochberg's (1988) sharper Bonferroni procedure.

Succinate dehydrogenase histochemistry. Succinate dehydrogenase (SDH), complex II of the electron transport chain, may also be affected in AD if our CO findings are attributable to a nonspecific effect on mitochondrial enzymes. To further characterize the specificity of our findings we chose to analyze the activity of this mitochondrial enzyme in the region with the largest reduction of CO activity, the PCC.

SDH histochemistry was performed on 40  $\mu$ m frozen sections adjacent to those used for CO histochemistry. The procedure was as follows: (1) slides were placed directly from  $-20^{\circ}\text{C}$  into 100 mm PBS with 10% (w/v) sucrose. Three baths of 5 min each with graded temperature from  $\sim$ 4–22°C were used to bring the slides to room temperature. (2) Slides were then transferred to PBS alone for 5 min. (3) Slides were incubated for 8 min at 37°C in the dark in the following reaction solution: 48 mm sodium succinate, 1.059 gm of EDTA (disodium salt), 0.046 gm of sodium azide, 0.859 gm of nitroblue tetrazolium (NBT), and 0.236 gm of 1-methoxyphenazine methylsulfate (mPMS) in 700 ml of double-distilled water. (4) The reaction was stopped with 10% formalin in PBS for 30 min at 22°C. (5) Slides were dehydrated in graded ethanol concentrations, 30, 50, 70, 95, 95, 100, and 100% in double-distilled water for 5 min each. (6) Sections were cleared in xylene, with three changes, 5 min each, and coverslipped with Permount.

Through the action of mPMS, an exogenous electron carrier, SDH catalyzes the dielectronic reduction of NBT to an insoluble product,

formazan, which can be visualized as a dark blue stain. Within a certain time course of the reaction, the accumulation of formazan does not affect the forward rate of reaction significantly. Therefore, endpoint measurements can be assumed to be linear with respect to reaction time up to 10 min, as we have determined empirically for our experimental conditions (data not shown).

The stained sections were analyzed for optical density with the same imaging system used for the CO analysis. Areas of measurement proximal to those used in the CO analysis were used. Deviations from the procedure above include that a laminar breakdown was not done; instead macroscopic average intensity measures encompassing  $\sim 2~\rm mm^2$  across all layers were made. The CO analysis was repeated in this manner to verify that no differences resulting from experimental method would skew these results, and mean CO results were the same with both methods. Data presented are mean  $\pm$  SE, and means were tested for a significant difference using a two-tailed t test.

Heme oxygenase-1 immunohistochemistry. Heme oxygenase-1 (HO1) is a putative antioxidant enzyme induced by oxidative stress (Premkumar et al., 1995). HO1 catalyzes the breakdown of heme motifs to form among other products, biliverdin, which is reduced to bilirubin, a molecule of known antioxidant properties. To determine whether our CO findings are attributable to a nonspecific effect of oxidative stress, HO1 immunohistochemistry was performed in the posterior cingulate cortex.

Frozen tissue sections from AD and control subjects were stained for the presence of HO1-reactive cells using on-the-slide immunohistochemistry. A polyclonal antibody, rabbit anti-rat HO1 (1:4000; 16 hr at 4°C; StressGen Biotech.Corp., Victoria, British Columbia, Canada) followed by biotinylated goat anti-rabbit IgG (1:200) and avidin-biotin-HRP (Vector Elite ABC kit; Vector Laboratories, Burlingame, CA) served to label the protein. Sections were then incubated in a chromagen solution, 3,3'-diaminobenzidine tetrahydrochloride (0.05%) in the presence of  $\rm H_2O_2$  (0.0015%), for 8 min at 22°C. Sections were dehydrated in graded ethanol concentrations, cleared in xylenes, and coverslipped.

Using an analog camera attached to an Olympus Optical (Tokyo, Japan) BX40 bright-field microscope at  $40\times$  magnification, images were captured and imported into JAVA software for analysis. An area of measurement across all layers ( $\sim$ 2 mm²) proximal to that used for CO and SDH histochemistry was delineated using the software. A pixel-by-pixel frequency histogram of ODs was generated, and a background threshold was created for each subject by selecting only those OD >1 SD above the mean. The object-counting feature of the software was then used to count the number of stained objects (cells) in this selected range. Adjacent stained pixels were deemed one object, and single pixel objects were eliminated from the overall count. The actual area sampled was recorded, and counts were normalized to cells per square millimeter. There were no significant group differences in the area sampled (data not shown). Data presented are mean  $\pm$  SE, and means were tested for a significant difference using a two-tailed t test.

This protocol assumes that HO1 is not reorganized from a diffuse to punctate spatial distribution between the control and pathological cases. This assumption is supported by the work of other groups, which indicate that HO1 immunoreactivity is coincident with reactive astrocytosis and colocalizes with glial fibrillary acidic protein, suggesting that it might be present in glial cells responding to neurological insult (Smith et al., 1994). If this were true in the AD cases, it would result in a greater number of immunoreactive objects identified in the above protocol. Admittedly, HO1 is an indirect measure of oxidative insult, but this analysis does provide us with some indication that our findings are not those of generalized transcriptional downregulation.

Cell counting in cresyl violet-stained sections. To determine whether the CO activity decreases simply reflect cellular degeneration, counts of NissI-stained cells were obtained from the same sections and regions used to demarcate the laminas for CO analysis as described in the section of the CO methods. Areas of measurement across all layers ( $\sim$ 2 mm²) used for the CO, SDH, and HO1 analyses were used and counted using the microscope setup and counting protocol as in the HO1 analysis. Counts were again normalized to cells per square millimeter. As in that analysis, there were no significant group differences in area sampled (data not shown). This form of counting, although not suited for indicating the actual number of cells, is sufficient to rule out group differences resulting from pathological degeneration in the sampled area (Arendt et al., 1983). Data presented are mean  $\pm$  SE, and means were tested for a significant difference using a two-tailed t test.

Amyloid plaque and neurofibrillary tangle histopathology. To investigate whether CO activity was related to AD histopathology a final series was

stained with thioflavin S for a survey of amyloid plaque and neurofibrillary tangle pathology in the PCC. Slide-mounted sections were fixed in 4% formalin (v/v) for 30 min, followed by pretreatment in 50% chloroform-50% ethanol for 1 hr. The slides were passed through graded alcohols (10 sec each) to distilled water (5 min), and then to 0.1% thioflavin S (Sigma, St. Louis, MO) in distilled water (10 min). Slides were differentiated in 80% ethanol (10-12 sec), quickly rinsed in distilled water, and coverslipped with an aqueous medium. Pathology was visualized on a Zeiss Universal fluorescent microscope using a 10× objective. Severity was semiquantitatively assessed on a seven-point scale ranging from 0 (none) to 6 (frequent), using the CERAD criteria as a guideline (Mirra et al., 1991). Cored/neuritic plaques, diffuse plaques, and neurofibrillary tangles were assessed separately, with each subject receiving one score for each. Scores were correlated on a subject-by-subject basis with laminar and mean CO activity in the AD patients using Pearson product moment correlations, and their probability of being different from zero was calculated.

#### **RESULTS**

## Subjects and samples

The subject information (age, PMI, brain weight) was compared between groups using two-tailed t tests. In the analysis of the PCC, the nondemented control group (n=13) consisted of seven males and six females with a mean  $\pm$  SEM age of  $81.2 \pm 2.1$  years, a mean PMI of  $4.1 \pm 0.7$  hr, and a mean brain weight of  $1255 \pm 48$  gm. The AD group (n=15) consisted of seven males and eight females with a mean age of  $83.4 \pm 1.4$  years, a mean PMI of  $4.2 \pm 1.0$  hr, and a mean brain weight of  $1060 \pm 40$  gm. AD patients' reports included confirmation of AD histopathology and, for 12 of the 15 subjects, years since diagnosis of AD ( $6.25 \pm 0.9$  years).

In the analysis of the PMC, the nondemented control group (n=12) consisted of seven males and five females with a mean age of  $80.7 \pm 1.7$  years, a mean PMI of  $3.2 \pm 0.2$  hr, and a mean brain weight of  $1240 \pm 51$  gm. The AD group (n=12) consisted of six males and six females with a mean age of  $84.8 \pm 1.3$  years, a mean PMI of  $4.0 \pm 0.8$  hr, and a mean brain weight of  $1051 \pm 48$  gm. Years since diagnosis with AD =  $6.25 \pm 0.9$  years (identical subset of subjects as in PCC).

The AD subjects across both areas had decreased brain weight relative to their age-matched controls (Student's two-tailed t=2.7 in PMC, 3.2 in PCC; p<0.05). No significant differences existed in age across the groups in either area. No difference was found in the PMI within the PCC subjects. In PMC, the AD patients demonstrated a PMI significantly longer than controls (t=-2.5; p<0.05); however, the actual time difference between the group means was inconsequential (<1 hr).

APOE status was available for the majority of the subjects. It may be of interest to point out that the only control subject to carry an  $\epsilon 4$  allele also had the lowest mean CO value (57 U); however, in the AD patients, there were no significant differences in CO activity between  $\epsilon 4$  carriers and noncarriers in any of the six layers, even before correction for multiple comparisons (n = 12; Student's two-tailed t test; p > 0.19).

The mean CO value was used to assess the contribution of agonal disease state in the control group. Agonal state (as assessed via cause of death when available) had no apparent effect on measured CO activity in the controls. Those who died from relatively acute conditions (e.g., cardiac arrest, stroke) showed similar activity (n = 5;  $78 \pm 11$  U) to those with more chronic agonal conditions (e.g., cancer, chronic obstructive pulmonary disease, degenerative joint disease; n = 5;  $104 \pm 12$  U; p > 0.13; two-tailed t test).

The mean CO value was used to assess the effect of gender within the groups. Female AD patients (n = 8) demonstrated significantly decreased CO activity relative to male AD patients

Table 2. Mean (SEM) of laminar cytochrome oxidase activity units (micromoles per minute per gram of tissue wet weight) in human postmortem posterior cingulate (area 23) and primary motor cortex (area 4) in AD patients and age-matched controls

	Posterior cingu	ılate cortex		Primary motor	Primary motor cortex			
	Control	AD	% Difference	Control	AD	% Difference		
Layer I	85 (8)	52 (3)**	-39%	86 (7)	68 (4)	-21%		
Layer II	88 (7)	60 (3)**	-32%	103 (7)	84 (4)	-18%		
Layer III	95 (9)	71 (4)*	-25%	115 (6)	100 (4)	-13%		
Layer IV	87 (8)	66 (3)*	-24%	102 (5)	91 (6)	-11%		
Layer V	83 (7)	63 (3)*	-24%	95 (6)	86 (5)	-9%		
Layer VI	73 (6)	55 (3)*	-25%	86 (6)	76 (5)	-12%		
Mean	85 (7)	61 (2)**	-28%	98 (6)	84 (4)	-14%		

Significantly lower than control (\*p < 0.05; \*\*p < 0.01; two-tailed Bonferroni-corrected).

(n=7) with 55  $\pm$  3 and 68  $\pm$  2 U, respectively, in the PCC (Student's two-tailed t test, -3.3; p < 0.01; -19%). There was no similar decrement in the PMC or in the control groups in either area. We also performed a multivariate analysis to conclude that "gender" was an independent and significant factor. We had no AD severity data, but we performed this analysis using ANCOVA to covary out brain weight within the AD group and compare across gender using the layer CO means, after excluding one outlier identified in the regression with a Studentized residual of -3.45. This analysis resulted in a highly significant effect of gender in the ANCOVA (p=0.007; n=7 per group).

#### Cytochrome oxidase laminar analysis

The results of the laminar analysis are presented in Table 2. Light microscopic examination of the CO histochemistry in the PCC and PMC showed that our procedure stains very few cell bodies and that the stain was essentially found exclusively in neuropil distributed across all the cortical laminas. Because CO is a marker of excitatory synaptic activity, the predominant neuropil staining is in line with the majority of the excitatory synapses in these cortical areas being axodendritic as opposed to axosomatic (Wong-Riley et al., 1998).

In the PCC, each layer (I-VI) showed significant CO activity decrements in the AD patients relative to the controls (p < 0.05). Layers I and II showed relatively more hypometabolism (p < 0.01), and layer I demonstrated a statistically greater decrement relative to layers III, IV, V, and VI in the AD group (Mann–

Whitney U tests; p < 0.05). Averaging across the laminas for each subject revealed a significant overall mean activity decrease in the AD PCC of 28% (p < 0.01) relative to the age-matched controls. A similar but nonsignificant trend was seen in the PMC. None of the PMC layers were significantly different between AD and controls.

Correlation matrices were constructed for normalized CO activity in each group and area (Table 3). Correlations between layers were found different from zero at p < 0.05 in both areas and both groups. Comparing the groups using an r-to-z transformation highlighted three significant (p < 0.05, corrected for multiple comparisons) changes in the correlative pattern in the PCC, indicating that the relationships between the laminas were altered by the disease process. The patterns of significant correlations were nearly identical across groups in the PMC with no correlations significantly different between groups after correction for multiple comparisons.

To investigate the influence of the patients' age and the duration over which they had been diagnosed with AD, those measures were correlated against CO activity in the PCC. There was no relationship between age and CO activity in AD or controls in the limited age range we studied. However, the duration of AD was inversely related to the severity of the decline of CO activity in layer I of the PCC (r = -0.68; p = 0.01), i.e., the longer the disease duration, the lower the CO activity (Table 4). No significant correlations were found in PMC.

Table 3. Interlaminar correlation matrices for normalized CO data in PCC and PMC

		Posterior	Posterior cingulate cortex							Primary motor cortex					
		I	II	III	IV	V	VI	I	II	III	IV	V	VI		
	I	1.00						1.00							
	II	0.76*	1.00					0.95*	1.00						
Control	III	-0.72*	-0.68*	1.00				-0.26	-0.19	1.00					
	IV	-0.94*	-0.83*	0.68*	1.00			-0.89*	-0.85*	0.41	1.00				
	V	-0.65*	-0.37	0.37	0.42	1.00		-0.73*	-0.81*	-0.29	0.53	1.00			
	VI	-0.25	-0.53	-0.16	0.34	-0.03	1.00	-0.84*	-0.82*	-0.17	0.59*	0.72*	1.00		
	I	1.00						1.00							
	II	0.57*	1.00					0.62*	1.00						
AD	III	-0.10	0.53*,**	1.00				-0.19	0.15	1.00					
	IV	-0.59*	-0.70*	0.09	1.00			-0.86*	-0.75*	-0.18	1.00				
	V	-0.73*	-0.91*,**	-0.51	0.51	1.00		-0.86*	-0.76*	-0.18	0.85*	1.00			
	VI	-0.63*	-0.73*	-0.68*	0.21	0.86*,**	1.00	-0.59*	-0.81*	-0.55	0.77*	0.77*	1.00		

<sup>\*</sup>Correlation different from zero at two-tailed p < 0.05.

<sup>\*\*</sup>Correlation significantly different from aged control (two-tailed p < 0.05; Bonferroni-corrected).

Table 4. Pearson correlations between cytochrome oxidase activity and subject age and duration of disease in AD patients

	Years						
Laminae	Age $(n = 15)$	Duration $(n = 12)$					
I	0.13	-0.68*					
II	0.01	-0.41					
III	0.00	-0.25					
IV	-0.06	-0.34					
V	-0.14	-0.21					
VI	-0.18	-0.16					

n. Number of subjects.

#### Succinate dehydrogenase analysis

SDH, the other mitochondrial electron transport enzyme measured, showed no significant differences between AD and controls in the PCC region showing the CO decrement. The SDH mean  $\pm$  SEM activity (OD units  $\times$  100) was 2.81  $\pm$  0.17 in controls and 2.56  $\pm$  0.11 in AD patients (p=0.24). However, CO activity (OD units  $\times$  100) reanalyzed in exactly the same way as SDH activity showed a highly significant group difference (p=0.0017) between controls (1.64  $\pm$  0.06) and AD patients (1.39  $\pm$  0.03).

### Heme oxygenase-1 analysis

HO1, the other oxidoreductase enzyme measured that is induced by oxidative stress, showed no significant differences between AD and controls in the PCC region showing the CO decrement. The HO1 mean  $\pm$  SEM expression (immunostained cell bodies per mm²) was 202  $\pm$  11 in controls and 187  $\pm$  7 in AD patients (p=0.30).

# Cell counting analysis

The number of Nissl-stained cells, counted to determine whether cell loss could account for the CO decrement in the sampled PCC region, showed no significant differences between AD and controls. The mean  $\pm$  SEM number of cells (Nissl-stained cell bodies per square millimeter) was 483  $\pm$  27 in controls and 478  $\pm$  14 in AD patients (p=0.87). We sampled the most histologically intact areas of the PCC to specifically rule out that any CO decrements could be accounted for by nonspecific cell loss or artifacts in the sampled areas. Thus, the point we made is that our sampled areas differ in CO but do not differ in cell body counts; it is not whether overall PCC degenerative changes may exist in AD as compared with controls.

### Histopathological analysis

To investigate the influence of the neuropathological burden on CO activity in the PCC, the three semiquantitative measures of thioflavin-stained cored/neuritic plaques, diffuse plaques, and neurofibrillary tangles were correlated against the laminar and mean CO activity measures in the AD patients. No significant correlations were found (p > 0.05). In spite of significantly (p < 0.05) greater histopathology in the AD group in every measure (cored/neuritic plaques: control, 1.54  $\pm$  0.14, AD, 3.86  $\pm$  0.06; diffuse plaques: control, 1.23  $\pm$  0.15, AD, 3.79  $\pm$  0.14; tangles: control, 0.15  $\pm$  0.04, AD, 3.21  $\pm$  0.19), there were no relationships between subject CO activity and histopathological measures on a subject-by-subject basis.

We also performed a principal components analysis as used by Vogt et al. (1998). Although we did not have laminar histopathol-

ogy scores and thus could not define Vogt's subtypes, we performed the principal components with the CO scores. This indicated no subtypes; that is, all six layers contributed significantly to the first principal component. A distinctive finding of our study is the lack of correlation between traditional AD histopathology and CO activity. Thus, it is not surprising that the CO findings do not resemble Vogt's subtypes based on histopathology.

#### DISCUSSION

The results suggest that prominent reductions in energy metabolism seen in the PCC in PET studies of probable AD cases may be caused by CO decrements that are more severe in the superficial laminas. This study was the first to quantify CO activity histochemically in PCC frozen samples from confirmed dementia cases of the AD-type and age-matched nondemented controls. As anticipated, CO activity showed a greater decrement in PCC than PMC, in agreement with PET studies showing prominent hypometabolism in the PCC in AD patients (Minoshima et al., 1997). A reduced CO activity is consistent with an energy metabolic dysfunction in the PCC of AD patients. However, our control data suggest this CO decrement in PCC may not be mainly attributable to nonspecific factors such as loss of cells or general decrements in other oxidoreductase enzymes. For example, the other respiratory enzyme measured, SDH, is part of the electron transport chain in the inner mitochondrial membrane (complex II), and nonspecific local effects on CO (complex IV) should have affected SDH activity in a similar way. Similarly, it appears unlikely that other nonspecific effects such as oxidative stress and oxygen utilization in the PCC may explain our CO results. For example, HO1 induction should have been affected because this enzyme incorporates oxygen to heme under conditions of oxidative stress, as demonstrated in other brain regions in AD patients (Smith et al., 1994). Although our data from control analyses cannot rule out all possible explanations, together with the CO and PET data, the data on number of Nissl-stained cells, HO1immunostained cells, and SDH histochemical activity suggest that AD patients exhibit a preferential CO decrement in the PCC. Table 1 also included six patients with premature death (heart failure, pneumonia, cerebral vascular accident) during the course of their AD, suggesting that our CO findings likely do not represent a simple end-stage phenomenon. Indeed, metabolic decrements in PCC have been shown to exist very early in the course of AD (Minoshima et al., 1994, 1997; Reiman et al., 1996), and longitudinal studies show PCC dysfunction as a clinical predictor for later AD (Johnson et al., 1998; Kogure et al., 2000) that is correlated with severity of symptoms (Alsop et al., 2000).

Specificity of the CO decrement is also suggested by the lack of correlation between CO activity and histopathological measures. No relationship was found between CO activity and the presence of amyloid plaques and neurofibrillary tangles in the PCC. This dissociation is important because the early hypometabolism seen in PCC in PET studies is greater than that found in other regions, although traditional AD histopathology is lower in PCC than other regions such as the temporal cortex (Braak and Braak, 1998). In AD temporal cortex, Hatanpaa et al. (1998) quantified CO subunit III mRNA in neurons in close proximity to neuritic plaques. They found no reduction in mRNA levels in the proximal neurons or their processes compared with more plaquedistant neurons. However, Hatanpaa et al. (1996) found levels of CO mRNA decreased by 26% in neurons bearing early-stage neurofibrillary tangles compared with tangle-free neurons. Unsurprisingly, tangle-bearing neurons do not produce as much CO,

<sup>\*</sup>Different from zero at two-tailed p = 0.01.

and likely other enzymes and proteins, as tangle-free neurons. Obviously any energy hypometabolism in these neurons may be simply secondary to the abnormal intracellular tangles. However, this did not appear to be the case in the PCC where the CO decrement was not associated with plaques or tangles. Thus, both the PET and CO data suggest that histopathology alone cannot account for PCC hypometabolism. The lack of correlation between the presence of AD neuropathological changes and CO decreases in PCC is important because it questions the linkage between traditional AD histopathology and functional disruption in AD.

Two principal mechanisms may explain decreases in CO activity in AD brains: one is a nonspecific, secondary mechanism caused by reduced energy demand, and the other is a specific, primary mechanism caused by CO catalytic impairment. The first one may be secondary to neurofibrillary tangles or any process that may downregulate CO activity. For example, reduced neuronal activity and synaptic loss would downregulate CO and other enzymes of oxidative phosphorylation. One way to evaluate the contribution of this secondary effect is to measure other enzymes of oxidative phosphorylation, as done with SDH in this study. Clearly this secondary reduction in CO activity is not specific to AD (Kish et al., 1999) and can be manipulated in experimental animals by denervation or functional decreases in neuronal activity (Wong-Riley, 1989). This secondary mechanism does not appear to be the only one in PCC because a neuronal downregulation should have affected SDH activity or some of the other measures besides CO activity.

The second mechanism that may compromise CO activity in certain brain regions more than others may involve a primary CO catalytic impairment. We speculate that this primary mechanism appears likely to contribute to our findings in PCC for several reasons (Gonzalez-Lima et al., 1998ab). First, a catalytic abnormality in CO has been found in AD brains (Parker et al., 1994b) as well as peripheral AD tissues where it is much less likely to have resulted from secondary neuronal downregulation or another nonspecific cause. This primary CO impairment in AD cells cannot be attributable to nonspecific neuron atrophy or loss because it has been found in non-neural cells, such as platelets (Parker et al., 1990, 1994a), fibroblasts (Curti et al., 1997), and skeletal muscle (Gonzalez-Lima et al., 1998b). A CO decrement in these various peripheral tissues indicates that a CO impairment may exist in addition to any local neuropathology or neuronal atrophy in the AD brain. However, denervation and neuropathology affecting synaptic function in any given region are likely to secondarily enhance any primary CO decrement in that region. Second, a primary CO impairment in AD would not affect all cells equally; vulnerability would hinge on the reliance of the cell on aerobic metabolism and possibly its ability to ameliorate and/or withstand oxidative stress (Shoffner and Wallace, 1994). Thus, a CO impairment in AD may affect preferentially certain organs (like brain), cell types (like pyramidal neurons), and regions (like PCC) that are more dependent on CO for oxidative energy metabolism (Gonzalez-Lima et al., 1998ab). For example, systemic administration in animals of sodium azide, a CO activity inhibitor, have been shown to produce deficits in maze learning and memory but no motor deficits (Bennett and Rose, 1992; Bennett et al., 1992a,b), as well as different degrees of CO inhibition in different brain regions (Cada et al., 1995). Preferential vulnerability to CO impairments in some brain regions may also be related to their differential expression of CO genes (Chandrasekaran et al., 1994). Therefore, perhaps the most reasonable explanation for the preferential hypometabolism observed in the PCC in PET studies and in the current study may be a synergy between a primary CO catalytic abnormality and a secondary neuronal downregulation caused by atrophy and denervation of synapses. Separating the contribution of each of these factors to the overall decrease in CO activity may be difficult, but effort should be dedicated to this end because CO could be an attractive target for therapeutic intervention.

This study also showed that CO decrements in AD were localized mainly to layer I. This is a synapse-rich molecular layer with predominant dendritic neuropil as opposed to cell bodies, a fact that may contribute to its vulnerability. Aging is the major risk factor for AD; so it is interesting that layer I appears most vulnerable to aging effects. Peters et al. (1998) showed a 30–60% reduction in synapse density in layer I in old monkeys, particularly indicating that spiny synapses that belong to the apical dendritic tufts of pyramidal cells are degenerating in old age. In the PCC, pyramidal neurons most vulnerable to atrophy in AD have their tufts of apical dendrites in layer I (Brun and Englund, 1981). Pyramidal cells also show the greatest degree of neurofibrillary tangle histopathology in AD (Hof et al., 1990; Van Hoesen, 1990; Braak and Braak, 1998). A combination of layer 1 aging effects (Peters et al., 1998) and AD-vulnerable pyramidal neurons with dendrites in layer I may also contribute to the preferential CO decrements in layer I in AD compared with age-matched controls. The observed CO decrement in layer I was correlated with disease duration, as expected of events relevant to AD progression. It would also be beneficial to assess the relationship between CO activity and dementia severity. To our knowledge, no such work has been done to date, and no severity information was available for our subjects. However CO decrement in the PCC was correlated with disease duration, and disease duration is accompanied by a progressive increase in symptom severity and PCC hypoperfusion (Alsop et al., 2000), reinforcing the possibility that dementia severity could be related to CO decrements.

Finally, CO activity was ~20% lower in the PCC of female compared with male AD subjects. This gender effect was AD-specific in that the control subjects did not demonstrate such a disparity and region-specific in that it was not found in the PMC in either group. Demographic studies support the idea that females have an increased vulnerability to AD (Launer et al., 1999; Letenneur et al., 1999). It is hoped that CO may soon become a useful marker of metabolic vulnerability in subjects at risk for AD using new *in vivo* noninvasive imaging (Heekeren et al., 1999).

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