A Mitocentric View of Alzheimer’s Disease Suggests Multi-Faceted Treatments

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Abstract. Alzheimer’s disease (AD) is defined by senile plaques made of amyloid-β peptide (Aβ), neurofibrillary tangles made of hyperphosphorylated tau proteins, and memory deficits. Thus, the events initiating the cascade leading to these end points may be more effective therapeutic targets than treating each facet individually. In the small percentage of cases of AD that are genetic (or animal models that reflect this form of AD), the factor initiating AD is clear (e.g., genetic mutations lead to high Aβ1-42 or hyperphosphorylated tau proteins). In the vast majority of AD cases, the cause is unknown. Substantial evidence now suggests that abnormalities in glucose metabolism/mitochondrial function/oxidative stress (GMO) are an invariant feature of AD and occur at an early stage of the disease process in both genetic and non-genetic forms of AD. Indeed, decreases in brain glucose utilization are diagnostic for AD. Changes in calcium homeostasis also precede clinical manifestations of AD. Abnormal GMO can lead to plaques, tangles, and the calcium abnormalities that accompany AD. Abnormalities in GMO diminish the ability of the brain to adapt. Therapies targeting mitochondria may ameliorate abnormalities in plaques, tangles, calcium homeostasis, and cognition that comprise AD.

Keywords: Alzheimer’s disease, amyloid, calcium, mitochondria, oxidative stress, neurofibrillary tangles, senile plaques

INTRODUCTION

Alzheimer’s disease (AD) is defined by senile plaques and neurofibrillary tangles and progressive cognitive changes including memory loss. Since these features are the defining characteristics of AD, hypotheses to explain AD must address what induces the alterations in the processing of amyloid-β protein precursor (AβPP) that lead to plaque formation and the hyperphosphorylation of tau protein that leads to tangles. Other abnormalities that accompany AD and occur earlier in the disease process are plausible candidates to be upstream events that initiate plaque and tangle formation as well as the cognitive loss. Considerable evidence suggests that the changes in glucose metabolism/mitochondrial function/oxidative stress (GMO) and calcium homeostasis precede plaques, tangles, and clinical manifestation of AD in humans and in mouse models. The changes in GMO are plausibly linked in a causative manner to the other abnormalities that define AD. Hence, the early alterations in GMO, which can induce multiple abnormalities, seem like more desirable therapeutic targets than the reversal of the individual facets that may result from abnormalities in GMO.

LINK OF GLUCOSE METABOLISM, MITOCHONDRIAL FUNCTION, AND OXIDATIVE STRESS IN BRAIN FUNCTION

The brain has a high rate of glucose metabolism and normal brain function depends on glucose metabolism. Even mild reductions in the oxygen or glucose available to the brain diminish brain function including cognition. Novel alternative pathways of glucose utilization and oxygen utilization do not occur in the brain,
but the coupling of neuronal properties to metabolism is unique. For example, the energy requirements for maintenance of high ionic gradients across membranes and an interaction of metabolism with neurotransmitters are unique to brain. Six neurotransmitters are derived from metabolic pathways and three use molecular oxygen for their synthesis. Furthermore, other aspects of neurotransmission depend upon energy metabolism. For example, any reduction in oxygen availability to the brain diminishes the release of acetylcholine and reduces cholinergic function. Furthermore, neurotransmitter receptors are modified by oxidants. Thus, even mild interruptions in metabolism or oxidative stress alter brain function [1].

Free radicals (compounds with an unpaired electron) or reactive oxygen species (ROS) are a normal part of metabolism. The multiple sources of ROS in cells have been reviewed extensively. The major producer of ROS is thought to be the electron transport chain (ETC) of the mitochondria, particularly complex II and III. Other reactions within the tricarboxylic acid (TCA) cycle including the pyruvate dehydrogenase complex (PDHC), the α-ketoglutarate dehydrogenase complex (KGDHC), or malate dehydrogenase (MDH) can also produce free radicals under appropriate conditions [2,3]. KGDHC may also serve as ROS sensors [4]. Any of the many oxygenases or dehydrogenases in cells are linked to electron transfer and can lead to free radical production. For example, monoamine oxidase can be a major producer of oxidants [5]. Together these reactions represent a high proportion of all reactions in a biological system.

As part of their role in normal metabolism, ROS serve as signaling molecules. Many physiological reactions in all tissues including the ETC are modulated by ROS [6]. ROS can modify neurotransmitter receptors or ion transporters with profound effect on the function of these proteins [1]. ROS communicate between metabolic pathways. For example, production of ROS by the ETC can inactivate KGDHC of the TCA cycle to diminish the production of NADH [7]. ROS produced by monoamine oxidase can also inactivate KGDHC [5]. ROS can even be regulatory within protein complexes. For example, KGDHC consists up of three protein subunits. The third protein of the complex produces ROS that regulates the first protein of the same complex [8].

Oxidative stress occurs under conditions in which production of free radicals or ROS exceeds the ability of the cell or brain to buffer them. Since the ETC chain uses most of the oxygen in neurons, oxidative stress is often associated with the ETC. Any interruption of ETC normal function increases ROS production. For example, in cell systems either increasing or lowering oxygen increases ROS production [2,8]. Impairment of the ETC by a variety of means leads to excess levels of ROS. For example, adding azide to block respiration increases ROS production and inactivates KGDHC [7]. Reducing oxygen and/or glucose to the brain also increases ROS [9]. The excess oxidative stress can modify DNA, RNA, lipids, and proteins. The damaged molecules include those that can be regulated by ROS. For example, KGDHC can be irreversibly inactivated by a variety of oxidants so that recovery is not possible [10]. Thus, oxidative stress diminishes the ability of the brain to adapt.

In vivo studies support the suggestion that interruption of normal metabolism increases free radical production. For example, reducing the activity of the KGDHC and PDHC increases oxidative stress [11]. Mild impairment of metabolism by thiamine deficiency (TD) also increases oxidative stress [12]. ROS also increase in response to hypoglycemia. Increased lipoperoxidation levels are observed before the onset of the hypoglycemic-induced isoelectric period, while 3-nitrotyrosine residues in proteins and nitrotyrosine-positive cells occur with glucose reperfusion. These changes occur only in vulnerable brain regions, suggesting a correlation between oxidative damage and vulnerability to hypoglycemic neuronal death in selective brain regions. The results demonstrate that a pro-oxidant state is promoted in certain brain regions during hypoglycemia and after the glucose reperfusion phase, which may result from the activation of several oxidative stress pathways and may relate to subsequent cell death [13]. Thus, one challenge in unraveling the pathophysiology of AD is to understand the source of the free radicals that lead to damage in AD.

GLUCOSE METABOLISM/MITOCOCHONDRIAL FUNCTION/OXIDATIVE STRESS ARE CLOSELY LINKED TO CALCIUM REGULATION

Calcium, like ROS, is a normal signaling molecule that is closely linked to metabolism. In isolated nerve endings, hypoxia diminishes calcium uptake, but increases cytosolic calcium and diminishes calcium dependent release of acetylcholine [1]. In neurons, intracellular calcium increases in response to depolarization or activation of a variety of receptors. The increased calcium activates numerous calcium dependent
enzymes in the cytosol, and elevates calcium in the mitochondria and endoplasmic reticulum (ER). The increased calcium in the mitochondria activates a variety of processes, including KGDHC, and can promote ROS production. ROS increase calcium in hypoxic tissues [14]. Calcium in the ER is involved in multiple processes including ER stress, which is involved in cell death processes. Stabilizing the ER calcium can protect against ischemia-induced cell death [15]. The calcium transport systems of the ER are also sensitive to oxidative stress [16].

Calcium, mitochondria, and the ER are closely linked morphologically and functionally. Considerable cross talk of cell death proteins occurs between the ER and mitochondria [17]. Calcium released from the ER can elevate mitochondrial calcium without increasing cytosolic calcium, which suggests a direct morphological link. Double label experiments of ER and mitochondria support this suggestion [18]. Presenilins, which are mutated in one familial form of AD, may be part of this link. Presenilins are enriched in ER membranes associated with mitochondria and appear to provide a bridge between the two organelles [18]. This link of mitochondria and ER may underlie the AD-related abnormalities that are caused by mutations in presenilin-1.

METABOLIC CHANGES IN AD SUGGEST MITOCHONDRIAL DEFICITS ARE EARLY AND CRITICAL

Imaging studies of living patients

Measurements of the cerebral metabolic rate for glucose suggest that metabolic changes precede the onset of histopathological or clinical features. The ability to reliably image glucose metabolism in living people with fluoro-deoxyglucose (FDG) by positron emission tomography (PET) in a non-invasive manner allows temporal studies of living patients in tests of the pathophysiology of AD. The rate of brain glucose utilization predicts whether the patients will progress from mild cognitive impairment (MCI) to AD. Over an 18-month period, patients who went from MCI to AD had lower glucose utilization in the right temporoparietal cortex than those who did not convert [19]. The baseline hippocampal FDG PET predicts decline from normal to AD with 81% accuracy and from normal to MCI with 71% accuracy. Greater rates of decline of hippocampal and cortical FDG-PET are found in AD as compared to individuals without cognitive decline. Thus, FDG-PET during normal aging predicts cognitive decline in advance of the clinical diagnosis [20]. Other studies demonstrate that glucose utilization in the entorhinal cortex also accurately predicts the conversion from normal to MCI. Among those who declined, the baseline glucose utilization in the entorhinal cortex predicted longitudinal memory and temporal neocortex metabolic reductions. At follow-up, those who declined showed memory impairment and hypometabolism in temporal lobe neocortex and hippocampus. These data suggest that an entorhinal cortex stage of brain involvement can be detected in normal elderly and predicts future cognitive deficits and reductions in brain metabolism [21]. Changes in glucose metabolism may also link the apolipoprotein E 4ε allele (APOE4) predisposition to the cognitive changes in AD. Among those subjects who declined from MCI to AD, APOE4 carriers show marked reductions in temporal cortex metabolism. Thus, a multiplicity of studies show that glucose metabolism declines in early stages of AD [21].

Changes in glucose metabolism decades before the onset of AD was demonstrated by examining patients whom have a predisposition due to the presence of APOE4. Patients with AD have abnormally low rates of cerebral glucose metabolism in posterior cingulate, parietal, temporal, and prefrontal cortex. Cognitively normal, late-middle-aged carriers of the APOE4 allele have abnormally low rates of glucose metabolism in the same brain regions as patients with probable AD. The young APOE4 carriers have abnormally low rates of glucose metabolism bilaterally in the posterior cingulate, parietal, temporal, and prefrontal cortex. Thus, carriers of a common AD susceptibility gene have functional brain abnormalities in young adulthood, several decades before the possible onset of dementia [22].

The addition of FDG-PET to a clinical diagnosis provides useful information that can affect the likelihood of detecting AD pathology and may suggest therapeutic options [23]. Recent studies compared three promising biomarkers of AD including CSF amyloid-β (Aβ), amyloid PET tracer, and glucose metabolism. The authors concluded that “evidence that FDG-PET abnormalities precede any cognitive symptoms in individuals who later progress to AD is probably the strongest” [24].

Results with autopsy brain suggest a molecular basis for the decline in metabolism in AD

Measures in living individuals provide a temporal profile of glucose changes but cannot provide as much
insight into mechanism as studies of autopsy brains. The results suggest that both transcriptional and post-translational modifications of key metabolic pathways are likely critical.

**Protein changes**

Activity measurements of enzymes of metabolic pathways in autopsied brains provide insight into the cause of the decline in glucose utilization with AD. A key enzyme of the pentose phosphate shunt, transketolase, declines 45% (Fig. 1) [25]. The TCA cycle is the main catabolic converter of glucose to energy in the form of NADH. To determine whether changes in the TCA cycle may underlie the decline in glucose utilization in AD, the activities of the enzymes involved the TCA cycle were determined. The changes in the activities with AD form a discrete pattern. The dehydrogenases that are also decarboxylases are diminished: PDHC (−41%), isocitrate dehydrogenase (ICDH) (−27%), and KGDHC (−57%). On the other hand, the dehydrogenases are increased: succinate dehydrogenase (SDH) (mitochondrial complex II) (+44%) and MDH (+54%) (Fig. 1). The highest correlation with the decline in clinical dementia rating scale is with PDHC ($r = 0.77$) [26].

The protein that shows AD related changes which has been studied in the greatest detail is KGDHC. The activity of KGDHC declines in both genetic and non-genetic forms of AD. In most cases of AD, the protein levels (i.e., immunoreactivity) of KGDHC are not altered, which suggests post-translational modifications cause the reduction in activity [27]. In familial AD patients in which the AD is caused by a mutation in Aβ/PP processing, the protein levels of KGDHC subunits as detected by the same antibodies suggest reduction in protein levels (i.e., a loss of immunoreactivity [28]). The results are consistent with post-translational inactivation of KGDHC by different ROS in different forms of AD because some ROS alter KGDHC immunoreactivity whereas others do not. The extreme sensitivity of KGDHC to oxidants and the widespread oxidant damage in AD brains suggests that the deficits in AD may be oxidant mediated.

Changes in gene expression in autopsy brain support the suggestion that alterations in glucose metabolism are critical in the pathophysiology of AD

Transcriptional changes also contribute to the reduction in brain glucose utilization in AD, and these changes may be mediated by oxidants [29]. Gene microarrays of controls and AD reveal that down-regulation of genes involved in protein folding/metabolism/transport, energy metabolism, and signaling pathways accompany AD [30]. Of the 51 members of the glycolytic, TCA, oxidative phosphorylation, and associated pathways, qPCR indicates that 15 are down-regulated in AD [31]. A genome-wide transcriptomic approach found alterations in the expression of 80 metabolically relevant nuclear genes in posterior cingulate cortex. AD cases have significantly lower expression of 70% of the nuclear genes encoding subunits of the mitochondrial ETC in posterior cingulate cortex, 65% of those in the middle temporal gyrus, 61% of those in hippocampal CA1, 23% of those in entorhinal cortex, 16% of those in visual cortex, and 5% of those in the superior frontal gyrus [32]. mRNA levels of 11 mitochondrial-encoded genes are altered in patients with early and definitive AD. Results reveal down regulation of genes in Complex I, whereas gene expression of Complex III and IV show increases mRNA in patients with early and definitive AD [33]. Genes related to mitochondrial energy metabolism and apoptosis are also upregulated in 2-month old Tg2576 mice, which bear a double mutation in Aβ/PP. The same genes were upregulated at 5 and 18 months [34]. The interpretation of these results and their consequences on brain function is difficult since these diverse changes only reflect steady-state mRNA levels.
Structural changes in mitochondrial also support their role in the disease process

Morphometric analysis shows that the area of intact mitochondria is significantly decreased in AD, whereas there is no difference between the area of damaged mitochondria in AD or control cases [35].

ANIMAL MODELS OF PLAQUES AND TANGLES SUGGEST METABOLIC DEFICITS OCCUR IN EARLY STAGES OF THE DISEASE PROCESS

Changes in mitochondria precede other pathological changes even in transgenic mice that are engineered to make plaques and tangles. Mitochondrial Aβ increases at 9 months in the triple Tg-AD mice, whereas mitochondrial function is altered as early as three months of age including the following: decreased respiration, decreased PDHC protein level and activity, increased oxidative stress as manifested by increased H$_2$O$_2$ production and lipid peroxidation. Even embryonic neurons derived from the hippocampus of the triple transgenic AD mouse model (3xTg-AD) exhibit decreases in mitochondrial respiration and increased glycolysis [36]. In other mouse models of AD, the changes in mitochondrial function also occur before plaque formation. Isolated mitochondria from double Swedish and London mutant AβPP transgenic mice show pronounced mitochondrial dysfunction including a drop of mitochondrial membrane potential and reduced ATP-levels at 3 months when elevated intracellular, but not extracellular Aβ deposits are present [37], and there are no reports on presence of mitochondria Aβ at such an early age.

Whether Aβ has to be present in the mitochondria to alter mitochondrial function is not clear. Aβ is a very reactive compound and a multitude of papers show that Aβ can alter multiple cell and mitochondrial functions when it is added outside the cells or mitochondria, especially at high concentrations (e.g., 10 µM). Even nM concentration of Aβ added outside cells alters mitochondrial function and oxidative activity [38]. Indeed, Aβ added to cells inhibits PDHC and KGDHC [39], which are diminished in brains from patients with AD. Aβ (10 µM) also blocks the entry of nuclear encoded proteins into the mitochondria [40] and alters expression of fission proteins. Aβ can also alter the mitochondrial permeability transition [41]. Aβ is present in mitochondria in brains from patients with AD by immunocytochemistry [42] and in transgenic mice.

All of the experiments in transgenic mice that demonstrate mitochondrial Aβ are done in mice that express high levels of mutant human AβPP. For example, the Tg2576 mouse expresses mutant human AβPP 5.56 to 5.76 times the level of endogenous mouse AβPP [43]. Thus, these mice are designed to make abnormal Aβ production the initiating cause. Even in these mice, the data are not clear if Aβ must be present in the mitochondria to get mitochondrial dysfunction. One study demonstrated an increase in Aβ within mitochondria in the J-20 transgenic mouse strain. The results show a very small amount of Aβ in mitochondria at 4 months and a marked increase in mitochondrial Aβ at 8 months. Amyloid plaques occur in this model around 5–7 months [44]. At 8 months, no differences are seen in the mitochondrial Aβ, respiratory control ratio by isolated mitochondria, protein levels of complex I, II, III, and IV or oxygen consumption between control and J-20. At 12 months, mitochondrial Aβ is quite high and small differences in respiratory control ratio as well as small reductions in complex III or IV are present [42]. However, the measures of mitochondrial function and oxidative stress that usually precede plaque formation by months have not been studied in the J-20 line. In Tg2576 mice, plaques are not present at two months of age. However, soluble intraneuronal Aβ deposits are present, but in only very low amounts (about 10% of those at 12 months) [45]. In addition, increased H$_2$O$_2$, but no significant increase in carboxyls or cytochrome oxidase occurs at two months of age [45]. In this strain, very few Aβ deposits occur in 8 month old mice, but by 6 months Aβ occurs in mitochondria [45]. Earlier studies showed increases in markers of oxidative stress in urine and blood of these mice at 8 months of age [46]. Many Aβ plaques are present at 17 months but no significant changes in H$_2$O$_2$ production or cytochrome oxidase are apparent [45]. Thus, the data are consistent that mitochondrial changes precede plaque formation and perhaps the Aβ accumulation in the mitochondria.

The mechanism for Aβ entry into mitochondria has been examined. The results from rat mitochondria show that Aβ is transported into mitochondria via the translocase of the outer membrane (TOM) machinery. The import is independent of the mitochondrial membrane potential [47].

AD models in drosophila also have early mitochondrial changes

Expression of wild-type or arctic form of Aβ in a small group of neurons in the adult fly allows study of
both function and structure of both axon and presynaptic terminals. Depletion of presynaptic mitochondria is the earliest detected phenotype. Diminished axonal transport of mitochondria, accumulation of Aβ intra-cellularly, decreased synaptic vesicles, increased large vacuoles, and elevated synaptic fatigue follow. The results suggest early depletion of presynaptic and axonal mitochondria, which lead to other presynaptic deficits and behavioral deficits [48]. The loss of mitochondria at the synapse may be due to reduced transport of mitochondria down the axon. In the Aβ42 fly brain, mitochondria are reduced in axons and dendrites, and accumulate in the somata without severe mitochondrial damage or neurodegeneration. The organization of microtubule or global axonal transport is not significantly altered at this stage [49]. Thus, even in flies engineered to make more Aβ, the mitochondrial changes are very early.

OXIDATIVE STRESS IS AN EARLY EVENT IN AD

AD patients

Oxidative stress appears to be an early change in AD. Although markers of oxidative stress cannot be measured in vivo in brain, measures of oxidative stress can be found in urine and occur at very early stages of AD. Isoprostanes are significantly higher in cerebrospinal fluid, plasma, and urine of subjects with MCI and AD compared with cognitively normal elderly subjects [50], suggesting these are early events in AD.

Elevations of markers of ROS damage to proteins, nucleic acids, carbohydrates, and lipids in autopsy brains from AD patients are well-documented [51–55]. Oxidative damage in AD brain is more pervasive than plaques and tangles [56]. Significant increases of an oxidized nucleoside derived from RNA, 8-hydroxyguanosine, oxidized amino acids, and nitrotyrosine occur in vulnerable neurons of patients with AD [57,58]. To determine whether oxidative damage is an early- or end-stage event in the process of neurodegeneration, the relationship between neuronal 8-hydroxyguanosine and nitrotyrosine and histological and clinical variables (i.e., Aβ plaques, neurofibrillary tangles, duration of dementia, and APOE4 genotype) was determined. The results suggest oxidative damage is quantitatively greatest early in the disease and declines with disease progression. Increases in Aβ deposition are associated with decreased oxidative damage. These relationships are more significant in patients with APOE4. Neurons with tangles show a 40%-56% decrease in relative 8-hydroxyguanosine levels compared with neurons free of tangles. The results suggest that increased oxidative damage is an early event in AD that decreases with disease progression and lesion formation [59].

Animal models

Metabolic changes/oxidative stress occur in early stages of the disease in animals that model various aspects of AD. Reduction in the metabolic enzymes that are diminished in AD brains increases oxidative stress. For example, TD in mice or rats leads to early reductions in the activity of the thiamine dependent enzymes that are diminished in AD. These include enzymes that catalyze key steps in the pentose shunt (transketolase), entry of carbon into the TCA cycle (PDHC) and of the TCA cycle (KGDHC). The reduction in the activities of these enzymes lead to increased oxidative stress in a manner that mimics several aspects of the oxidative stress observed in AD including hydroxynoneal, nitrotyrosine, redox active iron, microglia activation, and increased heme oxygenase [60]. More recent studies show that a reduction in one of the three subunits of KGDHC and PDHC decrease KGDHC and PDHC activities by one half and increases common markers of oxidative stress [11].

Most animal models that are intended to mimic AD are designed to produce plaques or tangles. For example, the Tg2576 mouse express mutant human AβPP 5.56 to 5.76 times the level of endogenous mouse Aβ/PP [43]. Even in these models, the oxidative stress is an early abnormality. Isoprostanes are specific and sensitive markers of in vivo lipid peroxidation. To determine whether Aβ deposition in vivo is associated with increased lipid peroxidation, the isoprostane levels were determined in the Tg2576 transgenic mouse model of plaque formation. Levels of urinary isoprostanes are higher in Tg2576 than in wild type animals as early as 8 months of age and remain high. A similar pattern occurs in plasma, cerebral cortex, and hippocampus. In contrast, a surge of Aβ levels as well as Aβ deposits in Tg2576 mouse brains occurs at 12 months of age [46]. Mitochondrial dysfunction in the 3xTg-AD mouse model, which has mutations leading to plaques and tangles, is also associated with higher levels of ROS [37].

3xTg mice with the human transgenes AβPP(Swe), PS1(M146V), and tau(P301L) also show early oxida-
tive deficits. At four months, these mice show cognitive deficits and intracellular amyloid oligomers, and at six months hyperphosphorylated tau and extracellular amyloid deposits appear. In striking contrast, glutathione is lower in the 3xTg neurons beginning at one month and continues to be low at 2, 4, and 6 months. NAD(P)H levels are low at 1 month and become lower at 4, 6, and 8 months. Although levels of 8-hydroxyguanine per cell do not increase until six months, cytochrome C immunoreactivity in mitochondria decreases at 4 and 6 months [61]. A proteomic analysis of this mouse reveals a massive deregulation of 24 proteins, one-third of which are mitochondrial proteins mainly related to complexes I and IV of the oxidative phosphorylation system [62]. Thus, considerable evidence suggests that mitochondrial dysfunction and oxidative stress occur early in animal models of AD.

Changes in Calcium in Endoplasmic Reticulum and Mitochondria in Living Cells from AD Patients

Abnormalities in calcium dynamics have been implicated in AD for many years [63]. Calcium dynamics are difficult to assess in brain. Reductions in calcium uptake were initially shown in fibroblasts from AD patients [64]. Subsequent studies reveal that the bombesin (or bradykinin) releasable calcium stores (BRCS) from the ER are exaggerated in fibroblasts from both sporadic AD patients and patients bearing presenilin mutations [65]. Recent studies show that the changes in BRCS in patients bearing presenilin mutations are because presenilins control calcium leak channels and the AD causing mutations block the leakage [66]. However, in AD fibroblasts bearing AβPP mutations, BRCS are diminished [28]. Mitochondrial calcium is also altered in fibroblasts from AD patients [67]. Disturbances of intracellular calcium homeostasis are early events in the pathogenesis of AD in humans and mouse models. In humans, changes in ER calcium occur in patients’ fibroblasts before the patients are symptomatic [68]. Changes occur in transgenic mice long before the pathology. Caffeine evokes a larger response in ER calcium in mutant presenilin-1 knock-in neurons from mice with either a presenilin mutation or in 3xTg mice [69].

Impaired Metabolism/Oxidative Stress Can Induce AD-like Changes in Mitochondria and AβPP Processing

If mitochondrial abnormalities and oxidative stress are upstream effects in AD, then impairing metabolism or oxidants should be able to produce AD-like effects. As described below, different forms of mitochondrial impairment or oxidative stress can mimic AD-like changes. However, one particular oxidant does not produce all of the changes. Similarly, one particular antioxidant does not block all of the effects. Thus, stimulating general mitochondrial function or a broad range of antioxidants may be the most effective strategy for treating AD.

In vitro evidence

Changes in mitochondrial enzymes.

Select oxidants can induce specific changes in metabolic enzymes that are reminiscent of AD and the changes can be very selective. KGDHC has a central role in metabolism and is among the proteins that are most sensitive to oxidants, and measures in AD brain suggest it has undergone post-translational modification. KGDHC is sensitive to diverse oxidants in a variety of cell free and cellular systems including H₂O₂ [7,70–73], hydroxynonenal (HNE) [74], chloroamine [70], sodium hypochlorite [70], t-butylhydroperoxide [75], and acrolein [54]. Catecholamines enhance inactivation of the E3 subunit by the copper Fenton system [76]. Knockout of SOD2 in mice increases ROS production and inactivates E1k of KGDHC [77]. Increases in cellular ROS and diminished KGDHC are seen following hyperoxia in CHO cells [78,79], inhibition of respiration in microglia [7], and elevated MAO expression [5]. Oxidant inactivation of KGDHC by NO can mimic changes in non-genetic forms of AD (i.e., reduced activity with no change in immunoreactivity). On the other hand, peroxynitrite leads to changes reminiscent of those in the brains from the AβPP670/671 family (i.e., reduced activity and diminished immunoreactivity) [7]. Thus, specific oxidants lead to inactivation of KGDHC by different mechanisms and each reflects AD with different causes.

Oxidants can also lead to the selective increase in the other TCA cycle enzymes in a manner that is reminiscent of AD brain. For examples, in cells in which KGDHC is compromised, H₂O₂ increases both message and protein for MDH (Fig. 2) [80]. Activity of MDH is elevated in AD brains (see Figs 1, 2).
Fig. 2. Oxidant treatment can induce diverse changes in metabolic enzymes that resemble the changes in AD brain [80,136]. Cells with a reduction in one of the three subunits of KGDHC (E2k) were treated with H$_2$O$_2$ (see the left half of the figure). KGDHC activity declined while MDH activity increased, which is reminiscent of what happens in AD brains (right half of the figure). Message levels show a similar response to oxidants.

Changes in ER calcium

The molecular basis of the changes in BRCS from the ER is well established for cells from patients bearing presenilin-1 mutations (see above). The cause of the changes in other patients (i.e., the vast majority) is not established. The observations that select oxidants can induce these alterations suggest they may be causative in the majority of AD patients, which do not bear presenilin-1 mutations (Fig. 3A). Different oxidants selectively induced various ROS in distinct patterns and selectively modify BRCS [16]. Of the several oxidants tested, tert-butyl-hydroperoxide is the most specific for exaggerating BRCS (i.e., a change reminiscent of those in patients bearing presenilin-1 mutations). By contrast, hypoxanthine/xanthine oxidase diminishes BRCS, which is reminiscent of the changes that occur in fibroblasts from patients bearing A$_\beta$PP670/671 mutations [28]. On the other hand, NO did not alter BRCS. Thus, these results are consistent with the hypothesis that abnormalities in selective cellular ROS cause AD-related changes in intracellular calcium regulation [16].

A$\beta$ production and plaque formation

Interference with energy metabolism and/or oxidative stress alters A$_\beta$PP processing and A$\beta$ production. Induction of oxidative stress by multiple approaches increases A$\beta$ production and plaque formation. The major enzyme that increases A$\beta$ from A$_\beta$PP is $\beta$-secretase (BACE1). Energy deprivation induces phosphorylation of the translation initiation factor eIF2$\alpha$, which in turn increases the translation of BACE1 [81].

In cultured cells, inhibition of ETC by sodium azide or uncoupling of the ETC by the mitochondrial uncoupler carbonyl cyanide m-chlorophenylhydrazone increases amyloidogenic fragments [82]. Pro-oxidant stimuli significantly increase generation of HNE, which increases production of BACE protein [83]. The stimulation of BACE expression by oxidative stress involves the c-jun N-terminal kinase/c-jun pathway [84]. H$_2$O$_2$ significantly increases BACE1 promoter activity, facilitates $\beta$-secretase activity and A$\beta$ generation. As emphasized previously, oxidants will also decrease KGDHC and increase ER calcium stores in a manner that is reminiscent of cells from AD patients. Thus, upregulation of BACE1 transcription by oxidative stress may contribute to the pathogenesis of AD [85].

Mitochondrial fission and fusion

Mitochondrial fission and fusion may be critical to the pathophysiology of AD. Proteins associated with fusion and fission are reduced (i.e., Drp1, OPA1, Mfn1, and Mfn2) or increased (Fis1) in AD. Manipulations of mitochondrial fission and fusion proteins in neuronal
cells to mimic their expression changes in AD cause 
AD-like changes in mitochondrial fission and fusion.
Oligomeric Aβ causes AD-like changes in mitochon-
drial fragmentation and reduces mitochondrial density 
in neuronal processes. Oligomeric Aβ induces synaptic 
change (i.e., loss of dendritic spine and postsynaptic
density proteins) that correlates with abnormal mito-
chondrial distribution. Dynamin-like protein 1 (DLP1) is 
a regulator of mitochondrial fission and distribution.
DLP1 overexpression prevents oligomeric Aβ-induced 
synaptic loss [86]. NO produces mitochondrial fission, 
in part via S-nitrosylation of dynamin-related protein 1 (Drp1) (SNO-Drp1). Preventing nitrosylation of Drp1 
by cysteine mutation abrogates these neurotoxic events.
SNO-Drp1 is increased in brains of AD patients. Thus, 
nitrosylation of Drp 1 may contribute to the pathogene-
sis of AD [87]. More than 80% M17 cells overexpress-
ing AβPPswt mutant (AβPPswt M17 cells) display 
alterations in mitochondrial morphology and distribu-
tion. AβPP elevated ROS levels, decreased mitochon-
drial membrane potential, reduced ATP production, 
and also caused differentiation deficiency upon retinoic 
acid treatment. Levels of DLP1 and OPA1 are signif-
icantly decreased whereas levels of Fis1 are signifi-
cantly increased in AβPPwt and AβPPswt. The stud-
ies suggest that AβPP, through Aβ production, caus-
es an imbalance of mitochondrial fission/fusion that 
results in mitochondrial fragmentation and abnormal 
distribution [88]. Abnormal mitochondrial distribution 
characterized by elongated mitochondria that accu-
ulated in perinuclear areas in 19.3% of fibroblasts from 
non-genetic forms of AD, which was in marked con-
trast to their normally even cytoplasmic distribution 
in the majority of human fibroblasts from normal sub-
jects (> 95%). DLP1 levels are decreased significan-
tly in sporadic AD fibroblasts. Overexpression of wild-
type DLP1 in sporadic AD fibroblasts rescued these 
mitochondrial abnormalities. Antioxidant treatment re-
versed the AD related changes in DLP1 and in mito-
chondria suggesting that elevated oxidative stress is 
likely the potential pathogenic factors that cause DLP1 
reduction [89].

In vivo changes

Amyloidogenic processing of AβPP

Interruption of metabolism/mitochondrial function 
induces oxidative stress and promotes plaque forma-
tion. Experimental TD is an established model for re-
ducing the activities of thiamine-dependent enzymes 
in brain. TD reduces three key enzymes that are di-
minished in autopsy brains: transketolase, PDHC, and 
KGDHC. TD diminishes thiamine-dependent enzymes 
throughout the brain, but produces a time-dependent 
selective neuronal loss, glial activation, inflammation, 
and oxidative abnormalities including elevated HNE in 
select brain regions. In Tg19959 transgenic mice over-
expressing a double mutant form of AβPP, TD exac-
terates AD-like pathology and enlarges the area oc-
cupied by plaques in cortex, hippocampus, and thala-
mus by 50%, 200%, and 200%, respectively. TD in-
creases BACE1 protein levels by 43% and Aβ levels 
by about three fold. Thus, the induction of mild im-
pairment of oxidative metabolism, oxidative stress, and 
inflammation by TD alters metabolism of AβPP and/or 
Aβ and promotes accumulation of plaques (Fig. 4) [90].
Much more restricted mitochondrial deficits can also 
exacerbate plaque formation. The mitochondrial dihy-
drolipoyl succinyltransferase enzyme (DLST) is a key 
subunit specific to KGDHC (E2k). Experiments test-
ed whether partial genetic deletion of DLST could ac-
celerate the onset of AD pathogenesis. A transgenic 
mouse model of amyloid deposition was crossed with 
DLST+/- mice in which KGDHC activity is reduced 
by about 40%. DLST deficiency increases Aβ plaque 
burden, Aβ oligomers, and nitrotyrosine levels and ac-
celerates spatial learning and memory deficits [91].

Interruption of metabolism by a variety of other 
means, including insulin, 2-deoxyglucose, 3-
nitropropionic acid, and kainic acid, can also promote 
plaque formation in plaque competent mice. Cerebral 
BACE-1 rapidly increases approximately 150% and 
the enzyme remains elevated for at least seven days. 
Aβ1–40 levels are increased approximately 200% of 
control at 7 days [92]. BACE1 is also induced in as-
sociation with oxidative stress in the brains of mice 
subjected to cerebral ischemia/reperfusion [84].

Diminishing antioxidant defense systems also in-
creases plaque formation. Knockout of one allele of 
manganese superoxide dismutase (MnSOD), a critical 
antioxidant enzyme, elevates oxidative stress. Cross-

est of MnSOD deficient mice with plaque competent 
mice increases brain Aβ levels and Aβ plaque burden 
in Tg19959 mice. These results indicate that oxidative 
stress can promote the pathogenesis of AD [93]. Sim-
ilarly, genetic combination of plaque competent mice 
with heterozygous SOD2 knockout shows that mito-
chondrial SOD2 deficiency exacerbates amyloid plaque 
burden [94].

Many of the effects of oxidants may be mediated 
by activation of BACE-1, the rate-limiting enzyme for 
Aβ production. TD increases BACE-1 [90]. Ener-
Fig. 4. Mild impairment of oxidative metabolism exaggerates plaque formation in plaque competent mice. TD diminishes the activities of PDHC, KGDHC, and transketolase, which are also diminished in AD. Just ten days of TD leads to a large exacerbation of plaque formation throughout the brain [90].

Inhibitors of energy metabolism increase levels of eIF2α-P, BACE1, Aβ, and amyloid plaques. Preventing eIF2α phosphorylation blocks the energy-deprivation-induced increase in BACE1 [81].

**Promotion of cell death by AβPP fragments**

Altered AβPP processing has many implications besides plaque formation. These other actions can occur soon after metabolic interruption and can be driven by diminished metabolism and oxidative stress. Abnormal cleavage of AβPP leads to production Aβ and various carboxy terminal fragments of AβPP (CTFs). Although the precise functions of the AβPP CTFs are unknown, they alter membrane currents and calcium homeostasis [95,96], stimulate mitogen activated protein kinase pathways, activate the transcription factor NF-κB, and elevate inducible nitric oxide synthase (iNOS) [97]. Some AβPP CTFs accumulate in the nucleus and bind with Fe65 [97] and CP2 to alter transcription [98]. The translocation of CTF to the nucleus has also been implicated in neuronal death in viro. C-terminal fragments of AβPP translocate to the nucleus of the neurons destined to delayed degeneration following neurotoxic striatal lesions [99]. Genetic knockouts of AβPP attenuate microglial activation and enhance neuron survival in substantia nigra compacta after axotomy [100]. Nuclear AβPP fragments also play a role in the progressive loss of dopaminergic substantia nigra pars compacta neurons following unilateral medial forebrain bundle transaction [99]. Thus, multiple approaches suggest that nuclear translocation of C-terminal fragments of AβPP are important in cell death.

Alteration in AβPP metabolism and nuclear translocation of CTFs of AβPP have been implicated in neuron death following mild impairment of oxidative metabolism. Neurons that will die on day nine of TD accumulate CTFs of AβPP in nuclei of within three days of TD. Western blot analysis of nuclear fractions reveals a 61% increase in CTF12 levels and a 214% increase in CTF15 levels by TD. These findings demonstrate that changes in AβPP metabolism occur in early stages of TD, and they may play an important role in TD-induced selective neuronal loss [101].
Interference with mitochondrial function/energy metabolism promotes tau phosphorylation

AD brains contain neurofibrillary tangles that are composed of abnormally hyperphosphorylated tau protein. Disturbed metabolism/oxidative stress can promote tau phosphorylation.

Carbonyl cyanide m-chlorophenylhydrazone, an uncoupler of mitochondrial oxidative phosphorylation, increases by more than 10-fold the proportion of fibroblasts reacting immunocytochemically with antibodies to paired helical filaments and by 157-fold the proportion of cells reacting with the Alz-50 monoclonal antibody [102]. Glucose deprivation in cultured hippocampal neurons can result in antigenic alterations similar to those seen in AD neurofibrillary tangles [103]. Mitochondrial SOD2 deficiency exacerbates levels of Ser-396 phosphorylated tau. This hyperphosphorylation of tau can be prevented with antioxidant treatment [94]. These observations suggest that the oxidative abnormalities previously described in tissues from patients with AD may contribute to the accumulation of abnormal cytoskeletal materials in this disorder.

Induction of in vivo alterations of glucose metabolism in mice by starvation or intraperitoneal injections of either insulin or deoxyglucose leads to abnormal tau hyperphosphorylation with patterns resembling those in early AD brains [104]. A possible mechanism linking the changes is protein O-GlcNAcylation, a common post-translational modification of nucleocytoplasmic proteins with β-N-acetyl-glucosamine and a process regulated by glucose metabolism. In AD, O-GlcNAcylation is markedly decreased, which negatively correlates with phosphorylation of tau protein. Hyperphosphorylated tau contains 4-fold less O-GlcNAcylation than non-hyperphosphorylated tau, which resembles changes of O-GlcNAcylated and phosphorylated tau in rodent brains with decreased glucose metabolism induced by fasting. Thus, impaired brain glucose metabolism may lead to abnormal hyperphosphorylation of tau and neurofibrillary degeneration via down regulation of tau O-GlcNAcylation in AD [105].

Caspase activation provides another link between tangle formation and impaired mitochondrial function. Inhibition of KGDHC leads to release of cytochrome c and activation of caspase before detectable changes in mitochondrial membrane potentials [106]. Recent studies indicate that caspase activation precedes and leads to tangle formation [107]. Thus, mild impairment of oxidation can lead to caspase activation which may in turn lead to tangle formation.

Thus, the abnormal glucose metabolism that accompanies AD can be linked to tangle formation. In AD patients, the decrease in two major brain glucose transporters (GLUT1 and GLUT3) correlate to the increase in the hyperphosphorylation of tau and to the density of neurofibrillary tangles in human brains [108]. Tangles are also present in Wernicke-Korsafoff patients who like AD patients have reductions in PDHC, KGDHC, and transketolase [25].

MITOCHONDRIAL THERAPIES

The data described above demonstrate that changes in GMO occur early in AD and that that oxidants or impaired metabolism can induce many AD-related abnormalities including changes in metabolic enzymes, ER calcium, A/PP processing, and tangle formation that define AD. However, antioxidant therapies of AD have not been successful. This may be because the multifaceted nature of the changes induced by oxidants. Different oxidants may lead to each of the abnormalities. Thus, a therapy must reverse all of the oxidant-induced changes or remove the source of the oxidants. Stabilization of mitochondrial function or reactivation of mitochondrial proteins may achieve this goal.

Understanding the role of KGDHC in neurodegeneration may help develop new therapeutic strategies. Blocking either upstream or downstream effects will likely be beneficial. Diminishing the oxidative stress related to TD protects neurons [109,110]. Lipoic acid, an essential cofactor for KGDHC, exerts antioxidant activity through a number of mechanisms involving its thiol-disulfide redox couple or metal binding properties. Lipoic acid supplements have been partially effective in aged rats [111,112], in Parkinson’s disease models [112], and in AD [112]. The antioxidant trolox dramatically increases KGDHC activity in cultured cells [72] as does N-acetylcysteine (unpublished results). Lipoamide, lipoic acid, glutathione, and cysteine protect KGDHC from HNE [74]. Inactive KGDHC from H2O2-treated mitochondria can be reactivated with dithiothreitol or glutaredoxin [74,113]. Activation of metabolic pathways may also bypass KGDHC deficiencies. Activation of PDHC extends life span in mouse models of Huntington’s disease [11]. Treatment of AD patients with thiamine or thiamine derivatives [114–116], or bypassing the KGDHC catalyzed step with malate, may be beneficial [117].

Specific oxidants may lead to specific deficits, and reversal of the abnormality may require specific antiox-
oxidants. For example, numerous oxidants were tested for their ability to mimic the exaggerated ER calcium that occurs in families bearing presenilin-1 mutations and in sporadic AD. We postulate that in cells from sporadic forms of AD oxidants induce the change. tert-butyl-hydroperoxide increases BRCs calcium without altering cytosolic calcium (i.e., a mimic of AD changes) (Fig. 3A) [16]. Furthermore, specific antioxidants can block the oxidant effects on ER calcium (Fig. 3A) [118]. This same antioxidant is able to bring an AD line back to control (Fig. 3B). Thus, for this one variable in a culture dish, an antioxidant can reverse one aspect of AD.

Similarly, several studies using plaques as endpoints have shown beneficial effects in vivo of diminishing oxidative stress. In Tg19959 transgenic AD mice, partial deficiency of the mitochondrial antioxidant enzyme MnSOD exacerbates amyloid pathology. The offspring of Tg19959 mice crossed with MnSOD-overexpressing mice have a 50% increase in catalase protein levels, a 50% decrease in levels of oxidized protein, and a 33% reduction in cortical plaque burden compared to Tg19959 littermates. Thus, MnSOD overexpression lowers plaque formation or increases plaque degradation, and markedly attenuates the phenotype in a transgenic AD mouse model [119]. Immunoreactive iNOS is detected in brains of mice with AD-like disease resulting from transgenic expression of mutant human Aβ and presenilin-1. AβPP- or AβPP/PS1-double transgenic mice were bred with iNOS(+/+) or iNOS(−/−) mice. Deficiency of iNOS substantially protected the AD-like mice from premature mortality, cerebral plaque formation, increased Aβ levels, protein tyrosine nitration, astrocytosis, and microgliosis [120]. Treatment with the antioxidant resveratrol can also diminish plaque formation. Resveratrol, a polyphenol found in red wine, peanuts, soy beans, and pomegranates, possesses a wide range of biological effects. Resveratrol diminishes plaque formation in a region specific manner. The largest reductions in the percent area occupied by plaques occur in medial cortex and hypothalamus. Furthermore, resveratrol reduces brain glutathione by 21% and increases brain cysteine by 54%. These changes may be mediated by changes in the vasculature since resveratrol is not detectable in brain under these conditions [121].

A difficulty in developing therapies is that multiple targets and multiple oxidants appear to be involved in AD. An understanding of which radicals are altered is critical to the disease, since a single antioxidant may not work. For example, treatment with α-tocopherol (vitamin E) prevents HNE activation of BACE in neuronal cell lines [83]. However, Trolox (a close analogue of vitamin E) does not protect KGDHC against oxidant challenge [122]. This lack of effect occurs even though the Trolox quenches the ROS production. The results suggest that KGDHC is more sensitive to oxidants than the probe that was used in these studies. α-keto-β-methyl-n-valeric acid diminishes the H2O2 effects on BRCs, while trolox and dimethyl sulfoxide exaggerate the response. Thus, rather than just blocking the downstream events such as BACE1, efforts should be made to block the oxidants that lead to activation. These same oxidants (or related factors) may lead to the abnormalities in the other facets of the disease.

Another critical question about the use of antioxidants is that some of the responses that are viewed as negative may be protective. For example, treatment of mitochondria with H2O2 leads to glutathionylation and inactivation of KGDHC. This inactivation diminishes NADH production, which would reduce subsequent free radical production by the ETC. As the H2O2 declines, KGDHC is de-glutathionylated and reactivated [123]. By analogy, the reduction of KGDHC activity in AD brain may be protective in the short term. However, if it is not reversed it can lead to damaging effects including apoptosis. Regardless of that, BACE1 activation may also be protective initially [124]. Thus, diminishing the causative factor of activation seems a better strategy than just blocking activation. For example, HNE, which increases in AD brains and in animal models of AD including TD, inactivates KGDHC and activates BACE1, which may be protective in the short term [125]. Blocking HNE production would reverse two AD-related changes and diminish the need for the protective response.

The multiple facets of AD suggest that treatments which are less specific and act on mitochondria may correct multiple facets of the disease and be more effective than agents that just block one facet of the disease (Fig. 5). Proposed treatments include: creatine [126], lipoic acid [127], CoQ10 (ubiquinone) [128], Idebonone (synthetic analogue of CoQ10), dimebon [129], Mito Q (CoQ10 targeted to mitochondria) [130], SS31 [131], a cocktail of resveratrol, glucose, malate [117], PGC1-α activator [132], stimulation of sirtuins (NAD-dependent histone deacetylase proteins) [133], activators of nrf2/ARE [134], and PPAR-γ agonists [135]. Thus, a variety of treatments have been proposed to act by protecting mitochondria or stabilizing mitochondrial function to allow the brain to adapt better to disease.
SUMMARY

GMO appear central to many facets of AD (Fig. 5). In those few families bearing mutations in AβPP in which the primary gene mutation promotes Aβ production, it seems likely Aβ must initiate the change. The majority of animals of AD model the situation in these few families (i.e., a gene mutation in AβPP) leads to overproduction of Aβ. However, in all other cases of AD (i.e., the vast majority), the change may be initiated and propagated by mitochondrial deficits. Even in the families with gene mutations in AβPP, mitochondria and oxidative stress appear to be integral parts of the pathophysiology and appropriate therapeutic targets. The changes in mitochondrial function can be plausibly linked to the cognitive deficits as well as the senile plaques and neurofibrillary tangles which are the signature of AD. Any screen of therapeutics should not just look at one facet of AD such as abnormal AβPP processing, but should test whether the treatment ameliorates multiple facets of the disease.

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