Apo E4 Alleles and Impaired Olfaction as Predictors of Alzheimer’s Disease

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Abstract

Alzheimer’s disease (AD) is the most common form of dementia that affects more than 5 million Americans. It is the only disease among the 10 causes of death that cannot be slowed or cured, thus raising the need for identification of early preclinical markers that could be the focus of preventative efforts. Although evidence is escalating that abnormalities in olfactory structure and function precede AD development and early cognitive impairments by one or more decades, the importance of olfaction is largely overlooked in AD, and such testing is not routinely performed in neurology clinics. Nevertheless, research using the olfactory model, has begun to advance our understanding of the preclinical pathophysiology of AD. Notably, an interesting series of studies is beginning to illuminate the relationship between Apolipoprotein E (ApoE) ε4 polymorphism and olfactory dysfunction and late-onset Alzheimer’s disease. In this article, we reviewed present research on the significance of ApoE and olfaction to AD, summarized current studies on the associations and mechanisms of ApoE and olfactory dysfunction, and highlighted important gaps for future work to further advance the translational application of the olfactory paradigm to early, preclinical diagnosis and treatment of AD.

Keywords: Apolipoprotein E; Alzheimer’s disease; Olfactory bulb; Olfactory cortex; Olfactory function; Odor evoked response potentials

Introduction

ApoE4 is a risk factor for AD

Numerous genetic studies have revealed that inheritance of ApoE4 allele increases the risk and rate of progression of late-onset Alzheimer disease, LOAD [1-3]. The human APOE gene exists as three polymorphic alleles—ε2, ε3 and ε4—which have population frequencies of 8.4%, 77.9% and 13.7%, respectively [4]. However, the frequency of the ε4 allele is dramatically increased to ~40% in patients with AD (4). Additionally, 65-80% of AD patients have at least one ApoE4 allele. Inheritance of a single ApoE ε4 variant increases a person’s risk of developing AD by a factor of three in men and four in women, and having two copies of the ε4 allele increases risk by up to 15-fold compared to persons without the ε4 variant [4,5]. Furthermore, ApoE4 inheritance decreases the age of onset of AD [1,6-8].

Apolipoprotein E (ApoE) mediates lipid transport from one tissue or cell type to another [9,10], thus participates in the regulation of lipid homeostasis. In peripheral tissues, ApoE is mainly produced by the liver and macrophages, and mediates cholesterol metabolism in an isoform-dependent manner. ApoE4 is associated with hypercholesterolemia, which is a risk factor for atherosclerosis, coronary heart disease and stroke [9,10]. In the CNS, ApoE is mainly produced by astrocytes, but also can be expressed by oligodendrocytes, and ependymal layer cells [11,12]. Cholesterol is transported to neurons by ApoE via ApoE receptors, which are members of the low-density lipoprotein receptor (LDLR) family [13]. Increasing evidence suggests that under diverse pathophysiological conditions, CNS neurons also express ApoE, although at lower levels than astrocytes [14-17]. The cellular origin of ApoE appears to influence its effects on AD pathology [18,19].

Neurofibrillary tangles and amyloid plaques, two neuropathological hallmarks of AD, are increased in brain samples from ApoE4 carriers as compared to non-ApoE4 carriers [3,7]. Both plaques and tangles appear earlier in ApoE4 carriers as compared to non-carriers of ApoE4. In addition, AD patients with ApoE4 genotype showed widespread degeneration of neurons in areas of the brain related to learning and memory, compared to non-ApoE4 patients [20]. Although various hypotheses have been proposed to explain the relationship between ApoE4 and AD, the mechanism by which ApoE4 leads to AD in humans, if at all, is still unclear. A comprehensive study of the pathophysiological effect of ApoE4 on AD progression is hampered by ethical limitations to sampling of brain tissues in living persons as they progress through preclinical to clinical stages of the disease. The early involvement of the more accessible olfactory pathways by AD biology, therefore offers some hope that the olfactory paradigm can be applied to investigative efforts aimed at elucidating preclinical pathophysiology of disease progression in people with genetic risk of AD. This review examines the scientific premise in support of the validity of the olfactory paradigm for investigation of early and progressive AD in ApoE4 carriers. To provide a template for interpreting data on ApoE-related changes in olfaction in AD, we first provide an overview of basic neurobiology of olfaction and a brief description of methods used for assessment of olfaction. This is followed by the review of extant data that examined whether the olfactory system reliably reflects the effect of apoE variations on AD vulnerability.

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Neurobiology of olfaction

The olfactory system enables us to perceive smell from the environment, and flavors from food. Loss of olfaction is linked to many neurodegenerative diseases, including Parkinson’s disease (PD), fronto-temporal dementia (FTD) and AD [21,22]. The olfactory system begins with the olfactory mucosa, a pseudostratified columnar epithelium in the posterior region of the nasal cavity, from which odor information is projected into the olfactory bulb (OB) at the base of the brain (Figure 1). In all mammals, nasal airflow carries odors into contact with olfactory receptors (ORs) located on the cilia of olfactory receptor neurons (ORNs) in the nasal olfactory mucosa. ORNs express only one OR type [23], and ORNs expressing the same OR innervate up to two glomeruli per OB [24]. Odorant binding with an OR triggers a G-coupled protein–mediated intracellular signaling cascade, ultimately producing an action potential [25,26]. The ORs possess unique chemosensory tuning properties [27,28], that provide a first step at which the olfactory system can sort the limitless number of odorants it may encounter. Action potentials in the ORNs transmit odorant information into discrete zones in the OB whose activation is dictated by nasal airflow [29-31]. These zones in the OB form a “spatial map” of odorant information [32-35] and are modulated by local glomerular layer neurons. This type of organization is believed to be important for the most basic aspects of olfactory perception, including odor perception and discrimination [36-38]. The OB is a six-layer structure in which the sequential stages of odor information processing take place (Figure 1). The axons of ORNs form the olfactory nerve layer of the OB [36-38]. Secondary olfactory neurons, called mitral cells (MCs) located in the mitral cell layer of OB, and tufted cells (TCs) located in the external plexiform layer, all innervate OB glomeruli. Another major cell population in the OB are the granule cells, which are axon-less interneurons organized in patchy aggregated rows in the most central cell layer of the OB (interneuron). The apical dendrites of granule cells form synapsis with MCs and TCs. Granule cells also receive centrifugal input from some secondary olfactory structures and display broader odor-tuning characteristics than the upstream MCs and TCs [39]. Granule cells are mostly GABAergic and glutamatergic. In many mammalian species granule cells are constantly renewed by neurogenesis during adulthood [40-43]. The activity of MCs, TCs, and interneurons in the OB is subject to neuromodulation [44]. The OB receives dense noradrenergic projections from the locus ceruleus, cholinergic input from the horizontal limb of diagonal band of Broca, and serotonergic afferents from the medial and dorsal raphe nuclei. Axons from MC and TC converge to form the lateral olfactory tract, whose distal projections innervate a variety of secondary olfactory structures, including the anterior olfactory nucleus (AON), piriform cortex, olfactory tubercle, the lateral entorhinal cortex, and paramygdaloid complex (Figure 2). These secondary olfactory structures are regarded as the primary olfactory cortex. Cells in the AON cells are highly responsive to odor stimulation [45] and they are believed to aid in intra-nasal odor localization [46]. The piriform cortex is well-known for its role in modifying the processing of odors based upon experience and learning [47-51]. Olfactory tubercle neurons are reliably activated by tasks that assess odor valence, motivated behaviors, and acquisition of rewards, suggesting important roles of the olfactory tubercle in guiding hedonic and valence-dependent responses to odors [52,53]. Neurons within these secondary olfactory structures project into tertiary olfactory structures, which include the orbitofrontal cortex, the insular cortex, and the dorsal hippocampus [54]. Of particular relevance to AD, the entorhinal cortex innervates the hippocampus via the perforant pathway [55]. Additionally, thalamic regions receive olfactory information from several of the secondary olfactory structures, including the AON, piriform cortex, and olfactory tubercle. Olfactory information is also transmitted to the hypothalamus via the amygdaloid complex [56]. These foregoing discussions suggest that odor experience is intertwined with motivated behavior, emotion and cognition through overlap neural circuitry.

Psychophysical Tasks of Olfaction

Generally, psychophysical tasks of olfaction are based upon the presentation of odors to a test subject, followed by examination of the subject’s responses to questions about some characteristics of the odorants presented. Olfactory psychophysical tasks most commonly
measured in studies of olfaction in aging include odor identification, odor memory, odor discrimination and odor threshold sensitivity tasks [57-59]. Other methods which have been employed for clinical assessment of olfaction include surveys of subjective rating of olfactory ability and quality of life [60] and assessments of odor familiarity [61] and hedonics [62].

Odor identification tasks

During odor identification tests, odorants are presented sequentially to test subjects at suprathreshold concentrations, and subjects are required to identify each odor from a list of descriptors. This forced-choice procedure is aimed at minimizing subjects' response bias. Odor identification tests are considered the most advanced type of test depiction of higher-order cortical functions among other psychophysical tests [63-65]. However, a major problem of odor identification test is that it correlates with the verbal abilities of the subject and has a strong cultural precondition, as not all odors are known equally well in various cultural groups. The most widely used identification tests include the 40-odorant University of Pennsylvania Smell Identification Test (UPSIT) [59] and the Sniffin' Sticks Identification Test [58]. However, several other tools are available, such as the 3-item Quick Smell Identification Test, Q-SIT [66]; the 12-item Brief Smell Identification Test, B-SIT [67]; the Smell Diskettes Olfaction Test [68]; the San Diego Odor Identification Test, (SDOIT), which consists of 8 common household odorants presented in opaque jars and includes a picture board to assist odor identification [20]; the Brief (Cross-Cultural) Smell Identification Test [67]; and the Scandinavian Odor Identification Test [69]. Additionally, odor identification can be assessed using flow-dilution olfactometers, such as the T&T Olfactometer [70].

Odor discrimination tasks

In odor discrimination procedures, different odorants are presented to subjects at suprathreshold concentrations, and subjects have to determine which of the odorants smell different [71]. This task examines the ability to distinguish between odors, not to recognize or identify them. In clinical applications discrimination tests are used in combination with identification and threshold tests. The Sniffin' Sticks Test (SST) discrimination task involves a triple-forced-choice procedure. Per triplet, two distracter pens encompass identical smells, while the respective third pen (the clue) contains a different odor. The number of correctly identified clues represents the discrimination score [71].

Odor threshold tasks

The principle behind the threshold tests is that a subject is repeatedly exposed to ascending and descending concentrations of the same odorant and is required to identify the least detectable concentration for the index odor (usually n-butanol or phenyl-ethyl-alcohol) [72]. The widespread use of the latter odorant in odor threshold applications is based on the premise that it more selectively activates olfactory receptors than the trigeminal receptors in the nose [72]. This task is normally associated with the peripheral part of the olfactory pathway [73], and can be measured by means of the T&T olfactometer [70], the Threshold Test of the Sniffin’ Sticks Extended Test [71], and the Smell Threshold Test (STT) [74].

Odor memory tasks

Perhaps due to their relative complexity, odor memory tasks were introduced more recently, compared to threshold and identification tasks. Most clinical applications of odor memory use a recognition task carried out in two stages: the acquisition and the recognition stages. During the acquisition stage, the subject is asked to smell a small set of common household odors at intervals of 30 seconds to control for odor adaptation. This is followed by the recognition stage whereby the subject would be required to recognize the previously presented odors among distractor odors [75]. The total number of correctly recognized, percent of odors in the acquisition stage that was correctly identified, and recognition errors are outcome variables of odor memory tests commonly used in clinical studies [75,76].

Neuroimaging and Physiological Measures of Olfaction

Neuroimaging procedures, including structural and functional magnetic resonance imaging, positron emission tests (PET), single-photon emission computed tomography (SPECT), and electroencephalography (EEG), have been used in neuroscience research to characterize the neurobiology of olfactory processing. The imaging modalities provide good spatial localizations of regions relevant to olfaction, while the EEG applications reveal the sequence of neuronal activations with high temporal resolution [77]. However, due to its relative simplicity, neurobiological investigations of the relationship between ApoE and olfaction have almost exclusively employed EEG methods.

To characterize EEG patterns of olfactory perception, researchers either place electrodes intranasally to acquire odor-induced electrophysiological activity locally in the olfactory epithelium (i.e., electro-olfactography, EOG), or acquire cortical activations during odor exposure, through placement of electrodes on the scalp (i.e., odor event-related potentials, OERPs). These methods provide more objective measures of olfactory function, independent of patients' response bias. Compared to EOG, OERPs are more commonly assessed in clinical populations, and their absence is often a strong indicator of olfactory loss [78]. Odor event-related potentials (OERPs) result from the sequential activation of different brain areas, beginning from olfactory bulbs and tracts and involving the orbitofrontal and insular cortices, along with rostrum-medial regions of the temporal lobe [79]. In most OERP designs, three scalp electrodes placed along the midline – frontal (Fz), central (Cz), and parietal (Pz) – regions allow for examination of relative activations of the cortical fronto-centro-parietal regions by olfactory stimuli [80]. Odor event-related potentials consist of a series of positive and negative voltage deflections, which are related to a set of underlying components. Most components are referred to by a letter (N/P) indicating polarity (negative/positive), followed by a number indicating either the latency in milliseconds or the component's ordinal position in the waveform. For instance, a negative-going peak that often occurs about 100 milliseconds after a stimulus is presented is often called N100 or N1. Typically, N1 is followed by a positive peak, known as the P200 or P2. The specified latencies for OERP components are often quite variable. For example, the P300 (P3) component may exhibit a peak anywhere between 250 ms – 700 ms [81]. These OERP parameters are important variables in studies investigating the influence of ApoE polymorphisms in olfactory functions and in AD.

ApoE4 association with olfactory dysfunction in AD

An overview of results from studies investigating the association between ApoE polymorphisms and olfactory functions is depicted in Table 1. As shown, deficits in odor fluency, odor identification, odor recognition memory, and odor threshold sensitivity have all been associated with inheritance of the ApoE4 genotype in several studies [20,82-86]. These impairments in olfactory psychophysics are observed early in the course of AD, even before the onset of clinical dementia.
<table>
<thead>
<tr>
<th>Study</th>
<th>Sample (n)</th>
<th>Age (years)</th>
<th>Measure</th>
<th>Findings</th>
<th>Other notes</th>
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<tbody>
<tr>
<td>Wang et al. [85]</td>
<td>E4+ (25); E4- (33)</td>
<td>E4+ =70.70 ± 7.36; E4- =71.67 ± 6.28</td>
<td>Cross-cultural Smell Identification Test (CC-SIT)</td>
<td>Significant differences in OERPs between E4+, and E4-. Significant main effect for congruency (F (1,16)=27.08, p &lt;0.001). Significant 4-way interaction between LR, DV, congruency and ApoE4 (F (1,16)=9.41, p=0.007). E4+ had a larger difference in amplitude between congruous and incongruous odor-image pairs at right ventral electrode sites than E4-, t (9)=3.97, p=0.003. ApoE4 carriers had a significantly smaller ERP amplitude difference than non-carriers (F (1,17)=5.12, p=0.037).</td>
<td>(\downarrow) smell identification in ApoE4 group</td>
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<td>Corby et al. [86]</td>
<td>Young and middle-aged subjects with E4 polymorphisms: E4+ (20); E4- (20); E4+ Young=23.9+2.8 Middle age=50.6+2.4. E4- Young 22.2+3.0 Middle age=49.9+3.2.</td>
<td></td>
<td>Chemosensory tests with the butanol odor threshold; SDOIT; OERPs.</td>
<td>Significant effects of ApoE status for P3 latency collapsed across age (F(1,38)=21.91, p=0.001, n2=0.38), with ApoE4+ participants demonstrating shorter latencies than ApoE4+ participants. In the young group olfactory P3 latency was the most significant predictor ((\beta=7.69, p=0.01)) resulting in overall classification rate of 75% (Sensitivity=80%, Specificity=70%). In the middle age group olfactory P3 latency was also the most significant predictor ((\beta=12.54, p=0.001)) resulting in overall classification rate of 80% (Sensitivity=80%, Specificity=80%).</td>
<td>OERP is sensitive to very subtle changes in the brain associated with the ApoE4 allele, even at much younger ages than previously demonstrated. Additionally, the OERP is more sensitive to these changes than traditional tests of olfactory functioning.</td>
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<td>Handley et al. [87]</td>
<td>Sibling group (24); E+ (10); E- (14). Control group (47) E+ (33); E- (14). Control group: 73.17 (61 – 87)</td>
<td>Sibling group mean age (range): 74.08 (59 – 88).</td>
<td>Odor identification performance test</td>
<td>Lower odor identification scores for the sibling group (4.17 \pm 2.20) compared to control group (6.60 \pm 2.22), F(1, 67) 10.42, p&lt;0.01. No difference in odor identification accuracy between E4+ (5.17 \pm 2.44), and E4- groups (5.09 \pm 2.44), p=ns. The group (\times) ApoE4 status interaction was significant, F(1, 67) 6.10, p&lt;0.05. The sibling E4+ group had the poorest mean odor identification scores, lower than control E4+ ((t(22)= -4.12, p&lt;0.01)) and control E4- ((t(44)= -2.33, p&lt;0.05)).</td>
<td>Odor identification deficits may reflect early disease progression in individuals at increased risk for developing the disease.</td>
</tr>
<tr>
<td>Calhoun-Haney and Murphy [83]</td>
<td>Year 1 (baseline): E4+ (22); E- (28). Year 4 (follow-up): E4+ (22); E- (28).</td>
<td>Year 1: E4+ =73; E4- =71; Year 4: E4+ =77; E4- =75.</td>
<td>Butanol odor threshold test; SDOIT</td>
<td>At baseline there was no significant difference between performance on odor identification between the two allele groups, ((F, (1,48)=2.9, p=0.09)), although there was the trend toward poorer performance in the E4+ group. Allele status significantly affected performance on the odor identification measure at Year 4 follow-up, ((F, (1,48)=20.0, p=0.0005)), with performance significantly poorer in E4+ than in E4- individuals. E4+ individuals significantly declined in odor identification performance over time ((F(1,21)=22.9, p=0.0005)), but E4- individuals did not ((F(1,27)=0.41, p=0.52)). There was no main effect of time ((F, (1,48)=0.39, p=0.54)) or allele status ((F, (1,48)=1.7, p=0.20)) on odor threshold. Detection of odor at higher dilution steps indicates better sensitivity. Neither the E4+ nor E4- group showed a significant decline in global cognitive performance over time.</td>
<td>Study demonstrates that normal non-demented elderly adults who carry the E4+ allele and are thus at risk for AD, showed a significant decline over a 4-year time period in performance on odor identification but not on odor threshold, picture identification or the DRIS.</td>
</tr>
</tbody>
</table>
Confirmed AD (38):

Patients n=18; E4+ (10)

Control group (38):

E4+ (23); E4- (15)

Control group (38):

E4+ (24); E4- (14)

Control group (38):

E4+ (19); E4- (19)

MCI patients (converters and nonconverters to AD during 1-9 years follow up):

Converters (39); Nonconverters (109); Controls (63)

Confirmed AD (38):

E4+ (23); E4- (15)

Control group (38):

E4+ (19); E4- (19)

Probable AD (38):

E4+ (24); E4- (14)

Confirmed AD (38):

E4+ (38); E4- (21)

E4+ (15)

AD and Controls (% carriers of E4 alleles):

Patients n=18; E4+ (37.3%)

Controls n=30; E4- (20.8%)

Kjelvik et al. [89]

Wetter and Murphy [84]

Bacon et al. [86]

Sundermann et al. [87]

Gilbert and Murphy [84]

Gilbert and Murphy [84]

ApoE4+ Individuals demonstrated delays in the processing of olfactory information compared to those who are E4-. Olfactory latency appears to be more sensitive for detecting olfactory deficits than the psychophysical measures utilized in this study.

Clinical sample of cognitively impaired, normal-demented patients exhibiting memory impairment shows that ApoE4- and olfactory dysfunction can be combined for prediction of conversion rate to AD.

The analysis of false positive errors in recognition memory for olfactory stimuli may be useful in identifying early deficits in cognition.

The remote memory for olfactory and visual stimuli was not impaired in non-demented E4+ individuals compared to E4 positive controls. These data suggest that the areas of the brain that are affected in non-demented E4+ individuals are not significantly affected in non-demented individuals genetically at risk for AD.

Compliance in olfactory identification was also associated with the volume of several brain structures, particularly hippocampus, more than scores on memory tasks in aMCI and AD.

Olfactory tests distinguished patients with aMCI and early AD from healthy control individuals, and suggested that patients with greater olfactory impairment have increased brain atrophy.
| Corby et al. [97] | Young and middle-aged subjects genotyped for ApoE | ApoE+: Young=22.2 ± 2.0; Middle=49.9 ± 3.2; ApoE-: Young=23.9 ± 2.8; Middle=50.6 ± 2.4 | Butanol odor threshold; SDOIT; OERP's | DRS scores, butanol thresholds, and odor ID scores did not differ between age groups, or ApoE groups, and there were no interaction effects (p>0.05). Young participants produced significantly larger P3 amplitudes than middle age participants (F(1,36) = 7.96, p=0.01, η2 = 0.16). No significant effects of ApoE status on visual or olfactory amplitude (p>0.05), and ApoE status did not interact with age (p=0.05). Significant effects of ApoE status for P3 latency collapsed across age (F(1,36)=21.91, p<0.001, η2=0.38), with ApoE+ participants demonstrating shorter latencies than ApoE+ participants; Significant olfactory P3 latency in young subjects with E4+ participants producing longer P3 latencies than E4- participants (p=0.05, η2=0.31). For middle age participants analysis demonstrated longer P3 latencies (p=0.01, η2=0.47) for E4+ participants compared to E4- participants. In the young group olfactory P3 latency was the most significant predictor (y2=12.54, p=0.001) resulting in overall classification rate of 75% (Sensitivity=80%, Specificity=70%). In the middle age group olfactory P3 latency was also the most significant predictor (y2=12.54, p=0.001) resulting in overall classification rate of 80% (Sensitivity=80%, Specificity=80%). Study demonstrates that OERP seems to be sensitive to very subtle changes in the brain associated with the ApoE4 allele, even at much younger ages than previously shown. The OERP appears to be more sensitive to these changes than traditional tests of olfactory functioning. |
| Landis et al. [77] | 60-66 age group (E4+): n=1121 (29.96); 72-78 age group: n=691 (29.67); 81-87: n=339 (22.44); 90+ age group: n=129 (18.52) | 60-66 age group: 63.06 ± 2.90; 72-78 age group=75.14 ± 2.99; 81-87 age group=83.92 ± 2.38; 90+ age group=91.95 ± 2.40 | Olfactory memory comprised of episodic odor recognition memory and odor identification, based on the Sniffin' Sticks test battery. Other measures: verbal and visual episodic and semantic memory | E4+ had an effect on olfactory memory (standardized estimate=0.08, p=0.01), such that the presence of an E4 allele was associated with poorer olfactory memory. No interaction effects between APOE and age were observed with regard to olfactory memory and verbal or visual episodic memory (p>0.10). This study found that olfactory memory is more sensitive to effects of age and APOE genotype compared to episodic and semantic memory. |
| Murphy et al. [20] | Nondemented older persons E4+ (7); E4- (20) | E4+ =74.00 ± 5.34; E4- =79.43 ± 7.74 | Butanol odor threshold; SDOIT | SDOIT scores were significantly lower in E4+ versus E4 group (p=0.006). Odor threshold task did not reach significance between groups. Study shows the dissociation between odor threshold and odor identification very early in the disease process, even before clinical signs of dementia. |
| Longitudinal study of episodic memory decline | Decline group n=110; E4+ (45) and E4- (65). No decline group n=977; E4+ (279) and E4- (698) | Decline group=65.6 ± 9.9. No decline group = 68.9 ± 10.8 | SOIT | Odor identification scores were lower in participants with episodic memory decline (main effect, F(1,1083)=7.480, p=0.006). Odor identification scores were lower in participants with E4 (main effect, F(1,1083)=4.467, p=0.035). Olfaction was significantly impaired in participants with both E4 and an ongoing episodic memory decline. |
| Longitudinal study of global cognitive decline | E4+ (143); E4- (353) | 73.3 ± 7.1 | A version of SOIT | An odor identification deficit, in combination with older age and ApoE4 predicted decline in global cognitive function. Olfactory deficit can dissociate between benign and non-benign global cognitive development in nondemented, very old E4-carriers, who are at high risk of developing dementia. However, further longitudinal assessment are needed to resolve whether a combination of olfactory deficit, E4, and high age predicts clinical dementia over a more extended time frame. |
The effect of ApoE4 on odor identification in the 75–80 age range is not driven by individuals who receive a dementia diagnosis within a 5-year period after olfactory assessment. The present results suggest that the ApoE gene plays a significant role for the integrity of the olfactory system in non-demented, elderly individuals.

Odor identification performance decreased most strongly in older participants with ApoE4; ApoE4 × age interaction was significant (p=0.033). Although several demographic, cognitive, and health variables affected odor identification, this results suggest that there was a unique effect of the ApoE4 × age interaction on odor identification. Predictor variables accounted for 21.3% of the explanatory variance in odor identification performance.

Study shows a positive linear relationship between adiposity and prolonged olfactory latencies in older adults. When analyzed separately, this relationship remained significant only in older adults who were positive for the ApoE4 allele. There were no significant differences in amplitude between groups.

Latencies were longer in older adults than in young adults, F(1, 49)=106.54, p<0.001. Young adults who were positive for the ApoE4 allele also showed shorter latencies than older adults with the ApoE4 allele, F(1, 23)=236.05, p<0.001; only the ApoE4 positive older adults maintained significant correlations between both measurements of adiposity and P3 latency at Pz, i.e., BMI [r(14)=0.629, p<0.05], waist circumference [r(14)=0.625, p<0.05]

Higher occurrence of olfactory dysfunction among Irish individuals at genetic risk of dementia.

Participants with DS possessing at least one E4 allele performed significantly poorer on the odor identification test compared to those without the E4 allele. The mean number of odors identified in the E4+ group was 4.4, compared to 5.7 in the E4- group, F (1, 33)=4.51, p=0.04.

Individuals with DS carrying the ApoE4 allele, exhibit significantly greater deficits in odor identification than those who are negative for the allele.

The SDOIT test revealed no significant main effects or interaction effects involving ApoE status collapsed across age groups (F(1,54)=8.9 ± 1.9 and 10.1 ± 1.2, p<0.05). Odor ERP task revealed a main effect of ApoE status collapsed across age groups (F(1,54)=8.9 ± 1.9 and 10.1 ± 1.2, p<0.05).

Varying patterns of brain activation were observed over the post-stimulus epoch for E4- versus E4+ individuals on topographical maps. Individuals with the E4 allele demonstrated different ERP peak latencies during identification of olfactory but not visual stimuli.

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### Table 1: ApoE olfaction studies.

<table>
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<tr>
<th>Author (year)</th>
<th>Animal model</th>
<th>Age (months)</th>
<th>Measure</th>
<th>Findings</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nathan et al. [101]</td>
<td>ApoE KO mice; WT C57BL/6 strain</td>
<td>4 months</td>
<td>BFP test; OC test; OCTA test.</td>
<td>ApoE KO mice performed poorly in all three tests compared to WT mice, while they learned the tasks at a rate comparable to WT mice. Latency to find the buried pellet was significantly longer in ApoE KO mice than WT mice. ApoE KO mice did not differentiate the odorant and failed the avoidance test.</td>
<td>ApoE deficiency in ApoE KO mice leads to a deficit in olfactory function, suggesting an important role for ApoE in the olfactory system.</td>
</tr>
<tr>
<td>Nathan et al. [103]</td>
<td>ApoE KO mice; WT C57BL/6 strain</td>
<td>2-3 months</td>
<td>OE Lesion; IHC of OB tissue (0, 3, 7, 21, 42, and 56 days post-lesion)</td>
<td>Slow OMP recovery in the OB in ApoE KO compared to WT mice. Recovery of glomerular area was similarly slower. GAP43 accumulation and restore in the OB were lower in KO mice.</td>
<td>Olfactory nerve regeneration is significantly slower in KO mice, suggesting ApoE participates in olfactory nerve regeneration.</td>
</tr>
<tr>
<td>MC Asey et al. [113]</td>
<td>ApoE KO mice; WT C57BL/6 strain</td>
<td>2-4 months</td>
<td>Ovariectomy. Estradiol replacement. IHC of Olfactory tissues (5, 14, 28 and 49 days after OVX and pellet replacement).</td>
<td>GFAP concentrations were higher in the E2-deprived mouse but did not increase in the E2-replaced group at 49 days. Syn and ApoE concentrations were significantly ↑ by 15% and 25%, respectively, in the E2-replaced compared to the vehicle-replaced group at 5 days, but by 14 days concentrations were equivalent.</td>
<td>Estradiol is able to suppress reactive gliosis. In addition, E2 replacement in OVX mice is associated with transiently higher levels of ApoE and Syn.</td>
</tr>
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</table>
The perikarya and processes of sustentacular (Sus) cells expressed ApoE-like immunoreactivity. The endothelial cells of blood vessels were intensely stained for ApoE in the lamina propria. Cells forming Bowman’s gland also immunostained for ApoE. The ApoE staining in the nerve fascicles was less intense, but was uniformly distributed throughout the core of the nerve bundles. Ensheathing glia, surrounding the nerve fascicles also stained heavily for ApoE.

ApoE is expressed in the adult OE and lamina propria at strategic locations where it could facilitate the differentiation, maturation and axonal growth of the ORN, perhaps by recycling lipids from degenerating ORN for use by growing axons.

Five days of E2 replacement significantly ↑ LRP expression in the hippocampus, OB and neocortex but not in cerebellum. In contrast, E2 treatment ↓ LRP expression in OB.

Hormone therapy (HT) modification of both ApoE and LRP could have wide-spread effects on cellular function given LRP’s manifold signaling functions.

In the absence of ApoE, synaptic recovery in whole bulb samples is substantially delayed compared to WT mice.

ApoE expression in the OE is highly regulated during the entire course of OE reconstitution post injury, and ApoE deficiency in ApoE KO mice leads to delayed recovery of mature OMP+ cells in the reconstituting OE.

ApoE production increases in the injured OE to facilitate maturation of olfactory sensory neurons.

ApoE may be required to elicit the complete effect of estradiol on Syn upregulation.

 Estradiol and ApoE act synergistically to minimize the loss of mature sensory neurons and synapses following ovariectomy.

The OE cultures derived from ApoE KO mice have significantly ↓ neurons with shorter neurite outgrowth than cultures from WT mice. Treatment with either purified human ApoE2 or with human ApoE3, but not ApoE4, significantly ↑ neurite outgrowth. The differential effects of human ApoE isoforms on neurite outgrowth were abolished by blocking the LRP with lactoferin and RAP.

ApoE2 and ApoE3 stimulate neurite outgrowth in OE cultures by interacting with the LRP ApoE4, the isofrom associated with AD, failed to promote neurite outgrowth, suggesting a potential mechanism whereby apoE4 may lead to olfactory dysfunction in AD patients.

Note: AD Alzheimer’s disease, BFP buried food pellet, BrdU bromodeoxyuridine, GAP 43 growth associated protein, GFAP Glial fibrillary acidic protein, IB immunoblotting, IHC immunohistochemistry, LRP low-density lipoprotein (LDL) receptor related protein, OB olfactory bulb, OC odor choice, OCTA odor cued taste avoidance, OE olfactory epithelium, OMP olfactory marker protein, ORN olfactory receptor neuron, O VX over victimized, PCX piriform cortex, RAP receptor-associated protein, Syn synaptophysin (a synaptic marker).

Table 2: ApoE mouse studies.
Non-demented carriers of ApoE4 polymorphism showed significant decline in olfactory processing as compared to individuals without ApoE4 allele [87,88]. Importantly, patients with increased brain atrophy have greater olfactory impairment [89], indicating that olfactory function is further diminished as AD progresses. In family studies, siblings of AD probands had lower olfactory identification scores compared to siblings of control probands [90]. Moreover, within families, siblings with ApoE4 allele showed greater deficits in odor identification tests than siblings without ApoE4 allele. This finding of cosegregation between ApoE4 and olfactory identification deficits suggests that odor identification deficits may reflect early disease expression in individuals at increased risk for developing the disease [90]. Data from longitudinal studies provide additional supporting evidence of the association between ApoE4 inheritance and poor scores in olfactory tests, but more importantly highlights the predictive effect of baseline ApoE4 status on progression of olfactory loss over a 4-year period [83]. A recent study suggests that domains of high-order olfactory functioning, like odor identification and remote memory measured by odor familiarity ratings, may be more impaired in AD E4/E4 homozygotes compared to E3/E4 heterozygotes and E3/E3 homozygotes [91]. These deficits give insight into how the presence of two E4 alleles may differentially affect the progression of AD. Down Syndrome involves trisomy of chromosome 21, where the gene for amyloid precursor protein (APP) is located. Down Syndrome (DS) represents premature aging disorder and autopsy studies show that by age 40, the brains of almost all individuals with DS have significant levels of plaques and tangles, which are definitive neuropathological features of AD. These consistent brain changes notwithstanding, development of AD is not uniform in DS [92]. Interestingly, individuals with Down Syndrome (DS) who are carriers of ApoE4 allele, exhibit significantly greater deficits in odor identification than those who are negative for the allele [93]. It implies therefore, that olfactory tests and ApoE4 genotyping may contribute to improved prediction of AD risks in DS populations.

Olfactory event-related potentials (OERPs) have also demonstrated high sensitivity to subtle changes in olfactory functioning, and to AD and ApoE status [94-96]. A growing number of clinical studies have shown that brain changes associated with ApoE4 allele are captured much earlier in age through OERP recordings than through psychophysical tests of olfaction, thereby suggesting the potential utility of OERP for identification of preclinical stages of AD [82-97-99].

Mechanistic studies linking ApoE4 to olfactory function

Despite consistent evidence of the robust relationship between olfactory dysfunction, ApoE4 inheritance and AD, the mechanism underlying these relationships are not fully understood. However, murine models of ApoE are beginning to illuminate our understanding of the role of ApoE in olfactory structure and function. The results of mechanistic studies in mice and from in vitro biochemical assays are highlighted in Table 2. ApoE deficiency in apoE KO mice leads to deficits in several tasks of olfactory function, suggesting an important role of ApoE in the mice olfactory system [100,101]. Previous studies showed that ApoE is expressed at high levels by a variety of cell types in the olfactory epithelium (OE). In particular, high expression of ApoE in the basal cells and adjoining lamina propria of the OE suggests that ApoE may play a role in the differentiation, maturation and axonal growth of ORNs, perhaps by recycling lipids from degenerating ORN for use by growing axons [102]. Indeed, studies of ApoE KO mice provide support that ApoE plays an important role in olfactory nerve regeneration [103]. Another group of investigators examined the effects of ApoE isoforms on neuronal differentiation and neurite outgrowth in OE explant cultures [104]. They discovered that ApoE2 and apoE3 stimulate neurite outgrowth in OE cultures by interacting with the lipoprotein receptor, LRP. ApoE4, the isoform associated with AD, failed to promote neurite outgrowth, signifying a potential mechanism whereby ApoE4 may lead to olfactory dysfunction in AD patients [104]. Moreover, Nathan et al. [101] investigated the involvement of ApoE in propagating regeneration of OE cells by inducing OE lesions in ApoE and WT mice [101]. The results revealed that ApoE expression in the OE is highly regulated during the entire course of OE reconstitution post injury, and that ApoE deficiency in ApoE KO mice leads to delayed recovery of mature OMP+ cells in the reconstituting OE [105]. Glomerular Synaptophysin (Syn) density, measured by immunohistochemistry, was lower in KO mice at all time points following the lesion [105]. This lower concentration of whole bulb Syn paralleled the slower recovery of glomerular area in KO mice. In the absence of ApoE, synaptic recovery in whole bulb samples was significantly delayed compared to WT mice [106]. This study highlights the important role of ApoE in neuronal differentiation. It is noteworthy that ApoE has also been shown to modulate other molecular factors that are important for neurogenesis, including WNT2 and granulin [107-109]. Some studies have identified a close relationship between estrogen and apolipoprotein E (ApoE) in the central nervous system, and neuroprotective the role of hormone therapy (HT) in several neurological disorders. Estrogen and ApoE function synergistically to minimize the loss of mature sensory neurons and synapses in OB, and OE following ovariectomy [110-113].

Gaps in current studies and future directions

Current diagnosis of Alzheimer’s disease (AD) is based on clinical examination, neuropsychological testing and brain imaging; however, a definite diagnosis can only be made by postmortem examination. Although brain imaging and cerebrospinal fluid biomarkers are applied in patients with mild or questionable symptoms to increase the level of diagnostic certainty, and peripheral bio-fluids are largely investigated, no definitive diagnostic tests are available yet. Biomarkers that reliably predict development of AD would greatly assist preventative and management treatments. This review demonstrates the potential relevance of olfactory system for both biomarker and pathophysiology studies of AD progression. However, many questions must still be answered. It is unclear how to identify and differentiate age-related olfactory changes and olfactory dysfunction caused by disease. Smell loss is also associated with other neurodegenerative disorders, like Parkinson’s disease. Olfactory testing would need to be used with other biomarkers, specific to each disease, or olfactory changes in each disease have to be better specified. While the role of ApoE in olfactory neurogenesis has been reasonably demonstrated in basic studies, it remains uncertain how ApoE promotes AD amyloid and tau pathology in the olfactory system, especially in OB tissue. There are numerous studies examining interactions between ApoE4 protein and Amyloid or Tau proteins in cortical and hippocampal tissues [114]. ApoE is thought to be involved in plaque formation, but how exactly ApoE is involved in pathogenesis of AD is not well understood. The hypothesis gaining widespread support is that ApoE is involved in deposition or clearance of Abeta by direct protein-to-protein interaction. When associated with lipid, ApoE4 bound preferentially to an intermediate aggregated form of Abeta and had higher avidity than did ApoE2 or ApoE3 [115]. The tenability of this hypothesis in olfactory tissues has not been studied. Moreover, mechanistic correlations between ApoE and olfaction in AD to date are performed in animal models, not in humans. There are substantial differences between olfactory systems in rodents and...
humans. Further research is necessary to clarify these uncertainties. Patient-derived olfactory neurons offer an excellent alternative tool to study correlations between ApoE4 and olfactory impairment. Moreover, these olfactory cells can be collected from people at high risk (e.g. ApoE4 carriers), particularly those who display progressive impairments in psychophysical and physiological tests of olfaction. With future accomplishment of these goals in mind, it is reasonable to anticipate that combination of psychophysical, neuroimaging, electrophysiological and molecular studies of olfactory tissues may hold promise for characterization preclinical stages of the disease in people at risk, such as the case of ApoE4 inheritance.

Conclusion

This review summarizes the research showing that ApoE4 is a significant player in olfactory impairment, an early AD symptom. Furthermore, we discussed the mechanistic studies that have been evaluated in ApoE KO mice. Our review stresses the importance of olfactory function as a biomarker of AD, and a potential useful test for prediction of AD development in those with genetic (i.e., ApoE4) risk. Olfactory tests should be incorporated in the assessment of populations at high risk for dementia, like ApoE4 alleles for early recognition of AD and successful intervention.

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