Alzheimer's disease care and management: Role of information technology

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Abstract:
Alzheimer's disease (AD) an ailment that is supposed to affect people in old age. There are evidences that it might affect others also. The number of elders is increasing as the average life expectancy is increasing. AD afflicts its patients with the dementia and AD might increase in malignance over time. People with cognitive disabilities can be overwhelmed through cognitive prosthetics. With the help of information technology we can enhance the quality of life. Significant achievements are possible with an interdisciplinary approach that includes genomic, genetic, technological and therapeutic measures. The combination and coordination of Bioinformatics facilitates generation of various diagnostic tools for the people who are suffering from Alzheimer's disease. These tools help the care providers also. In this article, we emphasize the literature regarding the use of technology and its methodologies to improve the quality of care for the people with Alzheimer's disease.

Key words: information technology; bioinformatics; alzheimer’s disease; electronic medical records

Background:
Alzheimer's disease (AD) was first discovered in 1907 by German neurologist Alois Alzheimer. [1] Alzheimer's disease becomes one of the major hurdles for further survival of elders thereby a many number of people may suffer from AD in the next few decades. Molecular genetics reached human genetics about 1976, when the first human genes were cloned [2] Transgenic methods, 'knock-outs' and ‘knock-ins’ began in about 1986, and in about 1996, database searching became a fruitful way to do genomic research [3] The term ‘genome’ refers to an organism's complete set of genes and chromosomes. The term genomics describes the scientific discipline of mapping, sequencing, and analyzing genomes. [4]

The fact that most diseases do not follow a simple inheritance patterns has led to a significant challenge in the genetic dissection of the complex traits of diseases such hypertension, Alzheimer's disease, schizophrenia and diabetes. Alzheimer’s (AD) affects its patients with a dementia that increases in malignance over time: the older an AD patient is, the worse the dementia is. Dementia is a result of the loss of neurons in the brain that assist in engagement of intellectual activities. The loss of neurons specifically affects the hippocampus, which is a central region for memory operation, and the cerebral cortex. The cerebral cortex is also involved in memory functions, but also works to accomplish reasoning and language functions. A big difference between a normal brain and a brain afflicted with AD is the presence of protein clusters inside and between neurons. The clusters inside are known as neurofibrillary tangles, which consist of a protein named tau. Another type of protein known as beta-amyloid is the protein that exists between neurons. While the presence of tau seems to be proportionate to the degree of dementia experienced by the patient, indicating a possible connection to the cause of AD, it is not unique to AD the way beta-amyloid plaques are in their unique concentrations. These beta amyloid proteins that cluster between neurons are accompanied by the immune system's microglia, reactive inflammatory cells that are thought to remove already damaged neurons and/or the amyloid plaques themselves. [1] They originate from the beta-amyloid precursor protein (bAPP) when a bAPP fragment that is 99 amino acids is cut by gamma-secretase, creating a beta-amyloid peptide, while the amyloid plaques themselves are present in most old people, the high concentration of them in the hippocampus and cerebral cortex in AD patients suggests a role in the neuronal degenerative process.

Memory loss is the most common and well known symptom for Alzheimer's disease. Other symptoms include loss of cognitive abilities, judgment, thinking and disorientation to place and time. Identification loss, depression, confusion, anxiety, fear, frustration, paranoia are also symptoms for Alzheimer's disease. The above symptoms may have different effects on different people. Currently medicines that are available for Alzheimer's disease slow down its progression or help control the
symptoms such as anxiety or sleeplessness. However, there is no available cure for Alzheimer's disease. While the curative approach is certainly crucial to combating the effects of AD, one avenue we might consider looking down is a focus on supplemental measures.

The development of new technologies that could help AD patients cope with loss of mental function might be appropriate, given the nature of the ailment. Developments in information technology could be offer assistance to AD patients in a way that could supplement the loss of biological function with mechanical functions. For example, a computer could be used to keep records of family members to help remind the patient about his or her past. While a desktop PC seems somewhat impractical for this, a computer small enough to fit into someone's eyeglasses, coupled with voice and image recognition technology, could provide AD patients with the kinds of information they need to continue to function. This, along with drugs to at least slow the process, could provide a treatment that could restore a quality of life to the patient in a way that is currently unavailable.

A substantial number of the IMI patients reflected regional hypometabolism similar to AD, suggesting that IMI is likely an early stage in progressive dementia. A large percentage of IMI patients converted clinically to AD within three years of initial study, though they observed impaired memory functioning well before a clinical diagnosis of AD could be made. In addition to potential clinical utility, IMI and PET represent an opportunity to study dementia in relation to brain chemistry at a time when brain pathology is in the process of development.

Genomics of Alzheimer's disease
Genome analysis may be divided into structural and functional genomics. Structural genomics is an initial phase of genome analysis, and has a clear end point which is the construction of high-resolution genetic, physical, and transcript maps of an organism (its complete DNA sequence). This genotypic approach focuses on understanding how genotypic variation gives rise to phenotypic variation, relying on physical and genetic maps and easily-typed DNA sequence polymorphisms. The expression approach (functional genomics) relies on the large collection of partially sequenced cDNA clones. The benefits of the information arising from the accumulation of human gene sequences includes developing systematic ways of finding genes of interest, and their functions; hence 'functional genomics'. The genes cloned and their corresponding DNA sequences provide the tools for comprehensive characterization of the expression patterns of this entire set of genes, and for systematic experimental investigations of the functional properties of their products. Thus, functional genomics, which represents a new phase of genome analysis, makes use of the structural genomics information. The investigation is primarily a systematic approach to elucidate the genome and its functions.

Pathologically, Alzheimer's disease (AD) is associated with generalized degeneration of the cerebral cortical and hippocampal neurons. Cholinergic neurons in the basal forebrain which project to cortex and hippocampus appear to be particularly vulnerable, and to an extent, so are serotonergic and noradrenergic afferents to cortical regions. The extracellular deposition of peptide fragments (amyloid-beta) from the larger membrane precursor protein (APP) is typical in affected brain tissue. Intracellular accumulations of tau-proteins (tangles) are present in many cortical and cortico-limbic regions. [5]

Genetic studies have led to the identification of three genes in which mutations can cause AD: the β-amyloid precursor protein gene located on chromosome 21, presenilin 1 (PS1) located on chromosome 14 and presenilin 2 (PS2) located on chromosome 1. [6, 7, 8] In addition, the E4 allele of the apolipoprotein E (apoE) gene is a risk factor for AD. While mutations associated with APP are extremely rare, the 50 or so mutations associated with PS1 may explain up to half of all cases of early-onset AD. A study which investigated the association of two candidate genes (PS1 and α1-antichymotrypsin (ACT)) with the risk of sporadic Alzheimer's disease on chromosome 14 reported that the frequency of the ACT*A allele was significantly higher in AD patients than in controls and the stratification of the ACT data by PS1 genotypes showed that the risk associated with the ACT*A allele was confined to PS1*1 carriers only. [9]

The two-site haplotype data for PS1 and ACT indicated that the A1 haplotype, carrying the ACT*A and PS1 alleles, was more frequent in Alzheimer's disease patients, and these results may also suggest that there is a possible synergistic effect of these two loci on the risk of AD. In contrast to early-onset AD, there is to date only one genetic factor indisputably linked with late-onset forms of this disorder; the E4 allele of apolipoprotein E. [10] Differences in ApoE genotyping appear to explain differences in patients' responses to drug therapy. With tacrine, a better response was seen in patients with the ApoE E2 or ApoE E3 allele than in those carrying the ApoE E4. The ApoE E4 allele has an inverse relationship with residual brain choline acetyltransferase (the acetyl-choline synthesizing enzyme) activity, and it appears that patients with this genotype may not have sufficient acetylcholine to benefit from a drug which acts as an inhibitor of acetylcholinesterase. However, patients with the ApoE E4 genotype appear to have a better response than other AD patients to treatment with another drug, Servier's S12024 (morpholinyl-2 methoxyl-8 tetrahydro-1, 2, 3, 4 quinoline) which is currently in phase II clinical trials. In fact, this drug had no detectable effect in patients with the other ApoE genotypes. S12024 does not appear to affect the cholinergic system, but rather to facilitate brain noradrenergic and vasopressinergic activity, and increases vasopressin synthesis and release in a dose-dependent manner. There may be a balance between cholinomimetic and vasopressinergic pathways, according to ApoE E4 allele presence or absence. [11]
to design of clinical trials, the important observation may be that alleles that appear to be conclusively associated with a therapeutically relevant phenotype can be used to select a subgroup of patients for clinical trials.

A polymorphic site need not be part of the target for the drug; it only needs to be associated with a response to the treatment. In responsive patients, the selective treatment could be more effective, and associated with fewer or less severe side effects. Furthermore, pre-emptive genotyping aimed at drug-associated genes could mean that fewer drug candidates would fail to reach the market place because of poor toxicity/efficacy profiles in the general population. For example, genotyping of early-onset AD is likely to include the two PS1 and PS2 genes involved in this disease. Predictive and diagnostic tests for PS1 mutations and diagnostic tests for ApoE alleles are already commercially available and other tests are being developed. Thus, genetic testing for AD exists for clinical use, and is likely to be used more often to stratify patients in Alzheimer's disease research, both in trials of preventive products and in tests of new pharmacological treatments. Therefore, predictive and diagnostic genetic testing for these highly penetrant mutations such as PS1 or PS2 may be appropriate for adults from families with a clear autosomal dominant pattern or inheritance, particularly those with a family history of early onset of symptoms. Testing is an option that could be discussed and that could reasonably be accepted or declined by the patients. However, the application of the ApoE test raises concerns, because although the E4 allele is associated with an increased risk of AD, its predictive value for individuals is quite limited. [11] The small increase in diagnostic confidence provided by ApoE genotyping does not justify the burdens of testing; such testing may have value in AD research, but its widespread clinical use is premature until practical benefits outweigh its costs.

**Information technology-Alzheimer’s disease**

Information technology role on Alzheimer's disease has already begun. In 2003, Intel entered into to a consortium with the Alzheimer's association, granting $1 million in Information technology research to be directed towards AD patients. Technology such as sensor networks is being used to study the habits of Alzheimer's disease patient behavior in hopes of finding ways to learn more about AD and to make it more livable. This is an example of how Information technology can work for AD patients. Scientist Hans Moravec has suggested that someday, entire human brains and the consciousnesses they hold will be able to be downloaded into a computer. This would certainly avoid the problem of neuronal deterioration, but it's possible that by the time, we have the technology to move minds into machines, we will know enough about AD to make it a livable or curable illness.

The early stage of AD primarily causes memory problems. At this stage a person can live independently and only requires assistance in remembering certain tasks. Cognitive prosthetics are helpful in aiding a person to remember such tasks. A pager was used to help a person remember the tasks during the early stage of AD. The Pager was able to record 80 letter alphanumeric message and then display the message at scheduled time. The person was able to perform the tasks independently within a week by using the pager. The functional memory of the person had improved after using the pager for six months.

The majority of people with AD usually fit into the category of mild to moderate disease progression. At this stage it is common for patients with AD to move-in with a caregiver. During these stages, persons with AD are prone to forgetting their way home or wandering in the streets. The use of technology for people with AD provides hope that they can live on their own for longer period. Global Positioning System (GPS) is the technology that may prove to be very helpful in these stages. GPS is a tracking device that can be used to identify the location of the patients. A patient can wear a GPS unit and a caregiver can be notified when the patient wanders out of the designated area. Currently GPS devices used to locate Alzheimer’s patients are being used with personal locater devices.

During the moderate to late stages of AD, behavioral disturbances need for more care. At this stage, motes can be programmed to inform caregiver of all the conditions of the patient. Sensors could be placed in the person's bed to monitor weight loss. A combination of sensors placed in chairs and infrared tags detected by cameras could inform the caregiver, if the AD patient has fallen down or is sitting in the chair.

Information technology can be used to help the caregivers with the responsibilities of monitoring the AD patients as well as informing the caregivers about AD and answering their questions. For example, a telephone-based intervention has been used to provide help and support to caregivers. The Project is known as REACH (Resource for Enhancing Caregiver Health) used an interactive voice response (IVR) to provide support and answers to caregivers. It is necessary for the caregiver to receive the support they need in order to perform their job easily. The World Wide Web can easily solve the problem by providing knowledge and support to the caregivers. They can get the validity of online information from the online service called Alzonline.net. [12]

Healthcare providers and caregivers are responsible for constantly monitoring the patient's condition. It is not possible for many caregivers to constantly monitor the patient's health because of lack of medical knowledge or equipment. [13] An important part of care giving consists of taking the patients to the physician, this can add further stress to the care giver and affect the accuracy of physician's judgment about the patient's condition. A healthcare provider might have a better idea regarding a patient’s condition if they could monitor the patient in a natural setting. A telehealth system could be used in order
for the healthcare provider to monitor the patients in their home setting.

Telehealth technology can be useful for rural caregivers. They need not drive long distances. Telehealth application can be used to monitor patients at home and eliminate unnecessary travel. A number of telehealth applications have been used to monitor the patients. Information technology can play a very important role to improve the condition of the people with Alzheimer's disease. It can provide a caregiver with information and support. On the other hand it can engage the patients in many different activities to reduce the caregivers stress. A combination of telemedicine, telecommunication projects and technologies can be used to monitor patients at home and eliminate unnecessary travel. A number of telehealth applications can be used to monitor patients at home and eliminate unnecessary travel. It can provide a caregiver with information and support. On the other hand it can engage the patients in many different activities to reduce the caregivers stress.


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