Ascertainment of Informative Alzheimer Disease Families from the IMAGE Project Registry for Genetic Linkage Analysis Studies


ABSTRACT: Genetic linkage analysis requires the identification and documentation of large families with many affected members present, preferably in more than one generation. The IMAGE Project has been establishing a population-based Alzheimer disease (AD) registry in the Saguenay - Lac-Saint-Jean region of the Province of Quebec. The population of this region has a well-documented ancestry, with reliable genealogical records (since 1842) computerized by SOREP. We have recently begun to investigate the pedigrees of selected probands (definite, probable and possible) from the IMAGE registry in order to identify informative pedigrees for genetic linkage analysis. Interviews were carried out with close relatives of the probands (at least one informant per sibship) to identify secondary AD cases. The questionnaires used pertain to the accuracy of genealogical records, to family medical history and to a retrospective diagnosis of AD for people with cognitive deficits. By these means, we have documented a large extended pedigree in which a total of 15 individuals with cognitive deficits were ascertained over three generations. Of these cases, 7 are still living and there is autopsy confirmation in another one. Computer simulations using the program SIMLINK revealed that this is a potentially informative family for linkage analysis. Horizontal extension of the pedigree to second cousins of the proband is now being carried out. This will render the family IMAGE/1 even more informative in genetic linkage analysis studies.

RESUME: Identification de familles informatives atteintes de la maladie d'Alzheimer a partir du registre du projet image pour des études génétiques de liaison Il est nécessaire d'identifier et de documenter de grandes familles incluant plusieurs membres atteints, de préférence sur plus d'une génération, pour procéder à des études de liaison. Le projet IMAGE est en train d'établir un registre de patients atteints de la maladie d'Alzheimer à partir de la population du Saguenay-Lac-St-Jean, une région de la province de Québec. Les origines ancestrales de cette population sont bien connues et SOREP possède des données généalogiques informatisées fiables sur cette population. Nous avons récemment commencé à investiguer les pedigrees de certains cas index (cas certains, probables et possibles) à partir du registre IMAGE, dans le but d'identifier les pedigrees susceptibles d'être informatifs pour l'analyse de liaison. Nous avons procédé à des entrevues avec les proches parents des cas index (au moins un informateur par fratrie) afin d'identifier les cas secondaires de maladie d'Alzheimer. Les questionnaires utilisés investiguent l'exactitude des données généalogiques, l'historique médicale familiale et un diagnostic rétrospectif de maladie d'Alzheimer chez les individus manifestant un déficit cognitif. Nous avons ainsi identifié une famille étendue comptant un total de 15 individus, répartis sur trois générations, manifestant un déficit cognitif. Parmi ces cas, 7 sont encore vivants et nous possédons un rapport d'autopsie confirmant la présence de la maladie d'Alzheimer chez un autre cas. Une simulation par ordinateur au moyen du programme SIMLINK a montré que cette famille était potentiellement informative pour l'analyse de liaison. Nous procédons actuellement à l'extension horizontale du pedigree, incluant les cousins(ines) au 2e degré du cas index, ce qui va rendre la famille IMAGE/1 encore plus informative pour les études de liaison.


While the etiology of Alzheimer disease (AD) is unknown, various hypotheses have been put forward to account for its development. Factors most often cited include transmissible agents, environmental toxins and genetic defects. Thus far, the risk factor most consistently identified, besides advanced age, is having an affected blood relative. However, the genetic basis...
of AD remains to be elucidated: the mode of transmission of the disease is uncertain and the percentage of familial cases in the general population is not known.

One of the best ways of establishing the involvement of genetic factors in the etiology of a disease is to show an association or a linkage of specific genetic markers with the disease trait. Due to inherent problems in establishing a pre-mortem diagnosis and to the late onset of the disease, genetic analyses have so far proven difficult to perform with Alzheimer cases. As a consequence of the late manifestation of AD, parental samples are rarely available for analysis, and few affected individuals can be sampled, even within the same generation. Furthermore, a definite diagnosis in most cases can be established only at autopsy so that there may be up to a 10-15% error rate in the diagnosis of living affected individuals. Ascertainment of cases from earlier generations is rarely based on well-documented medical records but relies mainly on retrospective family history, with its ensuing uncertainties. Some of these problems can be circumvented by analysing extended, multigeneration pedigrees in which an autosomal dominant transmission is established and where the affected individuals have been well characterized in terms of the clinical course of the disease and their neuropsychological deficits, and in whom neuropathological confirmation of AD was obtained through autopsy. The probands we have identified through the IMAGE Project comply with these requirements. [IMAGE is an acronym for Investigations de la Maladie d’Alzheimer: Epidémiologie et Génétique].

THE IMAGE PROJECT AND GENETIC STUDIES

The IMAGE Project was launched to provide a venue for a comprehensive study of the disease and to overcome the absence of an adequate animal or experimental model. Hence, a population-based registry of AD cases is currently being established in Saguenay - Lac-Saint-Jean (SLSJ), in the Province of Quebec (Figure 1), with the aim of screening as systematically as possible for all living AD cases and of following them longitudinally. The field network operations of IMAGE have been described in detail elsewhere. The objective of IMAGE is to accumulate and integrate data in a multidisciplinary fashion in order to gain better insight into the nature of AD. Ongoing projects within IMAGE are concerned with the clinical/neuropsychological/neuropathological correlations of AD, the population genetics of the field of study, the identification of risk factors involved in the development of AD, the socio-geographical distribution of cases over the area, the study of care-taking and case management in the field, and the molecular genetics of AD in large extended families.

The IMAGE registry is unique of its kind and its strength lies in the possibility for its investigators to access a computerized data-base of all the parish registries of the region, currently dating back to 1842, in collaboration with the SOREP consortium in Chicoutimi [SOREP stands for the Centre interuniversitaire de recherches sur les populations]. The SOREP computerized data-base contains data taken from church and civil records, censuses and various other sources (e.g. social, medical). The

![Figure 1 — The Saguenay-Lac-Saint-Jean (SLSJ) territory, selected as field of study for the IMAGE Project. SLSJ is located north of the city of Quebec; it covers an area of approximately 22,500 km², with a population of approximately 290,000 people.](image-url)
data, representing more than one million records, are divided between a central data-base and numerous sectorial files which have been linked through software that permits computerized reconstruction of families and genealogies. This data-base has information on place and date of birth, place and date of death, as well as residential and professional histories of the population. Anyone having spent his or her life in SLSJ appears on average 10 times in the SOREP records. This data-base will soon be extended into the Charlevoix area just south of SLSJ. Since it is estimated that 70-80% of the gene pool of SLSJ comes from Charlevoix, the IMAGE Project will be in a position to trace the ancestors of cases back to the seventeenth century. In practical terms, this ability to establish ascending and descending genealogies enables us to construct very wide pedigrees, e.g. up to second and third cousins of the AD probands.

Beyond the well-documented ancestry and the computerized genealogical records, the IMAGE Project provides further important advantages. Within the region, large families with an average of 10 to 12 children/generation are the norm rather than the exception. This familial structure makes it easier to find affected living individuals in more than one generation, with the children of the older members of the sibships being approximately the same age as the younger members. Furthermore, large families have closely spaced siblings and additional family members might also be diagnosed as having AD during a short period of time (2 to 5 years).

The whole region of SLSJ was mostly populated from immigrants coming from Charlevoix; indeed, between 1838 and 1911, 75% of the immigrants came from that region. Thus, even if AD turns out to be a heterogeneous disease, one could expect to find “pockets” of homogeneity in terms of the genetic background of the IMAGE cases, that is, familial AD patients with the same genetic defect even in relatives more distant than the second degree. A probable “founder effect” for AD has been described in the descendants of the Volga Germans.11 A similar founder effect in SLSJ would facilitate linkage studies, allowing the work to be carried out from a much larger familial base.

INVESTIGATION OF FAMILY MEDICAL HISTORY

We have recently begun to investigate the pedigrees of selected probands (definite, probable and possible) from the IMAGE registry in order to identify informative pedigrees for genetic linkage analysis. Two or more of the following criteria were applied, in combination, in the choice of families to be evaluated: 1) affected subjects with large sibships; 2) one or more probable AD cases (based on NINCDS-ADRDA criteria12); 3) neuropathological confirmation13 (also Ball, personal communication) on at least one case; 4) probable cases in two generations; 5) numerous first, second or third degree relatives having reached the age of risk. Most of the families we have investigated so far have fulfilled at least three of the above criteria.

Interviews were carried out with close relatives of the probands to identify secondary cases of AD. For each proband, one sibling and one member of each first cousin sibship were interviewed. In addition, any nephews or nieces who had reached the age of risk for AD were also investigated. Each informant was asked to provide information about all his/her first degree relatives (parents and siblings and, if necessary, about unmarried aunts and uncles).

In the first questionnaire administered, the informants were asked to verify the pedigree structure of their own sibship and to supply information concerning missing individuals. Although all the persons who have resided in the SLSJ region should be entered at least once in the SOREP genealogical registry, it is conceivable that some relatives of AD probands might have spent their entire lives outside the region. Thus, we were able to confirm and complete the pedigree structure obtained from the genealogical records of SOREP.

The second questionnaire pertaining to the family medical history was derived from a questionnaire developed and validated by Breitner.14 At this stage of the project we were mainly interested in the presence of memory loss, dementia, or AD in the relatives of the proband. If, through this questionnaire, a relative was identified as having cognitive deficits, the caregiver or the best possible informant was identified for this individual. If the suspected dementia case was not in the IMAGE registry, either the case and/or the caregiver was asked permission to undertake the IMAGE registry diagnostic assessment protocol.

If the suspected dementia case was deceased, the person who best knew the deceased near the end of his/her life was administered a questionnaire to establish a retrospective diagnosis of AD. This questionnaire, based on clinical experience, is divided into two sections: the first one seeks to identify exclusion criteria, while the second one concerns the presence of characteristic cognitive and functional impairments and their rate of progression. The questions in the first section pertain to causes of

Figure 2 — Pedigree structure of the AD family IMAGE/1. Proband is identified by an arrow.
Table 1: Simulation Analysis to Estimate the Power of Demonstrating Linkage with Pedigree IMAGE/1 using a Di-allelic Marker Locus (0.71, 0.29)

<table>
<thead>
<tr>
<th>True recombination fraction</th>
<th>Estimated mean and maximum lod scores</th>
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<tbody>
<tr>
<td></td>
<td>MEAN</td>
</tr>
<tr>
<td>0.00</td>
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<tr>
<td>0.05</td>
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<tr>
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<tr>
<td>0.15</td>
<td>.601</td>
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<tr>
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<td>.447</td>
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Simulation was performed with the SIMLINK program. One thousand replications of the pedigree shown in Figure 2 were analyzed.

Dementia other than AD (e.g. stroke, severe head injury, psychiatric diseases, other neurologic diseases, alcoholism, and drug abuse). In the second part, questions bear upon the presence of memory, language and orientation problems, as well as the extent of difficulty in performing the activities of daily living. The answers to this questionnaire were evaluated by a neurologist who then rendered a diagnosis of either probable AD, dementia of other cause, or absence of dementia.

To date we have completed the evaluation of six extended pedigrees. For all the families, we were able to contact at least one informant, per sibship, who was still residing in SLSJ. When required, we were also able to verify the presence of suspected dementia in an individual with additional informants. Although the information we had initially on some families suggested that they might provide interesting material for linkage analysis studies, full ascertainment in a short period of time allowed us to show that these were families with a low informative potential and that they need not be a priority. Only a set-up such as the one provided by the IMAGE-SOREP association (rapid genealogical reconstruction, families mostly confined to a small geographical territory, etc.) allows such a quick and rapid evaluation.

Identification of an Informative Pedigree

The following criteria were used for the selection of potential families for participation in genetic linkage studies: 1) clear evidence of autosomal dominant transmission; 2) clinical diagnosis of probable AD; 3) at least one neuropathological confirmation per family; 4) the presence of several affected and unaffected individuals available for genotyping and who are willing to cooperate.

Of the six documented pedigrees, we have identified one large family (IMAGE/1) which fulfills the criteria set above. The proband is a probable AD case in a family presenting with a history of cognitive deficits in his father and in his sister. Investigation of the family medical history has shown that a total of 15 individuals over three generations have or have had cognitive deficits (Figure 2). In the generation of the proband, there is one definite, three probable and two possible cases, as well as two putative cases who have yet to see a neurologist. In addition, four of the deceased cousins of the proband were diagnosed as AD by the retrospective diagnosis questionnaire. To date, of the 37 individuals in the generation of the proband surviving
beyond the age of 60, 12 have been affected by cognitive deficits. The mean age of onset in this family is 69.7±8.0 years.

We have assessed the informativeness of this pedigree for genetic linkage studies by performing computer simulations using the program SIMILINK. This program randomly assigns appropriate genotypes to all individuals available for sampling and then uses the LIPEP program to calculate lod scores. This exercise enables the investigator to estimate the power afforded by a given family in subsequent analyses. The results of these analyses can be used to determine the minimum number of families required for linkage studies.

The SLSJ area, even if only 5% of all cases are familial in nature and among the descendants, the ability to link together all these sibships will render the family IMAGE/1 even more informative.

Based on an estimated collection of 500 AD cases in the SLSJ area, even if only 5% of all cases are familial in nature and appropriate for linkage studies, we expect to be working from a base of 10 to 20 families. Given that the families described above are the norm rather than the exception in SLSJ, the population of the IMAGE registry provides a formidable resource to molecular genetics studies.

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REFERENCES


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