Canadian Consensus Guidelines on Use of Amyloid Imaging in Canada: Update and Future Directions from the Specialized Task Force on Amyloid imaging in Canada

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ABSTRACT: Positron emission tomography (PET) imaging of brain amyloid beta is now clinically available in several countries including the United States and the United Kingdom, but not Canada. It has become an established technique in the field of neuroimaging of aging and dementia, with data incorporated in the new consensus guidelines for the diagnosis of Alzheimer disease and predementia Alzheimer’s disease–related conditions. At this point, there are three US Food and Drug Administration– and European Union–approved tracers. Guided by appropriate use criteria developed in 2013 by the Alzheimer’s Association and the Society of Nuclear Medicine and Molecular Imaging, the utility of amyloid imaging in medical practice is now supported by a growing body of research. In this paper, we aimed to provide an update on the 2012 Canadian consensus guidelines to dementia care practitioners on proper use of amyloid imaging. We also wished to generate momentum for the industry to submit a new drug proposal to Health Canada. A group of local, national, and international dementia experts and imaging specialists met to discuss scenarios in which amyloid PET could be used appropriately. Peer-reviewed and published literature between January 2004 and May 2015 was searched. Technical and regulatory considerations pertaining to Canada were considered. The results of a survey of current practices in Canadian dementia centers were considered. A set of specific clinical and research guidelines was agreed on that defines the types of patients and clinical circumstances in which amyloid PET could be used in Canada. Future research directions were also outlined, notably the importance of studies that would assess the pharmaco-economics of amyloid imaging.
Since 1989, four Canadian Consensus Conferences on the Diagnosis and Treatment of Dementia (CCCDDT) have been held. Of those, the 4th CCCDDT1 focused on updating previous diagnostic approaches to Alzheimer’s disease (AD) taking into account revised diagnostic criteria proposed by the International Working Group5,6 and the recommendations made by the National Institute on Aging - Alzheimer Association (AA) workgroups.7-9 The focus of CCCDDT4 and of two accompanying papers10,11 was largely on neuroimaging and other biomarkers; nine recommendations on amyloid positron emission tomography (PET) were made, most of which are now outdated. In this paper, we aimed to provide updated Canadian guidelines to dementia care practitioners on proper use of amyloid imaging as formulated by the Specialized Task force on Amyloid imaging in Canada (STAC). We also wished to generate momentum for the industry to submit a new drug proposal to Health Canada so that this revolutionary technique could become part of our clinical armamentarium under precise and well-defined indications.

AMYLOID IMAGING: FROM 2004 TO TODAY

Since Klunk’s publication on Pittsburgh compound B (PiB) in 2004,12 PET using amyloid ligands has revolutionized AD research, leading to improved models of disease pathogenesis, providing evidence for a long preclinical disease phase, and stimulating therapeutic trials aimed at delaying or preventing the symptomatic phase of AD.13,14 For example, amyloid imaging has served as a secondary outcome measure in AD clinical trials with disease-modifying agents such as the anti amyloid monoclonal antibodies bapineuzumab and solanezumab.15,16 Brain amyloid reduction and slowing of cognitive decline were found after 1 year of treatment with aducanumab, a human immunoglobulin G1 monoclonal antibody against a conformational epitope found on amyloid beta (Aβ) 17 Finally, its added value also lies in subject selection for clinical trials given an approximate 15% amyloid negative rate.

Beyond research, amyloid imaging has demonstrated great potential as a diagnostic tool largely because it allows in vivo detection of amyloid plaques, a core pathologic feature of AD.12 It is now an established technique in the field of neuroimaging of aging and dementia, with data incorporated in the most recent consensus guidelines for the diagnosis of AD7 and predementia AD-related conditions.8,9 There currently are three US Food and Drug Administration and European Union–approved, fluorine-18–labeled, tracers available for clinical use: florbetapir since 2012,18 flutemetamol since 2013,19 and florbetaben since 2014.20 Converging evidence on the diagnostic utility of that technique has rapidly accumulated.21-29 Still, no tracer has been approved for clinical use in Canada so far.

AMYLOID PET IN COGNITIVELY NORMAL ELDERLY INDIVIDUALS AND VARIOUS CLINICAL POPULATIONS

Cognitively Normal Elderly Individuals

Cognitively normal elderly individuals (CNs) show elevated PiB binding in 10% to 34% of cases, a proportion similar to observed rates of amyloid pathology in autopsy studies.30 Increasing age and the presence of the apolipoprotein E ε4 allele are the major predictors of PiB positivity in CN.31,32 Recent findings in persons without dementia or mild cognitive impairment33,34 suggest that amyloid deposition is associated with very subtle cognitive deficits, especially among apolipoprotein E ε4 carriers.35 Interestingly, recent data suggest that the cooccurrence of Aβ and neurodegeneration (hippocampus volume and glucose metabolism) is associated with cognitive decline in CNs.36 The significance of a positive amyloid scan in CNs still remains uncertain, but cross-sectional studies have shown “AD-like” brain changes (hippocampal and tempo-parietal atrophy),37 whereas early longitudinal data have strengthened the notion that many (although probably not all) are in a “preclinical” phase of AD.9 Recently, a group of researchers compared the ability of molecular markers for AD, including amyloid imaging and cerebrospinal fluid (CSF) biomarkers (Aβ1-42, tau, ptau181, tau/Aβ1-42, ptau181/Aβ1-42), to predict time to incident cognitive impairment among cognitively normal adults aged 45 to 88 years and followed for up to 7.5 years. Results indicated that all AD biomarkers predicted incident cognitive impairment and supported the hypothesis that biomarkers signal underlying AD pathology at least several years before the appearance of dementia symptoms.38 From a diagnostic perspective, the significant number of amyloid-positive CNs emphasizes that amyloid positivity is not synonymous with AD, and that amyloid scans cannot replace a detailed clinical evaluation. At present, there is no clinical indication for amyloid imaging in CNs, though this will remain an area of active research in coming years, particularly with the advent of amyloid-lowering therapies, which might be most effective if initiated in the presymptomatic disease stage.39,40

Mild Cognitive Impairment

Current data in mild cognitive impairment (MCI)8,33,34 indicates that amyloid imaging provides prognostic information, presumably by identifying patients with underlying AD pathology.31 As a group, 52% to 87% of MCI subjects show elevated PiB binding in a regional distribution similar to that observed with AD.30 In longitudinal studies, 1-year conversion rates to AD range from 33% to 47% in PiB-positive MCI subjects, whereas virtually no conversions are seen in PiB-negative subjects.32 In a longitudinal study,43 the authors compared baseline amyloid deposition between MCI converters and nonconverters in 31 subjects followed over 3 years. The conversion rate was 82% in those with increased PiB uptake, but only 7% in PiB-negative subjects. Results from the Australian Imaging Biomarkers and Lifestyle study of aging on 87 participants with MCI (age, 73.7 ± 8.27) showed that 59% had progressed to probable AD over 3 years.44 Multivariate analysis showed β-amyloid imaging as the variable most strongly associated with progression. Almost all amnestic MCI subjects with a positive amyloid scan developed AD. Altogether, the literature clearly suggests that PiB-positive amnestic MCI patients are likely to have early AD, and amyloid imaging will
help in risk stratification and selection of patients who may benefit from experimental disease-specific therapies. Similar to normal aging, positivity of both Aβ and neurodegeneration biomarkers in MCI can further stratify risk of imminent conversion to dementia.

Alzheimer’s Disease

Most studies in AD have found high (90% or greater) PiB-PET positivity, with a pattern that closely mirrors the distribution of plaques found at autopsy. Tracer binding is diffuse and symmetric, with high uptake consistently found in prefrontal cortex, precuneus and posterior cingulate cortex, followed closely by lateral parietal and lateral temporal cortex, and striatum. Studies in atypical clinical presentations of AD have shown that amyloid deposition is more common in the logopenic variant of primary progressive aphasia than in nonfluent or semantic variants, supporting the hypothesis that logopenic variant of primary progressive aphasia is often associated with underlying AD. Others have detected high PiB binding in patients with posterior cortical atrophy, a visuospatial/biparietal clinical syndrome often caused by AD. Much like 18F-fluorodeoxyglucose-PET (FDG-PET), amyloid imaging will probably not add value to the diagnostic workup of patients with straightforward clinical AD, but is likely to be useful in patients with atypical complex presentations or early age-of-onset dementia (see the following two sections).

The Frontotemporal Lobar Degeneration Spectrum of Disorders

Considering that the frontotemporal lobar degeneration spectrum of disorders (FTLD) and AD are the leading causes of early age-of-onset dementia, that distinguishing the two during life can be clinically challenging but is also important, and that Aβ plaques are not part of the FTLD pathologic spectrum, several authors have argued for a valuable role of amyloid imaging in the differential diagnosis of these conditions. Small case series have reported low rates of PiB positivity (0% to 15%) and flurbetaben positivity (9%) in FTLD. Recently, results from the largest study currently available on the diagnostic utility of amyloid PET in FTLD showed, in 62 AD and 45 FTLD patients, that PiB visual reads had a higher sensitivity for AD than FDG-PET, with similar specificity. PiB outperformed FDG in classifying patients with known histopathology, and visual reads showed higher inter-rater reliability and agreement than for FDG, suggesting it was the more accurate and precise technique.

Complex, Atypical Patients With an Uncertain Diagnosis

The clinical diagnosis of AD has only moderate sensitivity and specificity when compared with the pathological cause of dementia as determined at autopsy. Misdiagnosis rates are even higher in complex, atypical patients with an uncertain diagnosis, approaching 30%. A growing body of literature supports the clinical utility of amyloid imaging for the differential diagnosis of atypical patients with an uncertain diagnosis. This has major implications for a cohort of individuals who are often younger than 65 years of age and still active in the workforce. Indeed, several dementia experts have argued that an accurate diagnosis helps direct therapy (i.e. avoid unnecessary or undesired cholinesterase inhibitors or memantine prescriptions), determine a better care plan (which considers patient safety and minimizes the risk of preventable complications), and enable patients to participate in legal and financial planning.

Recently, a group of Canadian researchers investigated the clinical utility of amyloid PET in the differential diagnosis of atypical cases and its impact on caregivers in the context of a tertiary memory clinic. Using the amyloid tracer 11C-NAV4694, they prospectively scanned 25 patients (mean age, 59.3 years; standard deviation [SD], 8.8; and mean Mini-Mental State Examination, 21.6; SD 6.2) with an atypical syndrome as determined by dementia experts. All patients had a full workup (i.e. history, examination, blood tests, neuropsychology, magnetic resonance imaging [MRI], and FDG-PET), yet no certain diagnosis could be arrived at following that investigation. Amyloid PET was either positive or negative based on qualitative and quantitative reads by two qualified independent expert nuclear medicine specialists. Physicians rated whether amyloid PET was associated with a change in diagnosis and altered management. They also reported their degree of confidence in diagnosis before and after amyloid PET. Caregivers were met 3 months after having been told of the diagnosis and completed a 21-item Likert scale questionnaire along with a 1-hour interview designed to assess the impact of the amyloid scan. The cohort was 48% amyloid positive and 52% amyloid negative. Inter-rater reliability was 100%. Amyloid PET was associated with a diagnostic change in 36% (9/25) of cases (24% changed from AD to non-AD and 12% from non-AD to AD). There was a significant increase (40%) in diagnostic confidence following the scan. Altogether, this study corroborated recent findings and suggested an additive role for amyloid PET in atypical cases with an unclear diagnosis despite the detailed workup of a tertiary memory clinic. Amyloid PET increased diagnostic confidence and generated significant alterations in management in almost three-quarters of cases. Furthermore, the overall process was very well received by caregivers, reducing anxiety and depressive symptomatology as well as increasing quality time spent with their loved ones.

Other Clinical Conditions

Other clinical conditions studied with amyloid PET include vascular cognitive impairment, cerebral amyloid angiopathy, Parkinson’s disease dementia, and dementia with Lewy bodies (DLB). In one study on vascular cognitive impairment, authors found that 69% of patients were PiB-negative. High PiB binding rates were found in nondemented patients with cerebral amyloid angiopathy. Most studies showed higher amyloid plaques in DLB than in Parkinson’s disease dementia or nondemented PD patients and, in some, PiB positivity was associated with more rapid disease progression. The high frequency of plaques and high rates of positive scans in DLB suggest that amyloid PET is unlikely to be helpful in differentiating DLB from AD.

AMYLOID PET IN CLINICAL PRACTICE TODAY: ARE WE READY YET?

The translation from the research setting into the clinic has progressed steadily. For example, a group recently determined the sensitivity and specificity of amyloid PET with flutemetamol using neuropathologically determined neuritic plaque levels and showed high sensitivity and specificity in an end-of-life population. In an effort to guide clinicians, the Society of Nuclear Medicine and Molecular Imaging (SNMMI) and the AA jointly published criteria for the appropriate use of amyloid
PET. Over the past 3 years, almost 10 reports on the practical clinical applications of amyloid imaging have been published, and their conclusions indicated a significant role in orienting treatment (i.e. deciding whether to initiate or discontinue AD symptomatic medications) and a positive impact on caregivers. Finally, much is expected from the soon-to-be launched Imaging Dementia – Evidence for Amyloid Scanning (IDEAS) Study, a $100 million open-label longitudinal cohort effort on approximately 18,500 US Medicare beneficiaries. In this venture, diagnostically uncertain cases of MCI and atypical dementia will be scanned to determine whether knowledge of amyloid status leads to significant changes in patient management and if this translates into improved medical outcomes.

In light of the outstanding worldwide momentum surrounding the utility of amyloid imaging in clinical practice, and the fact that Canada lags behind on approval of the technique by health authorities, we first aimed to provide updated Canadian guidelines to dementia care practitioners on proper use of amyloid imaging. Second, we wished to generate momentum for the industry to submit a new drug proposal to Health Canada.

METHODS

The STAC met in Montreal, QC, Canada on May 14, 2015, to update the 2012 Canadian consensus guidelines, which incorporated nine recommendations on amyloid PET, most of which are now outdated. The meeting included all members of the STAC (a group of local, national, and international dementia experts and imaging specialists; see Appendix A) as well as consensus meeting participants (clinical and academic; see Appendix A), and leading representatives from the molecular imaging and imaging software industries. Technical and regulatory considerations pertaining to Canada were discussed with a member of Health Canada who attended the meeting. Peer-reviewed and published literature between January 2004 and May 2015 was searched before the meeting. A survey of current diagnostic practices in Canadian dementia centers was also presented to allow focused discussions on Canadian medical practices. In brief, we discussed indications and, just as importantly nondications, of amyloid imaging based on clinical and nonclinical scenarios with variables including symptoms (typical and atypical), clinical settings, clinical contexts, evidence of cognitive deficits, family history, knowledge of AD genetic risk, and age.

Although previous CCCDDTs had used the evidence grading system developed by the Canadian Task Force on Preventive Health Care, for this iteration we attempted to follow, where possible, the Grading of Recommendations Assessment, Development and Evaluation system in keeping with current recommendations for the conduct of consensus conferences. Each participant was allowed to take part in the discussion. Consensus was defined as 80% or more conference participants being in agreement with a recommendation. Partial consensus was defined as 60% to 79% being in agreement. Recommendations reaching consensus are listed in Tables 1 and 2. Recommendations reaching only partial or no consensus are only mentioned in the Discussion section.

RESULTS

A set of updated guidelines were agreed on that define the types of patients and clinical circumstances in which amyloid PET could be used in Canada (Table 1). Future research directions were also outlined, notably the importance of studies that would assess the pharmaco-economics of such diagnostic procedure (Table 2).

Survey of the Availability and Use of Biomarkers in Canada

Three months before the meeting, a survey was sent to all Canadian dementia centers. A total of 27 respondents (British Columbia, 3; Alberta, 3; Saskatchewan, 1; Ontario, 7; Quebec, 10; Maritimes, 3) provided answers mainly to two questions: (1) How many early-onset atypical dementia cases do you see per month, and (2) Which advanced diagnostic techniques do you use in practice beyond clinical history, physical examination, standard laboratory tests, and basic computed tomography imaging (i.e. MRI, hippocampal volumetry, molecular imaging, CSF Aβ42 and tau). Results indicated that an average of 5.8 (SD: 5.4) early-onset atypical dementia cases per clinician were seen per month. More than 80% use MRI. Only clinicians from Quebec use FDG-PET in clinical practice, whereas single-photon emission computed tomography is used by a majority of clinicians outside Quebec, because of regionally specific provincial reimbursement issues. Only 20% of clinicians use CSF measures. Less than 11% prescribe acetylcholine-esterase inhibitors to atypical cases with uncertain diagnoses. Finally, in 85% of cases, clinicians reported that knowledge of the amyloid status of their atypical patient would change their therapeutic approach. Some clinicians also reported sending patients to the United States for an amyloid scan.

DISCUSSION

Amyloid PET is now an established neuroimaging technique with data incorporated in the consensus guidelines on AD and predementia AD-related conditions. So far, three different fluorine-18–labeled agents have been approved for clinical use in a variety of jurisdictions around the world. Despite these major advancements, Canada is not yet one of those jurisdictions. Publication of appropriate use criteria by the AA and the SNMMI has paved the way for other countries to adopt a standardized model reinforcing proper use of amyloid imaging in medical practice. This paper is derived from discussions of the STAC, a group of local, national and international dementia experts and imaging specialists who revisited the scenarios in which amyloid PET could be used appropriately in Canada. The final product is an updated set of guidelines to the 2012 CCCDDTs4 effort, which also factors in the results of a survey of current practices in Canadian dementia centers. Furthermore, it is tailored to the Canadian reality and wishes to promote future development of amyloid imaging in our country. We hope this paper will generate momentum for the industry to submit a new drug proposal to Health Canada so that regulatory bodies approve the technique and discussions about provincial reimbursement can begin.

Canadian Challenges

This consensus effort allowed members to realize that only two Canadian provinces (Alberta and Quebec) have access to neurological FDG-PET examinations, a well-established molecular imaging technique in the field of dementia. Indeed, use of FDG-PET has been supported by a wealth of literature (see Bohnen et al for a review). In the diagnosis of AD, authors have showed that FDG-PET is superior to a baseline clinical evaluation and similar to an evaluation performed 4 years later. The addition of
Table 1: Recommendations for clinicians on behalf of the Canadian Consensus Conference on the Use of Amyloid Imaging

1. Amyloid imaging represents a promising technique in the evaluation of dementia for which much has been learned over the past decade. It is not currently approved for clinical use in Canada. When it becomes available to Canadian clinicians, it must not be considered a routine test:
   A. In accord with Appropriate Use Criteria for Amyloid PET, we recommend its use in patients with objectively confirmed cognitive impairments in whom there is diagnostic uncertainty after a comprehensive clinical evaluation (mental status testing, laboratory tests, and structural brain imaging using MRl), and in whom knowledge of Alzheimer’s disease status is expected to provide a more precise diagnosis and alter management;
   B. Clinicians who wish to obtain amyloid imaging should refer patients to dementia centers with an expertise in this technique, i.e. dementia experts; with substantial clinical experience and practice in dementia care who work in conjunction with nuclear medicine specialists qualified in amyloid imaging;
   C. We strongly recommend against the use of amyloid imaging in cognitively normal individuals or for the initial investigation of cognitive complaints.

2. Physicians should be cautious about interpreting the significance of amyloid test results, i.e. used in isolation this test cannot diagnose Alzheimer’s disease; MCI or differentiate normal from abnormal aging. When faced with such situations, we recommend they consult with dementia centers with an expertise in this technique.

3. At present, there is no clinical indication for amyloid imaging in:
   A. Attempting to differentiate AD from other aβ-associated dementia (e.g. dementia with Lewy bodies, cerebral amyloid angiopathy);
   B. Attempting to differentiate between AD clinical variants (e.g. classic amnestic AD vs. posterior cortical atrophy or logopenic variant of primary progressive aphasia);
   C. Attempting to differentiate between the various clinical presentations associated with frontotemporal lobar degeneration spectrum of disorders (e.g. behavioral variant frontotemporal dementia vs progressive supranuclear palsy) to try to define the underlying pathology;
   D. Staging the severity of a dementing syndrome.

4. Patients with MCI represent a highly heterogeneous cohort for which amyloid imaging may be appropriate under specific circumstances. As a general rule, amyloid PET could be considered in MCI patients for whom the dementia expert has determined that greater certainty about the underlying pathology would alter management (e.g. knowledge of amyloid burden in an individual <65 years old with confounding circumstances such as depression or other medical disorders, and for whom safety issues at work could have major consequences). In such a case, determination of a positive amyloid status could lead to the diagnosis of MCI resulting from AD, as opposed to a nondegenerative condition, and have significant repercussions for future care and planning.

5. The actual process of undergoing an amyloid scan and the implications associated with disclosure of the results should be taken very seriously because this can be highly stressful for patients and families. To maximize safety and effectiveness of disclosing results, we recommend adopting parts of the sequence recently developed by Harkins et al in cognitively normal older adults participating in AD prevention studies. This format includes an educational session with clinical scenarios before the scan, assessment of mood and willingness to receive the results, and a formal face-to-face disclosure session in which results are discussed along with their diagnostic and prognostic implications.

Table 2: Recommendations for research and translational development to clinical care on behalf of the Canadian Consensus Conference on Use of Amyloid Imaging

1. In research settings with amyloid imaging capabilities, investigators should be encouraged to develop projects to better define the clinical and research uses of this technique and evaluate its readiness for translation to clinical care.

2. Trial designers are strongly encouraged to use this technique to (1) decrease the heterogeneity of their mild cognitive impairment population; (2) identify a cohort that is likely to respond to a drug with antiamyloid properties; and (3) study patients that are likely to convert to AD in a relatively short time frame.

3. Testing and longitudinal follow-up of asymptomatic individuals or patients with subjective cognitive impairments not meeting mild cognitive impairment criteria, or at-risk individuals (e.g. gene mutation carriers, family history of AD, apolipoprotein E ε4) should be restricted to research.

4. Future research should explore (1) the natural evolution of amyloid burden and its role in the pathophysiology of AD and other dementias, (2) its use as a potential surrogate marker for anti-amyloid therapies, (3) pharmacoeconomics issues of amyloid imaging, (4) positron emission tomography-pathology correlations, and (5) the links between amyloid imaging with cerebrospinal fluid AD biomarkers as well as downstream markers of neurodegeneration.

AD = Alzheimer’s disease; MCI = mild cognitive impairment; MRI = magnetic resonance imaging; PET = positron emission tomography.

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AD = Alzheimer’s disease.
One could argue that amyloid imaging, when approved by regulatory bodies, will be in no different situation than FDG-PET because its use will also depend on provincial reimbursement policies. In 2013, our American colleagues faced a similar situation with regards to reimbursement for amyloid imaging. The Centers for Medicare and Medicaid Services (CMS) National Coverage Decision on amyloid PET imaging in dementia and neurodegenerative disease (CAG-00431N) ruled not to cover the scans because “the evidence is insufficient to conclude that the use of amyloid PET imaging is reasonable and necessary for the diagnosis or treatment of illness or injury or to improve the functioning of Medicare beneficiaries with dementia or neurodegenerative disease.” CMS questioned the ability of amyloid PET to lead to improved health outcomes, such as avoidance of futile treatment or tests, improving or slowing the decline of quality of life, and survival. However, CMS did find sufficient evidence that the use of amyloid PET is promising: (1) to exclude AD in narrowly defined and clinically difficult diagnoses and (2) to enrich clinical trials seeking better treatments or prevention strategies for AD. The soon-to-be launched IDEAS study (discussed previously) was developed partly in response to this decision and it is hypothesized that it will demonstrate that knowledge of amyloid status leads to significant changes in patient management and improved medical outcomes, particularly in diagnostically uncertain cases of MCI and atypical dementia.

We believe time has come for Canadian dementia experts to take leadership in defining the role of molecular imaging in the differential diagnosis of dementia. Reimbursement issues surrounding the two PET techniques currently available should be discussed with appropriate provincial health authorities. Both techniques (FDG-PET and amyloid imaging) are now supported by a solid body of evidence, and politicians properly informed of their benefits should be open to fund programs where FDG-PET and amyloid PET are reimbursed in specific clinical situations along the guidelines detailed previously (e.g. differential diagnosis of AD vs FTLD).

**Amyloid Imaging versus CSF**

Amyloid imaging is not the sole biomarker which may signal underlying AD pathology. A decade or more before the appearance of dementia symptoms, and possibly before amyloid accumulation is detectable by PET, CSF changes can appear and correlate with brain atrophy in cognitively normal elderly. As suggested in the CCCDTD4 paper, authors have compared amyloid imaging with CSF AD biomarkers in the same study and found that CSF Aβ1-42 analyzed consecutively in routine clinical practice at an accredited laboratory can be used with high accuracy to determine whether a patient has normal or increased cortical Aβ deposition and so can be valuable for the early diagnosis of AD. Other groups replicated these findings using cross-sectional and longitudinal designs. Interestingly, abnormal flutemetamol retention levels correlated with disease stage in patients with mild cognitive symptoms, but this was not the case for CSF Aβ1-42.

The utility of CSF in diagnosing unclear dementing syndromes such as those on the FTLD spectrum is less clear. Although tau fibrils and aggregates are pathological hallmarks of several FTLD subtypes, total CSF tau (t-tau) appears to be a general marker of neurodegeneration, whereas phosphorylated tau (i.e. p-tau-231 and p-tau-181) are useful in discriminating between AD and frontotemporal dementia. Increased ratio of tau/Aβ1-42 can also distinguish AD from FTLD, and a low CSF p/t-tau ratio may distinguish FTLD-TDP from FTLD-tau. Another study suggests that total and p-tau in CSF were elevated in primary progressive aphasia relative to the behavioral variant frontotemporal dementia.

At the moment, however, CSF variability across techniques and centers is such that it limits proper confident interpretation of the results. Because of the absence of appropriate laboratory infrastructure in Canada, or consensus as to where the samples should be sent for analysis, Aβ1-42, t-tau, and p-181-tau have no clinical utility in Canada (not recommended for clinical practice), although they are part of research protocols in observational and therapeutic studies. Current international efforts to standardize CSF AD biomarkers, notably with more reliable enzyme-linked immunosorbent assay techniques, are currently under way.

**ACTION PLAN**

An action plan was developed modeled on appropriate use criteria by Johnson and colleagues. These recommendations cover issues that could be disseminated to Canadian health care professionals and dementia organizations through knowledge translation activities: (1) who should be referred for an amyloid scan; (2) education (patients and families, health care professionals, dementia organizations); (3) amyloid PET scanning technique, interpretation (visual vs quantitative), translation into a clinical decision; and (4) proper disclosure of results.

**Who Should Be Referred for an Amyloid Scan?**

This decision should rely on the dementia expert guided by current Canadian guidelines. A dementia expert is a physician with substantial clinical experience and practice in dementia care. Expertise in dementia is acquired through formal training and clinical experience in neurology, psychiatry, and geriatric medicine; however, not all dementia experts have expertise in amyloid imaging and/or work in conjunction with a nuclear medicine specialist (NMS) qualified in amyloid imaging, hence the recommendation to refer to a dementia center with expertise in this technique, when appropriate.

**Education (Patients, Families, Health Care Professionals, Dementia Organizations)**

Dementia specialists in Canada are committed to disseminating information to the public and dementia organizations (e.g. information on amyloid imaging already exists on the Alzheimer Society’s website at http://www.alzheimer.ca), to assisting colleagues in appropriate use of amyloid imaging, and in providing clarifications on how to incorporate amyloid PET in medical practice. Knowledge translation activities should be organized in all Canadian provinces to introduce the most recent guidelines on amyloid imaging.

**Amyloid PET Scanning, Interpretation, and Translation Into a Clinical Decision**

**Amyloid PET Scanning**

Imaging procedures should be performed by qualified nuclear medicine technologists and NMS with national certification in nuclear medicine and appropriate qualification in amyloid
imaging. It should be performed in an imaging facility certified by Canadian accrediting agencies. Procedure guidelines for amyloid PET (SNMMI and European Association of Nuclear Medicine) should be followed.

**Interpretation**

The nuclear medicine team (i.e. the technologist and the NMS) performing the scan must be familiar with brain anatomy and must have adequate specific training in amyloid PET interpretation because amyloid PET imaging can be technically challenging and should be performed with strict attention to quality control.²⁹,⁶¹ Web-based instruction programs have been developed and validated and should be completed successfully by all amyloid imaging teams before reading scans. Interpretation of amyloid PET imaging should be communicated to the referring physician by the NMS by way of a written report according to a standard diagnostic imaging practice as outlined in the SNMMI General Imaging Guideline. At this time, the final reading should indicate whether Aβ was present (amyloid positive) or was not present (amyloid negative). The protocol for the qualitative read that determines positivity or negativity must be standardized⁵⁹ and must conform to a specific guideline provided by the manufacturer. Indeterminate results may arise as a result of technical or physiological factors and should be reported as such.

**Translation Into a Clinical Decision**

The NMS report should not equate amyloid positivity with AD dementia (amyloid positivity is not synonymous with AD, and amyloid scans cannot replace a detailed clinical evaluation). Upon receiving the NMS report, the dementia expert should proceed to a comprehensive review of the clinical assessment (history, physical examination) and test results (laboratories, neurocognitive testing, MRI, FDG-PET) and incorporate amyloid results into a clinical decision process, always considering that amyloid imaging is an evolving modality and that image interpretation criteria, clinical significance of positive and/or negative scans, and technical imaging considerations are evolving.²⁵

**Proper Disclosure of Results**

That moment can be highly stressful for patients and families. To maximize safety and effectiveness of disclosing results, we recommend consulting the sequence recently developed by Harkins et al.⁶⁵ in cognitively normal older adults participating in AD prevention studies. This process included: (1) an educational session, in which participants receive verbal and written information covering what is known and unknown about amyloid imaging, including possible results and their meaning; (2) screening for anxiety and depression to determine suitability to receive amyloid imaging information; (3) checking comprehension and recognizing distress; (4) conducting imaging on a separate day from consent, and disclosing results on a separate day from imaging; (5) proceeding to disclosure in person, with time for questions (at disclosure, physicians should assess mood and willingness to receive results); and (6) offering resources for support (brochures, follow-up call). The latter were developed for normal older adults participating in AD prevention studies. This may not be entirely possible in clinical practice, but can serve as general principles to guide proper disclosure.

**Future Challenges**

Amyloid imaging has been approved for widespread clinical use by leading health authorities in the United States and the United Kingdom. Despite this, several unknowns about its diagnostic utility remain and future studies should particularly focus on (1) its sensitivity and specificity as compared with pathology in practice-based settings (as opposed to the hospice studies), (2) technical and patient factors that could lead to false positives and false negatives, (3) the relative contribution of both diffuse and neuritic plaques’ binding to the in vivo signal, (4) interpretation of the test as a dichotomous result versus assessing binding intensity and spatial distribution, (5) inter- and intra-rater reliability of visual and/or quantitative interpretations, (6) the optimal quantitative threshold for defining a positive scan for each of the available tracers, (7) whether the threshold for PiB positivity should be adjusted based on demographic factors such as age or genetic variables, and (8) cost effectiveness issues. Such issues are relevant for any diagnostic test, and should be addressed as research continues to target key variables associated with amyloid imaging. Recent longitudinal initiatives such as the IDEAS study should help answer several of these questions.

**Conclusions**

Amyloid imaging represents a major breakthrough in the evaluation of dementia that will doubtlessly translate into better clinical care and ultimately help guide the development of molecular-based therapies for these devastating illnesses. An impressive body of research has already been generated in the field, and studies of practical clinical applications are emerging with a specific indication in patients with objectively confirmed cognitive impairments where diagnostic uncertainty remains even after a comprehensive clinical evaluation in a tertiary memory clinic. This technique should always remain an adjunct imaging tool that is part of a comprehensive clinical evaluation in which a dementia expert determines that having a more accurate clinical diagnosis will alter management. Fundamentally, amyloid imaging detects a brain histological state, and is not a clinical diagnosis. Used in isolation, it cannot diagnose AD, MCI, or differentiate normal from abnormal aging. Clinical availability of new tracers in Canada would represent a major advancement for the many Canadians affected by an unclear dementicing condition who suffer in silence while being exposed to unnecessarily prolonged delays before diagnosis, repeated and pointless visits and diagnostic tests with inferior sensitivity and specificity than amyloid imaging, and inappropriate treatments or lack thereof when indicated. In the end, we wish this effort to generate momentum for the industry to submit a new drug proposal to Health Canada so that regulatory bodies approve the technique and approval of provincial reimbursement can follow its proper course.

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REFERENCES


APPENDIX A: MEMBERS

A. Specialized Task Force on Amyloid imaging in Canada (STAC)

1. Organizing Committee Members
   Serge Gauthier, MD (chair)
   Robert Laforce Jr, MD, PhD
   Pedro Rosa-Neto, MD (co-chair)
   Jean-Paul Soucy, MD

2. Invited International Leaders
   Bruno Dubois, MD
   Gil D. Rabinovici, MD

B. Consensus meeting participants

   Clinical and Academic
   Howard Chertkow, MD
   Guy Lacome, MD
   Alain Robillard, MD
   Sylvia Villeneuve, PhD

   C. Member from Canadian Regulatory Bodies (Health Canada)
   Agnes Klein, MD

D. Leading representatives from the molecular imaging and imaging software industries
   Susan De Santi, PhD
   Benoît Galarneau
   Rick Hiatt
   René Rebeaud
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APPENDIX B: RELATIONSHIPS WITH INDUSTRY AND MANAGEMENT OF CONFLICTS OF INTEREST

The group rigorously attempted to avoid any actual, perceived, or potential conflicts of interest that might have arisen as a result of an outside relationship or personal interest of the writing committee members. We reviewed our own industry relationship policies to ensure that the ensuing process adhered to current standards. Members were required to provide disclosure statements of all relationships that might be perceived as real or potential conflicts of interest. These statements were reviewed by the chair and senior author of this paper.