Advances in functional neuroimaging in dementias and potential pitfalls

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Abstract

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Introduction

Neuroimaging is continuously advancing at a rapid rate and has progressed from excluding relatively uncommon secondary causes (stroke, tumor) to assisting with early diagnosis and subtype of dementia. Structural imaging has given way to functional, metabolic and receptor imaging. An emerging focus is the appreciation of human connectomics, neuro-opathies and their imaging counterparts.14 Fluorodeoxyglucose positron emission tomography (18FDG-PET) brain, single photon emission computed tomography (SPECT) brain, PET Pittsburgh Compound B (PIB) and PET receptor (Dopa) imaging are becoming more widespread and gaining acceptance as part of the routine work up (Table 1).18FDG-PET brain in particular has been an important tool in assisting in the early diagnosis of mild cognitive impairment (MCI) type of case as well as in differentiating different types of dementia, with frontotemporal disorders (FTD) and Alzheimer’s disease (AD) already being an Food and Drug Administration asserted indication for example.1 Furthermore, basic neuroscience is increasingly delineating newer molecular phenotypes (beta amyloid, alpha synuclein and TDP-43) and subcategories which are particularly complex with FTD (TDP 43, tau, FUS, C90RF2). The pathologies take effect at the synaptic and neuronal level and are the earliest events in these disease states. Neuroimaging is increasingly able to decipher these earlier changes long before the clinical state even emerges or the extensive atrophy seen on structural scans. As late or end stages are barely amenable to significant intervention, the earlier detection is key.

Currently PET brain imaging offers the most accurate diagnostic method for the 5 main dementia categories as well as a number of the subcategories2 (Table 2). A number of patterns have emerged that reliably differentiate the major dementia subtypes. This is important for treatment and prognostic decision making. For example the posterior cortical atrophy syndrome (Benson syndrome), an AD variant progresses very slowly with largely retained cognitive function, at the same time dominated by complex visual impairments including visual agnosias and Balint’s syndrome (Figure 1).3 With FTD (frontotemporal hypoemetabolism) (Figure 2) and AD (temporal, parietal, posterior cingulate) the footprint of hypometabolism is relatively easily identified (Figure 3). However with Parkinson’s disease with dementia (PDD) and Diffuse Lewy Body Disease (DLBD), the differentiating features even on PET scanning are less distinct and considerable overlap in the patterns are seen and consequently confusion may arise. This is particularly the case with PDD, DLDB and cortical basal ganglioc (CBG) disease. PDD and DLDB are particularly challenging as there is no clear PET pattern as for example with AD and FT. This is perhaps not surprising as DLDB and AD share features at a clinical, neuroimaging, pathological and pharmacotherapeutic level.4 Furthermore, Parkinson’s disease (PD), PDD, DLBD and AD represent a pathological spectrum with loss of both cholinergic neurons and dopaminergic neurons as the most prominent neurotransmitter perturbation initially.5 Adding to the complexity, there is overlap with AD and cognitive vascular disorders (CVD), the frontal variant of Alzheimer’s and the frontal variant of FTD that present added diagnostic challenges in individual cases. Even more importantly, diagnosis and differentiation may not be possible by clinical means alone especially in the pre-symptomatic phase when no cognitive impairment exists particularly in the context of presumed high cognitive reserve. However intervention and treatment may yield the most significant results at this stage.6

In addition to the 4 most common dementia syndromes discussed, newer dementia etiologies continuously added and expand the diagnostic work up. Autoimmune dementias and the prefrontal atrophy secondary to chronic stimulation of the pain matrix (chronic pain syndrome) is also regarded as a neurodegenerative condition today. Brain atrophy and as such neurodegeneration has been associated with various chronic pain conditions including fibromyalgia, posttraumatic headache, complex regional pain syndrome (CRPS), and chronic back pain.7-10 The potential cause and effect relationship of the atrophy with pain continues to be elucidated. In relation to dementias, memory and pain processing share close anatomic relationships. Numerous functional neuroimaging studies using positron emission tomography (PET)11-13 and functional magnetic resonance imaging (fMRI)14,15 have demonstrated that the anterior cingulate cortex (ACC) and posterior cingulate cortex (PCC) have key roles in processing of pain perception.16 Lesional studies have corroborated these findings.17,18 The ACC has not been demonstrated to be involed in coding stimulus intensity or location but participates in both the affective and attentional components of pain sensation and response. Increased blood flow is also found in the posterior parietal and prefrontal cortices and is thought to reflect networks for attention and memory activated by painful stimulation. Memory and pain processing networks are physically close in the cingulate cortex. Episodic memory retrieval is predominately in the caudal portion of the PCC while pain processing occurs in the rostral portion of the PCC.19 Given that patients with Alzheimer’s disease and other neurodegenerative dementias are noted to report less pain and may receive less analgesics that comparable normal peers, consideration has been provoked that the dementia related neurodegeneration process may be affecting pain processing pathways. However, studies have demonstrated heightened magnitude and duration of activity in the ACC and other pain processing areas during pain stimulus in Alzheimer’s disease patients.20 As such

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the perception of pain is not diminished in Alzheimer’s, and should raise awareness of clinical care providers to insure adequate analgesia for dementia patients with painful conditions.

PET brain scan imaging and cerebrospinal fluid (CSF) biomarkers are becoming increasingly important to diagnose dementia syndromes especially in the setting of overlap syndromes. Learning to identify metabolic uptake patterns by cartographic analysis as well as semi-quantitative methods may be crucial for correct diagnosis in the face of conflicting clinical as well as neuro-radiological findings. The four major neurodegenerative diseases include AD, DLBD, FTD and CVD in approximate order of frequency. AD and FTD

Table 1. Spectrum of current functional imaging modalities.

| Magnetic resonance imaging multimodality |
|-----------------------------------------|
| Magnetic resonance imaging (routine series) |
| TI/T2 fluid attenuated inversion recovery, gradient echo sequence, magnetic resonance angiogram largely to detect degree of concomitant vascular disease, atrophy pattern and other secondary pathologies |
| Magnetic resonance imaging - diffusion tensor imaging |
| Fiber tract pathology especially in traumatic brain injury, multiple sclerosis |
| Magnetic resonance imaging - quantitative atrophy estimation |
| At least 5 different patterns of the major dementia syndromes (Seeley WW et al.) |
| Magnetic resonance imaging - perfusion |
| Perfusion as a reflection of hypometabolism, similar to single photon emission computed tomography (perfusion) and positron emission tomography (metabolism) patterns of abnormality |
| Magnetic resonance spectroscopy |
| Biochemical analysis, choline, lactate particularly useful in brain tumor diagnosis |

| Single photon emission computed tomography |
|-------------------------------------------|
| Hypoperfusion (in vascular or hypometabolism) |
| Hyperperfusion for example with ictal foci |
| Position emission tomography brain |
| Frontotemporal disorders |
| Alzheimer’s disease |
| Diffuse lewy body disease |
| Parkinson’s |
| Progressive supranuclear palsy |
| Huntington’s |
| Cortical basal ganglionic cognitive vascular disorders |
| Intrinsic state connectivity maps |
| Default mode |
| Salience network |
| Parkinson’s (increased connectivity of the basal ganglionic and thalamocortical loops) |
| Cortical basal ganglionic |
| Cognitive vascular disorders |

| Quantitative electrophysiological and magnetoencephalographic |
|---------------------------------|
| Alzheimer’s disease - reduced connectivity of alpha and beta in frontotemporal and frontoparietal regions |
| Frontotemporal disorders - uncertain |
| Parkinson’s - increased connectivity of alpha and beta locally and globally |
| Diffuse lewy body disease - reduced connectivity alpha range locally and globally |

Figure 1. 18FDG PET brain: posterior cortical atrophy syndrome with marked posterior cortical hypometabolism.

Figure 2. FTD end stage with extensive bifrontal cortical atrophy (top) and 18FDG PET brain of early non fluent aphasial subtype (down), demonstrating left inferior frontal hypometabolism (arrow) in context of normal structural MRI scan.
Review (Figure 2) have distinct and easily identifiable patterns and CVD is guided primarily by stroke related imaging. The situation with DLDB can be difficult due to the overlap of neuropathologies. A correct diagnosis is nevertheless vital to guide care for the patient. Refining the diagnosis may assist with appropriate treatment and in avoiding serious side effects not the least of which include neuroleptic sensitivity such as neuroleptic malignant syndrome or catatonia. People with DLDB generally respond better to acetylcholinergic (Ach) therapy in terms of improved alertness, less fluctuation in cognition and memory than do people with AD. The complex clinical and neuroradiological patterns of the dementia syndromes no doubt foster misdiagnosis. The neuroradiological findings of DLDB may be initially confused the diagnosis of AD diagnosis. Hallucinations, a hallmark of DLDB can occur with AD as well as other dementias. A recent report of a specific 5HT - 2A antagonist specific for visual hallucinations with DLDB is further reason for precise diagnostic accuracy translating into therapeutically effectiveness. There are already several recognized metabolic features of PET scanning that have been identified with DLDB, namely i) diffuse glucose hypometabolism in entire cortex including the occipital region typical feature of DLDB and distinctive from AD, ii) lateral occipital hypometabolism (Figure 3) which may have the highest sensitivity and iii) posterior cingulate preserved metabolism or posterior cingulate island sign, (PCIS) which may have the highest specificity (Figure 4). However, the so called PCIS appears to reflect the much more pronounced occipital hypometabolism relative to the also reduced (but to a lesser extent) or sometimes normal metabolism of the posterior cingulate cortex.

With AD on the other hand, hypometabolism is seen very early in the medial portions of the parietal lobes as well as the posterior cingulate region. The difficulty posed by the fluctuating symptomatology adds to the diagnostic dilemma. In addition, the uncertainty of hallucination origin, whether drug induced, due to DLBD or AD is a frequent frustration shared by both patients and physicians treating these conditions. The advent of biomarker assisted diagnosis has already initiated new diagnostic criteria for AD that enable a pre-mortem diagnosis that is heavily reliant on positive CSF and PET findings. The trend of clinical, PET and CSF diagnostic features rendering increased diagnostic accuracy will likely translate similarly in the DLDB-Parkinsonism complex as it has already done for example with MCI and AD. With the advent of MR perfusion scanning giving similar information to PET brain scanning, this modality may become more desirable in view of its availability and lack of radiation. A continuously expanding therapeutic armamentarium for DLDB both in terms of receptors for symptom alleviation as well as disease modifying therapies all bode for precise diagnostic accuracy.

Emerging imaging modalities: default mode network or intrinsic connectivity networks imaging and other (molecular) networks

The Default Mode Network (DMN) can be imaged by functional MRI (resting state without activation procedures) and reflects the basal or default mode activity of the brain. It links particular brain regions that includes the posterior cingulate, the precuneus, lateral parietal, lateral temporal, medial frontal areas. DMN impaired connectivity has already been shown in AD, FTD, schizophrenia, epilepsy, autism later life depression. The DMN is active during rest and becomes less active during cerebral task engagement. It is implicated in the pathophysiology of AD as the distribution of the DMN is similar to the fibrillar amyloid deposition in patients with AD (amyloid PET scanning). The A-beta deposition overlaps considerably with the DMN and the tau deposition overlaps with the DMN component that is concerned with episodic memory. It has been surmised that over-activity of DMN (posterior cingulate, later parietal, medial

![Figure 3. Transaxial 18FDG PET brain revealing bilateral predominantly occipital hypometabolism (arrows) in a patient with DLDB.](image1)

![Figure 4. Sagittal 18FDG PET brain revealing preserved mid to posterior cingulate metabolism (arrows) with temporoparietal hypometabolism.](image2)

| Table 2. Positron emission tomography brain patterns in dementias. |
|---------------------------------------------------------------|
| **Dementia subtype**                                           | **18Fluorodeoxyglucose positron emission tomography hypometabolism pattern** |
| Alzheimer                                                     | Relatively symmetric parietotemporal, medial temporal, posterior cingulate, frontal association cortex to lesser degree |
| Alzheimer’s disease variant (Posterior cortical atrophy syndrome) | Occipital hypometabolism predominates |
| Frontotemporal disorders behavioral variant                   | Frontal and anterior temporal hypometabolism |
| Parkinson’s disease with dementia                             | Temporo-parietal, may be similar to AD |
| Diffuse lewy body disease                                     | Occipital and temporal hypometabolism |
| Cognitive vascular disorders                                  | Cortical and subcortical, singular or multifocal, correlating with structural imaging abnormality |
| Corticobasal degeneration                                     | Global reduction in metabolism as well as asymmetric prefrontal, premotor, sensorimotor superior temporal, parietal hypometabolism with thalamic hypometabolism contralateral to limb apraxia |
| Huntington’s                                                  | Caudate nucleus hypometabolism, frontal association cortex to a lesser degree |
| Progressive supranuclear palsy                                | Caudate nucleus, putamen, thalamus, pons, superior and anterior frontal cortex |

Note that FDG PET increases diagnostic accuracy beyond that derived from clinical evaluation.
frontal) in younger life may lead to a metabolic impairment predisposing people to amyloid deposition in later life.34 The DMN is known to subsurface several key memory processes including episodic encoding, retrieval, autobiographical, metamemory processes, moral decision making and theory of mind. Petrella et al. reported lower connectivity in DMN in patients with MCI who subsequently were diagnosed with AD over a 2-3 years period.32 This type of functional connectivity MRI (fc-MRI), is an attractive tool because MRI scanners with blood oxygen level dependent (BOLD) capability are widely available and fc-MRI is non invasive, radiation free, can be repeated multiple times and have short acquisition time of 5-8 min.35

The quest for neuroimaging biomarkers for diagnosing pre-clinical disease may soon be realized albeit with perhaps a complementary role played by FDG-PET, Pittsburgh compound B PET (PIB PET), PET neurotransmitter imaging and fc-MRI of the DMN and perhaps other networks such as the salience (for FTLD) and attentional networks.36-39 With intrinsic connectivity networks the pattern for AD and FTLD is fairly clear. However for the PDD and DLDB the pattern is less obvious and there may also be hyperactivity in the BG as opposed to the usual underactivity seen in the other neurodegenerative conditions (Table 3).

Neurotransmitter and neurotransmitter receptor position emission tomography (and sometimes single photon emission computed tomography) imaging

In the hopes of guiding therapy more accurately, the cholinergic (nicotinic receptors) and dopaminergic systems have been investigated in this regard. An increase in 11C nicotinic binding sites as well as associated cognitive improvement were reported after 3 months of rivastigmine for AD patients.36 At times neurodegenerative conditions coexist and unraveling the most pertinent neurotransmitter systems at fault is useful. This has been shown for acetylcholine involvement in Parkinson's with and without dementia by 11C methyl-4piperidyl acetate (MP4A). In this study, the dopaminergic system was also measured using 18F fluorodopa (FDOPA) which revealed decreased uptake in the striatum and MP4A was decreased in the Parkinson's group with dementia.37

Cortical atrophy patterns

Cortical atrophy patterns are generally discernible at a later stage of the process and perhaps the least sensitive (Table 4). Nevertheless, a recent pivotal study showed 5 different neurodegenerative syndromes and their atrophy patterns corresponding to 5 different intrinsic functional connectivity networks.29 In particular the salience network has been shown to correlate with frontotemporal lobe dementia.40 This relatively novel approach of brain analysis, called connectomics by some is showing promising results. Assessment of brain connectomics is regarded as an area of priority in future cognitive research (The Human Connectome Project).41

The importance of considering cognitive reserve status by functional imaging in conjunction with cognitive or neuropsychological testing

No direct relationship exists between the extent of pathology and clinical manifestation of the underlying disease or damage for that matter. Katzman et al., reported on 10 elderly normal women with advanced AD pathology supported this premise, speculating that their brains had more cognitive reserve.42 Cognitive reserve is considered to include: i) brain reserve capacity (correlate - hardware, brain size, neural count or synaptic count; ii) cognitive reserve (correlate - software). Attempting to cope with brain damage using cognitive compensatory approaches. Higher education, bilingualism, literacy and participation in hobbies for example, allow people to withstand brain damage better. Cognitive reserve in turn has been divided into: i) Neural reserve: cerebral networks less susceptible to disruption due to greater inherent efficiency; ii) neural compensation: post brain damage, additional or nonconventional networks are deployed to compensate for brain damage.43

Functional imaging studies support the neural reserve and neural compensation reflecting individual compensatory differences to pathology. For example, two people with the same cognitive impairment may have markedly different degrees of underlying AD pathology. This is clearly important for the diagnosis of preclinical Alzheimer's disease, as mild cognitive impairment (MCI) patients may have both minimal pathology or more extensive pathology. The cognitive reserve (CR) hypothesis, is used to describe this variability and is considered an important part of the assessment therefore. Clinical evaluation alone cannot be relied on and biomarkers (whether CSF analysis of tau and amyloid beta 1-42 or metabolic imaging) will be part of the work up.44

Since the proposal of the CR hypothesis, this has been recently supported using the PIB and 18FDG PET in relation to education in mild AD. In this study, 12 high educated (15 or more years) and 13 low educated patient with the same degree of cognitive deterioration were evaluated with PET brain scanning using both [11C] PIB and 18F Fluorodeoxyglucose as ligands. The high-educated people showed increased PIB uptake in the lateral frontal cortex as well as lower glucose metabolic rate in the temporoparietal cortical regions compared to low educated people.

How may we utilize the various imaging modalities today and in the near future?

A likely hierarchical approach to using surrogate neuroimaging in cognitive patients may be as follows may take the following format: i) Resting State Network Imaging (DMN,

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**Table 3. Intrinsic connectivity network patterns in dementias.**

| Dementia subtype                      | Intrinsic connectivity pattern                           |
|---------------------------------------|---------------------------------------------------------|
| Alzheimer                             | Default mode network shows reduced connectivity         |
| Frontotemporal lobe disorder behavioral variant | Salience network shows reduced connectivity            |
| Parkinson’s                           | Basal Nuclei-thalamocortical loops show increased connectivity |
| Diffuse lewy body disease             | Uncertain at present but may show ascending brainstem projection system |
| Cortical basal ganglionic              | Uncertain                                               |

**Table 4. Atrophy patterns in dementia subtypes.**

| Dementia subtype                     | Atrophy pattern                                      |
|---------------------------------------|------------------------------------------------------|
| Alzheimer                             | Temporoparietal, medial hippocampus, precuneus       |
| Frontotemporal disorders behavioral variant | Frontotemporal                                      |
| Parkinson’s with dementia             | Temporoparietal                                      |
| Diffuse lewy body disease             | Parietal atrophy but no hippocampal atrophy. Caudate atrophy currently debatable |
| Vascular cognitive disorder (vascular cognitive impairment, vascular dementia) | Nil specific, subcortical leukoaraiosis as opposed to periventricular rimming leukoaraiosis is frequent |
| Progressive supranuclear palsy        | Midbrain atrophy (Hummingbird, Penguin signs)        |
| Cortical basal ganglionic              | Pronounced fronto-parietal atrophy, often asymmetric, corpus callosum atrophy |

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Salience and others) by f-MRI; ii) beta amyloid accumulation assessed by PET brain PIB (also CSF assays); iii) the subsequent synaptic dysfunction assessed by FDG-PET brain; iv) finally, neuronal loss follows, as assessed by volumetric MRI.

**Conclusion**

Since the revised AD criteria now include biomarker testing, neuroimaging (particularly PET) has gained acceptance. However, the conditions with most ambiguous results on PET scanning (PDD and DLBD), certain neuroradiological identification such as the PCIS may be an important feature in improving confidence in the discrimination between AD and DLBD in clinical overlap syndromes and support differing therapeutics. For all the other conditions, PET reliably discerns the differing syndromes. The advent of ICN imaging looks differing therapeutics. For all the other conditions, PET reliably discerns the differing syndromes. The advent of ICN imaging looks promising to enable an even earlier and more precise diagnosis of the ever expanding array of dementia syndromes.

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