Multimodality Imaging in Primary Progressive Aphasia

M. Roitman, G.C. Chiang, M.L. Gordon, and A.M. Franceschi

ABSTRACT

SUMMARY: Primary progressive aphasia is a clinically and neuropathologically heterogeneous group of progressive neurodegenerative disorders, characterized by language-predominant impairment and commonly associated with atrophy of the dominant language hemisphere. While this clinical entity has been recognized dating back to the 19th century, important advances have been made in defining our current understanding of primary progressive aphasia, with 3 recognized subtypes to date: logopenic variant, semantic variant, and nonfluent/agrammatic variant. Given the ongoing progress in our understanding of the neurobiology and genomics of these rare neurodegenerative conditions, accurate imaging diagnoses are of the utmost importance and carry implications for future therapeutic triaging. This review covers the diverse spectrum of primary progressive aphasia and its multimodal imaging features, including structural, functional, and molecular neuroimaging findings; it also highlights currently recognized diagnostic criteria, clinical presentations, histopathologic biomarkers, and treatment options of these 3 primary progressive aphasia subtypes.

Neurodegenerative disorders encompass an assortment of clinically and histopathologically diverse conditions, typically slowly progressive and associated with gradual neurologic dysfunction. While precise mechanisms leading to their development continue to be elucidated, these disorders can be broadly grouped into categories based on similar histopathology, including tauopathies (eg, some cases of frontotemporal lobar degeneration [FTLD], corticobasal degeneration, progressive supranuclear palsy), transactive-response DNA-binding protein 43 (TDP-43) accumulation (eg, other cases of FTLD, amyotrophic lateral sclerosis), β-amyloid accumulation (eg, Alzheimer disease [AD]), and α-synucleinopathies (eg, Parkinson disease, dementia with Lewy bodies, multiple system atrophy), noting mixed pathologies in some cases.1 Primary progressive aphasia (PPA), clinically characterized by language-predominant impairment, has been histopathologically linked to both underlying FTLD (eg, semantic variant PPA [svPPA] and nonfluent/agrammatic variant PPA [nfvPPA]) and AD-type pathology2 (eg, logopenic variant PPA [lvPPA]) (Fig 1). The microtubule-associated protein τ, encoded by the MAPT gene, has been shown to pathologically aggregate when in an abnormal hyperphosphorylated form (p-τ) and result in extracellular neurofibrillary tangles that contribute to neurodegeneration. Tauopathies are characterized by the predominance of abnormally deposited alternatively spliced τ isoforms (eg, 3-repeat [3R] versus 4-repeat [4R]-tauopathies), which may be seen in some cases of PPA.3 Deposits of TDP-43, a cellular protein encoded by the TARDBP gene and with 4 described subtypes (A, B, C, D), is an additional frequently identified pathologic substrate seen in cases of PPA.4 While trends have been reported between these histopathologic entities and PPA subtypes (lvPPA: AD pathology; svPPA: TDP-43; nfvPPA: 4R-τ), no absolute association exists and inconsistencies regarding the frequency of molecular alterations for each variant have been described across studies, likely related to varied diagnostic approaches.5

While a progressive neurodegenerative disorder characterized by language-predominant impairment has been a recognized entity dating back to the 19th century, inconsistencies regarding its
Inclusion and exclusion criteria for PPA diagnosis

| Criteria | Inclusion | Exclusion |
|----------|-----------|-----------|
| Most prominent clinical feature | Language difficulty | Pattern of deficits is better accounted for by other nondegenerative nervous system or medical disorders |
| Aphasia should be the most prominent deficit at symptom onset and for the initial phases of the disease | Prominent initial episodic memory, visual memory, and visuoperceptual impairments |
| Prominent initial behavioral disturbance |

Logopenic Variant PPA

LvPPA is the most recently described of the 3 PPA variants, first reported in 2004 by Gorno-Tempini et al. Clinically characterized by word-finding difficulties and lapses in conversation, LvPPA has been reported to exhibit histopathologic findings and biomarkers that overlap with AD pathology (eg, \( \beta \)-amyloid \([A\beta]\) and neurofibrillary tangles) and, therefore, is considered a clinical variant of AD.

As a whole, PPA is a rare entity with an estimated prevalence of 3–7 cases per 100,000, often occurring in late middle life (mean age of disease onset, 62.34 years) with an average delay between first symptoms and diagnosis of 3.21 years in 1 study. Given the novelty of lvPPA as a clinical entity, its precise prevalence is not definitively known. However, a 2016 retrospective analysis of a cohort from a tertiary center \((n = 97)\) in patients with language deficits and CSF biomarkers from the French AD databank, performed to better understand PPA demographics, revealed lvPPA as the most common variant of PPA \((51%, 49/97)\) with a slight female predominance \((57%, 28:21, \text{female/male})\). Within this group, lvPPA was more frequently associated with an AD CSF profile \((85%)\) than nfvPPA \((35%)\) or svPPA \((20%)\), contributing to the present day notion of lvPPA as a clinical variant of AD.
In a study investigating CSF fluid biomarkers (including Aβ42, τ, p-τ) in 13 patients with lvPPA, 62% (8 of 13) demonstrated a profile indicative of AD pathology (lvPPA+), while 38% (5 of 13) had a non-AD profile (lvPPA−). Subsequent analyses demonstrated that those in the lvPPA+ group exhibited more advanced imaging findings compared with those in the lvPPA− group, including more extensive hypometabolism and larger regions of involvement throughout the inferior parietal and superior and middle temporal cortices. Such heterogeneity of pathologies identified in this study may reflect a “logopenic aphasia complex,” with at least 2 existing lvPPA subvariants.

PPA may be inherited in an autosomal dominant manner, most commonly associated with mutations in the progranulin (GRN) gene on chromosome 17. While the presence of a GRN mutation does not necessarily lead to PPA, a language disorder often emerges in many patients with this mutation. The clinical symptoms of this language disorder can vary widely, with heterogeneity even among family members with the same GRN mutation. However, among patients with nonamyloid PPA with GRN mutations, lvPPA was found to be the most frequent linguistic variant. In 1 study, 42% of patients with lvPPA were found to carry the apolipoprotein E ε4 allele, known to confer an increased risk of sporadic AD and in keeping with our understanding of lvPPA as a clinical variant of AD, compared with 26% of patients with svPPA and 20% of patients with nfvPPA. Other predisposing gene variants or mutations found to be associated with lvPPA include TREM2, TOMM40, APP, PS1 and PS2, and MAPT.

While the aforementioned evidence supports underlying AD pathology in most patients with lvPPA, not all cases of lvPPA are attributed to AD pathology. In 1 postmortem analysis of 99 patients with lvPPA, 76% had primary AD pathology, while FTLD-TAR DNA-binding protein (FTLD-TDP) and FTLD-τ pathologies were identified in 14% and 5% of patients, respectively. To date, no pharmacologic options have been shown to improve or protect against declining function in lvPPA. However, supportive care measures such as speech-language therapy have demonstrated efficacy in improving communication. Given that underlying AD pathology is associated with most lvPPA cases, use of emerging anti-amyloid therapies in lvPPA may be investigated.

### Structural Imaging

The presence of specific regional patterns of atrophy or metabolic impairment is the key neuroimaging diagnostic feature for each of the 3 PPA variants (Table 2). Structural imaging, including CT and MR imaging, can be used to identify these classic patterns of focal atrophy. Included in the 2011 Gorno-Tempini et al diagnostic criteria, MR imaging can be used to identify asymmetric, classically left-sided widening of the Sylvian fissure, indicative of the posterior peri-Sylvian and temporoparietal atrophy seen in lvPPA. The posterior aspect of the left superior temporal gyrus, corresponding to the expected Wernicke area, is typically involved. This finding, particularly when progressive over multiple examinations and in conjunction with a clinical history of progressive word-finding difficulty, should raise the possibility of underlying lvPPA (Fig 2). Notably,
while most patients are left-hemispheric language dominant, involvement of the right hemisphere has been reported and hypothesized to occur in left-handed individuals or those with a history of developmental learning disabilities (eg, dyslexia). The preferred structural imaging technique for the diagnosis of PPA is MR imaging, due to its superior soft-tissue resolution and ability to precisely localize anatomic atrophy. Advanced MR imaging techniques, such as voxel-based morphometry analysis and DTI, may be used to demonstrate focal GM atrophy (Fig 3) and WM alterations, respectively. In particular, cortical volumetric software such as the FDA-approved NeuroQuant (https://www.cortechs.ai/products/neuroquant-ct/) and Icometrix (https://www.icometrix.com/) are increasingly being used in routine clinical assessment of various neurodegenerative disorders. In a study investigating the utility of MR imaging in differentiating PPA variants, MR imaging demonstrated a high specificity for the characteristic atrophy patterns of lvPPA (95%) and nfvPPA (91%), noting a low sensitivity for both (43% for lvPPA; 21% for nfvPPA). Therefore, while the presence of left posterior peri-Sylvian or temporoparietal region atrophy is highly suggestive of lvPPA, its absence does not exclude the diagnosis. In a prospective study investigating 130 patients with neurodegenerative aphasia, of whom 52 had lvPPA, GM loss was identified in patients with lvPPA, more commonly on the left and greatest in the posterior temporal lobe extending to the frontal and parietal regions. Fractional anisotropy and mean diffusivity analyses within this cohort revealed left-greater-than-right bilateral WM involvement, greatest in the posterior left temporal WM and extending into the anterior temporal, frontal, parietal, and occipital WM, as well as involving the bilateral superior and inferior longitudinal fasciculi and inferior occipitofrontal fasciculus. The posterior superior temporal and inferior parietal cortices have been shown to play a role in phonologic loop functions. Therefore, involvement of these regions and WM tracts in the superior and inferior longitudinal fasciculi likely account for the poor repetition, naming, and comprehension seen in patients with lvPPA.

Atrophy may also be identified anteriorly with involvement of the hippocampi, among other structures, with the overall extent and pattern of atrophy varying widely among individual patients. Most important, the presence of progressive atrophy with time supports the diagnosis of PPA, a critical observation worthy of mention when interpreting such structural imaging examinations.

Functional Imaging. fMRI, which can be performed with task-based paradigms or in a resting state, uses blood oxygen level-dependent contrast to identify areas of brain activation on the basis of oxygen extraction. While fMRI is not in routine clinical use for the diagnosis of PPA and limited literature exists regarding its specific findings in lvPPA, fMRI has been reported to demonstrate functional changes in patients with svPPA (formerly referred to as semantic dementia), to be discussed later in this review. Such advanced imaging techniques could be useful in
identifying aberrant and compensatory language network changes and may prove useful in guiding future therapeutic trials. Cerebral perfusion, which has been linked to cognition and neuronal activity, can be measured with MR imaging using the noncontrast arterial spin-labeling (ASL) technique. While limited data exist regarding use of ASL for lvPPA specifically, ASL has been shown to identify hypoperfusion patterns in other dementia subtypes before symptom onset and also correspond to disease-specific regions of hypometabolism identified on [18F] FDG-PET. However, limitations of the ASL technique for clinical use include its low SNR on conventional 1.5T field strength MR imaging, with 3T field strength MR imaging preferred for use in the ASL technique, as well as technical issues with quantification. Further studies assessing the potential role of ASL for patients with lvPPA as well as its performance compared with [18F] FDG-PET are warranted.

Molecular Imaging Molecular imaging allows in vivo identification and quantification of cerebral metabolism, abnormal deposition of β-amyloid and τ, and the presence of brain inflammation, important neuroimaging biomarkers that may improve early diagnosis and assist in assessing neurodegenerative disease progression. Currently available molecular imaging modalities include SPECT and PET, with a number of investigational radiotracers on the horizon. Advantages of PET imaging include superior spatial and contrast resolution compared with SPECT, though it is a more expensive examination and less widely available. Limitations include attenuation correction and motion artifacts, which may cause inaccurate anatomic coregistration. Molecular neuroimaging has been shown to be useful in the diagnosis of PPA. [18F] FDG-PET assesses cerebral glucose metabolism, which is abnormally reduced in neurodegeneration due to synaptic dysfunction and neuronal loss. The characteristic pattern of hypometabolism in lvPPA includes asymmetric involvement of the left posterior peri-Sylvian and left lateral temporoparietal regions, mirroring previously described regions of atrophy, specifically involving the left inferior parietal lobule and left posterior superior and middle temporal gyri, including the expected Wernicke area (Broadman area 22) (Figs 3 and 4). SPECT, which demonstrates regional hypoperfusion similar to the metabolic alterations on [18F] FDG-PET is infrequently used in clinical practice due to technical disadvantages and poorer accuracy.

Amyloid PET is a valuable examination for the diagnosis of AD and other neurodegenerative disorders demonstrating Aβ-pathology. In a meta-analysis of 1251 patients from 36 dementia centers, Aβ-positivity was identified in 86% of patients with lvPPA with evidence of AD pathology identified in 76% of those who
underwent a postmortem examination (Fig 5).\textsuperscript{17} $\alpha\beta$-positivity was also seen in a minority of those with nfvPPA (20%) and svPPA (16%); however, $\alpha\beta$-positivity was thought to represent a concomitant age-related process in these patients rather than being attributable to their PPA syndrome. In another study, 88% (46 of 52 patients) meeting the criteria for lvPPA demonstrated $\alpha\beta$-positivity, with low $\alpha\beta$-positivity rates in patients not meeting criteria for lvPPA (10%, 13 of 130 patients).\textsuperscript{18} In a study evaluating amyloid metabolism in PPA, 100% (4/4) of patients with lvPPA demonstrated elevated cortical Pittsburgh compound B uptake versus 16% (1/6) of patients with nfvPPA and 20% (1/5) of those with svPPA.\textsuperscript{19} Thus, amyloid PET imaging can be useful in distinguishing lvPPA from nfvPPA and svPPA, noting that comorbid age-related $\alpha\beta$-pathology may occur in each of these entities and an amyloid PET scan with positive findings does not equate to a diagnosis of lvPPA.

In addition to $\beta$-amyloid plaques, $\tau$ neurofibrillary tangles are a hallmark pathologic finding in AD.\textsuperscript{20} $\tau$-targeting PET tracers have been used for molecular imaging in PPA, most commonly $[^{18}\text{F}]$ flortaucipir (AV-1451), an FDA-approved, first-generation $\tau$ PET ligand.\textsuperscript{21,22} In a study investigating use of $[^{18}\text{F}]$ AV-1451 in PPA, patients with lvPPA exhibited striking uptake throughout the neocortex, most notably in the left temporoparietal region, compared with controls and subjects with other PPA variants, confirming the use of this radiotracer in distinguishing PPA subtypes.\textsuperscript{23} In a case series of patients with typical amnestic AD and atypical variants (posterior cortical atrophy, lvPPA, and corticobasal syndrome), all patients demonstrated region-specific distribution of $[^{18}\text{F}]$ AV-1451, indicating that $\tau$ PET can serve as a key biomarker linking molecular AD neuropathologic conditions with clinically significant neurodegenerative syndromes.\textsuperscript{24}

The use of PET tracers that target the translocator protein 18 kDa (TSPO) have also been explored in PPA, with the goal of characterizing the role of microglial activation and associated neuroinflammation in the pathogenesis of PPA. TSPO, originally named the peripheral benzodiazepine receptor, is an 18-kDa outer mitochondrial membrane protein, which has been found in disease-relevant areas across a broad spectrum of neurodegenerative diseases.\textsuperscript{24} Histopathologic studies have demonstrated asymmetric distribution of activated microglia in PPA, including high microglial densities in the superior temporal and inferior frontal gyri of the language-dominant hemisphere, consistent with postmortem and/or in vivo atrophy distribution.\textsuperscript{25} Patterns of microglial activation revealed variation favoring areas of increased atrophy in regions associated with language function, demonstrating concordance among patterns of microglial activation, atrophy, and clinical PPA phenotype.\textsuperscript{25} These findings support the potential use of TSPO PET in the evaluation of PPA subtypes.

**Semantic Variant PPA**

svPPA, previously referred to as “semantic dementia” in case reports throughout the 20th century, is a heterogeneous clinical syndrome characterized by progressive loss of knowledge about words and objects, including a fluent dysphasia with severe anoma, reduced vocabulary, and prominent impairment of single-word comprehension, which progresses to a stage of virtually complete dissolution of the semantic components of language.\textsuperscript{26} The criteria for the clinical diagnosis of svPPA are detailed in Table 2. Notably, a right temporal lobe–predominant variant of svPPA featuring impaired facial recognition (ie, prosopagnosia) and changes in affect and social behavior, in addition to semantic memory impairment, has also been described.\textsuperscript{34} SvPPA is currently recognized within the clinical spectrum of FTLD (Fig 1), with underlying histopathology most often attributable to TDP-43 type C, characterized by long dystrophic neurites and identified in >80% of patients with svPPA.\textsuperscript{5,35,36} Some cases have been associated with other histopathologic findings, including FTLD-TDP types A and B, FTLD-\(\tau\) (particularly 3R-\(\tau\) [Pick bodies] and 4R globular glial tauopathy), as well as AD pathology.\textsuperscript{5,35,36} SvPPA is almost always sporadic, rather than familial, and is the least heritable compared with other FTLD syndromes, with only 2%–4% of cases demonstrating an autosomal dominant pattern and suspected family history identified in 2%–17% of patients.\textsuperscript{26,37,38} Genetic disorders linked to familial FTLD-TDP include those associated with mutations in the GRN gene (FTLD-TDP type A), expansions in the chromosome 9 open reading frame 72 (C9orf72) gene (FTLD-TDP types A or B), and mutations in the valosin-containing protein (VCP) gene (FTLD-type D).\textsuperscript{39} In the absence of a strong family history of svPPA, an underlying genetic abnormality is considered unlikely.\textsuperscript{35} The precise prevalence of svPPA is not definitively known because diagnosis requires extensive clinical expertise and available data has largely been derived from tertiary care research centers referrals, likely not representative of the general population. However, the 2016 retrospective analysis from the French AD databank identified 26% (25 of 97) of PPA cases with CSF biomarkers to be svPPA, with a mean age of disease onset at 59.5 years and an average delay of 4.5 years between first symptom and diagnosis, a male predominance (68%; 8:17, female/male), and underlying AD pathology in only 20% of cases.\textsuperscript{40} The estimated svPPA prevalence based on the French AD databank cohort was 0.8 per 100,000 individuals, increasing with patient age. With regard to FTLD, its overall prevalence varies widely across studies, ranging from 2 to 31 per 100,000 individuals with an estimated true point prevalence of 15 to 22 per 100,000 and an incidence of 2.7 to 4.1 per 100,000 in those younger than 70 years of age.\textsuperscript{40} One series assessing 353 consecutive patients with FTLD identified 18.7% as having svPPA.\textsuperscript{41} An epidemiologic study investigating FTLD syndromes in 2 UK counties with a population of 1.69 million yielded an estimated svPPA prevalence of 1.2 per 100,000.\textsuperscript{42}

No disease-modifying medications are currently available for the treatment of svPPA. A variety of psychotropic medications have been used to manage associated behavioral symptoms, though evidence of efficacy from randomized clinical trials is lacking. Speech-language therapy has been shown to slow progression of anoma and may even offer a protective benefit to lexical items not yet lost, noting that treatment is suspected to be most beneficial at early stages of disease, supporting the advantage of an early diagnosis.\textsuperscript{43} A double-blind, sham-controlled, randomized clinical trial of transcranial direct current stimulation in patients with svPPA is ongoing.\textsuperscript{44}
Structural Imaging. The most common imaging features associated with svPPA include regional atrophy predominantly involving the left temporal lobe, most marked anteriorly involving the temporal pole (Fig 6). While focal atrophy is typically more pronounced on the left side, patients presenting with right-dominant temporal atrophy have been described in the literature. A study involving voxel-based morphometry in patients with svPPA reported that those with prosopagnosia had bilateral temporal lobe GM volume loss with greater involvement to the right, while those without prosopagnosia had predominantly left anterior temporal lobe volume loss. A meta-analysis of voxel-based morphometry studies investigating svPPA identified reduced GM volume in the bilateral fusiform and inferior temporal gyri, extending to the medial portion of the temporal lobes with involvement of the amygdala and parahippocampal gyri, as well as the left temporal pole, middle temporal gyrus, and caudate nucleus. Surface-based analysis of patients with svPPA identified marked cortical thinning in the left temporal lobe, particularly at the temporal pole; entorhinal cortex; and parahippocampal, fusiform, and inferior temporal gyri, with similar-yet-less extensive involvement of the contralateral cortex. Similarly, a longitudinal investigation mapping the progression of GM atrophy in predominantly left-versus-predominantly right temporal lobe variants of svPPA identified significant progression of GM atrophy in both the affected and contralateral temporal regions. Voxel-based morphometry has also identified asymmetric regional reduction in the temporal, periventricular, and callosal WM in patients with svPPA. DTI has been used to study structural connectivity changes on the whole-brain level in patients with svPPA, with reports of reduced fractional anisotropy and increased diffusivity in the anterior temporal lobe extending dorsally and posteriorly into the ventral frontal regions. Tractography has similarly been implemented in svPPA, identifying disruptions of structural connectivity related to GM atrophy, most severely affecting the WM tracts connecting the temporal regions with the frontal, parietal, and occipital regions (ie, uncinate fasciculus, arcuate fasciculus, superior longitudinal fasciculus, and inferior longitudinal fasciculus).

Functional Imaging. Task-based fMRI studies have reported that patients with svPPA compared with healthy controls demonstrate decreased activation in the mid-fusiform and superior temporal gyri; increased activation in the intraparietal sulcus, inferior frontal gyrus, and left superior temporal gyrus/sulcus; and lack of activation in the anterior temporal lobe. Similarly, resting-state fMRI studies in patients with svPPA have demonstrated reduced functional connectivity in the language and executive networks, with extensive disruptions between the anterior temporal lobe and a broad range of brain regions across the temporal, frontal, parietal, and occipital lobes. Magnetoencephalographic imaging has also been implemented to investigate whole-brain resting-state functional connectivity, identifying significant hyposynchrony of α and β frequencies within the left temporoparietal junction in patients with svPPA.

Regarding ASL MR imaging, a study investigating the prognostic value of regional CBF as measured by ASL MR imaging in patients with svPPA reported that ASL MR imaging may be sensitive to functional changes not identified on structural MR imaging, potentially serving as a prognostic biomarker marker of disease progression. Further studies assessing the potential role of ASL in patients with svPPA as well as its performance compared with [18F] FDG-PET are warranted.

Molecular Imaging. [18F] FDG-PET and SPECT can be performed to demonstrate characteristic asymmetric hypometabolism/hypoperfusion predominantly affecting the anterior temporal regions, most
commonly involving the left temporal pole (Figs 8 and 9). While svPPA is often associated with TDP-43 pathology, no available PET radioligand exists to date. Given the presence of AD pathology in some cases of svPPA, reported in up to 20% of cases within the 2016 French AD data-bank retrospective cohort,14 amyloid PET may result in an amyloid-positive examination in a minority of svPPA patients. While pathology is infrequently reported in patients with svPPA, studies with radioligands that demonstrate an affinity for $\tau$, including $[^{18}F]$ THK-5351 and flortaucipir ($[^{18}F]$ AV-1451), have demonstrated asymmetric retention in the temporal lobes in a pattern consistent with the expected distribution of TDP-43 pathology, hypothesized to reflect off-target binding to monoamine oxidase as a consequence of inflammation in the affected regions.51,56-58 A study investigating the novel $\tau$ PET tracer $[^{18}F]$ PI-2620, which has a low affinity for monoamine oxidase, demonstrated slightly elevated uptake involving the anterior and lateral temporal lobes in 1 of 2 subjects with svPPA, without elevated uptake in the other subject [Fig 10].59 There are no studies to date of PET ligands with an for inflammatory biomarkers (eg, monoamine oxidase B, TSPO/peripheral benzodiazepine receptor, or cyclooxygenase) in svPPA.60

Nonfluent/Agrammatic PPA

nvfPPA, previously referred to as “progressive nonfluent aphasia” in a series by Grossman et al61 and “PPA with agrammatism” by Mesulam,62 is the most diverse of the 3 PPA subtypes. Classically, patients present with a progressive language-predominant disturbance characterized by agrammatism in language production and apraxia of speech, with abnormally short (ie, telegraphic) phrases that tend to lack function words.63 Patients with nvfPPA have reduced verb production and diminished complexity in terms of grammar use,63 as well as difficulty with complex coordination of muscle groups involved in articulation of speech sounds (ie, apraxia of speech) and distortion of prosody (ie, rhythm, stress, and intonation of speech).64 The rate of word production in patients with nvfPPA has been reported to be less than one-third of the rate in healthy adults.65 Single-word

FIG 8. $[^{18}F]$ FDG-PET (A), axial TI (B), and PET MR imaging (C) views demonstrate an abnormal FDG distribution pattern with markedly decreased tracer uptake in the temporal lobes, particularly in the left temporal pole. There is corresponding advanced cortical atrophy with a “knife-blade” appearance in the left anterior temporal lobe on the axial TI sequence.

FIG 9. $[^{18}F]$ FDG-PET cortical surface maps demonstrate an abnormal FDG distribution pattern with severe left and moderate right hypometabolism in the anterior temporal lobes (A), with corresponding disproportionate cortical atrophy, particularly pronounced in the left temporal pole visualized on brain CT images (B), findings further supported by semiquantitative FDG-PET analysis using $z$ scores calculated in comparison with findings in age-matched cognitively healthy controls, semiquantitative FDG-PET analysis demonstrate markedly decreased values in the temporal lobes including the temporal poles (left > right) (B).
comprehension is notably preserved, though patients with nfvPPA may experience difficulty understanding sentences with complex syntax, such as those with relative clauses (eg, “he met a man who knew his brother”) or those with passive voice (eg, “the snake was bitten by the mongoose”).62,64 Complete criteria for a clinical diagnosis of nfvPPA, as defined by Gorno-Tempini et al,6 are described in Table 2.

Although typically considered a tauopathy, nfvPPA is the most heterogeneous of the PPAs, with a variety of other associated underlying pathologies. While 4R-τ is the most commonly reported underlying pathology, 1 postmortem series identified 23% of patients with nfvPPA exhibiting 3Ra-τ pathology (Pick bodies) and a minority with underlying TDP-43 or AD-type pathology.65 Patients with apraxia of speech and parkinsonism are more often associated with having a tauopathy than TDP-43 pathology.5,66 Similar to its pathologic heterogeneity, the clinical spectrum of nfvPPA has been described as the most diverse of the PPA subtypes, with a number of variant nfvPPA subsyndromes reported.11

NfvPPA is also the most heritable of the PPAs, with approximately 30% of patients reporting a positive family history.67 Mutations in all of the major genes associated with FTLD (eg, GRN, MAPT, C9orf72) have been identified in nfvPPA. Therefore, genetic screening should be considered in patients with a relevant family history. Additional detailed phenotyping of the genetic forms of nfvPPA will be required to improve on the continuously evolving understanding of PPA variants within the spectrum of FTLD.

The precise prevalence of nfvPPA is not definitively known because diagnosis requires extensive clinical expertise and available data have largely been derived from tertiary care research center referrals, likely not representative of the general population. However, the 2016 retrospective analysis from the French AD database identified 24% (23 of 97) of PPA cases with available CSF biomarkers as nfvPPA, with a mean age of disease onset at 60.9 years, an average delay of 2.3 years between first symptom and diagnosis, slight male predominance (52%; 11:12, female/male), and underlying AD pathology in 35% of cases.5 In 2 series of 353 patients with FTLD, 24.6% of cases were identified as nfvPPA.41 An epidemiologic study investigating FTLD syndromes in 2 UK counties with a population of 1.69 million yielded an estimated nfvPPA prevalence of 1.5 per 100,000.42

No pharmacologic option exists to improve or protect against declining function for patients with nfvPPA. However, speech-language treatment has demonstrated efficacy and structured oral reading has been proposed as a treatment method for apraxia of speech in nfvPPA.68 Additionally, transcranial direct current stimulation over the left posterior peri-Sylvian region and the Broca area has also been investigated as a potential treatment technique for nfvPPA.69 Supportive care remains the mainstay of PPA treatment, noting that patients with nfvPPA may experience dysphagia and should consult with a dietician or speech therapist for consideration of assisted feeding.11 Early detection of deficits, particularly physical, is critical to optimize outcomes related to changes in functional status. Support groups can also be extremely helpful, both for patients and their caregivers. Patients with nfvPPA who experience limited verbal output but with preserved comprehension may also benefit from alternative forms of communication devices.

Structural Imaging. The most common imaging feature associated with nfvPPA is regional atrophy predominantly involving the inferior frontal, opercular, and insular regions of the dominant hemisphere (Broadmann area 44/45; Broca area), most commonly on
Longitudinal progression of atrophy in nfvPPA has been reported to involve the posterior frontal regions, supplementary motor area, insula, striatum, inferior parietal regions, and underlying WM. Furthermore, atrophy typically progresses from the frontal operculum to the supplementary motor complex through the frontal aslant tract, which plays a role in the initiation and execution of movements, particularly articulation, and ultimately to the basal ganglia and supramarginal gyrus. The loss of integrity of the frontal aslant tract in nfvPPA is associated with distortion errors made by patients in spontaneous speech, as well as verbal fluency task performance. Therefore, nfvPPA is an example of a network disorder involving the circuit of regions and connections involved in speech production.

FIG 12. Axial CT (A), axial T2-weighted MR imaging (B), axial T1-weighted MR imaging (C), coronal CT (D), and coronal T1-weighted MR imaging (E) in a right-handed individual with apraxia of speech demonstrate asymmetric widening of the left Sylvian fissure with predominant left posterior frontoinsular atrophy (black arrows, A–E), suspicious for nfvPPA.

Such results suggest the possibility of fMRI serving as a useful imaging technique for the early detection of PPA, particularly nfvPPA, which may ultimately improve patient outcomes as disease-modifying therapies emerge in the clinical setting.

There are limited data regarding the use of ASL for nfvPPA specifically. Further studies assessing the potential role of ASL for patients with nfvPPA as well as its performance compared with $^{18}$F FDG-PET are needed.

Functional Imaging. A resting-state fMRI study demonstrated decreased functional connectivity between the left inferior frontal gyrus and posterior middle temporal gyrus in nfvPPA, even in patients without advanced atrophy.

Molecular Imaging. $^{18}$F FDG-PET and SPECT can be performed to demonstrate characteristic asymmetric hypometabolism-hypoperfusion predominantly affecting the left posterior frontal and peri-insular regions, including the left frontal operculum (Fig 13). Specifically, metabolic reduction in the left posterior frontoinsular region, including the inferior frontal gyrus, insula, and premotor and supplementary motor areas, is necessary to make an imaging-supported diagnosis of nfvPPA.

Regarding the cortical amyloid burden in nfvPPA, a study investigating $^{[1]}$C-Pittsburgh compound B in PPA subtypes demonstrated increased binding in only a few subjects with nfvPPA, in an uptake pattern similar to that of AD, including elevated tracer binding throughout the neocortex and striatum. A recent study investigating patients with PPA with discordant amyloid status (eg, nfvPPA with AD pathology) found that most cases exhibited FTLD-τ as the primary pathologic diagnosis with AD as an incidental age-related contribution. Specifically, 24 of 28 patients (86%) with svPPA and 28 of 31 patients (90%) with nfvPPA had negative amyloid PET findings, whereas 25 of 26 patients (96%) with logopenic PPA had scans with positive findings.

AJNR Am J Neuroradiol 43:1230–43 Sep 2022 www.ajnr.org

1239
amyloid-positive svPPA and nfvPPA cases with available postmortem data (2 of 4 and 2 of 3, respectively) all had a primary FTLD and secondary AD pathology diagnoses. Therefore, some patients with nfvPPA may have a PET examination positive for amyloid, and the presence of amyloid-positivity is not pathognomonic for any 1 PPA variant.

\[18F\] flortaucipir PET tracers have been used in nfvPPA, most commonly [18F] flortaucipir (AV-1451), a first-generation \( \tau \) PET ligand, which exhibits increased uptake in the left-greater-than-right frontal operculum and left middle and inferior frontal gyri (Fig 14).19 nfvPPA has also been shown to demonstrate less robust-but-focal uptake in the frontal WM and subcortical structures (Fig 15), regions known to be functionally impaired in nfvPPA, suggesting disease-specific binding to FTLD-4R \( \tau \).81 In nfvPPA, AV-1451 has been used to study \( \tau \) propagation in the left-hemispheric syntactic network, which comprises anterior frontal and posterior temporal nodes connected by the left arcuate fasciculus, with deposition greatest in the 2 nodes of the

FIG 13. [18F] FDG-PET cortical surface maps demonstrate an abnormal FDG distribution pattern with severe left-greater-than-right hypometabolism, most pronounced in the dorsal frontal lobes and left peri-insular region [A], with corresponding disproportionate cortical atrophy particularly pronounced in the left insular region visualized on brain MR views [B], findings further supported by semiquantitative FDG-PET analysis using \( z \) scores calculated in comparison with age-matched cognitively healthy controls, demonstrating markedly decreased values in the left > right peri-insular region, including in the pars opercularis and pars triangularis of the left inferior frontal gyrus, corresponding to the expected Broca area [B].

FIG 14. [18F] Flortaucipir in nfvPPA. A, On voxelwise comparison with healthy controls, agPPA demonstrates increased uptake in the left-greater-than-right frontal operculum; middle and inferior frontal gyri; and left superior frontal gyrus (pFWE < .05). B, The W score frequency map demonstrates elevated W scores above 1.65 in the bilateral middle frontal gyrus and frontal operculum in approximately two-thirds of patients scanned, with voxels above 1.65 in 8 of 11 patients in peak areas. C, ROI analyses reveals group differences in those with in nfvPPA compared with controls in the bilateral pars opercularis (left, \( P = .0001 \); right, \( P = .0018 \)); pars triangularis (left, \( P = .0016 \); right, \( P = .0029 \)); precentral gyrus (left, \( P = .003 \); right, \( P = .0012 \)); and superior frontal gyrus (left, \( P = .03 \); right, \( P = .045 \)). Adapted with permission from Tsai et al.81
syntactic network. The left arcuate fasciculus also demonstrated decreased fractional anisotropy in nfvPPA, particularly near the anterior node, suggesting propagation from node to connected node in human brain networks in the setting of neurodegenerative diseases, including PPA.

TSPO-targeting tracers have been explored in PPA, with the goal of characterizing the role of microglial activation and associated neuroinflammation in the pathogenesis of PPA. One study identified significantly increased mean [11C] PK-11195 binding in FTLD (n = 5, including 4 patients with nfvPPA and 1 patient with behavioral-variant frontotemporal dementia) in regions such as the left dorsolateral prefrontal cortex, right hippocampus, and parahippocampus.

CONCLUSIONS
PPA is a unique and complex spectrum of neurodegenerative disorders that requires a multidisciplinary approach to diagnosis, relying on the aggregate findings of clinical presentation, histopathologic biomarkers, and imaging features. A number of structural, functional, and molecular imaging modalities can support an accurate diagnosis, and neuroradiologists should be familiar with the classic imaging features of each PPA subtype, because prompt and accurate diagnosis may allow improved outcomes and intervention, particularly as disease-modifying therapies enter clinical practice.

Disclosure forms provided by the authors are available with the full text and PDF of this article at www.ajnr.org.

REFERENCES
1. Patel KP, Wymer DT, Bhatia VK, et al. Multimodality imaging of dementia: clinical importance and role of integrated anatomic and molecular imaging. Radiographics 2020;40:200–22 CrossRef Medline
2. Brown RKJ, Bohnen NI, Wong KK, et al. Brain PET in suspected dementia: patterns of altered FDG metabolism. Radiographics 2014;34:684–701 CrossRef Medline
3. Irwin DJ. Tauopathies as clinicopathological entities. Parkinsonism Relat Disord 2016;22:S29–33 CrossRef Medline
4. Montembeault M, Brambati SM, Gorno-Tempini ML, et al. Clinical, anatomical, and pathological features in the three variants of primary progressive aphasia: a review. Front Neurol 2018;9:692 CrossRef Medline
5. Spinelli EG, Mandelli ML, Miller ZA, et al. Typical and atypical pathology in primary progressive aphasia variants. Ann Neurol 2017;81:430–43 CrossRef Medline
6. Gorno-Tempini ML, Hillis AE, Weintraub S, et al. Classification of primary progressive aphasia and its variants. Neurology 2011;76:1006–14 CrossRef Medline
7. Mesulam MM. Primary progressive aphasia: a language-based dementia. N Engl J Med 2003;349:1535–42 CrossRef Medline
8. Mesulam M, Wieneke C, Rogalski E, et al. Quantitative template for subtyping primary progressive aphasia. Arch Neurol 2009;66:1545–51 CrossRef Medline
9. Sietek EJ, Narozniaka E, Brockhuis B, et al. Neuroimaging in the differential diagnosis of primary progressive aphasia: illustrative case series in the light of new diagnostic criteria. Polish J Radiol 2014;79:251–58 CrossRef Medline
10. Gorno-Tempini ML, Dronkers NF, Rankin KP, et al. Cognition and anatomy in three variants of primary progressive aphasia. Ann Neurol 2004;55:335–46 CrossRef Medline
11. Marshall CR, Hardy CJ, Volkmer A, et al. Primary progressive aphasia: a clinical approach. J Neurol 2018;265:1474–90 CrossRef Medline

AJNR Am J Neuroradiol 43:1230–43 Sep 2022 www.ajnr.org 1241
12. Grossman M. Primary progressive aphasia: clinicopathological correlations. *Nat Rev Neurol* 2010;6:88–97 CrossRef Medline

13. Butts AM, Machulda MM, Duffy JR, et al. Neuropsychological profiles differ among the three variants of primary progressive aphasia. *J Int Neuropsychol Soc* 2015;21:429–35 CrossRef Medline

14. Magnin E, Démonet JF, Wallon D, et al. on behalf of ePLM orators. Primary progressive aphasia in the Network of French Alzheimer Plan Memory Centers. *J Alzheimers Dis* 2016;54:1459–71 CrossRef Medline

15. Teichmann M, Kas A, Boutet C, et al. Deciphering logopenic primary progressive aphasia: a clinical, imaging and biomarker investigation. *Brain* 2013;136:3474–88 CrossRef Medline

16. Lundeen TF, Seibyl JP, Covington MF, et al. Signs and artifacts in amyloid PET. *Radiographics* 2018;38:2123–33 CrossRef Medline

17. Bergeron D, Gorno-Tempini ML, Rabinovici G, et al. Prevalence of amyloid-β pathology in distinct variants of primary progressive aphasia. *Ann Neurol* 2018;84:729–40 CrossRef Medline

18. Botha H, Duffy JR, Whitwell JL, et al. Classification and clinicoradiologic features of primary progressive aphasia (PPA) and apraxia of speech. *Cortex* 2015;59:220–36 CrossRef Medline

19. Rabinovici GD, Jagust WJ, Furst AJ, et al. AB amyloid and glucose metabolism in three variants of primary progressive aphasia. *Ann Neurol* 2008;64:388–401 CrossRef Medline

20. Beyer L, Brendel M. Imaging of tau pathology in neurodegenerative diseases: an update. *Semin Nucl Med* 2021;51:253–63 CrossRef Medline

21. Josephs KA, Martin PR, Botha H, et al. [18F]AV-1451 tau-PET and primary progressive aphasia. *Ann Neurol* 2018;83:599–611 CrossRef Medline

22. Mueller A, Bullich S, Barret O, et al. Tau PET imaging with 18F-PI-2620 in patients with Alzheimer disease and healthy controls: a first-in-humans study. *J Nucl Med* 2020;61:911–19 CrossRef Medline

23. Xia C, Makaretz SJ, Caso C, et al. Association of in vivo [18F]AV-1451 tau PET imaging results with cortical atrophy and symptoms in typical and atypical Alzheimer disease. *JAMA Neurology* 2017;74:427–36 CrossRef Medline

24. Werry EL, Bright FM, Piguet O, et al. Recent developments in TSPO PET imaging as a biomarker of neuroinflammation in neurodegenerative disorders. *Int J Mol Sci* 2019;20:1–21 CrossRef Medline

25. Kim G, Bobolkan K, Gelen T, et al. Atrophy and microglial distribution in primary progressive aphasia with transactive response DNA-binding protein-43 kDa. *Ann Neurol* 2018;83:1096–1104 CrossRef Medline

26. Hodges J, Patterson K, Oxbury S, et al. Semantic dementia: progressive fluent aphasia with temporal lobe atrophy. *Brain* 1992;115:1783–806 CrossRef

27. Mahalingam S, Chen MK. Neuroimaging in dementias. *Semin Neurol* 2019;39:188–99 CrossRef Medline

28. Masdeu JC. Neuroimaging of diseases causing dementia. *Neurol Clin* 2020;38:65–94 CrossRef Medline

29. Nasrallah IM, Wolk DA. Multimodality imaging of Alzheimer disease and other neurodegenerative dementias. *J Nucl Med* 2014;55:2003–11 CrossRef Medline

30. Villedame VL, Barkhof F, Garibotto V, et al. Molecular imaging approaches in dementia. *Radiology* 2021;298:517–30 CrossRef Medline

31. Clark DG, Charuvastra A, Miller BL, et al. Fluent versus nonfluent primary progressive aphasia: a comparison of clinical and functional neuroimaging features. *Brain Lang* 2005;94:54–60 CrossRef Medline

32. Collins JA, Montal V, Hochberg D, et al. Focal temporal pole atrophy and network degeneration in semantic variant primary progressive aphasia. *Brain* 2017;140:457–71 CrossRef Medline

33. Panegyres PK, McCarthy M, Campbell A, et al. Correlative studies of structural and functional imaging in primary progressive aphasia. *Am J Alzheimers Dis Other Dement* 2008;23:184–91 CrossRef Medline

34. Thompson SA, Patterson K, Hodges JR. Left/right asymmetry of atrophy in semantic dementia: behavioral-cognitive implications. *Neurology* 2003;61:1196–203 CrossRef Medline

35. Landin-Romero R, Tan R, Hodges JR. An update on semantic dementia: genetics, imaging, and pathology. *Alzheimers Res Ther* 2016;8:1–9 CrossRef Medline

36. Rohrer JD, Geser F, Zhou J, et al. TDP-43 subtypes are associated with distinct atrophy patterns in frontotemporal dementia. *Neurology* 2010;75:2204–11 CrossRef Medline

37. Seelaar H, Kamphorst W, Rosso SM, et al. Distinct genetic forms of frontotemporal dementia. *Neurology* 2008;71:1220–26 CrossRef Medline

38. Goldman J, Farmer J, Wood E, et al. Comparison of family histories in FTLD subtypes and related tauopathies. *Neurology* 2005;65:1817–19 CrossRef Medline

39. Pottier C, Ravenscroft TA, Sanchez-Contreras M, et al. Genetics of FTLD: overview and what else we can expect from genetic studies. *J Neurochem* 2016;138:32–53 CrossRef Medline

40. Onyike CU, Diehl-Schmid J. The epidemiology of frontotemporal dementia. *Int Rev Psychiatry* 2013;25:130–37 CrossRef Medline

41. Clerc MT, Guntan A. Frontotemporal lobar degeneration. *Dement Prevence, Risk Factors Manag Strateg* 2014;6:87–114 CrossRef Medline

42. Coyle-Gilchrist ITS, Dick KM, Patterson K, et al. Prevalence, characteristics, and survival of frontotemporal lobar degeneration syndromes. *Neurology* 2016;86:1736–43 CrossRef Medline

43. Henry ML, Reeson PM, Rapcsak SZ. Treatment for anemia in semantic dementia. *Semin Speech Lang* 2008;29:60–70 CrossRef Medline

44. Sanches C, Levy R, Benisty S, et al. Testing the therapeutic effects of transcranial direct current stimulation (tDCS) in semantic dementia: a double blind, sham controlled, randomized clinical trial. *Trials* 2019;20:1–15 CrossRef Medline

45. Chan D, Fox NC, Scähill RL, et al. Patterns of temporal lobe atrophy in semantic dementia and Alzheimer’s disease. *Ann Neurol* 2001;49:433–42 CrossRef Medline

46. Edwards-Lee T, Miller B, Benson D, et al. The temporal variant of frontotemporal dementia. *Brain* 1997;120:1027–40 CrossRef Medline

47. Josephs KA, Whitwell JL, Vemuri P, et al. The anatomic correlate of prosopagnosia in semantic dementia. *Neurology* 2008;71:1628–33 CrossRef Medline

48. Yang J, Pan P, Song W, et al. Quantitative meta-analysis of gray matter abnormalities in semantic dementia. *J Alzheimers Dis* 2012;31:827–33 CrossRef Medline

49. Rohrer JD, Warren JD, Modat M, et al. Patterns of cortical thinning in the language variants of frontotemporal lobar degeneration. *Neurology* 2009;72:1562–69 CrossRef Medline

50. Bramati SM, Rankin KP, Narvid J, et al. Atrophy progression in semantic dementia with asymmetric temporal involvement: a tensor-based morphometry study. *Neurobiol Aging* 2009;30:103–11 CrossRef Medline

51. Good CD, Scähill RL, Fox NC, et al. Automatic differentiation of anatomical patterns in the human brain: Validation with studies of degenerative dementias. *Neuroimage* 2002;17:29–46 CrossRef Medline

52. Yang Q, Guo QH, Bi YC. The brain connectivity basis of semantic dementia: a selective review. *CNS Neurosci Ther* 2015;21:784–92 CrossRef Medline

53. Guo CC, Gorno-Tempini ML, Gesierich B, et al. Anterior temporal lobe degeneration produces widespread network-driven dysfunction. *Brain* 2013;136:2979–91 CrossRef Medline

54. Ranasinghe KG, Hinkley LB, Beagle AJ, et al. Distinct spatiotemporal patterns of neuronal functional connectivity in primary progressive aphasia variants. *Brain* 2017;140:2737–51 CrossRef Medline

55. Olm CA, Kandel BM, Avants BB, et al. Arterial spin labeling perfusion predicts longitudinal decline in semantic variant primary progressive aphasia. *J Neurol Neurosurg Psychiatry* 2016;87:2212–17 CrossRef Medline

56. Kobayashi R, Hayashi H, Kawakatsu S, et al. [18F]THK-5351 PET imaging in early-stage semantic variant primary progressive aphasia: a report of two cases and a literature review. *BMC Neurol* 2018;18:109 CrossRef Medline

57. Bevan-Jones WR, Cope TE, Simon Jones P, et al. AV-1451 binding in vivo mirrors the expected distribution of TDP-43 pathology in the...
semantic variant of primary progressive aphasia. J Neurol Neurosurg Psychiatry 2018;89:1032–37 CrossRef Medline

58. Smith R, Santillo AF, Waldö ML, et al. F-Flortaucipir in TDP-43 associated frontotemporal dementia. Sci Rep 2019;9:1–10 CrossRef Medline

59. Mormino EC, Toueg TN, Azevedo C, et al. Tau PET imaging with 18F-P1-2620 in aging and neurodegenerative diseases. Eur J Nucl Med Mol Imaging 2021;48:2233–44 CrossRef Medline

60. Schain M, Kreisl WC. Neuroinflammation in neurodegenerative disorders: a review. Curr Neurol Neurosci Rep 2017;17:1–11 CrossRef Medline

61. Grossman M, Mickanin J, Onishi K, et al. The heritability and genetics of frontotemporal lobar degeneration. J Neurolinguistics 2009;22:370–83 CrossRef Medline

62. Grossman M. The non-fluent/agrammatic variant of primary progressive aphasia. Lancet Neurol 2012;11:545–55 CrossRef Medline

63. Olney NT, Spina S, Miller BL. Fronto temporal dementia. Neurology Clin 2017;35:339–74 CrossRef Medline

64. Grossman M, Mickanin J, Onishi K, et al. Treatment for apraxia of speech in nonfluent variant primary progressive aphasia. Brain 2013;136:77–88 CrossRef Medline

65. Wang J, Wu D, Chen Y, et al. Effects of transcranial direct current stimulation on language improvement and cortical activation in non-fluent variant primary progressive aphasia. Neurosci Lett 2013;549:29–33 CrossRef Medline

66. Mandelli ML, Vilaplana E, Brown JA, et al. Healthy brain connectivity predicts atrophy progression in non-fluent variant of primary progressive aphasia. Brain 2016;139:2778–91 CrossRef Medline

67. Mandelli ML, Vitali P, Santos M, et al. Two insular regions are differentially involved in behavioral variant FTD and nonfluent/agrammatic variant PPA. Cortex 2016;74:149–57 CrossRef Medline

68. Sajjadi SA, Sheikh-Bahaei N, Cross J, et al. Can MRI visual assessment differentiate the variants of primary-progressive aphasia? AJNR Am J Neuroradiol 2017;38:954–60 CrossRef Medline

69. Mandelli ML, Caverzasi E, Binney RJ, et al. Frontal white matter tracts sustaining speech production in primary progressive aphasia. J Neurol 2014;349:75–67 CrossRef Medline

70. Wilson SM, Galantucci S, Tartaglia MC, et al. Syntactic processing depends on dorsal language tracts. Neuron 2011;72:397–403 CrossRef Medline

71. Wilson SM, Dronkers NF, Ogar JM, et al. Neural correlations of syntactic processing in the nonfluent variant of primary progressive aphasia. J Neurolinguistics 2010;30:16845–54 CrossRef Medline

72. Miller ZA, Mandelli ML, Rankin KP, et al. Handedness and language learning disability differenti ally distribute in progressive aphasia variants. Brain 2013;136:3461–73 CrossRef Medline

73. Dick AS, Garcia D, Graziano P, et al. The frontal aslant tract (FAT) and its role in speech, language and executive function. Cortex 2019;111:148–63 CrossRef Medline

74. Bonakdarpour B, Rogalski EJ, Wang A, et al. Functional connectivity is reduced in early-stage primary progressive aphasia when atrophy is not prominent. Alzheimer Dis Assoc Disord 2017;31:101–16 CrossRef Medline

75. Minoshima S, Drzegza AE, Barthel H, et al. SNMMI procedure standard/EANM practice guideline for amyloid PET imaging of the brain 1.0. J Nucl Med 2016;57:1316–22 CrossRef Medline

76. Santos-Santos MA, Rabinovici GD, Iaccarino L, et al. Rates of amyloid imaging positivity in patients with primary progressive aphasia. JAMA Neurol 2018;75:342–52 CrossRef Medline

77. Tsai RM, Bejanin A, Lesman-Segev O, et al. 18F-Flortaucipir (AV-1451) tau PET in frontotemporal dementia syndromes. Alzheimers Res Ther 2019;11:1–18 CrossRef Medline

78. Pascual B, Funk Q, Zanotti-Fregonara P, et al. Multimodal 18F-AV-1451 and MRI findings in nonfluent variant of primary progressive aphasia: possible insights on nodal propagation of tau protein across the syntactic network. J Nucl Med 2020;61:263–69 CrossRef Medline

79. Cagnin A, Rossor M, Sampson EL, et al. In vivo detection of microglial activation in frontotemporal dementia. Ann Neurol 2004;56:894–97 CrossRef Medline

80. Madhavan A, Whitwell JL, Weigand SD, et al. FDG PET and MR imaging in logopenic primary progressive aphasia versus dementia of the Alzheimer’s type. PLoS One 2013;8:e62471 CrossRef Medline