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Neural regions underlying object and action naming: Complementary evidence from acute stroke and primary progressive aphasia

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Abstract

Background: Naming impairment is commonly noted in individuals with aphasia. However, object naming receives more attention than action naming. Furthermore, most studies include participants with aphasia due to only one aetiology, commonly stroke. We developed a new assessment, the Hopkins Action Naming Assessment (HANA), to evaluate action naming impairments.

Aims: Our aims were to show that the HANA is a useful tool that can (1) identify action naming impairments and (2) be used to investigate the neural substrates underlying naming. We paired the HANA with the Boston Naming Test (BNT) to compare action and object naming. We considered participants with aphasia due to primary progressive aphasia (PPA) or acute left hemisphere stroke to provide a more comprehensive picture of brain-behaviour relationships critical for naming. Behaviourally, we hypothesised that there would be a double dissociation between object and action naming performance. Neuroanatomically, we hypothesised that different neural substrates would be implicated in object vs. action naming and that different lesion-deficit associations would be identified in participants with PPA vs. acute stroke.

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Declaration of Interest

Declarations of interest: none

Methods & Procedures: Participants (N=138 with PPA, N=37 with acute stroke) completed the BNT and HANA. Behavioural performance was compared. A subset of participants (N=31 with PPA, N=37 with acute stroke) provided neuroimaging data. The whole brain was automatically segmented into regions of interest (ROIs). For participants with PPA, the image variables were the ROI volumes, normalised by the brain volume. For participants with acute stroke, the image variables were the percentage of each ROI affected by the lesion. The relationship between ROIs likely to be involved in naming performance was modelled with LASSO regression.

Outcomes & Results: Behavioural results showed a double dissociation in performance: in each group, some participants displayed intact performance relative to healthy controls on actions but not objects and/or significantly better performance on actions than objects, while others showed the opposite pattern. These results support the need to assess both objects and actions when evaluating naming deficits. Neuroimaging results identified different regions associated with object vs. action naming, implicating overlapping but distinct networks of regions. Furthermore, results differed for participants with PPA vs. acute stroke, indicating that critical information may be missed when only one aetiology is considered.

Conclusions: Overall, the study provides a more comprehensive picture of the neural bases of naming, underscoring the importance of assessing both objects and actions and considering different aetiologies of damage. It demonstrates the utility of the HANA.

Keywords

aphasia; stroke; primary progressive aphasia; object naming; action naming

Introduction

While aphasia is most commonly caused by stroke, other aetiologies including brain tumor, traumatic brain injury, infection, and neurodegenerative diseases such as primary progressive aphasia (PPA) can also result in aphasia. Studying individuals with each of these causes of aphasia can provide complementary information about the neural bases of language. For example, many people with post-stroke aphasia have been observed who have difficulty linking word sounds and word meanings after focal damage to left posterior superior temporal cortex, supporting Wernicke's classic observations that the area is critical for word comprehension (e.g., Hillis, Rorden, & Fridriksson, 2017). Similarly, individuals with semantic variant PPA exhibit impaired semantic memory along with disproportionate atrophy of the anterior temporal lobes, supporting the theory that those regions form a semantic hub (e.g., Lambon Ralph, Jefferies, Patterson, & Rogers, 2016), another component of the network underlying word comprehension. Converging evidence across diseases has also been reported (Faria et al., 2013). For instance, double dissociations between spoken and written naming have been observed in both stroke and PPA (e.g., Caramazza & Hillis, 1990; Ellis & Young, 1988; Gorno-Tempini et al., 2011; Hillis, Oh, & Ken, 2004; Miceli & Capasso, 1997; Rapp, Benzing, & Caramazza, 1997; Tainturier, Moreaud, David, Leek, & Pellat, 2001). However, results from different aetiologies are seldom compared directly. Comparing individuals with different aetiologies of damage can provide a more comprehensive picture of the brain-behavior relationships critical for

language. It could be that similar patterns are observed regardless of aetiology if the same critical regions are damaged in both. Alternatively, it could be that complementary information is observed from different aetiologies since different regions are more likely to be damaged by stroke versus neurodegenerative disease (or other lesion types). Both complementary and converging evidence from different aetiologies can be informative. Here, we compare the neural substrates of object and action naming in PPA and acute stroke.

Naming impairments and their neural bases are commonly investigated in individuals with aphasia, both post-stroke (e.g., Arévalo, Lu, Huang, Bates, & Dronkers, 2011; Faroqi-Shah et al., 2014; Forseth et al., 2018; Fridriksson et al., 2018; Fridriksson, Fillmore, Guo, & Rorden, 2015; Gleichgerrcht et al., 2016; Halai, Woollams, & Lambon Ralph, 2017; Hillis et al., 2018; Hillis, Tuffiash, Wityk, & Barker, 2002; Hope, Leff, & Price, 2018; Lacey, Skipper-Kallal, Xing, Fama, & Turkeltaub, 2017; McKinnon et al., 2018; Mirman et al., 2015; Newhart, Ken, Kleinman, Heidler-Gary, & Hillis, 2007; Parkinson, Raymer, Chang, FitzGerald, & Crosson, 2009; Piras & Marangolo, 2007; Python, Glize, & Laganaro, 2018; Schwartz, Faseyitan, Kim, & Coslett, 2012; Schwartz et al., 2009; Thye & Mirman, 2018; Tochadse, Halai, Lambon Ralph, & Abel, 2018; Tsapkini, Frangakis, & Hillis, 2011; Walker et al., 2011; Weiss et al., 2016) and in those with PPA (e.g., Beber et al., 2019; Benetello et al., 2016; Budd et al., 2010; Cotelli et al., 2016; Gesierich et al., 2012; Gleichgerrcht, Fridriksson, & Bonilha, 2015; Henry & Grasso, 2018; Kemmerer, Rudrauf, Manzel, & Tranel, 2012; Leyton, Hodges, Piguët, & Ballard, 2017; Marcotte et al., 2014; Mesulam et al., 2013; Meyer, Faria, Tippett, Hillis, & Friedman, 2017; Migliaccio et al., 2016; Mion et al., 2010; Race et al., 2013; Riello et al., 2018; Schwartz et al., 2012; Sebastian et al., 2018; Snowden et al., 2018; Thompson, Lukic, King, Mesulam, & Weintraub, 2012). Furthermore, individuals with stroke and primary progressive aphasia have been observed whose naming impairment is more severe for objects than actions, while others display the opposite pattern of performance (e.g., Bak & Hodges, 2003; Beber et al., 2019; Berndt, Mitchum, Haendiges, & Sandson, 1997; Damasio & Tranel, 1993; Hillis et al., 2006, 2004; Hillis, Tuffiash, & Caramazza, 2002; Thompson et al., 2012; Zingeser & Berndt, 1990). This finding suggests that the difference between object and action naming is not just degree of difficulty, with actions being harder to name than objects, as has sometimes been assumed. Rather, at least some neural substrates of object and action naming seem to be in separate locations that can be damaged independently. Converging evidence from individuals with stroke and primary progressive aphasia as well as functional neuroimaging supports this claim. Many studies indicate that left posterior frontal regions support action naming, while left temporal cortex supports object naming (e.g., Benetello et al., 2016; Cotelli et al., 2016; DeLeon et al., 2007; Foundas, Daniels, & Vasterling, 1998; Gesierich et al., 2012; Halai et al., 2017; Hart & Gordon, 1990; Hillis et al., 2018; Hillis, Tuffiash, Wityk, et al., 2002; Hurley, Paller, Rogalski, & Mesulam, 2012; Kemmerer et al., 2012; Leff et al., 2009; Leyton et al., 2017; Martin, Haxby, Lalonde, Wiggs, & Ungerleider, 1995; Mesulam et al., 2013; Meyer et al., 2017; Migliaccio et al., 2016; Newhart et al., 2007; Piras & Marangolo, 2007; Race et al., 2013; Schwartz et al., 2009; Snowden et al., 2018; Tranel, Adolphs, Damasio, & Damasio, 2001; Tranel, Kemmerer, Adolphs, Damasio, & Damasio, 2003; Tranel, Manzel, Asp, & Kemmerer, 2008; Tsapkini et al., 2011; Tyler, Russell, Fadili, & Moss, 2001; Walker et al., 2011; Weiss et al., 2016)

Assessment of action naming is important not only to detect all cases of impaired naming, but also to understand other aspects of production. For instance, grammatical sentence production requires verb retrieval. However, aphasia assessments often focus on naming of objects alone, making them insensitive for detecting certain language deficits and providing an incomplete view of the neural regions involved in naming. Here, we address this gap by presenting a new assessment to evaluate action naming, the Hopkins Action Naming Assessment (HANA). While this assessment has been used in conjunction with a battery of other assessments in some published reports (de Aguiar et al., 2020; Keator et al., 2019, Keator et al., 2020; Long, Sebastian, Faria, & Hillis, 2018; Odolil et al., 2020; Purcell et al., 2017; Race et al., 2013; Riello et al., 2018; Rofes et al., 2019; Sebastian et al., 2016; Sebastian et al., 2018; Tippett et al., 2020; Tsapkini et al., 2018; Unal et al., 2020), the current study presents the first focused description of the assessment and its norming. The HANA consists of 30 black and white images of actions. The items are matched in frequency and length to the 30-item version of the Boston Naming Test (BNT; Mack, Freed, Williams, & Henderson, 1992) (see Table 1). The HANA can be given in conjunction with the BNT, which is already in frequent clinical use, providing diagnostically useful information about the relative strengths of action and object naming while requiring minimal time to be added to the diagnostic session.

In the current study, we administered both the BNT and the HANA to individuals with acute left hemisphere stroke (N=37) and PPA (N=138). Our first aim was to show that the HANA is clinically useful for identifying action naming impairments, which are separate from the object naming impairments that can be diagnosed via the BNT. We hypothesised that a double dissociation between object and action naming would be observed, with some individuals showing deficits for objects but not actions and vice versa. Our other aim was to show that these assessments can be a useful tool for comparing the neural substrates associated with performance on object and action naming. To investigate this aim, we used neuroimaging data from a subset of the same participants (N=31 with PPA, N=37 with acute stroke). We hypothesised that different neural substrates would be implicated in object vs. action naming and that different lesion-deficit associations would be identified in participants with PPA vs. acute stroke. The last hypothesis is based on the premise that different regions of the brain are affected in PPA vs. stroke, allowing sufficient power to detect associations in different regions. For example, if few people with stroke have damage to the temporal pole, damage to that area is unlikely to be statistically associated with naming or other deficits, even if the left temporal pole is a critical node of the naming network. To assess as many areas as possible that might be important for naming, we included all variants of PPA, as different brain regions tend to be affected across variants (e.g., Gorno-Tempini et al., 2004, 2011), as well as across aetiologies (PPA vs stroke). The overall goal of the work presented here is to demonstrate the utility of the HANA and provide a more comprehensive picture of brain-behavior relationships critical for naming.

Materials and Methods

Participants

PPA—Participants (N=138, 73 females, mean age 68.7 ± 8.24 years, mean symptom duration 42.0 ± 26.64 months) who had been diagnosed with primary progressive aphasia were recruited from the senior author's outpatient neurology clinic in accordance with the policies of the Johns Hopkins Medical Institutions IRB. Diagnosis of PPA was based on presentation with progressive language impairment that remains the most prominent symptom in the absence of other cognitive, behavior, or personality changes (Gorno-Tempini et al., 2011; Mesulam, 1982). Information about the participants with PPA, including demographic information and scores on the tasks of interest, is presented in Table 2.

If possible, participants were classified as having logopenic (N=50), nonfluent (N=41), or semantic (N=33) variants using recent guidelines (Gorno-Tempini et al., 2011) on the basis of detailed language and cognitive assessments, history, comprehensive neurological examination, and available neuroimaging. However, fitting one of the classification types was not required for inclusion: 14 participants were unclassifiable (either because they did not have all of the core distinguishing features of any variant, as in cases of pure anomia, or because they had core features of more than one variant). As a group, the unclassifiable participants were significantly earlier in the course of PPA (mean symptom duration = 29.9 ± 18.30 months) than those with lvPPA (mean symptom duration = 42.3 ± 24.97 months; $t(28)=-2.06, p<.05$) or svPPA (mean symptom duration = 48.7 ± 32.45 months; $t(41)=-2.52, p<.05$); they may have progressed after these assessments to fit one of the classification types. Across classification types, participants were comparable on education and age, with the exception that participants with svPPA were significantly younger (mean age 65.5 ± 7.60 years) than participants with lvPPA (mean age 70.3 ± 7.70 years; $t(69)=-2.80, p<.05$) or unclassifiable PPA (mean age 70.4 ± 7.26 years; $t(26)=-2.12, p<.05$).

All 138 PPA participants completed the HANA; 135 also completed the BNT. A subset of 31 PPA participants (7 svPPA, 12 nfvPPA, 11 lvPPA, 1 unclassified; mean age 68.3 ± 8.68 years; 14 female) who had high resolution T1 weighted image (T1-WI) anatomical MRI scans within five months of the naming assessment were included in the neuroimaging analysis. The other PPA participants were excluded from this analysis because they did not have the required imaging.

Stroke—Participants (N=37, 14 females, mean age 56.5 ± 13.41 years) with acute left hemisphere ischemic stroke were enrolled at Johns Hopkins Hospital or Johns Hopkins Bayview Medical Center in accordance with the policies of the Johns Hopkins Medical Institutions IRB. All had MRI-confirmed strokes, with diffusion-weighted imaging (DWI) revealing the presence of unilateral left hemisphere infarct without hemorrhage on the initial scan. Participants were premorbidly fluent English speakers with normal or corrected-to-normal vision and hearing. They did not have history of dementia, previous symptomatic stroke, or other neurological disease affecting the brain, nor did they have reduced level of consciousness or ongoing sedation. All 37 acute stroke participants completed the HANA within five days of hospital admission; 34 also completed the BNT. All stroke participants were included in the neuroimaging analysis. Information about the participants with acute

stroke, including demographic information and scores on the tasks of interest, is presented in Table 3.

Controls—An additional twenty-six neurologically intact participants (20 female, mean age 64.0 ± 12.05 years) were recruited to serve as controls for behavioral testing.

Behavioural assessments

Object naming: Boston Naming Test (short form) [BNT]—Every other item from the original 60-item Boston Naming Test (Kaplan et al., 1983) was administered, following Mack et al. (1992). Participants named black and white line drawings of 30 objects. Uncued first responses were scored. The twenty-six controls tested here provided correct first responses to 22–29 items ($mean=26.3 \pm 2.48$).

Action naming: Hopkins Action Naming Assessment [HANA]—Participants named black and white line drawings or photographs of 30 actions. These items were matched in frequency and length to the names of the objects in the BNT (Table 1) using N-Watch (Davis, 2005) and the MRC Psycholinguistic Database (Coltheart, 1981). Frequency measures were drawn from those reported by Kuera and Francis (1967), the CELEX English linguistic database (Baayen et al., 1995), the Sydney Morning Herald corpus (Dennis, 1995), and Thorndike and Lorge (1942). Welch two sample two-tailed t-tests showed that the two assessments did not differ on any of the length or frequency measures (all p-values $> .05$). Pilot testing of the assessment showed that each picture had at least 75% name agreement among a group of twenty young controls. The older controls tested in the present study provided correct first responses to 17–30 items ($mean=23.9 \pm 3.46$). Actions generally have poorer name agreement compared to objects, as several action names can correctly describe most actions (e.g. yell, shout, scream). For both objects and actions, if the participant gave a name that was correct (synonymic) but not the target name, they were asked to give another name; but only the target, if named without a cue, was considered correct.

Behavioural Analysis

We dichotomised naming performance as normal or impaired relative to controls. The mean and standard deviation of controls was used to calculate standard scores. A standard score of $z < -1.645$ was considered significantly worse performance than controls ($p < .05$, one tailed) and classified as impaired. The pattern of intact vs. impaired performance on the object and action naming tasks was compared for all individual participants.

For each individual, we also compared accuracy of object vs. action naming using two-tailed Fisher's Exact Tests.

Neuroimaging Analysis

PPA—Anatomical information was acquired using a MPRAGE T1 sequence on a 3T scanner. Scans occurred within five months of the cognitive testing reported here (mean scan-assessment interval: scans 1.0 ± 2.32 months subsequent to assessment). Atlas-based analysis was used to segment each T1 MRI into 289 regions of interest (ROIs) using

the MRI Cloud platform (www.mricloud.org). This type of analysis allows the anatomical parcellation defined in an atlas template image to be applied to the brain of each individual participant so that multiple regions of interest (ROIs) can be automatically segmented. Such an analysis ameliorates drawbacks of voxel-based analysis, which suffers from the noise resulting from the assumption that normalization is accurate enough to identify the same voxel in a standardised brain and the individual participants', and of manual delineation of ROIs, which suffers from limited reproducibility and cannot be practically applied to whole brain analyses because of the amount of time necessary to apply such analyses (Faria et al., 2010). For each participant, the volume of brain tissue in each region was calculated in native space. Based on previous lesion and neuroimaging studies of naming, we selected 20 of the 289 regions resulting from the automatic parcellation as regions of interest (e.g., Benetello et al., 2016; Cotelli et al., 2016; DeLeon et al., 2007; Foundas, Daniels, & Vasterling, 1998; Gesierich et al., 2012; Gorno-Tempini et al., 2011; Halai et al., 2017; Hart & Gordon, 1990; Hillis et al., 2018; Hillis, Tuffiash, Wityk, et al., 2002; Hurley, Paller, Rogalski, & Mesulam, 2012; Kemmerer et al., 2012; Leff et al., 2009; Leyton et al., 2017; Martin, Haxby, Lalonde, Wiggs, & Ungerleider, 1995; Mesulam et al., 2013; Meyer et al., 2017; Migliaccio et al., 2016; Newhart et al., 2007; Piras & Marangolo, 2007; Race et al., 2013; Schwartz et al., 2009; Snowden et al., 2018; Tranel, Adolphs, Damasio, & Damasio, 2001; Tranel, Kemmerer, Adolphs, Damasio, & Damasio, 2003; Tranel, Manzel, Asp, & Kemmerer, 2008; Tsapkini et al., 2011; Tyler, Russell, Fadili, & Moss, 2001; Walker et al., 2011; Weiss et al., 2016). These regions consisted of the pars opercularis, pars orbitalis, and pars triangularis of the left inferior frontal gyrus; left precentral gyrus; left postcentral gyrus; left supramarginal gyrus; left angular gyrus; left superior temporal gyrus; right superior temporal gyrus; left superior temporal pole; right superior temporal pole; left middle temporal gyrus; right middle temporal gyrus; left middle temporal pole; right middle temporal pole; left inferior temporal gyrus; right inferior temporal gyrus; left fusiform gyrus; right fusiform gyrus; and left insula. The temporal lobe was subdivided such that the temporal pole consists of the regions of the superior and middle temporal gyri (labeled superior and middle temporal poles, respectively) lying anterior to a vertical plane through the anterior commissure, based on the anterior ending of the superior temporal sulcus. The superior and middle temporal poles were separated in the atlas template because they have different connectivity with other regions (Insausti et al., 1998; Kondo et al., 2003, 2005; Stefanacci et al., 1996). The volume for each ROI was normalised by cerebral volume, defined as total brain volume removing myelencephalon and cerebrospinal fluid (CSF), to control for relative regional atrophy. The ratio of cerebral to intracranial volume, defined as total brain volume removing myelencephalon but including CSF, was also calculated for use as a predictor in models to control for inter-individual brain size differences.

Acute stroke—Participants underwent MR imaging that included MPRAGE T1 anatomical scans as well as diffusion weighted imaging (DWI). Technicians blinded to the behavioral results used MRIcron (<https://www.nitrc.org/projects/mricron>) to trace areas of tissue disfunction, defined as areas of dense ischemia/infarct that were bright on DWI maps but dark on apparent diffusion coefficient (ADC) maps. Traced lesions were coregistered to T1 scans using SPM12 (<https://www.fil.ion.ucl.ac.uk/spm/software/spm12>). Lesioned tissue was replaced with tissue from the undamaged right hemisphere. The resulting

artificial T1 image was then segmented using the automated MRICloud pipeline (<https://braingps.mricloud.org/>) that relies on a highly accurate large deformation diffeomorphic mapping algorithm to transform images between native space and atlas space and a multi-atlas fusion label algorithm that allows the use of multiple atlases to reduce segmentation and mapping inaccuracies. Via this pipeline, each individual's scan was segmented into 289 regions of interest. The original lesion map was overlaid onto the resulting segmentation to calculate the percentage of tissue damaged in each of the 289 regions. As in the PPA analysis, we selected a smaller subset of regions for further analysis. From the list of ROIs used in the PPA analysis, seven were removed due to lack of variability: no participants had lesions in the six right hemisphere regions since unilateral left hemisphere stroke was an inclusion criterion, and none happened to have lesions in the left middle temporal pole. A minimum of five participants had lesions in the thirteen remaining ROIs used in the analyses of acute stroke data, consisting of the left inferior frontal gyrus pars opercularis, pars orbitalis, and pars triangularis; left precentral gyrus; left postcentral gyrus; left supramarginal gyrus; left angular gyrus; left superior temporal gyrus; left superior temporal pole; left middle temporal gyrus; left inferior temporal gyrus; left fusiform gyrus; and left insula. Total lesion volume was also calculated for use as a predictor in models to control for severity of stroke.

Relating neuroimaging and behavioural data—Least Absolute Shrinkage and Selection Operator (LASSO) regression (Tibshirani, 1996) was used to evaluate which ROIs were related to behavior on object and action naming. This method, which results in selection of a subset of covariates, was used instead of simpler multiple regression analyses as the neuroimaging data had high multicollinearity. That is, participants with damage in one region were highly likely to have damage in neighboring regions. LASSO regression is appropriate in this situation as it reduces over-fitting and provides for automated feature selection: when multiple highly correlated variables are entered into the model, only one is retained while the others are set to zero. This is helpful here where we have a large number of predictors relative to the sample size and where some of those predictors are closely related (Meinshausen & Yu, 2009). LASSO regression with standardised features was instantiated with the `glmnet` package in R (<https://cran.r-project.org/web/packages/glmnet/index.html>), using leave-one-out cross-validation to select the λ value that resulted in the minimum mean cross validated error. Inference testing and calculation of p-values for the features selected by the LASSO was conducted with the `selectiveInference` package in R (<https://cran.r-project.org/web/packages/selectiveInference/selectiveInference.pdf>). Separate analyses were conducted for each of the four combinations of object vs. action naming and PPA vs. acute stroke. Object naming models used number correct on the BNT as the dependent variable; action naming models used number correct on the HANA as the dependent variable. PPA models included cerebral to intracranial volume ratio as well as regional volume normalised by cerebral volume for each of the 20 ROIs as predictors. These models used one-tailed LASSO regression, considering only positive predictors where smaller regional volume (i.e. less healthy tissue) was associated with worse naming performance. Acute stroke models included total lesion volume as well as percent damage in each of the 13 ROIs as predictors. These models used one-tailed LASSO regression,

considering only negative predictors where larger lesion volume (i.e. less healthy tissue) was associated with worse naming performance.

Results

Behavioural Results

We hypothesised that there would be a double dissociation in performance for object vs. action naming. To investigate this, we first established that the assessments identified participants with naming impairments. We separately considered participants whose aphasia was due to PPA vs. stroke. Performance is shown in Figures 1 and 2.

On the BNT, the cut-off for normal performance was 23 based on the definition of normal performance as a standard score of $z > -1.645$. Correct naming of 22 or fewer items was considered impaired. Of the 135 participants with PPA who completed the BNT, 101 (75%) showed impaired performance on object naming. Likewise, 13 of the 34 (38%) with acute stroke were impaired on the BNT. On the HANA, the cut-off value derived from control performance was 19. Impaired performance on the HANA (i.e., correctly naming 18 or fewer items) was demonstrated by 89 of the 138 (64%) participants with PPA and 11 of the 37 (30%) with acute stroke who completed this assessment.

For a double dissociation to exist between two cognitive processes, there must be some individuals who demonstrate impaired performance on tasks tapping one function in the face of intact performance on tasks tapping the other function as well as other individuals who show the opposite pattern. Here, we would expect to find some individuals who show impaired performance on the BNT but not the HANA and vice versa to observe a double dissociation between object and action naming. We investigated this in two ways, the combination of which is stronger than either alone.

First, we looked at individuals' patterns of performance on the two tasks defining impaired vs. intact performance with the cut-offs described above. We identified eighteen participants with PPA (ten lvPPA, two nfvPPA, one svPPA, and four unclassifiable) and five with acute stroke who showed impaired performance on the BNT but intact performance on the HANA (mean difference 2.9 ± 4.12 more items correct on the HANA than the BNT; range 13 more items correct on the HANA than the BNT to 3 more items correct on the BNT than the HANA). We also identified the opposite dissociation of impaired action but not object naming, finding five participants with PPA (four lvPPA, one nfvPPA) and one with acute stroke who showed impaired performance on the HANA but not the BNT (mean difference 8.5 ± 3.27 more items correct on the BNT than the HANA; range 5–13 more items correct on the BNT than the HANA). Together, these patterns of performance constitute a double dissociation.

Second, we directly compared performance on the two tasks, regardless of status relative to cut-offs, using Fisher's Exact Tests to identify individuals who performed significantly better on one task than the other ($p < .05$). Such individuals would further support the finding of a double dissociation, ruling out the possibility that the findings reported above are an artifact of the specific cut-offs used to define intact vs. impaired performance. As expected,

we observed both participants who performed significantly better on the HANA than on the BNT (two PPA [two svPPA], one acute stroke; mean difference 11.3 ± 1.53 more items correct on the HANA than the BNT; range 10–13 more items correct on the HANA than the BNT) and participants who performed significantly better on the BNT than on the HANA (twelve PPA [four lvPPA, two nfvPPA, six svPPA], one acute stroke; mean difference 9.2 ± 2.08 more items correct on the BNT than the HANA; range 6–13 more items correct on the BNT than the HANA).

To provide even stronger evidence of a double dissociation, we note that some individuals fit both criteria that we considered: they demonstrated intact performance on one task but impaired performance on the other task relative to controls AND their performance was significantly better on one task than the other. Under these combined criteria, we identified two individuals with deficits on object naming but not action naming (one PPA, one acute stroke; 8 items correct on the BNT vs. 21 correct on the HANA and 16 items correct on the BNT vs. 27 correct on the HANA) and three individuals with deficits on action naming but not object naming (three PPA; 26 items correct on the BNT vs. 18 correct on the HANA, 25 items correct on the BNT vs. 13 correct on the HANA, and 25 items correct on the BNT vs. 12 correct on the HANA). These behavioral results support the hypothesis that there is a double dissociation between object and action naming.

Neuroimaging Results

We hypothesised that different neural substrates would be implicated in object vs. action naming and for those with acute stroke vs. PPA. To investigate these hypotheses, we used LASSO regression models, one for each of the four combinations of object vs. action naming and PPA vs. acute stroke. Results are summarised in Tables 4 and 5.

Models for participants with PPA included all 20 regions of interest as predictors, as well as the ratio of cerebral to intracranial volume to control for brain size differences. These were one-tailed analyses in which only positive predictors were selected, reflecting relationships in which greater volume was indicative of better naming. For the model of object naming, we found that the volumes of left supramarginal gyrus, left superior temporal pole, right middle temporal gyrus, left inferior temporal gyrus, right inferior temporal gyrus, and left fusiform gyrus were associated with performance on the BNT, as was the ratio of cerebral to intracranial volume. None of the covariates were independent, statistically significant predictors of object naming in PPA, although there was a trend for cerebral to intracranial volume ratio ($p=.096$) and left fusiform gyrus volume ($p=.081$) to be independently predictive. For the model of action naming, we found that the volumes of left superior temporal pole, right superior temporal pole, right middle temporal gyrus, left fusiform gyrus, and right fusiform gyrus were associated with performance on the HANA, as was the ratio of cerebral to intracranial volume. None of the covariates were independent, statistically significant predictors of action naming in PPA, although there was a trend for larger ratio of cerebral to intracranial volume ($p=.091$) to be independently predictive. Results are summarised in Table 4 and visualised in Figure 3. Models for participants with acute stroke removed the regions of interest in which no participants had lesions, including the six right hemisphere ROIs and left middle temporal pole. Therefore, 13 regions of interest were

included, as was total lesion volume to control for severity of stroke. One-tailed analyses were conducted in which only negative predictors were selected, reflecting relationships in which less damage was indicative of better naming. For the model of object naming, we found that larger percentage of tissue lesioned in the left angular gyrus and left middle temporal gyrus were associated with worse performance on the BNT. Greater damage to the left angular gyrus independently predicted object naming performance ($p < .001$). For the model of action naming, we found that larger percentage of tissue lesioned in the left angular gyrus and left insula were associated with worse performance on the HANA, as was larger total lesion volume. None of the covariates were independent, statistically significant predictors of action naming in acute stroke. Results are summarised in Table 5 and visualised in Figure 4.

Comparing the results of the object and action naming models provides a test of the hypothesis that different neural substrates are involved in object vs. action naming. For participants with PPA, we did find regions that were involved with both, namely left superior temporal pole, right middle temporal gyrus, and left fusiform gyrus, as well as the ratio of cerebral to intracranial volume. However, in line with our hypothesis, we also found regions that were associated with object naming only: left supramarginal gyrus, left inferior temporal gyrus, and right inferior temporal gyrus. There were also regions associated with action naming only: right superior temporal pole and right fusiform gyrus. For participants with acute stroke, we again found a region commonly associated with both object and action naming: left angular gyrus. There were differences in the neural substrates for the two types of naming, as left middle temporal gyrus was associated with objects but not actions, while left insula and total lesion volume were associated with actions but not objects.

Comparing the results of the models of PPA and acute stroke participants provides a test of the hypothesis that distinct regions of damage are statistically associated with deficits in naming when considering individuals from these different populations. Of the eight regions associated with object naming in either aetiology, none were identified in both stroke and PPA. Likewise, of the seven regions associated with action naming in either aetiology, none were identified for both stroke and PPA. The regions that were associated with both object and action naming in PPA were distinct from those that were associated with both types of naming in stroke. Thus, in line with our hypotheses, different neural substrates were identified for PPA and acute stroke.

Discussion

In the present study, we paired a novel assessment of action naming, the HANA, with an established assessment of object naming in order to demonstrate their combined utility and explore the neural bases of naming. Clinical evaluations of naming are often (although not always) limited by considering only object naming; we expanded the scope of our project to compare matched picture naming assessments of actions and objects. Studies of the brain regions underlying language in neurological populations are often limited by including only participants with damage due to one aetiology; we expanded the scope of our project to compare participants with language deficits due to PPA as well as acute left hemisphere stroke. Looking at both object and action naming in multiple aetiologies in the same study

affords us an unusual perspective from which we were able to investigate three hypotheses: (1) there is a behavioral double dissociation between object and action naming; (2) different neural substrates are involved in object vs. action naming; and (3) different lesion-deficit associations are identified in those with PPA vs. acute stroke. Our results confirmed these hypotheses.

First, we were able to demonstrate the utility of the HANA in identifying participants with action naming impairments. Of the 175 participants who completed the HANA, 100 were impaired relative to controls. Furthermore, there was a double dissociation such that some participants were impaired on object naming but not on action naming, while others were impaired on action naming but not object naming. This was the case regardless of whether we defined impairment dichotomously based on the performance of healthy controls or whether we looked at relative impairment defined as significantly better performance on one task than the other. Each of these methods has limitations. Using cut-offs based on control performance may lead to classifying individuals whose performance is similar across tasks as showing a dissociation because their scores on one task happen to fall just above the cut-off while their scores on the other happen to fall just below the cut-off. Using significance testing may lead to classifying individuals who perform well on both tasks as having a dissociation when their performance on one task is numerically better than the other even though both are in the range of healthy controls. Some participants may be identified as significantly better on the BNT than the HANA due to chance since the relatively small group of control participants found naming of objects easier than naming of actions (the cut-off for normal performance was 4 items higher for the BNT than the HANA). However, combining these methods mitigates these limitations, increasing confidence that true double dissociations were observed: some participants showed intact performance on one task relative to controls that was significantly better than their performance on the other task, which was classified as impaired relative to controls, while others showed the opposite pattern. This finding underscores the clinical importance of assessing and treating naming not only of objects but also of actions. Evaluating only one type may lead to missing functionally important deficits; treating object naming exclusively, which is typical of aphasia therapy, may have implications for sentence formulation. It was not the case that actions were always harder to name than objects as has sometimes been asserted. This result was not surprising in light of double dissociations previously reported in the literature. However, it does demonstrate that the HANA is a useful tool for evaluating action naming. As shown here, it is sensitive to action naming impairments in participants with aphasia and acute stroke. It is relatively quick to administer and score clinically. It is matched to a widely used standardised assessment of object naming, the Boston Naming Test, and performance on the two can be compared to evaluate naming of objects and actions.

Second, we showed that, beyond their clinical utility in diagnosing impairments, the HANA and BNT can be used to investigate the neural bases of naming. Different regions were identified as related to object naming performance vs. action naming performance in the different groups. Based on previous studies, we expected left temporal cortex to be critical for object naming and left posterior frontal regions to be critical for action naming. This was not the pattern that we uncovered: instead, damage to left temporal regions was implicated in poor performance on both object and action naming, and damage to left frontal regions

was implicated in poor performance on neither object nor action naming. This divergence from expectations may be due to inadequate power in specific areas or to the statistical methods applied. LASSO regression reduces multicollinearity by selecting only the most predictive variable among those that are highly correlated. Regions that are in fact critical may have been eliminated if they were strongly correlated with those selected by the model. Although the pattern of regions we identified did not match our hypotheses, we did find distinct networks of areas involved in each type of naming. In participants with PPA, left supramarginal gyrus, left inferior temporal gyrus, and right inferior temporal gyrus were associated with object naming but not action naming, while right superior temporal pole and right fusiform gyrus were associated with action naming but not object naming. The left hemisphere regions related to object naming have previously been identified as involved in naming (e.g., Damasio & Tranel, 1993; Daniele et al., 1994; DeLeon et al., 2007; Hart & Gordon, 1990; Migliaccio et al., 2016; Mirman et al., 2015; Race et al., 2013; Riello et al., 2018; Schwartz et al., 2012; Thye & Mirman, 2018). The identification of right hemisphere homologues of left hemisphere language regions (right inferior temporal gyrus, right superior temporal pole, and right fusiform gyrus) may indicate that participants initially can compensate by recruiting right hemisphere regions; it is only when atrophy increases bilaterally that naming deficits become more severe (e.g., Rohrer et al., 2013).

In participants with acute stroke, left middle temporal gyrus was associated with object naming but not action naming, while left insula and total lesion volume were associated with action naming but not object naming. The left middle temporal gyrus has previously been implicated in object naming (Faroqi-Shah et al., 2014; Fridriksson et al., 2018; Hillis, Tuffiash, Wityk, et al., 2002; Piras & Marangolo, 2007; Python et al., 2018; Riello et al., 2018; Schwartz et al., 2009; Thye & Mirman, 2018; Tochadse et al., 2018). Larger total lesion volume was associated with worse performance on actions but not objects, which may indicate that action naming was a more difficult, cognitively demanding task for many – although, as described above, not all – of the participants. Similarly, previous work has shown that ischemic strokes that involve the insula are larger than strokes that exclude the insula and therefore are associated with more common and persistent deficits (Kodumuri et al., 2016). Furthermore, the left insula has been associated with action naming in the past (e.g., Benetello et al., 2016; Hillis et al., 2006; Kemmerer et al., 2012; Race et al., 2013; Tranel et al., 2001).

Note that we did also identify common regions that were critical for both object and action naming, as damage was predictive of poor performance on both. For participants with PPA, these regions consisted of left superior temporal pole, right middle temporal gyrus, and left fusiform gyrus, as well as ratio of cerebral to intracranial volume. For acute stroke, greater damage to the left angular gyrus was related to poor performance on both. It makes sense that common regions would exist as many of the cognitive processes underlying naming of the two word classes are shared, and these regions have previously been identified in studies of naming (e.g., Benetello et al., 2016; Damasio & Tranel, 1993; Daniele et al., 1994; DeLeon et al., 2007; Faria et al., 2014; Faroqi-Shah et al., 2014; Foundas et al., 1998; Fridriksson et al., 2018; Gleichgerrcht et al., 2015, 2016; Gorno-Tempini et al., 2004, 2011; Halai et al., 2017; Hillis, Tuffiash, Wityk, et al., 2002; Hurley et al., 2012; Kemmerer et al., 2012; Lambon Ralph et al., 2016; Mesulam et al., 2013; Meyer et al., 2017; Migliaccio et

al., 2016; Mion et al., 2010; Newhart et al., 2007; Piras & Marangolo, 2007; Race et al., 2013; Raymer et al., 1997; Riello et al., 2018; Schwartz et al., 2009; Snowden et al., 2018; Thye & Mirman, 2018; Tranel et al., 2001, 2003, 2008; Tsapkini et al., 2011; Tyler et al., 2001; Walker et al., 2011).

Third, different regions were identified for participants with different aetiologies of brain damage. No single region was associated with both object and action naming in both PPA and acute stroke. Rather, all regions identified in the analyses were associated with naming only for participants with PPA or only for participants with acute stroke. This finding underscores the importance of considering converging evidence. For instance, many of the regions that we identified in the PPA participants were in the right hemisphere. It would not be possible to identify such regions in the typical participant with aphasia due to left hemisphere stroke since these regions are not infarcted. Similarly, some regions are less likely to be damaged in stroke than in PPA based on the blood supply of the brain. For example, the temporal poles are rarely damaged in stroke unless the infarct is quite large, while they are the predominant site of atrophy in semantic variant PPA. Examining the neural bases of cognition in only one of these populations may lead to missing important regions; comparing the two provides a more complete picture. One reason that there may be distinct regions associated with naming in the two aetiologies is that there are different distributions of underlying cognitive deficits in the two groups. For example, more patients with PPA are likely to have impaired naming due to impairments of conceptual semantics.

This study has several limitations. First, using a picture naming task as an assessment of action naming is complicated by the fact that depicting most actions necessarily involves depicting objects (e.g., the picture of “vacuum” includes a vacuum cleaner and a person using it). If a participant is impaired on object processing, they may also appear to be impaired on action processing because of this confound. Additionally, actions take place over time, which is difficult to depict in a static two-dimensional picture. However, we were able to find a double dissociation between the object and action naming tasks, indicating that, while this limitation may have weakened effects, it did not completely mask them. Future work may benefit from development of assessments where video clips of actions (and objects) are used as stimuli for naming instead of static pictures.

Next, it was not possible to directly compare the participants with PPA and acute stroke using exactly the same neuroimaging measures and thus we could not directly compare the two groups within the same statistical model. This is because it is impossible to determine the precise percentage of a region that is atrophied in PPA without having an earlier, premorbid scan of the same individual to compare with the later damaged scan. Since we did not have access to such data for all participants, we used a neuroimaging measure that it was possible to observe: volume of the region as a percentage of cerebral volume. It is more straightforward to measure the amount of damage in acute stroke as it is possible to visualise and trace the lesioned tissue on diffusion-weighted scans. Unlike the case of chronic stroke, there is no distortion of the brain shape due to encephalomalacia in acute stroke. In the future, studies could improve the assessment of the relationship between damage and behavior in PPA by relating longitudinal changes in brain volume over time from multiple scans to longitudinal changes in behavior over the same period (as in Faria,

Sebastian, Newhart, Mori, & Hillis, 2014). Longitudinal tracking of brain and behavior may be of particular interest in PPA because it is a chronic condition that may include changes reflecting some plasticity and reorganization as individuals attempt to adapt to neurodegeneration.

Another important limitation is the sample size that was included in the neuroimaging analysis. Although behavioral data were available for a relatively sizable sample of participants with PPA, contemporaneous neuroimaging data were not available for many of the participants. Similarly, the sample size was relatively small for the acute stroke participants. It is possible that the results were overfitted by the LASSO model. However, we preferred this statistical approach to simpler regression models because it addressed the multicollinearity inherently present in the data (e.g., participants with damage in one anatomical region are likely to have damage in nearby regions as well due to biological processes). While the LASSO model reduced multicollinearity by eliminating highly correlated variables, we do note that some information may have been lost in the process if more than one of the highly correlated regions makes an important contribution to naming. Therefore, we cannot make any claims about the regions we did not identify in the analyses: it is possible that they were not identified due to the type of model used or because there was inadequate power in terms of number of patients with atrophy or lesions in those areas. For example, typically only individuals with nfvPPA have frontal atrophy, whereas individuals with both svPPA and lvPPA have atrophy in temporal areas (e.g., Gorno-Tempini et al., 2004). However, the finding that distinct lesion-deficit associations were identified for objects vs. actions within each aetiology cannot be due to this power issue, because data from the same patients were used in detecting object and action naming deficits. It is likely that naming of objects and actions depends on partially overlapping neural networks; here we identified a subset of the critical regions for each and for both. We may have also missed key brain areas because we did not attempt to identify the white matter tracts critical for naming objects or actions. We plan to continue collecting data on these tasks, and data from larger samples in the future may be able to address these concerns.

Furthermore, the results show the importance of comparing individuals with aphasia due to different underlying causes of brain damage as complementary information can be obtained. However, this was limited to only two aetiologies: PPA and acute stroke. Expanding this work to other aetiologies such as tumor, focal traumatic brain injury, and infection (e.g., herpes simplex virus encephalitis), or to temporary lesions created with repetitive transcranial magnetic stimulation in future work is likely to provide further information to enrich understanding of the brain-behavior relationships critical for naming. Lesion-deficit association in each of these populations has challenges and limitations.

Conclusions

The present study introduced a new assessment, the Hopkins Action Naming Assessment, that can be used in conjunction with the Boston Naming Test not only to better diagnose naming impairments but also to investigate the neural bases of naming. Comparing the association between naming performance and damage to neural regions of interest in both PPA and stroke provides a more comprehensive picture of the brain-behavior relationships

critical for naming than could be obtained by studying either in isolation. Complementary information can be obtained by including analyses of both PPA and acute stroke, likely because different regions tend to be damaged in these populations. Likewise, as double dissociations between object and action naming performance were observed in both populations, identifying lesion-deficit associations for both word classes yields a more comprehensive picture of regions critical for naming.

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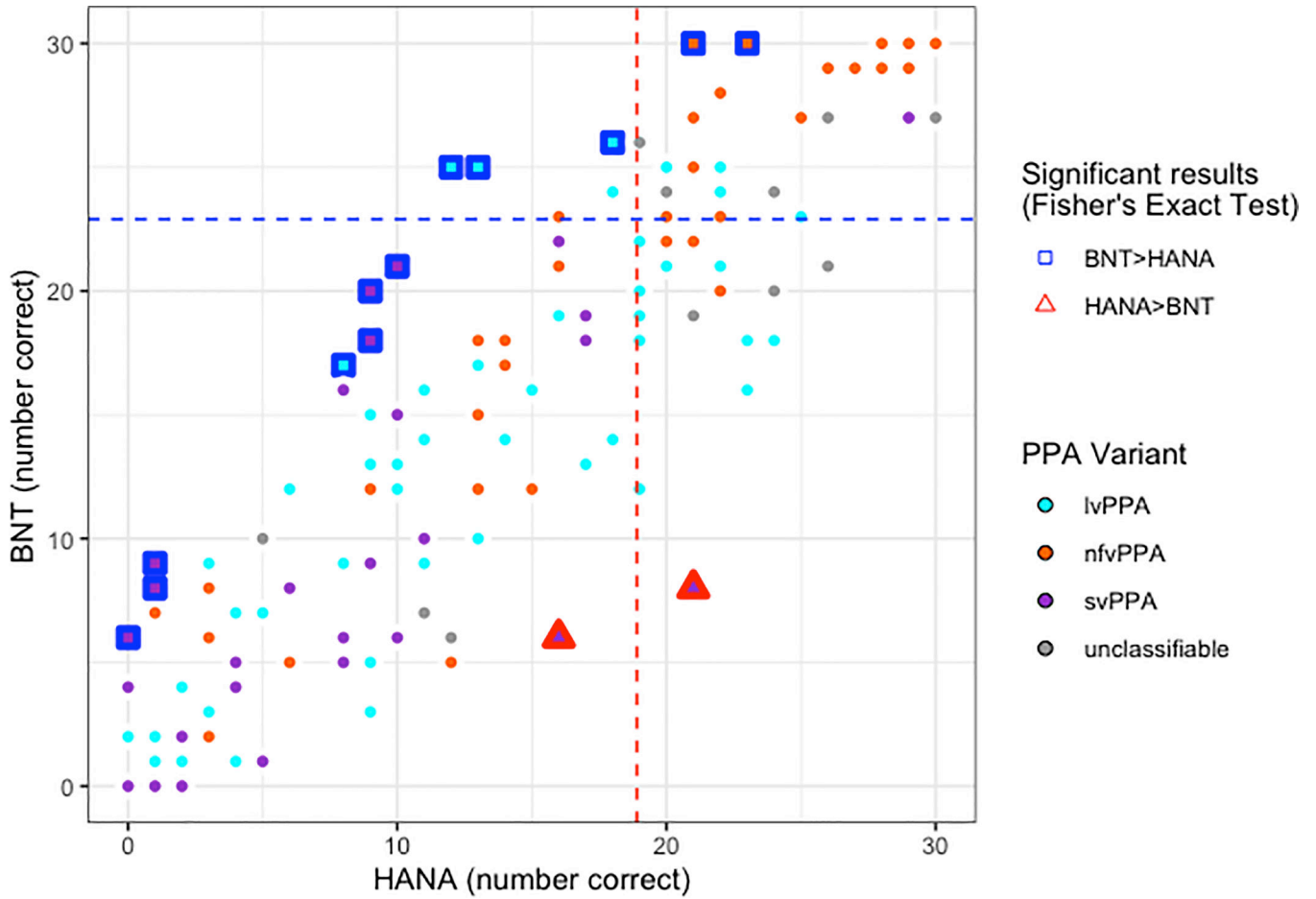


Figure 1.

Behavioural results for participants with PPA (N=135) who completed both the HANA and BNT assessments. Performance on the BNT is shown on the y-axis (maximum score=30) and performance on the HANA is shown on the x-axis (maximum score=30). Dot fill colours represent different PPA variants (cyan=lvPPA; orange=nfvPPA; purple=svPPA; gray=unclassifiable). Dashed lines represent cut-offs for normal performance at $z > -1.645$ relative to controls, blue for the BNT and red for the HANA. Participants who showed intact performance on the BNT but impaired performance on the HANA relative to controls appear in the upper left; those who showed the opposite pattern are shown in the lower right. Participants who performed significantly better on the BNT than the HANA according to Fisher's Exact Test are highlighted with blue squares. Participants who performed significantly better on the HANA than the BNT according to Fisher's Exact Test are highlighted with red triangles.

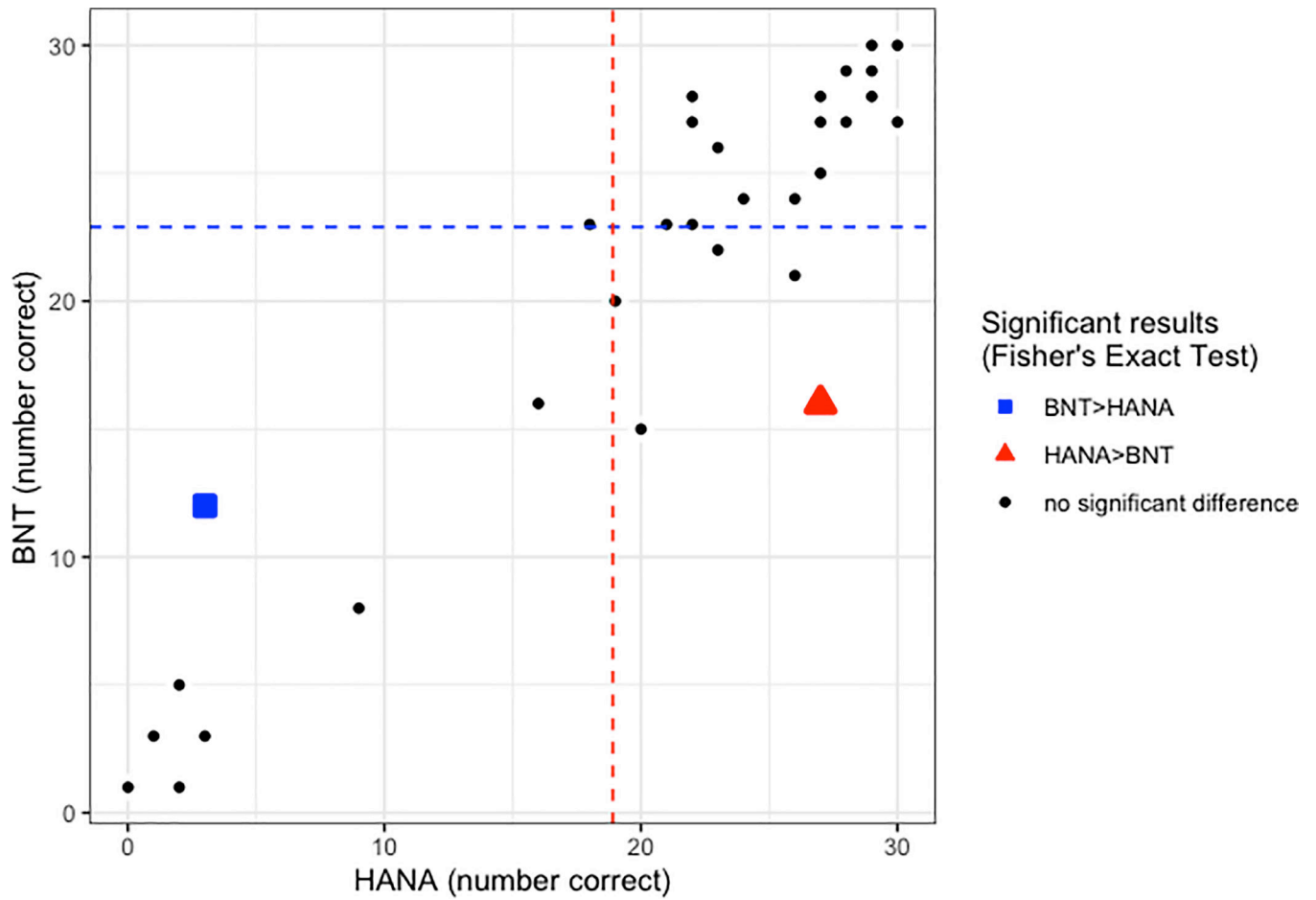


Figure 2.

Behavioural results for participants with stroke (N=34) who completed both the HANA and BNT assessments. Performance on the BNT is shown on the y-axis (maximum score=30) and performance on the HANA is shown on the x-axis (maximum score=30). Dashed lines represent cut-offs for normal performance at $z > -1.645$ relative to controls, blue for the BNT and red for the HANA. Participants who showed intact performance on the BNT but impaired performance on the HANA relative to controls appear in the upper left; those who showed the opposite pattern are shown in the lower right. The participant who performed significantly better on the BNT than the HANA according to Fisher's Exact Test is highlighted with a blue square. The participant who performed significantly better on the HANA than the BNT according to Fisher's Exact Test is highlighted with a red triangle.

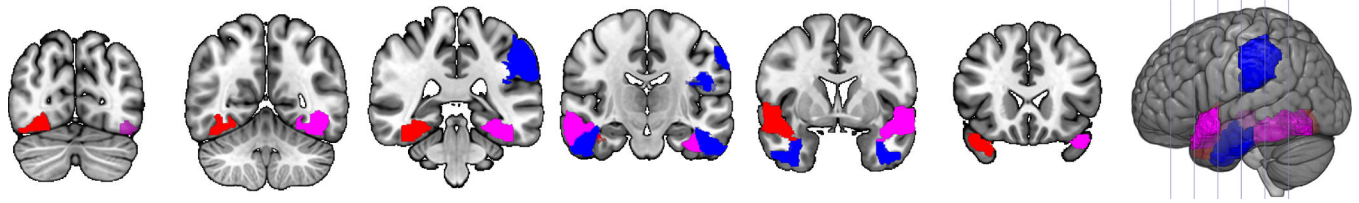


Figure 3. Neuroimaging results for PPA. Regions in red were associated with action naming performance. Regions in blue were associated with object naming performance. Regions in magenta were associated with naming of both objects and actions. Slices are $y=-73$, -55 , -36 , -17 , 2 , and 21 .

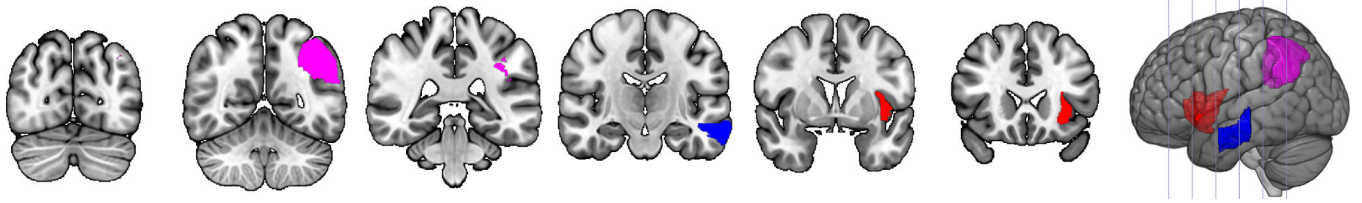


Figure 4. Neuroimaging results for acute stroke. Regions in red were associated with action naming performance. Regions in blue were associated with object naming performance. Regions in magenta were associated with naming of both objects and actions. Slices are $y=-73$, -55 , -36 , -17 , 2 , and 21 .

Table 1.

Length and frequency of the Boston Naming Test and Hopkins Action Naming Assessment items.

item	CELEX English linguistic database	Frequency			Length		
		Ku era & Francis	Sydney Morning Herald	Thorndike & Lorge	letters	phonemes	
<i>BNT items</i>							
bed	244.47	135	59.43	1236	3	3	
pencil	15.75	36	5.38	186	6	5	
whistle	9.66	4	8.87	211	7	4	
comb	5.70	6	1.37	96	4	3	
saw	387.88	353	159.85	2443	3	2	
helicopter	10.56	2	15.44	0	10	10	
octopus	1.45	1	2.43	9	7	7	
hanger	0.78	0	1.15	16	6	5	
camel	8.16	1	4.27	24	5	4	
pretzel	0.34	0	0	0	7	6	
racquet	0.78	1	3.11	0	7	5	
volcano	3.46	2	3.33	22	7	7	
dart	4.30	0	2.43	68	4	3	
globe	10.28	15	12.16	43	5	4	
beaver	2.07	3	1.32	28	6	5	
rhinoceros	1.56	3	0.94	4	10	9	
igloo	0.45	0	0.47	0	5	4	
dominoes	0.56	0	0.21	0	8	7	
escalator	1.34	0	1.49	0	9	9	
hammock	0.84	5	0.51	39	7	5	
pelican	0.78	0	3.41	5	7	7	
pyramid	3.97	2	3.92	17	7	7	
unicorn	0.67	0	0.77	0	7	7	
accordion	0.84	1	1.37	10	9	7	
asparagus	2.07	1	3.58	53	9	9	
latch	2.68	5	0.38	15	5	3	
scroll	3.13	0	1.45	14	6	5	
sphinx	0.84	1	0.64	4	6	6	
trellis	0.95	0	0.73	19	7	6	
protractor	0	0	0	0	10	10	
<i>mean</i>	24.211	19.2	10.014	152.1	6.6	5.8	
<i>SD</i>	81.6581	67.82	30.3284	487.95	1.92	2.19	
<i>range</i>	0–387.88	0–353	0–159.85	0–2443	3–10	2–10	
<i>HANA items</i>							
run	229.89	246	385.83	1643	3	3	

item	Frequency				Length	
	CELEX English linguistic database	Ku era & Francis	Sydney Morning Herald	Thorndike & Lorge	letters	phonemes
vacuum	15.14	22	9.09	48	6	6
suck	10.00	5	3.03	49	4	3
shave	5.87	6	3.11	62	5	3
point	403.35	402	400.67	1377	5	4
curtsy	0.78	0	0.30	0	6	5
crawl	7.82	11	3.28	168	5	4
hibernate	0.39	2	0.13	5	9	7
shear	0.73	40	0.68	38	5	3
underline	2.85	2	2.22	4	9	7
spill	4.19	1	11.39	74	5	4
peel	10.56	3	6.78	91	4	3
erase	2.12	1	1.41	24	5	4
sharpen	1.96	1	1.75	32	7	5
subtract	2.46	2	0.60	6	8	8
quack	0.56	9	0.73	10	5	4
juggle	0.84	0	2.60	10	6	4
extinguish	0.78	1	1.19	16	10	10
navigate	0.67	1	1.54	13	8	7
sled	0.78	0	1.07	26	4	4
carve	2.18	3	4.05	82	5	3
yawn	2.79	2	2.22	67	4	3
knit	2.51	11	4.31	117	4	3
dribble	1.34	0	0.81	8	7	5
knead	1.17	1	0.21	10	5	3
hypnotize	0	0	0	17	9	8
shred	1.96	3	1.28	79	5	4
prescribe	3.58	5	2.13	42	9	8
whisper	14.92	12	4.56	673	7	6
play	276.37	205	413.04	2606	4	3
<i>mean</i>	33.619	33.2	42.334	246.6	5.9	4.8
<i>SD</i>	94.4779	89.77	121.2887	590.27	1.91	1.97
<i>range</i>	0–403.35	0–402	0–413.04	0–2606	3–10	3–10
Welch two sample t-test results						
t	–0.41	–0.68	–1.42	–0.68	1.42	1.86
df	56.81	53.97	32.61	56.02	58.00	57.39
p-value	.681	.498	.166	.502	.162	.068

Notes: BNT= Boston Naming Test; HANA= Hopkins Action Naming Assessment; SD = standard deviation; df= degrees of freedom. Refer to the text for the normative sources of the frequency measures.

Table 2.

Demographic and naming assessment data for participants with PPA

participant	sex	age	education (years)	symptom duration (months)	scan-assessment interval (months)	BNT	HANA
<i>Participants with lvPPA</i>							
L01	female	79	16	30	-5	2	0
L02	female	73	18	108	0	3	9
L03	male	72	18	42	5	9	3
L04	female	76	18	12	1	10	13
L05	male	68	13	69	-2	16	11
L06	female	68	18	36	4	16	15
L07	male	76	12	31	0	17	8 *
L08	female	66	18	66	1	21	20
L09	male	51	12	39	3	23	25
L10	male	55	18	12	1	25	20
L11	male	72	20	24	0	25	22
L12	female	73	18	91		1	1
L13	female	72	16	48		1	2
L14	female	65	12	78		1	2
L15	female	73	12	68		1	4
L16	female	79	18	24		2	1
L17	male	85	18	78		3	3
L18	male	78	16	96		4	2
L19	female	72	14	60		5	9
L20	female	68	18	42		7	4
L21	female	72	18	36		7	5
L22	female	77	12	24		9	8
L23	male	75	18	10		9	11
L24	female	69	18	48		12	6
L25	male	71	20	42		12	10
L26	female	65	14	84		12	19
L27	male	69	19	60		13	9
L28	female	70	16	24		13	10
L29	male	75	16	30		13	10
L30	female	78	18	36		13	17
L31	female	54	15	36		14	11
L32	female	71	16	12		14	14
L33	female	68	12	24		14	18
L34	female	74	18	57		15	9
L35	female	69	12	33		16	23
L36	male	70	18	57		17	13
L37	male	52	12	12		17	13

participant	sex	age	education (years)	symptom duration (months)	scan-assessment interval (months)	BNT	HANA	
L38	male	64	18	24		18	19	
L39	male	73	20	12		18	23	
L40	female	68	18	36		18	24	
L41	female	70	12	15		19	16	
L42	female	90	16	24		19	19	
L43	male	75		18		20	19	
L44	male	80	12	6		21	22	
L45	female	70	14	48		22	19	
L46	female	69	16	36		24	18	
L47	female	65	18	35		24	22	
L48	male	60	13	35		25	12 *	
L49	male	73	20	77		25	13 *	
L50	female	56	16	72		26	18 *	
<hr/>								
	<i>mean</i>	70.3	16.1	42.3		13.8	12.5	
	<i>SD</i>	7.70	2.66	24.97		7.61	7.18	
	<i>range</i>	51–90	12–20	6–108		1–26	0–25	
<hr/>								
<i>Participants with <i>nfvPPA</i></i>								
N01	male	77	20	18		4	2	3
N02	female	79	14	35		0	7	1
N03	male	72	14	52		-4	8	3
N04	male	77	8	6		4	13	17
N05	female	65	16	48		4	15	13
N06	female	52		45		0	16	8
N07	male	85	12	24		3	17	13
N08	male	77	18	58		1	27	25
N09	male	65	18	24		2	29	29
N10	male	62	20	12		2	30	23 *
N11	male	73	16	22		0	30	29
N12	male	58	18	56		-2	30	30
N13	female	85	12	22		5	6	
N14	female	78	16	60		5	12	
N15	female	65	16	91		6	3	
N16	male	69	18	96		12	9	
N17	female	65	12	30		12	13	
N18	female	62	16	6		12	15	
N19	female	78	3	24		17	14	
N20	male	48	12	81		18	13	
N21	male	52	18	60		18	14	
N22	male	66	20	18		20	22	
N23	male	64	18	18		21	16	

participant	sex	age	education (years)	symptom duration (months)	scan-assessment interval (months)	BNT	HANA	
N24	female	67	16	31		22	20	
N25	male	76	18	42		22	21	
N26	female	74	12	30		23	16	
N27	male	81	16	36		23	20	
N28	female	71	18	59		23	22	
N29	male	73	18	48		25	20	
N30	male	61	16	54		25	21	
N31	male	74	12	24		25	21	
N32	male	65	18	36		25	21	
N33	female	82	12	54		27	21	
N34	female	77	18	97		28	22	
N35	female	67	16	84		29	26	
N36	male	63	16			29	27	
N37	male	67	19	46		29	28	
N38	male	70	19	10		29	28	
N39	male	52	18	12		30	21 *	
N40	male	69	15	28		30	28	
N41	male	59	14	12			28	
<i>mean</i>		68.8	15.7	40.2		20.4	18.1	
<i>SD</i>		9.14	3.46	24.93		8.46	8.05	
<i>range</i>		48–85	3–20	6–97		2–30	1–30	
Participants with svPPA								
S01	female	60	18	29		1	4	0
S02	female	68	16	68		0	6	0 *
S03	male	71	20	42		2	9	1 *
S04	female	77	12	36		0	9	9
S05	male	59	13	45		2	16	8
S06	female	62	18	48		4	27	29
S07	female	59	18	143		0		1
S08	female	59	12	60			0	0
S09	female	74	16	129			0	1
S10	male	58	16	97			0	2
S11	male	64	16	45			1	5
S12	male	64	16	120			2	2
S13	female	67	16	36			2	3
S14	male	65	13	24			4	2
S15	female	72	16	72			4	4
S16	female	52		12			5	4
S17	female	76	18	12			5	8
S18	female	68	16	68			6	3

participant	sex	age	education (years)	symptom duration (months)	scan-assessment interval (months)	BNT	HANA
S19	female	66	16	48		6	8
S20	male	74	18	30		6	10
S21	male	61	16	40		6	16 *
S22	female	61	18	31		8	1 *
S23	male	79	18	48		8	6
S24	female	68	16	45		8	21 *
S25	female	74	16	24		9	9
S26	male	55	18	36		10	11
S27	female	61	18	60		15	10
S28	male	79	12	24		18	9 *
S29	female	61	14	36		18	17
S30	female	73	20	12		19	17
S31	male	61	18	18		20	9 *
S32	female	62	12	48		21	10 *
S33	male	50	12	22		22	16
<i>mean</i>		65.5	16.0	48.7		9.2	7.6
<i>SD</i>		7.60	2.37	32.45		7.33	6.87
<i>range</i>		50–79	12–20	12–143		0–27	0–29

Participants with unclassifiable PPA

U01	female	62	16	41	0	27	30
U02	female	63	16	24		6	12
U03	female	69	18	30		7	11
U04	male	68	19	20		10	5
U05	female	55	16	48		18	14
U06	male	74	16	12		18	19
U07	female	68	12	72		19	21
U08	male	74	20	22		20	24
U09	male	80	16	12		21	26
U10	female	74		48		24	20
U11	male	68	18	36		24	24
U12	male	72	18	6		26	19
U13	female	78	12	12		27	26
U14	female	81	12	36			19
<i>mean</i>		70.4	16.1	29.9		19.0	19.3
<i>SD</i>		7.26	2.66	18.30		7.23	6.83
<i>range</i>		55–81	12–20	6–72		6–27	5–30

All participants with PPA

<i>mean</i>	68.7	15.9	42.0	15.2	13.7
<i>SD</i>	8.24	2.84	26.64	8.82	8.44

participant	sex	age	education (years)	symptom duration (months)	scan-assessment interval (months)	BNT	HANA
<i>range</i>		48–90	3–20	6–143		0–30	0–30

Notes: Bolded lines represent participants who contributed data to the neuroimaging analysis. SD = standard deviation; scan-assessment interval (months)= number of months between scan and assessment, with negative values indicating scan before assessment and positive values indicating assessment before scan; BNT= number of correct responses on the Boston Naming Test, maximum score of 30; HANA= number of correct responses on the Hopkins Action Naming Assessment, maximum score of 30.

* indicates a significant difference between performance on the BNT and HANA, Fisher's Exact Test $p < .05$.

Summary values are reported for participants with each variant of PPA as well as the whole group that contributed behavioral data; summary values are not reported for the scan-test interval as they apply only to the subset of participants with neuroimaging data.

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Table 3.

Demographic and naming assessment data for participants with acute stroke

participant	sex	age	education (years)	scan-stroke interval (days)	scan-admission interval (days)	scan-assessment interval (days)	BNT	HANA
A01	male	56	14	1	0	-1	1	0
A02	male	43	12	2	1	-3	1	2
A03	male	61	16	8	0	-5	3	1
A04	female	54	18	0	0	-2	3	3
A05	female	28	12	7	5	1	5	2
A06	male	53	12	1	0	-4	8	9
A07	male	65	12	1	0	-1	12	3 *
A08	female	61	9	3	3	1	15	20
A09	male	37	16	0	0	-4	16	16
A10	female	39	12	12	2	1	16	27 *
A11	male	87	20	0	0	-3	20	19
A12	female	65	14	8	1	-1	21	26
A13	male	48	11	1	0	-2	22	23
A14	female	49	12	2	0	-4	23	18
A15	male	52	12	0	0	-1	23	21
A16	female	53	14	7	6	4	23	22
A17	female	60	13	0	0	-2	24	24
A18	male	65	18	2	2	0	24	26
A19	male	49	15	4	0	-1	25	27
A20	male	46	12	4	3	0	26	23
A21	female	77		0	0	0	27	22
A22	female	32	14	5	4	1	27	27
A23	female	56	12	0	0	-2	27	28
A24	male	64	16	0	0	-4	27	30
A25	male	43	13	6	0	-3	28	22
A26	male	61	16	0	0	-2	28	27
A27	male	60	14	1	1	-2	28	29
A28	male	60	13	1	0	0	29	28
A29	male	57	14	1	0	-1	29	28
A30	male	35	14	1	0	-2	29	29
A31	male	70	13	0	0	-4	29	29
A32	male	55	14	0	0	-2	30	29
A33	female	60	14	0	0	-2	30	30
A34	male	60	14	2	2	1	30	30
A35	female	71		2	1	-2		0
A36	female	81	14	0	0	-5		3
A37	male	78	30	0	0	-1		29
<i>mean</i>		56.5	14.3	2.2	0.8	-1.5	20.9	19.8

participant	sex	age	education (years)	scan-stroke interval (days)	scan-admission interval (days)	scan-assessment interval (days)	BNT	HANA	
		<i>SD</i>	13.41	3.48	2.97	1.54	1.95	9.38	10.58
		<i>range</i>	28–87	9–30	0–12	0–6	–5–4	1–30	0–30

Notes: *SD* = standard deviation; scan-stroke interval (days) = number of days after onset of stroke symptoms that scan was obtained; scan-admission interval (days) = number of days after admission to hospital that scan was obtained. (This differs for some participants as compared to scan-stroke interval as participants sought treatment at different points relative to the onset of symptoms); scan-assessment interval (days)= number of days between scan and assessment, with negative values indicating scan before assessment and positive values indicating assessment before scan; BNT= number of correct responses on the Boston Naming Test, maximum score of 30; HANA= number of correct responses on the Hopkins Action Naming Assessment, maximum score of 30.

* indicates a significant difference between performance on the BNT and HANA, Fisher's Exact Test $p < .05$.

Table 4.

LASSO results—PPA.

	BNT		HANA			
	LASSO value	Adjusted Coefficient	p value	LASSO value	Adjusted Coefficient	p value
model intercept	-6.142×10^{-16}			-7.101×10^{-16}		
cerebral to intracranial volume ratio	0.303	0.327	0.096	0.416	0.515	0.091
left inferior frontal gyrus-pars opercularis						
left inferior frontal gyrus-pars orbitalis						
left inferior frontal gyrus-pars triangularis						
left postcentral gyrus						
left precentral gyrus						
left supramarginal gyrus	0.186	0.276	0.252			
left angular gyrus						
left superior temporal gyrus						
right superior temporal gyrus						
left superior temporal pole	0.212	0.292	0.1	0.092	0.155	0.575
right superior temporal pole				0.119	0.101	0.462
left middle temporal gyrus						
right middle temporal gyrus	0.072	0.161	0.494	0.030	0.161	0.567
left middle temporal pole						
right middle temporal pole						
left inferior temporal gyrus	0.020	-0.042	0.893			
right inferior temporal gyrus	0.054	0.064	0.462			
left fusiform gyrus	0.271	0.285	0.081	0.198	0.28	0.243
right fusiform gyrus				0.030	-0.022	0.716
left insula						

Table 5.

Lasso results—acute stroke.

	BNT		HANA	
	LASSO value	Adjusted Coefficient	LASSO value	Adjusted Coefficient
model intercept	1.367×10^{-16}		1.455×10^{-16}	
total lesion volume			-0.225	-0.167
left inferior frontal gyrus-pars opercularis				
left inferior frontal gyrus-pars orbitalis				
left inferior frontal gyrus-pars triangularis				
left postcentral gyrus				
left precentral gyrus				
left supramarginal gyrus				
left angular gyrus	-0.524	-0.555	-0.405	-0.487
left superior temporal gyrus				
left superior temporal pole				
left middle temporal gyrus	-0.121	-0.151	0.397	
left inferior temporal gyrus				
left fusiform gyrus				
left insula			-0.015	-0.134

Note:

* denotes significance at $p < .05$