

Florbetapir processing methods

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Summary

ADNI Florbetapir summary data are updated regularly and uploaded to LONI by our group. We use a native-space MRI scan for each subject that is segmented and parcellated with Freesurfer (version 4.5.0) to define cortical grey matter regions of interest (frontal, anterior/posterior cingulate, lateral parietal, lateral temporal) that make up a summary cortical ROI. We have also defined five reference regions (cerebellar grey matter, whole cerebellum, brainstem/pons, eroded subcortical white matter, and a composite reference region). We then coregister each florbetapir scan to the corresponding MRI and calculate the mean florbetapir uptake within the cortical and reference regions.

Florbetapir SUVRs can be created by averaging across the 4 cortical regions and dividing this cortical summary ROI by one of the five reference regions. Selection of a cortical region of interest and reference region depend on the goals of the analysis. We have provided two summary SUVRs in our dataset. The first is the cortical summary ROI divided by the whole cerebellum reference region (SUMMARYSUVR_WHOLECEREBNORM), which our group [1, 2] and others have used for cross-sectional florbetapir analyses. The second is the cortical summary ROI divided by a composite reference region

(SUMMARYSUVR_COMPOSITE_REFNORM), which we have evaluated recently for longitudinal florbetapir analyses and is described below.

Method

Acquisition of florbetapir and MRI image data from LONI

We download florbetapir data from LONI in the most fully pre-processed format (series description in LONI Advanced Search: "AV45 Coreg, Avg, Std Img and Vox Siz, Uniform Resolution"). Each subject's first florbetapir image is coregistered using SPM5 to that subject's MRI image (series description: ADNI 1 scans *N3;* and ADNI GO/2 scans *N3*) that was closest in time to the florbetapir scan. We use the MRI scan that is closest in time to the first florbetapir scan, but note that not all subjects have a concurrent MRI available on LONI so in some cases we use an MRI scan acquired at another visit. The baseline MRI is used to define regions for all subsequent florbetapir scans.



Figure 1: All cortical grey matter-only regions that can be used to create a composite florbetapir index (SUVR) are shown in red on an example subject's MRI. Main regions are frontal, anterior/posterior cingulate, lateral parietal, and lateral temporal cortex. Subregions within each of these main regions are listed to the left.

Freesurfer-defined cortical regions

We carry out Freesurfer processing to skull-strip, segment, and delineate cortical and subcortical regions in all MRI scans. We then extract florbetapir means from grey matter in





each subregion within 4 large regions (frontal, anterior/posterior cingulate, lateral parietal, lateral temporal – shown in Figure 1, subregions listed below). This process has been described in recent publications from our lab [3, 4]. Subregion and region means and subregion sizes are available in the LONI dataset and are NOT intensity normalized. Available subregions are listed here according to Freesurfer naming convention:

Frontal regions:

ctx-lh-caudalmiddlefrontal ctx-lh-lateralorbitofrontal ctx-lh-medialorbitofrontal ctx-lh-parsopercularis ctx-lh-parsorbitalis ctx-lh-parstriangularis ctx-lh-rostralmiddlefrontal ctx-lh-superiorfrontal ctx-lh-frontalpole ctx-rh-caudalmiddlefrontal ctx-rh-lateralorbitofrontal ctx-rh-medialorbitofrontal ctx-rh-parsopercularis ctx-rh-parsorbitalis ctx-rh-parstriangularis ctx-rh-rostralmiddlefrontal ctx-rh-superiorfrontal ctx-rh-frontalpole

Anterior/posterior cingulate regions:

ctx-lh-caudalanteriorcingulate ctx-lh-isthmuscingulate ctx-lh-posteriorcingulate ctx-lh-rostralanteriorcingulate ctx-rh-caudalanteriorcingulate ctx-rh-isthmuscingulate ctx-rh-posteriorcingulate ctx-rh-rostralanteriorcingulate

Lateral parietal regions:

ctx-lh-inferiorparietal ctx-lh-precuneus ctx-lh-superiorparietal ctx-lh-supramarginal ctx-rh-inferiorparietal ctx-rh-precuneus ctx-rh-superiorparietal ctx-rh-supramarginal

Lateral temporal regions:

ctx-lh-middletemporal ctx-lh-superiortemporal

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ctx-rh-middletemporal ctx-rh-superiortemporal

Calculation of florbetapir cortical summary values

To account for the varying sizes of the subregions listed above, we created weighted means for each of the 4 main cortical regions (frontal, anterior/posterior cingulate, lateral parietal, lateral temporal) using the florbetapir mean and volume of each subregion as follows: (subregion1_florbetapir_mean X subregion1_volume) + (subregion2_florbetapir_mean X subregion2_volume) + (subregionN_florbetapir_mean X subregionN_volume) / (subregion1_volume + subregion2_volume +... subregionN_volume)

Freesurfer-defined reference regions

We calculate means for five Freesurfer-defined reference regions (cerebellar grey matter, whole cerebellum, brainstem/pons, eroded subcortical white matter, and a composite reference region) that can be used to intensity normalize the cortical summary ROI or individual cortical regions. The composite reference region is a conventional (nonweighted) average of whole cerebellum, brainstem/pons, and subcortical WM regions, as proposed recently by Koeppe [5]. The only modification we have made to the Freesurfer delineation of these reference regions is that we eroded the subcortical white matter region away from cerebrospinal fluid and cortical grey matter in order to reduce spillover from signal in these regions into white matter. To do this, we smoothed a binarized a Freesurfer-defined white matter region mask to the same resolution as the PET data (effective 8X8X8mm^3) and then thresholded it at 0.70, resulting in an eroded subcortical WM region made up of voxels containing at least 70% white matter.



Figure 2: Reference regions included in our dataset are shown. The WM is eroded away from cortex to avoid partial volume effects. The composite reference region is an average of the whole cerebellum, brainstem/pons, and eroded WM regions. See Landau et al. JNM 2015.

Calculation of florbetapir SUVR

A florbetapir cortical summary measurement (SUVR) can be calculated by creating a conventional (nonweighted) average across the 4 main cortical regions (Figure 1) and dividing by one of the five reference regions (Figure 2).

Calculation of florbetapir cutoff for cross-sectional analyses

A cutoff for establishing amyloid positivity or negativity is specific to both the radiotracer and the image processing methods used[6, 7]. Even for a specific tracer and processing method, selection of an appropriate cutoff depends on the goals of the study or analysis.

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For this dataset, our current recommendation for cross-sectional florbetapir analyses is a **florbetapir cutoff of 1.11 using the whole cerebellum reference region**, which is equivalent to the upper 95% confidence interval above the mean of a group of young normal controls[8]. In addition, recent work by Clark and colleagues[9] showed that no patients with probable Alzheiemer's disease at autopsy had a florbetapir SUVR of greater than 1.10, based on Avid's template-based processing method. To determine the relationship between Avid-processed SUVRs and Freesurfer-processed SUVRs (both using a whole cerebellum reference region) for 325 ADNI florbetapir scans, we plotted SUVRs in the same group of ADNI subjects (Figure 3). We used the linear regression equation that resulted from this correlation to convert the Avid cutoff of 1.10 to an almost identical value of 1.11 in Freesurfer "units". The cutoff of 1.11 applies to the SUVR normalized by the whole cerebellum reference region ONLY and is NOT applicable to the individual subregions or the SUVR normalized by the composite reference region.

ADNI subjects can be categorized as amyloid positive or negative by applying this cutoff to the florbetapir composite SUVR value described above.



Comparison of 2 florbetapir processing methods: Avid's template-based processing and UC Berkeley's Freesurfer (native space) processing

Figure 3: Plot showing cortical AV45 uptake for 325 ADNI subjects analyzed using Avid's template-based process method and the Freesurfer processing stream outlined here. We used the regression equation for this associatior convert Avid's autopsy-validated cutoff of 1.10 from Clark et al. (JAMA 2011) to Freesurfer "units", resulting in a Freesurfer cutoff of 1.11.



Recommendations for longitudinal florbetapir analyses

Recent work in our laboratory and others[10, 11] has shown that reference regions containing subcortical eroded WM result in longitudinal florbetapir measurements that appeared to be more accurate[12]. Specifically, reference regions containing WM result in longitudinal florbetapir measurements that are less likely to decrease in a group of subjects who are expected to increase (see Figure 4 below and Landau et al. JNM 2015). Therefore our current recommendation is to use eroded WM or a <u>composite reference region, made up of whole cerebellum, brainstem/pons, and eroded subcortical white matter for longitudinal florbetapir analyses</u>. Transforming the whole cerebellum cutoff of 1.11 into composite reference region units using linear regression results in a <u>cutoff of 0.79</u>. We have provided amyloid positivity categorizations by the whole cerebellum reference cutoff and the composite reference cutoff in SUMMARYSUVR_WHOLECEREBNORM_1.11CUTOFF and SUMMARYSUVR_COMPOSITE_REFNORM_0.79CUTOFF, respectively.

Note that the Freesurfer analysis that defines regions used for florbetapir SUVRs is carried out on the baseline MRI scan only. (When an MRI scan was not available at the same timepoint as the florbetapir scan, we used the MRI that was closest in time to the florbetapir scan.)



Figure 4: Cortical SUVR Visit 1 and Visit 2 measurements using a number of candidate reference regions in a group of normal and EMCI subjects who were expected to increase over time (based on their CSF Aβ positive status at Visit 1). Use of reference regions containing eroded WM (bottom row) result in cortical SUVR change measurements that were less likely to decline in this group. See Landau et al. 2015 for more detailed information.



References

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Rev June 25 2015