Quality assessment of spatial transformation of locus coeruleus (LC) functional and structural magnetic resonance imaging (MRI) data

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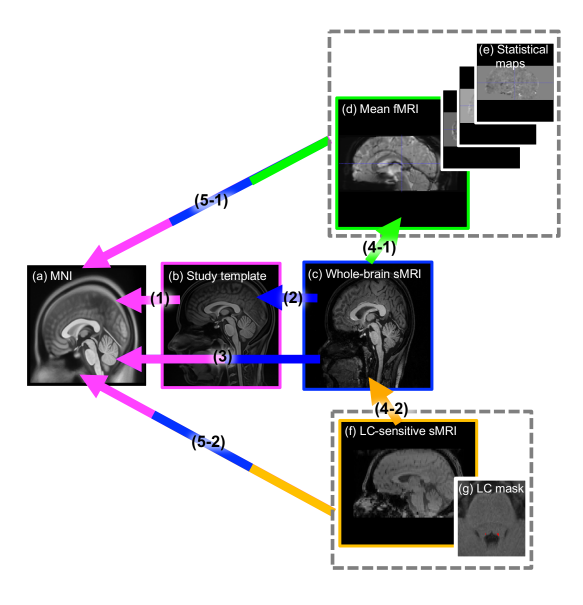
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1. Introduction

a. Summary of spatial transformation procedure



Previous to the landmark-based spatial transformation quality assessment, a series of transformations were executed on the mean functional images (d), whole-brain structural images (c), neuromelanin-sensitive structural images (f), and LC segmentations (g). As explained in detail in the section 2.5 of the main text, single-subject functional and structural images in native space (c-g) were transformed into MNI space (a). The code for the spatial transformation procedure described above is in the section 3.b of this manual.

b. Overview of landmark assessment procedure

After all subject datasets have been processed, the quality of the transformation was examined. This examination was done by selecting the landmarks in the MNI space, drawing them on the individual transformed mean functional images, and calculating Euclidean distances between the MNI-defined landmarks and their single-subject counterparts. Additionally, prior to the landmark assessment, a video that takes single-subject transformed functional or structural images per frame can be made and inspected to quickly identify abnormal registration. More detailed procedures will be explained in the walkthrough section.

c. The background and the goal of this procedure

The LC is so small that even a slight spatial deviation of a voxel size could significantly reduce the statistical power of group-level analyses. For more robust functional assessment of such structure, it is not only important to register and normalise each single subject functional image properly onto the group space, but also to control the quality of the transformed images and rectify any problems that arose from the spatial transformation. With the quality assessment protocol that are introduced in detail below, we aim to achieve more reliable and powerful voxel-wise analyses of the functional LC imaging data than existing approaches in the field.

2. Protocol

a. Preparation

i. Required data

The files that are needed for the assessment are:

- the MNI space to which all images are transformed
- transformed mean functional images of each subject

The MNI space can also be a group space depending on the analysis that will be done with the transformed functional data. In the example dataset, an MNI space created by Fonov et al (2011) was used.

ii. Required software

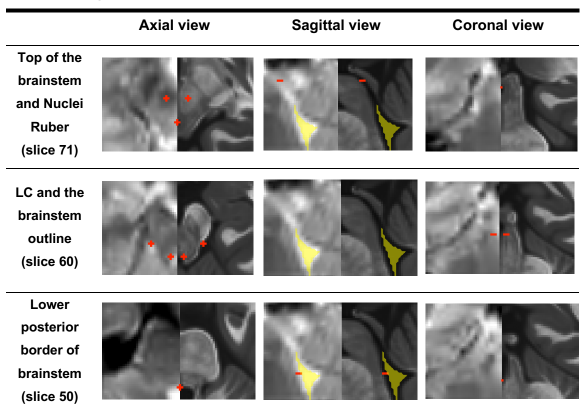
To draw the landmarks, we will use:

- ITK-SNAP (version later than 3.6.0-RC1; http://www.itksnap.org), or any segmentation-enabled brain image viewers
- MATLAB (version later than 2013b, Mathworks, Sherborn, MA, USA), or any computational programmes that can read Nifti images

The example MATLAB codes for generating the overlay video and statistical assessment is in section 4.

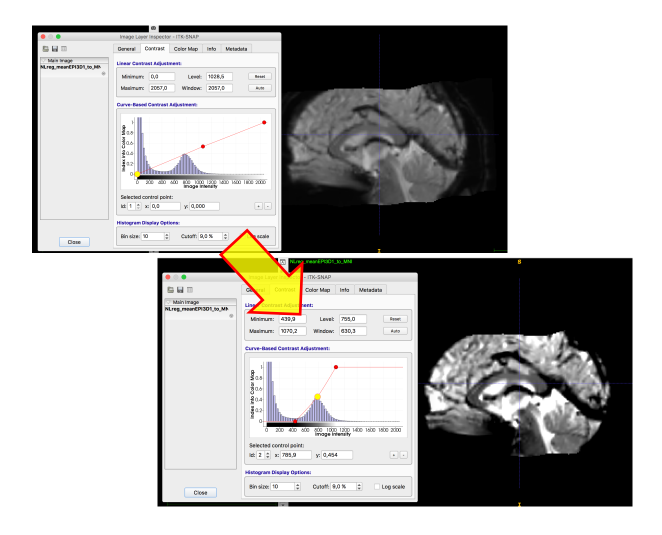
b. Procedure

i. Selecting landmarks



Selecting an appropriate set of landmarks depends on the region of interest (ROI). In this manual's ROI, LC, is located in the brainstem near the lateral floor of the fourth ventricle. Therefore, selecting landmarks in the upper brainstem that delineate the bounds of LC location is most helpful for the evaluation process. In addition, the landmarks should be clearly visible

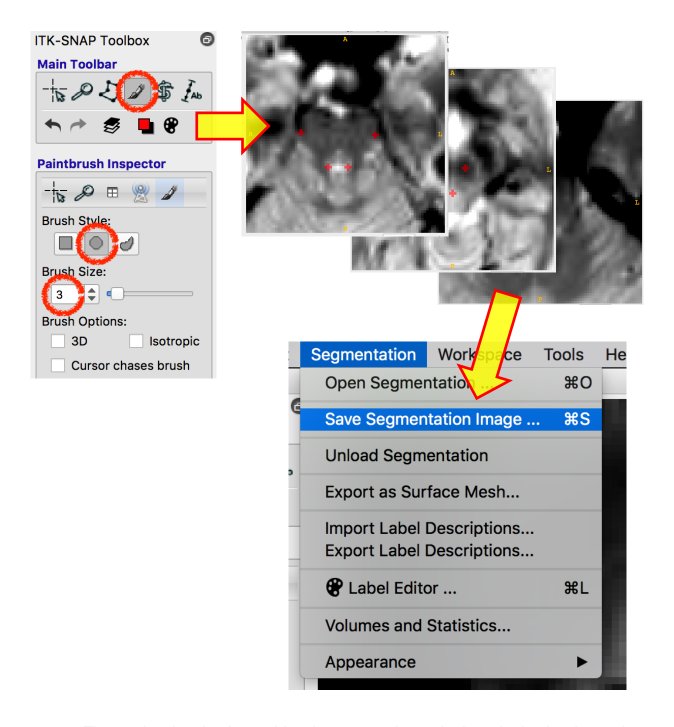
and anatomically distinguishable compared to the surrounding structures on the mean functional images, as these landmarks are drawn manually through visual inspection. Based on these criteria, the landmarks specified in the table above were selected in the main text. The top of the brainstem and nuclei Ruber were selected as a landmark to indicate the upper bound of the larger ROI area of the study, the brainstem. Also, Nucleus Ruber is distinctly visible in the functional images, which makes it an ideal candidate for a landmark. The LC and the outline of the brainstem was selected to delineate the bounds of LC activation. Lastly, the lower posterior border of brainstem was selected to indicate the lowest bound of possible LC location. Once the landmarks are selected, evaluators should familiarise themselves with them.



ii. Drawing landmarks

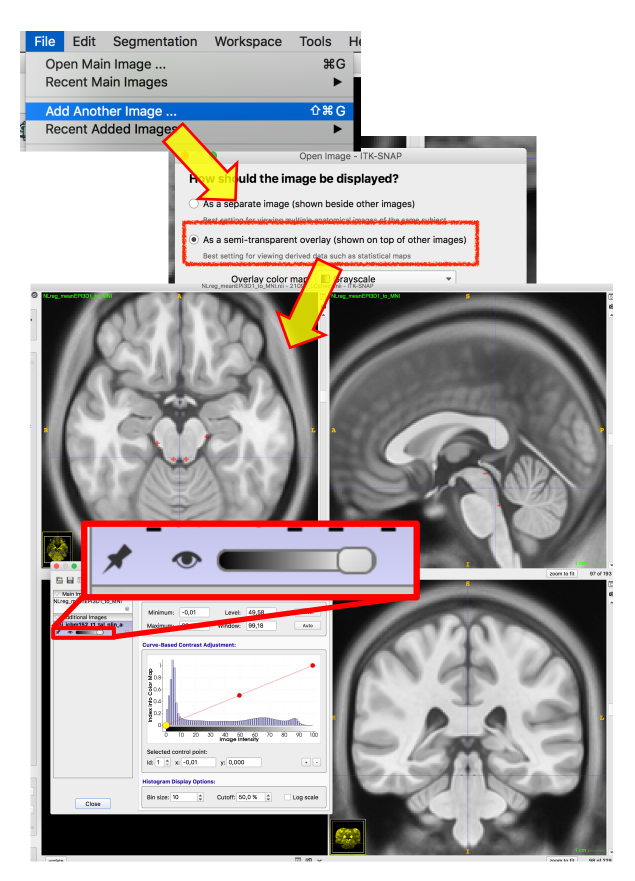
Next, evaluators inspect one of the transformed single-subject mean functional images. If necessary, adjust the contrast slider so that the fourth ventricle is as bright and distinguished as it can be. After adjusting the contrast of the image, magnify the image that you can draw the landmarks more precisely.





Then, using the size-3 round brush segmentation tool, place the landmarks on the transformed mean functional images in axial view. However, use the anatomical clues from other views to aid the landmark placement. For example, when drawing lower posterior border of the brainstem, the deepest groove of the fourth ventricle can be a guiding structure. When the assessment of the single-subject image is complete, save the landmarks as a segmentation image.

Optionally, after saving the image, evaluators could open the group space image used for transforming the mean functional images as a semi-transparent overlay. Once the image is overlaid, adjust the transparency setting to examine whether the landmark segmentations

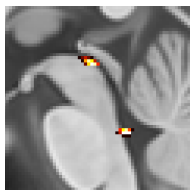


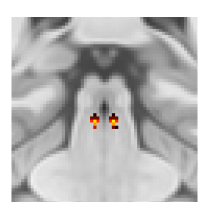
are in the right place. Take note of the subject ID if the landmarks were badly placed. However, DO NOT change the landmark image you already saved, as it indicates the quality of the functional image transformation.

iii. Aggregating single-subject landmarks for preliminary inspection

When the assessment of the whole dataset is complete, visual inspection of all landmarks aggregated in a same space is performed by executing the custom code from section 4 of this manual. By performing this step together with the frame-by-frame video of transformed images (the code to generate the video is in section 3-c, an example of this video is in the section 2.4 of the supplementary material), outliers are easily identified.







iv. Statistical quality assessment of aggregated landmarks of transformed functional images

Once all faulty transformation issues are addressed and rectified, statistical analyses of the quality assessment is performed to quantify the transformation precision. An example code for the analyses is in the section 3.d. Through this step, the spatial transformation can be further refined. The detailed background and example results are described in the main text.

v. Statistical quality assessment of transformed structural images

Additionally, the spatial transformation quality of the structural images can be assessed using the transformed LC segmentations. As specified in the section 2.5 of the main text and the section 1.a of this manual, the LC segmentations are transformed onto the MNI space using the transformation matrices and the deformation fields generated from warping neuromelanin-sensitive structural image (f) to the MNI space (step 5-2). The quantification of the assessment can be done by calculating Euclidean distance between the slice-wise centroids of the template LC mask in the MNI space, e.g. the meta LC mask by Dahl et al.

(2021) in this manual's case, and the transformed LC segmentations of each subject. An example code for the calculating the distance is in the section 3.e.

3. Codes

a. The spatial transformation pipeline (a shell script)

```
#!/bin/bash
# subject ID
ID=1001
# set up folders and group space images
folder=/mnt/work/yyi/temp/ED coreg/"${ID}"/
MNI=/mnt/work/yyi/temp/ED coreg/mni icbm152 t1 tal nlin asym 09c.nii
template=/mnt/work/yyi/temp/ED coreg/pilot template.nii.gz
# LC segmentation
mask=$(ls -t "${folder}"/data/LCmask "${ID}".nii.gz)
# ----- prepare images for trasnformation ----- #
# bias field correct T1 and EPI
N4BiasFieldCorrection -d 3 -v 1 -r 0 -i "${folder}"data/T1mean.nii -o
"${folder}"data/T1mean corrected.nii -s 2 -c [200x150x100x50,1e-6] -b 200
N4BiasFieldCorrection -d 3 -v 1 -r 0 -i "${folder}"data/meanEPI.nii -o
"${folder}"data/meanEPI corrected.nii -s 2 -c [200x150x100x50,1e-6] -b 200
# make an EPI mask (FSL)
/usr/local/fsl/bin/bet "${folder}"data/meanEPI corrected.nii "${folder}"data/meanEPI brain -f
0.5 -g 0 -n -m
# resample t1slab to T1 resolution (FreeSurfer)
mri convert -cs 1 -odt float -rl "${folder}"data/T1mean.nii -rt cubic "${folder}"data/t1slab.nii
"${folder}"data/t1slab 1mm.nii
# ------#
```

```
# ----- start the transformation ----- #
# study template -> MNI
antsRegistrationSyN.sh -d 3 -t s -f "${MNI}" -m "${template}" -o
"${folder}"NLreg_template_to_MNI_
# T1 -> study template
antsRegistrationSyN.sh -d 3 -t s -f "${template}" -m "${folder}"data/T1mean_corrected.nii -o
"${folder}"data/NLreg T1mean to template
# T1 -> MNI
antsApplyTransforms -d 3 -v 1 -n BSpline[4] -t
"${folder}"NLreg template to MNI 1Warp.nii.gz -t
"${folder}"NLreg template to MNI 0GenericAffine.mat -t
"${folder}"data/NLreg T1mean to template 1Warp.nii.gz -t
"${folder}"data/NLreg T1mean to template 0GenericAffine.mat -i
"${folder}"data/T1mean corrected.nii -r "${MNI}" -o
"${folder}"data/NLreg T1mean to MNI.nii
# T1 -> EPI
antsRegistrationSyN.sh -d 3 -t r -m "${folder}"data/T1mean corrected.nii -f
"${folder}"data/meanEPI corrected.nii -x "${folder}"data/meanEPI brain mask.nii.gz -o
"${folder}"data/coreg T1mean to meanEPI
# t1slab(resampled) -> T1
antsRegistrationSyN.sh -d 3 -t r -m "${folder}"data/t1slab 1mm.nii -f
"${folder}"data/T1mean_corrected.nii -o "${folder}"data/coreg_t1slab_to_T1mean_
# T1 -> MNI
antsApplyTransforms -d 3 -v 1 -n BSpline[4] -t
"${folder}"NLreg template to MNI 1Warp.nii.gz -t
"${folder}"NLreg template to MNI 0GenericAffine.mat -t
"${folder}"data/NLreg T1mean to template 1Warp.nii.gz -t
"${folder}"data/NLreg T1mean to template 0GenericAffine.mat -i
"${folder}"data/T1mean corrected.nii -r "${MNI}" -o
"${folder}"data/NLreg T1mean to MNI.nii
```

```
# EPI -> MNI
antsApplyTransforms -d 3 -v 1 -n BSpline[4] -t
"${folder}"NLreg template to MNI 1Warp.nii.gz -t
"${folder}"NLreg template to MNI 0GenericAffine.mat -t
"${folder}"data/NLreg_T1mean_to_template_1Warp.nii.gz -t
"${folder}"data/NLreg T1mean to template 0GenericAffine.mat -t
["${folder}"data/coreg T1mean to meanEPI 0GenericAffine.mat, 1] -i
"${folder}"data/meanEPI_corrected.nii -r "${MNI}" -o
"${folder}"data/NLreg meanEPI to MNI.nii
# LC segmentation -> MNI
antsApplyTransforms -d 3 -v 0 -n NearestNeighbor -t
"${folder}"NLreg template to MNI 1Warp.nii.gz -t
"${folder}"/NLreg template to MNI 0GenericAffine.mat -t
"${folder}"/data/NLreg T1mean to template 1Warp.nii.gz -t
"${folder}"/data/NLreg T1mean to template 0GenericAffine.mat -t
"${folder}"/data/coreg t1slab to T1mean 0GenericAffine.mat -i "${mask}" -r "${MNI}" -o
"${folder}"/data/NLreg LCmask to MNI.nii
# 1st-level stats images -> MNI : ! linear interpolation !
for I in {01..19}
do
      antsApplyTransforms -d 3 -v 1 -n Linear -t
"${folder}"NLreg template to MNI 1Warp.nii.gz -t
"${folder}"NLreg template to MNI 0GenericAffine.mat -t
"${folder}"data/NLreg T1mean to template 1Warp.nii.gz -t
"${folder}"data/NLreg T1mean to template 0GenericAffine.mat -t
["${folder}"data/coreg T1mean to meanEPI 0GenericAffine.mat, 1] -i
"${folder}"data/con 00${I}.nii -r "${MNI}" -o "${folder}"data/con 00${I} mni.nii
done
# ----- #
```

b. Generate Nifti landmark overlay image (a MATLAB script)

 $\$ Check the coregistration with EPI landmarks

% make a plot

```
clear;clc
```

```
%% preparation
% SET PATHS
path_root = '/path/to/your/landmark/images/';
path_save = '/path/to/the/folder/where/the/heatmap/will/be/saved/';
cd(path_save)
ids = [];% put your ID list
nid = length(ids);
%% make
cc = 0; % reset the counter
averplot = zeros(193,229,193); % pre-assign the dummy variable for plotting
the masks
figure; % open sketchbook
for i = 1:length(ids)
    averplotdum = zeros(193,229,193);
    name = num2str(ids(i));
    data = spm_read_vols(spm_vol([path_root name '/data/LCmask_check_' name
'.nii'])); % change here accordingly to your file name
    [x y z] = ndgrid(1:size(data,1), 1:size(data,2), 1:size(data,3));
    % get x/y/z level of LC masks
    ycoordz = y(find(data(:) == 1)); uycoordz = unique(ycoordz);
    ylevel = floor(median(uycoordz)); % take middle one
    xcoordz = x(find(data(:) == 1)); uxcoordz = unique(xcoordz);
    xlevel = ceil(median(uxcoordz)); % take middle one
    zcoordz = z(find(data(:) == 1)); uzcoordz = unique(zcoordz);
    zlevel = ceil(median(uzcoordz)); % take middle one
```

```
% right now, z position, which is an axial view, seems to be the most
    % appropriate slice to check the coregistration. if you want to take a
    % look at the other views, please change the script accordingly
    dumz = squeeze(data(:,:,zlevel));
    imagesc(dumz);title([name ' ' num2str(i) ]) ;% middle slice
    cc = cc+1;
    zstore(cc) = zlevel;
    averplotdum(:,:,zlevel) = dumz; % average points and squeeze them in
      dumy = squeeze(data(:,ylevel,:));
      imagesc(dumy);title([name ' ' num2str(i) ]);% middle slice
용
      cc = cc+1;
용
      ystore(cc) = ylevel;
      averplotdum(:,ylevel,:) = dumy;
용
      dumx = squeeze(sum(data(:,(xlevel-1):(xlevel+1),:),2));
      imagesc(dumx);title([name ' ' num2str(i) ]);% middle slice
용
용
      cc = cc+1;
용
      xstore(cc) = xlevel;
용
      averplotdum(xlevel,:,:) = dumx;
    averplot = averplot + averplotdum; % record the points
    imagesc(squeeze(averplotdum(:,:,zlevel)));hold on % and plot
      imagesc(squeeze(averplotdum(xlevel,:,:)));hold on % x position plot
      imagesc(squeeze(averplotdum(:,ylevel,:)));hold on % y position plot
    clear dumx dumy dumz
end
averplot = averplot/cc; % get average
%% generate the overlay image in 3D space
% one mask where all voxel > 0 are 1
averplot_nifti = averplot; averplot_nifti(find(averplot_nifti(:)>0)) = 1;
hdr = spm vol([path root /data/landmarks.nii']); % pick just any header from
a file
hdr.fname = [path_save 'landmark_heatmap.nii'];
```

```
hdr.dim = size(averplot_nifti);
hdr = rmfield(hdr,'pinfo');
hdr.nii = spm_write_vol(hdr,averplot_nifti);
```

c. Generate video of transformed mean functional or structural images (a MATLAB script)

```
%% Visual check for coregistered images
   make movie of epis to see whetehr OK aligned
%% preparation
clear;clc
% set paths
path root = '/path/to/your/transformed/images/'; % where are the files?
path save = '/path/where/the/video/will/be/saved/'; % where is the video
going to be saved?
% set variables
ids = [];
input prompt = {'duration per frame in seconds (default=0)'; 'name of the
file'; 'view(sagittal, coronal, axial)'; 'which slice should the video
capture?'};
defaults = {'0', 'null', 'sagittal', '96'};
input_answer = inputdlg(input_prompt, 'specify the properties of the video',
1, defaults);
vidframerate = str2num(input_answer{1,1});
vidformat = '.avi';
           = [input_answer{2,1} vidformat];
vidname
           = input_answer{3,1};
vidview
slicenum = str2num(input_answer{4,1});
cd(path_save)
            = VideoWriter(vidname);
if vidframerate ~= 0
   v.FrameRate = 1/vidframerate;
elseif vidframerate == 0
   v.FrameRate = 30; % it's super fast
end
%% start recording
```

```
open(v) % open the file
cc = 0; figure % open a sketchbook
for i = 1:length(ids)
        name = num2str(ids(i)); disp(num2str(ids(i)))
        data
                               spm_read_vols(spm_vol([path_root
                                                                        name
'/data/transformed_image.nii']));
        % position the slice - dimensions are: (x=sagittal, y=coronal,
z=axial)
        if strcmpi(vidview, 'sagittal')
            dum = squeeze(data(slicenum,:,:)); % sagittal
        elseif strcmpi(vidview, 'coronal')
            dum = squeeze(data(:,slicenum,:)); % coronal view
        elseif strcmpi(vidview, 'axial')
            dum = squeeze(data(:,:,slicenum)); % axial
        else
            error('check your view input')
        end
        dum = rot90(dum);% t1WB on template
      dum = zscore(dum);
        imagesc(dum);
        title(num2str(ids(i)))
        cc = cc+1;
        % Store the frame
        M(cc)=getframe(gcf); % leaving gcf out crops the frame in the movie.
end
writeVideo(v,M) % export the video
close(v)
close all;
```

d. Calculate the Euclidean distance of single-subject landmarks (a MATLAB script)

```
%% LANDMARK PLOTTING: create 3D plots of created heatmaps, and compare
clc;clear;close all
% set env
path_transformed = '/path/to/your/individual/landmarks/';
load('/path/to/your/subject/IDs/IDs.mat') % load IDs
% load individual landmark images
transformed landmarks single=[];
for subj=1:length(IDs)
    transformed_landmarks_single{subj,1}=spm_read_vols(spm_vol([
path_transformed 'landmark_' IDs{subj} '.nii']));
end; clear subj
% load the landmark image drawn on MNI
MNIlandmark =
spm read vols(spm vol(['/path/to/the/MNI/landmark/MNI landmark.nii'])); %
change here accordingly to your file name
MNIlandmark(find(MNIlandmark<1))=NaN;</pre>
% write down the coordinates of MNI landmarks
MNI.NucRuber=[];
MNI.NucRuber.left=[102 113 71];
MNI.NucRuber.right=[92 113 71];
MNI.TopBrainstem=[97 102 71];
MNI.OutlineBrainstem=[];
MNI.OutlineBrainstem.left=[109 103 60];
MNI.OutlineBrainstem.right=[85 103 60];
MNI.LC=[];
MNI.LC.left=[100 97 60];
MNI.LC.right=[94 97 60];
MNI.BottomBrainstem=[97 94 50];
disp('prep done')
%% identify median coords in the single landmarks
transformed landmark coords=[];
for subj=1:length(IDs)
    clear xs ys zs tmp idx2
    [xs,ys,zs] =
ind2sub(size(transformed landmarks single{subj}),find(transformed landmarks
_single{subj}~=0));
    tmp = [xs ys zs];
    [\sim,idx2] = sort(tmp(:,3)); tmp_sorted = tmp(idx2,:);
    slices = unique(tmp(:,3));
    slices = slices(end:-1:1);
```

```
%% the first slice, with three landmarks: top, NucRubs
    clear idx3 positionSlice positionX_sorted cluster1 cluster2 cluster3
    positionSlice = tmp sorted(find(tmp sorted(:,3)==slices(1)),:); % find
the points in the first slice
    [~,idx3] = sort(positionSlice(:,1)); positionX sorted =
positionSlice(idx3,:);
    % ruber right
    cluster1 = min(positionX_sorted(:,1));
    cluster med1 =
positionX sorted(find(positionX sorted(:,1)==cluster1+1),:);
    cluster_medpt{subj,1} = cluster_med1( find( cluster_med1(:,2)==ceil(
median(cluster_med1(:,2))) ),: );
    % top curve
    cluster2 = min(positionX sorted(positionX sorted(:,1)>cluster1+2,1));
    cluster med2 =
positionX sorted(find(positionX sorted(:,1)==cluster2+1),:);
    cluster_medpt{subj,2} = cluster_med2( find( cluster_med2(:,2)==ceil(
median(cluster_med2(:,2))) ),: );
    % ruber left
    cluster3 = min(positionX_sorted(positionX_sorted(:,1)>cluster2+2,1));
    cluster med3 =
positionX_sorted(find(positionX_sorted(:,1)==cluster3+1),:);
    cluster_medpt{subj,3} = cluster_med3( find( cluster_med3(:,2)==ceil(
median(cluster_med3(:,2))) ),: );
    transformed_landmark_coords{subj,1}.TopSlice = [];
    transformed landmark coords{subj,1}.TopSlice.NucRuber left =
cluster medpt{subj,3};
    transformed landmark coords{subj,1}.TopSlice.NucRuber right =
cluster_medpt{subj,1};
    transformed landmark coords{subj,1}.TopSlice.TopBrainstem =
cluster medpt{subj,2};
    clear cluster medpt
    %% the second slice, with four landmarks: brainstem outline and LC
    clear idx3 positionY positionX_sorted cluster1 cluster2 cluster3
    positionSlice = tmp sorted(find(tmp sorted(:,3)==slices(2)),:);
    [~,idx3] = sort(positionSlice(:,1)); positionX sorted =
positionSlice(idx3,:);
    % brainstem outline right
    cluster1 = min(positionX sorted(:,1));
    cluster med1 =
positionX sorted(find(positionX sorted(:,1)==cluster1+1),:);
    cluster medpt{subj,1} = cluster med1( find( cluster med1(:,2)==ceil(
median(cluster med1(:,2))) ),: );
    % LC right
    cluster2 = min(positionX sorted(positionX sorted(:,1)>(cluster1+2),1));
    cluster med2 =
positionX_sorted(find(positionX_sorted(:,1)==cluster2+1),:);
```

```
cluster medpt{subj,2} = cluster med2( find( cluster med2(:,2)==ceil(
median(cluster med2(:,2))) ),: );
    % LC left
    cluster3 = min(positionX sorted(positionX sorted(:,1)>cluster2+2,1));
    cluster med3 =
positionX_sorted(find(positionX_sorted(:,1)==cluster3+1),:);
    cluster medpt{subj,3} = cluster med3( find( cluster med3(:,2)==ceil(
median(cluster_med3(:,2))) ),: );
    % brainstem outline left
    cluster4 = min(positionX sorted(positionX sorted(:,1)>cluster3+2,1));
    cluster med4 =
positionX sorted(find(positionX sorted(:,1)==cluster4+1),:);
    cluster medpt{subj,4} = cluster med4( find( cluster med4(:,2)==ceil(
median(cluster med4(:,2))) ),: );
    transformed landmark coords{subj,1}.MidSlice = [];
    transformed landmark coords{subj,1}.MidSlice.OutlineBrainstem left =
cluster medpt{subj,4};
    transformed landmark coords{subj,1}.MidSlice.OutlineBrainstem right =
cluster medpt{subj,1};
    transformed landmark coords{subj,1}.MidSlice.LC left =
cluster medpt{subj,3};
    transformed_landmark_coords{subj,1}.MidSlice.LC_right =
cluster_medpt{subj,2};
    clear cluster medpt
    %% the third slice, with one landmark: bottom window brainstem
    clear idx3 positionY positionX sorted cluster1 cluster2 cluster3
cluster4
    positionSlice = tmp sorted(find(tmp sorted(:,3)==slices(3)),:);
    [~,idx3] = sort(positionSlice(:,1)); positionX sorted =
positionSlice(idx3,:);
    % brainstem bottom
    cluster1 = min(positionX_sorted(:,1));
    cluster med1 =
positionX_sorted(find(positionX_sorted(:,1)==cluster1+1),:);
    cluster_medpt{subj,1} = cluster_med1( find( cluster_med1(:,2)==ceil(
median(cluster_med1(:,2))) ),: );
    transformed_landmark_coords{subj,1}.BottomSlice = [];
    transformed landmark coords{subj,1}.BottomSlice.BottomBrainstem =
cluster medpt{subj,1}
    clear cluster_medpt
    fprintf('\n subject %s done\n', IDs{subj})
end;clear subj
disp('coordinates collected')
%% calculate euclidian distances in single subject level
```

```
varnames =
{'NucRuber left';'NucRuber right';'TopBrainstem';'OutlineBrainstem left';'O
utlineBrainstem_right';...
    'LC_left';'LC_right';'BottomBrainstem'};
for v1=1:length(varnames)
   eval([varnames{v1} '=[];'])
end
EuclideanDistances indv=[];
for subj=1:length(IDs)
    EuclideanDistances indv{subj,1}.NucRuber=[];
    EuclideanDistances indv{subj,1}.NucRuber.left =
sum((transformed landmark coords{subj,1}.TopSlice.NucRuber left-
MNI.NucRuber.left).^2).^0.5;
    EuclideanDistances indv{subj,1}.NucRuber.right =
sum((transformed_landmark_coords{subj,1}.TopSlice.NucRuber_right-
MNI.NucRuber.right).^2).^0.5;
    EuclideanDistances indv{subj,1}.TopBrainstem =
sum((transformed landmark coords{subj,1}.TopSlice.TopBrainstem-
MNI.TopBrainstem).^2).^0.5;
    EuclideanDistances_indv{subj,1}.OutlineBrainstem=[];
EuclideanDistances indv{subj,1}.OutlineBrainstem.left=sum((transformed land
mark coords{subj,1}.MidSlice.OutlineBrainstem left-
MNI.OutlineBrainstem.left).^2).^0.5;
EuclideanDistances indv{subj,1}.OutlineBrainstem.right=sum((transformed lan
dmark coords{subj,1}.MidSlice.OutlineBrainstem right-
MNI.OutlineBrainstem.right).^2).^0.5;
    EuclideanDistances indv{subj,1}.LC=[];
EuclideanDistances_indv{subj,1}.LC.left=sum((transformed_landmark_coords{su
bj,1}.MidSlice.LC left-MNI.LC.left).^2).^0.5;
EuclideanDistances_indv{subj,1}.LC.right=sum((transformed_landmark_coords{s
ubj,1}.MidSlice.LC right-MNI.LC.right).^2).^0.5;
    EuclideanDistances indv{subj,1}.BottomBrainstem =
sum((transformed_landmark_coords{subj,1}.BottomSlice.BottomBrainstem-
MNI.BottomBrainstem).^2).^0.5;
    % write as a table
    NucRuber_left(subj,1)=EuclideanDistances_indv{subj,1}.NucRuber.left;
    NucRuber_right(subj,1)=EuclideanDistances_indv{subj,1}.NucRuber.right;
    TopBrainstem(subj,1)=EuclideanDistances_indv{subj,1}.TopBrainstem;
OutlineBrainstem left(subj,1)=EuclideanDistances indv{subj,1}.OutlineBrains
tem.left;
OutlineBrainstem right(subj,1)=EuclideanDistances indv{subj,1}.OutlineBrain
stem.right;
    LC_left(subj,1)=EuclideanDistances_indv{subj,1}.LC.left;
    LC_right(subj,1)=EuclideanDistances_indv{subj,1}.LC.right;
BottomBrainstem(subj,1)=EuclideanDistances indv{subj,1}.BottomBrainstem;
```

```
spssID = cell2mat(cellfun(@str2num, IDs, 'UniformOutput',false));
T=table(NucRuber_left,NucRuber_right,TopBrainstem,OutlineBrainstem_left,Out lineBrainstem_right,...
    LC_left,LC_right,BottomBrainstem);
ED_export
=[spssID',NucRuber_left,NucRuber_right,TopBrainstem,OutlineBrainstem_left,OutlineBrainstem_right,...
    LC_left,LC_right,BottomBrainstem];
save(['/path/where/the/folder/the/data/will/be/saved/EuclideanDistance_land marks.mat'],'ED_export')
disp('ED_calc_done')
```

e. Calculate the Euclidean distance between the slice-wise centroids of a template LC mask and single-subject transformed LC segmentations (a MATLAB script)

```
%% calculate Euclidean distance: LC segmentations
clc;clear;close all
warning off
% paths
path transformed = '/path/to/the/transformed/LCsegmentations/';
IDs=[]; % subject IDs
% load images
% aggregated LC segmentations, transformed & binarized
LC_tf_bin = spm_read_vols(spm_vol([path_transformed
'LCmask_heatmap_binary.nii']));    LC_tf_bin(find(LC_tf_bin==0))=NaN;
% template MNI LC mask
MNImask =
spm_read_vols(spm_vol(['/path/to/the/template/LCMask/mni_icbm152/mni_icbm15
2 LCmetaMask MNI05 s01f plus50 bin.nii'])); MNImask(MNImask==0)=0;
% identify coordinates of each voxel
[x_MNI,y_MNI,z_MNI] = ind2sub(size(MNImask),find(MNImask~=0));
[x_tf,y_tf,z_tf] = ind2sub(size(LC_tf_bin),find(~isnan(LC_tf_bin)));
% coordinates in double
positions MNI = [x MNI,y MNI,z MNI];
[~,idx1] = sort(positions MNI(:,2));
slices MNI mask = unique(positions MNI(:,3)); clear idx1
%% draw and check the transformed & aggregated LC segmentations in 3D space
close all
S1 = repmat([170], numel(x tf), 1);
S2 = repmat([200], numel(x_MNI), 1);
hFig = figure();
axh = axes('Parent', hFig);
set(gca,'FontSize',25); hold on
hold(axh, 'all');
h2 = scatter3(x_MNI,y_MNI,z_MNI,S2,'d',...
'MarkerEdgeColor','g',...
'MarkerFaceColor',[0.4660 0.6740 0.1880]);
h1 = scatter3(x_tf,y_tf,z_tf,S1,'o',...
    'MarkerEdgeColor','r',...
'MarkerFaceColor',[1 0.2 0.2],'MarkerFaceAlpha',.5);
xlabel('X','FontWeight','bold')
ylabel('Y','FontWeight','bold')
zlabel('Z','FontWeight','bold')
xlim([88 107])
ylim([91 99])
zlim([46 64])
view(axh, -33, 22);
grid(axh, 'on');
set(gca, 'FontSize', 18);
legend(axh, [h1,h2], {'Transformed Masks', 'MNI Meta LC mask'});
%% find centroid voxel
```

```
% left LC
dummybase = zeros(193,229,193); % make a blank canvas
tmp MNI=[x MNI,y MNI,z MNI]; % tidy up the xyz coordinates from the
original mask
indL = tmp MNI(:,1) < mean(x MNI); % identify coordinates of left and right</pre>
coordsL = tmp MNI(indL,:,:);
indR = tmp MNI(:,1)>mean(x MNI);
coordsR = tmp MNI(indR,:,:);
leftLC=dummybase;
leftLC(sub2ind(size(leftLC),coordsL(:,1),coordsL(:,2),coordsL(:,3))) = 1;
rightLC=dummybase;
rightLC(sub2ind(size(leftLC),coordsR(:,1),coordsR(:,2),coordsR(:,3))) = 1;
[x_Left,y_Left,z_Left]=ind2sub(size(leftLC),find(leftLC~=0));
[x_Right,y_Right,z_Right]=ind2sub(size(rightLC),find(rightLC~=0));
centroid_L_mni =
[round(mean(x_Left)),round(mean(y_Left)),round(mean(z_Left))];
centroid R mni =
[round(mean(x_Right)),round(mean(y_Right)),round(mean(z_Right))];
centroid_both_mni =
[round(mean(x_MNI)),round(mean(y_MNI)),round(mean(z_MNI))];
%% extract centroid of single-subject masks
centroid L = []; centroid R =[];
for id = 1:length(IDs)
    clear tmp leftLC rightLC indL indR coordsL coordsR xi Left yi Left
zi Left ...
        xi Right yi Right zi Right
    LCsegs individual binary{id,1} =
spm read vols(spm vol([path transformed 'threshold025 ' IDs{id})
'_conjmask_mni.nii']));
    dummybase = zeros(193,229,193); % make a blank canvas
    [xi,yi,zi] =
ind2sub(size(LCsegs individual binary{id,1}),find(LCsegs individual binary{
id, 1 \sim = 0);
    tmp=[xi,yi,zi]; % tidy up the xyz coordinates from the original mask
    indR = tmp(:,1)>97;%mean(xi); % identify coordinates of
    coordsR = tmp(indR,:,:);
    indL = tmp(:,1)<97;%mean(xi);</pre>
    coordsL = tmp(indL,:,:);
    leftLC=dummybase;
    leftLC(sub2ind(size(leftLC),coordsL(:,1),coordsL(:,2),coordsL(:,3))) =
1;
    rightLC=dummybase;
    rightLC(sub2ind(size(leftLC),coordsR(:,1),coordsR(:,2),coordsR(:,3))) =
1;
    % tmp keren=kerenmask;%[x keren,y keren,z keren];
```

```
% leftLC = tmp keren;leftLC(x keren<mean(x keren),:,:)=0; % choose</pre>
everything that's on the left
    % rightLC = tmp_keren;rightLC(x_keren>mean(x_keren),:,:)=0; % choose
everything that's on the right
    [xi_Left,yi_Left,zi_Left]=ind2sub(size(leftLC),find(leftLC~=0));
    left_indivs{id,1} = [xi_Left,yi_Left,zi_Left];
    [xi_Right,yi_Right,zi_Right]=ind2sub(size(rightLC),find(rightLC~=0));
    right_indivs{id,1} = [xi_Right,yi_Right,zi_Right];
    centroid L(id,:) =
[round(mean(xi Left)),round(mean(yi Left)),round(mean(zi Left))];
    centroid R(id,:) =
[round(mean(xi_Right)),round(mean(yi_Right)),round(mean(zi_Right))];
    centroid both(id,:) =
[round(mean(xi)),round(mean(yi)),round(mean(zi))];
end
%% ED from the centroid point, slicewise analysis
% find centre of each slice
MNI_left = [x_Left,y_Left,z_Left];
MNI_right = [x_Right,y_Right,z_Right];
MNI slices L = unique(z Left);
MNI slices R = unique(z Right);
for slices = 1:length(MNI slices L)
    clear tmp zind
    % sort the left
    zind=z Left==MNI slices L(slices);
    tmp = MNI left(zind',:,:);
    if size(MNI_left(zind',:,:),1)==1
    Lcentrer_mni(slices,:) = [tmp(:,1:2) MNI_slices_L(slices)];
    Lcentrer mni(slices,:) = [mean(tmp(:,1:2)) MNI slices L(slices)];
end
for slices = 1:length(MNI slices R)
    clear tmp zind
    % sort the left
    zind=z_Right==MNI_slices_R(slices);
    tmp = MNI_right(zind',:,:);
    if size(MNI_right(zind',:,:),1)==1
    Rcentrer_mni(slices,:) = [tmp(:,1:2) MNI_slices_R(slices)];
    Rcentrer_mni(slices,:) = [mean(tmp(:,1:2)) MNI_slices_R(slices)];
    end
end
% now find centres of individual masks
Lcentre_indiv=[];Rcentre_indiv=[];
for id=1:length(IDs)
    clear maskL maskR slicesL slicesR sl
```

```
% first, left
    maskL=right_indivs{id,1}; % there's something wrong with this left and
right
    slicesL=unique(maskL(:,3));
    for sl=1:length(slicesL)
        clear tmp zind
        % sort the left
        zind=maskL(:,3)==slicesL(sl);
        tmp = maskL(zind',:,:);
        if size(tmp,1) == 1
            Lcentre indiv{id,1}(sl,:) = [tmp(:,1:2) slicesL(sl)];
            Lcentre_indiv{id,1}(sl,:) = [mean(tmp(:,1:2)) slicesL(sl)];
        end
    end
    % then, right
    maskR=left indivs{id,1}; % there's something wrong with this left and
right
    slicesR=unique(maskR(:,3));
    for sl=1:length(slicesR)
        clear tmp zind
        % sort the left
        zind=maskR(:,3)==slicesR(sl);
        tmp = maskR(zind',:,:);
        if size(tmp,1) == 1
            Rcentre_indiv{id,1}(sl,:) = [tmp(:,1:2) slicesR(sl)];
        else
            Rcentre_indiv{id,1}(s1,:) = [mean(tmp(:,1:2)) slicesR(s1)];
        end
    end
end
%% calculate Euclidian distance of centroids, slicewise
ED_slice_L=[]; ED_slice_R=[];
for id=1:length(IDs)
    % find the slices that matches
    clear Lslc_tmp A B
    Lslc tmp = Lcentre indiv{id,1}(:,3);
    indslc1=ismember(MNI_slices_L,Lslc_tmp,'rows');
    indslc2=ismember(Lslc_tmp,MNI_slices_L,'rows');
    A = Lcentrer mni(indslc1,:);
    B = Lcentre_indiv{id,1}(indslc2,:);
    for k=1:size(B,1)
        clear v
             = B(k,:) - A(k,:);
        ED_slice_L{id,1}(k) = sqrt(nansum(v .^ 2));
    end
    % find the slices that matches
    clear Rslc_tmp A B
    Rslc_tmp = Rcentre_indiv{id,1}(:,3);
    indslc1=ismember(MNI slices R,Rslc tmp,'rows');
    indslc2=ismember(Rslc_tmp, MNI_slices_R, 'rows');
```

```
A = Rcentrer mni(indslc1,:);
    B = Rcentre indiv{id,1}(indslc2,:);
    for k=1:size(A,1)
        clear v
            = B(k,:) - A(k,:);
        ED_slice_R{id,1}(k) = sqrt(nansum(v .^ 2));
end
% additional processing
ED_slice_mean_L = cellfun(@nanmean, ED_slice_L);
ED slice mean R = cellfun(@nanmean, ED slice R);
%% simple stats
means(1,1)=nanmean(ED_slice_mean_L);
means(1,2)=nanmean(ED_slice_mean_R);
stds(1,1)=nanstd(ED_slice_mean_L);
stds(1,2)=nanstd(ED_slice_mean_R);
medians(1,1)=nanmedian(ED slice mean L);
medians(1,2)=nanmedian(ED_slice_mean_R);
fprintf('\n left LC mean: %1.4f, STD: %1.4f, median: %1.4f \n right LC
mean: %1.4f, STD: %1.4f, median: %1.4f \n', ...
    means(1,1), stds(1,1), medians(1,1), means(1,2), stds(1,2),
medians(1,2)
```