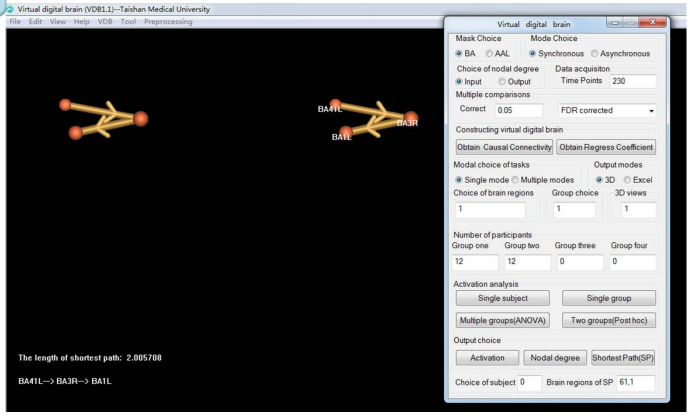
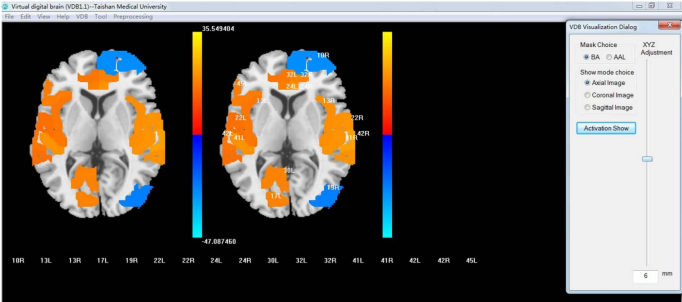
Virtual Digital Brain Manual





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1. Introduction

Please cite as '... was/were performanced using the virtual digital brain software package VDB1.1

(https://www.nitrc.org/projects/vdb/) while using the software to make publicized paper.

VDB1.1 is a 3D visualization tool of human brain, which is used to research neural activities of brain regions

evoked by the virtual stimulating signal (or the virtual task signal). In addition, this tool can also help

researchers to study causal relationships among brain regions, nodal degree and the shortest path length of the

directional brain network.

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2. Installation

Firstly, you need to download the software package from (https://www.nitrc.org/projects/vdb/) and unzip downloaded file. Copy all files in the folder "VDB1.1" to the directory: D: \\VDB, and then add the executable file VDB 1.1.exe in the folder "VDB" to the desktop shortcut.

3. Construction of virtual digital brain

3.1. Data preprocessing

Data were preprocessed using spm8(<u>http://www.fil.ion.ucl.ac.uk/spm/software/spm8/</u>). The performance is described as follows.

- (2) Spatial smoothing with a Gaussian kernel of a specified width is applied to the normalized functional images.
- (3) Normalized structural images are registered to the normalized functional images by applying rigid registration.
- (4) Those registered structural images are segmented into the white matter, gray matter, and cerebrospinal fluid images.
- (5) Preprocessed functional images of every subject are placed in a folders named as Subxxx(such as Sub001). All folders (Sub001, Sub002,···Subxxx) are combined into a big destination folder(for example, Detrend).

Preprocessed data mentioned above are further preprocessed using the following procedures.

(1) The removal of linear and quadratic trends. Open the software (i.e. run the VDB1.0.exe file in the folder "VDB" or the desktop) and click on the menu "Preprocessing", and then click on the option "Detrend", select the folder "Detrend" in the opened dialog. Finally, click on the button "Ok", the procedure will run and execute Detrend. Processed functional image files with the prefix "D" are stored in the folder "Detrend" (Figure 1).

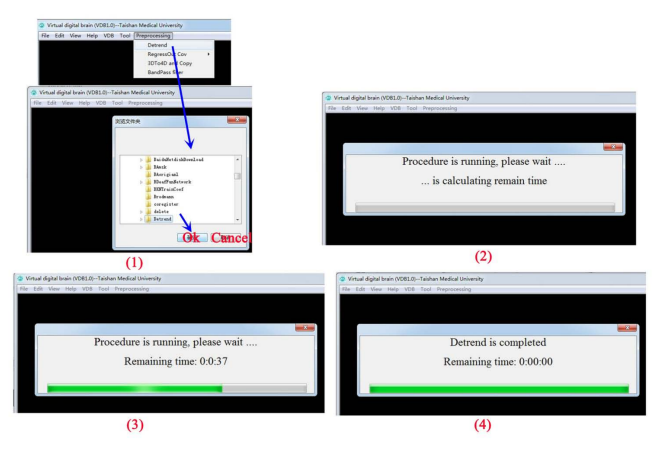


Figure 1. Detrend

- (2) Copy the white matter, cerebrospinal fluid and motion parameter image files of every subject to those folders Subxxx(such as Sub001, Sub002,…)in the folder "Detrend".
- (3) Regress out covariates including realignment parameters (motion parameters), the global mean signal, mean white matter signal, and the mean cerebrospinal fluid signal. Click on the option "RegressOut Cov" in the menu "Preprocessing". Click on the button "Starting regression" in the covariate regression dialog and select the folder "Detrend" in the opened dialog. Finally, click on the button "Ok", the procedure will run and execute covariate regression. Processed functional image files with the prefix "C" are stored in the directory: D: \DetrendCoved (Figure 2).

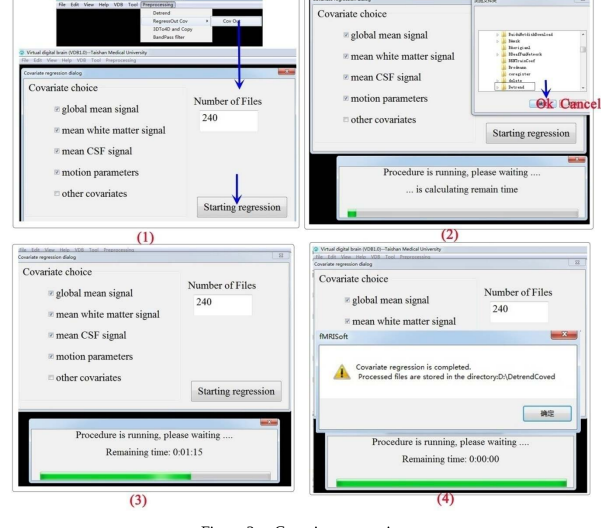


Figure 2. Covariate regression

(4) Integrating 3D images to 4D. As shown in Figure 3, firstly, click on the menu "Preprocessing", and then click on the option "3DTo4D and Copy". Select the folder "DetrendCoved" in the opened dialog, finally, click on the button "Ok", the procedure will run and execute data transform. Constructed 4D data are stored in the folder "DetrendCoved4D" (the directory:D: \\DetrendCoved4D).

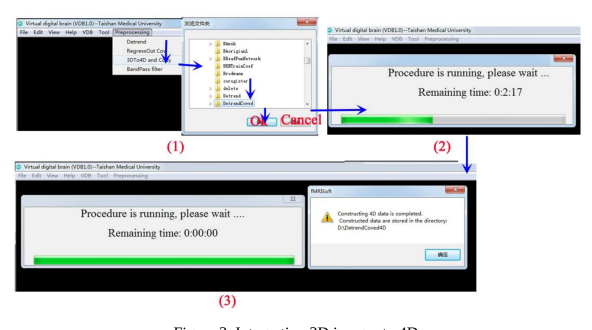


Figure 3. Integrating 3D images to 4D

(5) Band-pass temporal filter. As shown in Figure 4, firstly, click on the menu "Preprocessing", and then click on the option "Bandpass filter". Select the folder "DetrendCoved4D" in the opened dialog, finally, click on the button "Ok", the procedure will run and execute filtering. Filtered 4D data are stored in the folder "ConstructionVDB" (the directory:D: \\ConstructionVDB).

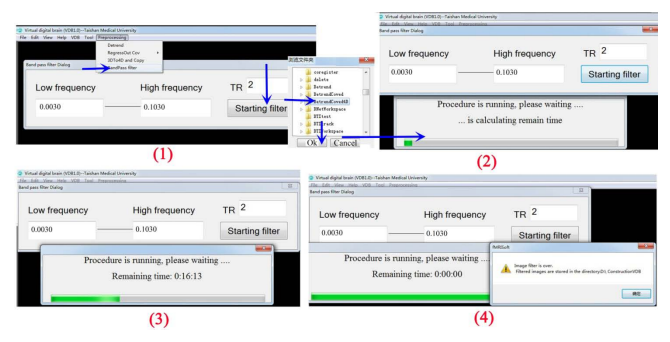


Figure 4. Band-pass temporal filter

3.2. Construction of brain causal network

The steps of construction are described as follows:

1. Firstly, open the software and click on the menu VDB, and then start the construction of the brain causal network (Figure 5).

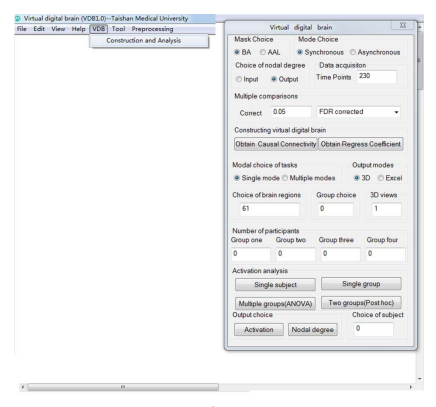


Figure 5. Construction of the brain causal network.

2. Select "BA" or "AAL" in the mask choice, "Synchronous" in the mode choice, "Time Points" in the data acquisition, and then click on the button "Obtain Causal Connectivity" and open the destination folder (Figure 6).

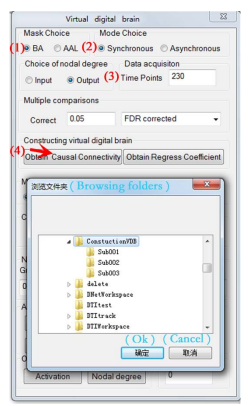


Figure 6. Opening the destination folder

3. Run the procedure and obtain the matrix of synchronous causal connectivity. The result is automatically stored in the directory: D: \\\VDB\\\CausalConnectivity(Figure 7).

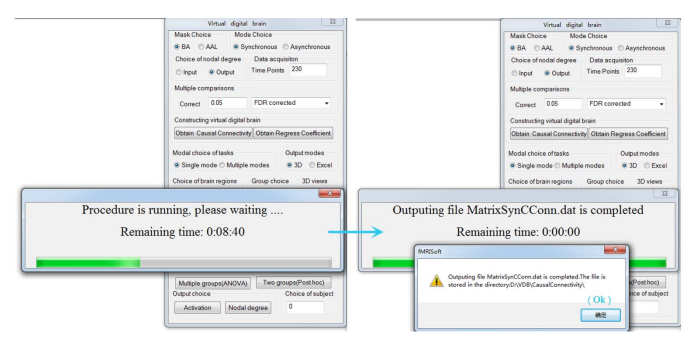


Figure 7. Obtaining the matrix of synchronous causal connectivity.

- 4. Select "Asynchronous" in the mode choice and repeat the steps 2 and 3. The matrix of asynchronous connectivity is also stored in the directory: D: \\\VDB\\\CausalConnectivity.
- 5. Select "BA" or "AAL" in the mask choice, "Time Points" in the data acquisition, corrected parameter in the multiple comparisons, and then click on the button "Obtain Regress Coefficient" and open the destination folder. Run the procedure and obtain the regress coefficient. The result is automatically stored in the directory: D:\\VDB\\RegressCoefficient.(Figure 8).

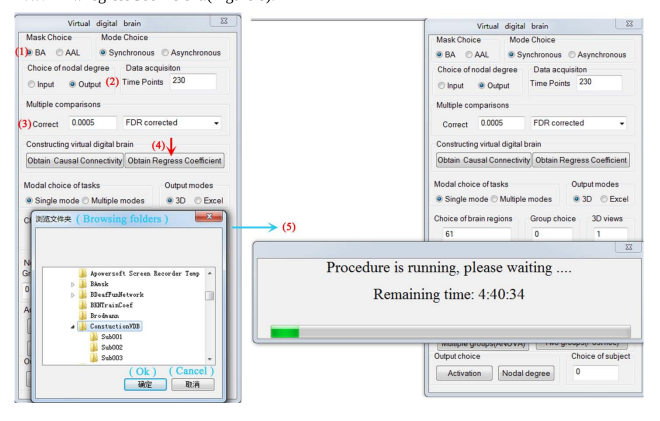


Figure 8. Obtaining the regress coefficient

4. Analysis of nodal degrees

4.1. Nodal degrees of the causal connectivity network

1. Nodal degrees of all subjects. Select "BA" or "AAL" in the mask choice, "Synchronous" or "Asynchronous" in the mode choice, "Input" or "Output" in the choice of nodal degree, corrected parameter in the multiple comparisons (this parameter must be equal to the value that has been used in the "Obtain Regress Coefficient" step), "Excel" in the output modes, and then fill "0" in the editor control "Group choice". In addition, the number of participants must be filled in these editor controls (Group one to four). Click on the button "Nodal degree" and obtain the nodal degrees of all subjects. The result is showed in an excel file (Figure 9).

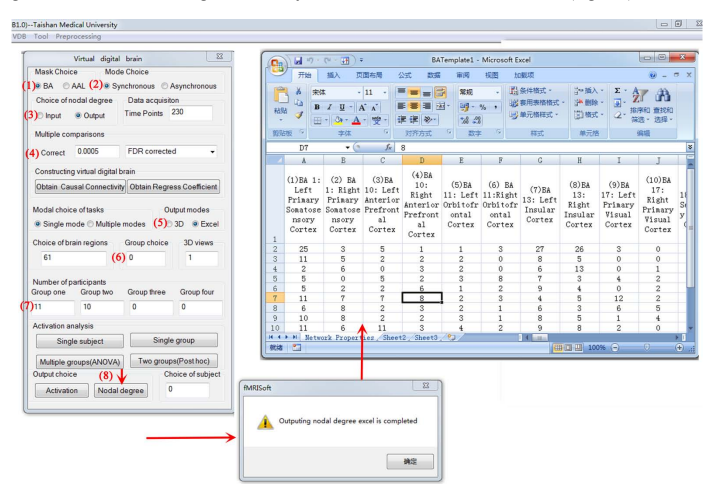


Figure 9. Nodal degrees of all subjects

2. Nodal degrees of the group (causal connectivity of every brain region). Select "BA" or "AAL" in the mask choice, corrected parameter in the multiple comparisons, "Excel" in the output modes, and then fill the code of group such as "1" or others (2-4) in the editor control "Group choice". In addition, the number of participants must be filled in these editor controls (Group one to four). Click on the button "Nodal degree" and obtain the

nodal degrees of the group. The result is showed in an excel file (Figure 10). Positive real numbers indicate the strengths of synchronous causal connectivity, and negative real numbers indicate the strengths of asynchronous causal connectivity. The real numbers of every row indicate the strengths of output causal connectivity corresponding to every node, and the real numbers of every column indicate the strengths of input causal connectivity corresponding to every node.

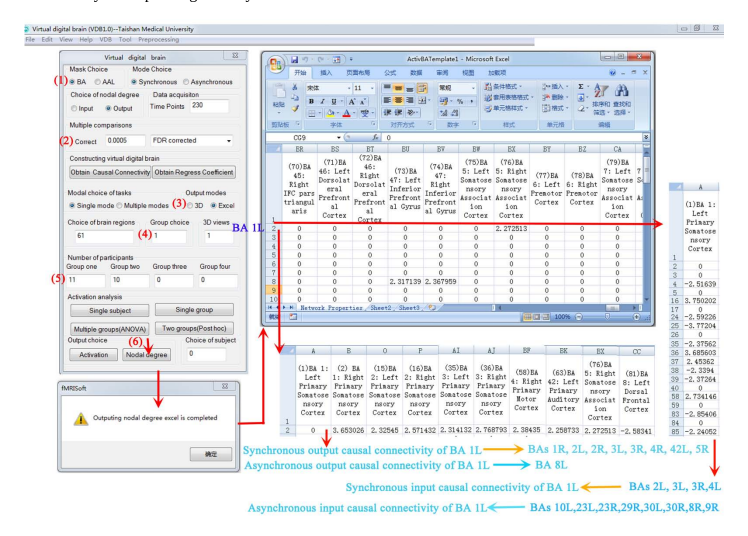


Figure 10. Nodal degrees of the group

3.Nodal degree of one brain region (3D visualization). Select "BA" or "AAL" in the mask choice, "Synchronous" or "Asynchronous" in the mode choice, "Input" or "Output" in the choice of nodal degree, corrected parameter in the multiple comparisons, and "3D" in the output modes, fill the index of displayed brain region in the "Choice of brain regions" control, and then fill the code of group such as "1" or others (2-4) in the editor control "Group choice", the index of 3D view in the editor control "3D views" (the index "1" indicates the superior view, "2" indicates the inferior view, "3" indicates the left view, "4" indicates the right view), and the

index of brain region in the editor control "Choice of brain regions" see also table 1 and table 2 for details. In addition, the number of participants must be filled in these editor controls (Group one to four). Click on the button "Nodal degree", and 3D visualization will run. The result is showed in the left of client area (Figure 11). The size of bar indicates the strength of causal connectivity, and the arrow indicates the direction of causal connectivity.

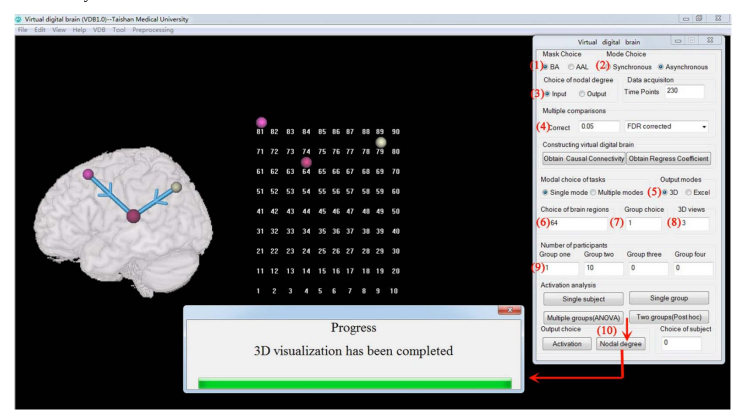


Figure 11. 3D visualization of the nodal degree.

Table 1 Indexes of brain regions and corresponding Brodmann areas

| Indexes | Index of Brodmann Areas |
|---------|-------------------------------------|
| 1 | 1 (L). Primary Somatosensory Cortex |
| 2 | 1 (R). Primary Somatosensory Cortex |
| 3 | 10 (L). Anterior Prefrontal Cortex |
| 4 | 10 (R). Anterior Prefrontal Cortex |
| 5 | 11 (L). Orbitofrontal Cortex |
| 6 | 11 (R). Orbitofrontal Cortex |
| 7 | 13 (L). Insular Cortex |
| 8 | 13 (R). Insular Cortex |
| 9 | 17 (L). Primary Visual Cortex |
| 10 | 17 (R). Primary Visual Cortex |
| 11 | 18 (L). Secondary Visual Cortex |
| 12 | 18 (R). Secondary Visual Cortex |

| 13 | 19 (L). Associative Visual Cortex |
|----|--|
| 14 | 19 (R). Associative Visual Cortex |
| 15 | 2 (L). Primary Somatosensory Cortex |
| 16 | 2 (R). Primary Somatosensory Cortex |
| 17 | 20 (L). Inferior Temporal Gyrus |
| 18 | 20 (R). Inferior Temporal Gyrus |
| 19 | 21 (L). Middle Temporal Gyrus |
| 20 | 21 (R). Middle Temporal Gyrus |
| 21 | 22 (L). Superior Temporal Gyrus |
| 22 | 22 (R). Superior Temporal Gyrus |
| 23 | 23 (L). Ventral Posterior Cingulate Cortex |
| 24 | 23 (R). Ventral Posterior Cingulate Cortex |
| 25 | 24 (L). Ventral Anterior Cingulate Cortex |
| 26 | 24 (R). Ventral Anterior Cingulate Cortex |
| 27 | 25 (L). Subgenual cortex |
| 28 | 25 (R). Subgenual cortex |
| 29 | 27 (L). Piriform Cortex |
| 30 | 27 (R). Piriform Cortex |
| 31 | 28 (L). Posterior Entorhinal Cortex |
| 32 | 28 (R). Posterior Entorhinal Cortex |
| 33 | 29 (L). Retrosplenial Cingulate Cortex |
| 34 | 29 (R). Retrosplenial Cingulate Cortex |
| 35 | 3 (L). Primary Somatosensory Cortex |
| 36 | 3 (R). Primary Somatosensory Cortex |
| 37 | 30 (L). Cingulate Cortex |
| 38 | 30 (R). Cingulate Cortex |
| 39 | 31 (L). Dorsal Posterior Cingulate Cortex |
| 40 | 31 (R). Dorsal Posterior Cingulate Cortex |
| 41 | 32 (L). Dorsal anterior Cingulate Cortex |
| 42 | 32 (R). Dorsal anterior Cingulate Cortex |
| 43 | 33 (L). Anterior Cingulate Cortex |
| 44 | 33 (R). Anterior Cingulate Cortex |
| 45 | 34 (L). Anterior Entorhinal Cortex |
| 46 | 34 (R). Anterior Entorhinal Cortex |
| 47 | 35 (L). Perirhinal cortex |
| 48 | 35 (R). Perirhinal cortex |
| 49 | 36 (L). Parahippocampal cortex |
| 50 | 36 (R). Parahippocampal cortex |
| 51 | 37 (L). Fusiform gyrus |
| | |

| 52 | 37 (R). Fusiform gyrus |
|----|---|
| 53 | 38 (L). Temporopolar Area |
| 54 | 38 (R). Temporopolar Area |
| 55 | 39 (L). Angular gyrus |
| 56 | 39 (R). Angular gyrus |
| 57 | 4 (L). Primary Motor Cortex |
| 58 | 4 (R). Primary Motor Cortex |
| 59 | 40 (L). SupramarginalGyrus |
| 60 | 40 (R). SupramarginalGyrus |
| 61 | 41 (L). Primary Auditory Cortex |
| 62 | 41 (R). Primary Auditory Cortex |
| 63 | 42 (L). Primary Auditory Cortex |
| 64 | 42 (R). Primary Auditory Cortex |
| 65 | 43 (L). Subcentral Area |
| 66 | 43 (R). Subcentral Area |
| 67 | 44 (L). IFC pars opercularis |
| 68 | 44 (R). IFC pars opercularis |
| 69 | 45 (L). IFC pars triangularis |
| 70 | 45 (R). IFC pars triangularis |
| 71 | 46 (L). Dorsolateral Prefrontal Cortex |
| 72 | 46 (R). Dorsolateral Prefrontal Cortex |
| 73 | 47 (L). Inferior Prefrontal Gyrus |
| 74 | 47 (R). Inferior Prefrontal Gyrus |
| 75 | 5 (L). Somatosensory Association Cortex |
| 76 | 5 (R). Somatosensory Association Cortex |
| 77 | 6 (L). Premotor Cortex |
| 78 | 6 (R). Premotor Cortex |
| 79 | 7 (L). Somatosensory Association Cortex |
| 80 | 7 (R). Somatosensory Association Cortex |
| 81 | 8 (L). Dorsal Frontal Cortex |
| 82 | 8 (R). Dorsal Frontal Cortex |
| 83 | 9 (L). Dorsolateral Prefrontal Cortex |
| 84 | 9 (R). Dorsolateral Prefrontal Cortex |

Table 2 Indexes of brain regions and corresponding AALareas

| Indexes | AAL Areas |
|---------|----------------|
| 1 | Precentral (L) |
| 2 | Precentral (R) |

| 3 | Frontal Sup (L) |
|----|------------------------|
| 4 | Frontal Sup (R) |
| 5 | Frontal Sup Orb (L) |
| 6 | Frontal Sup Orb (R) |
| 7 | Frontal Mid (L) |
| 8 | Frontal Mid (R) |
| 9 | Frontal Mid Orb (L) |
| 10 | Frontal Mid Orb (R) |
| 11 | Frontal InfOper (L) |
| 12 | Frontal InfOper (R) |
| 13 | Frontal Inf Tri (L) |
| 14 | Frontal Inf Tri (R) |
| 15 | Frontal Inf Orb (L) |
| 16 | Frontal Inf Orb (R) |
| 17 | RolandicOper (L) |
| 18 | RolandicOper (R) |
| 19 | Supp Motor Area (L) |
| 20 | Supp Motor Area (R) |
| 21 | Olfactory (L) |
| 22 | Olfactory (R) |
| 23 | Frontal Sup Medial (L) |
| 24 | Frontal Sup Medial (R) |
| 25 | Frontal Med Orb (L) |
| 26 | Frontal Med Orb (R) |
| 27 | Rectus (L) |
| 28 | Rectus (R) |
| 29 | Insula (L) |
| 30 | Insula (R) |
| 31 | Cingulum Ant (L) |
| 32 | Cingulum Ant (R) |
| 33 | Cingulum Mid (L) |
| 34 | Cingulum Mid (R) |
| 35 | Cingulum Post (L) |
| 36 | Cingulum Post (R) |
| 37 | Hippocampus (L) |
| 38 | Hippocampus (R) |
| 39 | ParaHippocampal (L) |
| 40 | ParaHippocampal (R) |
| 41 | Amygdala (L) |
| | 15 |

| 42 Anygdala (R) 43 Calcarine (L) 44 Calcarine (R) 45 Cuncus (L) 46 Cuneus (R) 47 Lingual (L) 48 Lingual (R) 49 Occipital Sup (L) 50 Occipital Mid (L) 51 Occipital Mid (R) 53 Occipital Inf (R) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (R) 61 Parietal Inf (R) 62 Parietal Inf (R) 63 SupraMarginal (R) 64 Angular (R) 65 Angular (R) 66 Angular (R) 67 Precueus (R) 69 Paracentral Lobule (R) 71 Caudate (R) 73 Putamen (L) 74 Pallidum (R) | | |
|--|----|------------------------|
| 44 Calcarine (R) 45 Cuneus (L) 46 Cuncus (R) 47 Lingual (L) 48 Lingual (R) 49 Occipital Sup (L) 50 Occipital Mid (L) 51 Occipital Mid (R) 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (R) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (R) 61 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (R) 64 SupraMarginal (R) 65 Angular (R) 67 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (R) 73 Putamen (R) 74 Putamen (R) 75 Pallidum (R) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) <th>42</th> <th>Amygdala (R)</th> | 42 | Amygdala (R) |
| 45 Cuneus (R) 47 Lingual (L) 48 Lingual (R) 49 Occipital Sup (L) 50 Occipital Sup (R) 51 Occipital Mid (L) 52 Occipital Mid (R) 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (R) 61 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (R) 66 Angular (R) 67 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (R) 73 Putamen (R) 74 Putamen (R) 75 Pallidum (R) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R | 43 | Calcarine (L) |
| 46 Cuneus (R) 47 Lingual (L) 48 Lingual (R) 49 Occipital Sup (L) 50 Occipital Mid (L) 51 Occipital Mid (R) 52 Occipital Mid (R) 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (R) 73 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschi (L) <th>44</th> <th>Calcarine (R)</th> | 44 | Calcarine (R) |
| 47 Lingual (R) 48 Lingual (R) 49 Occipital Sup (R) 50 Occipital Mid (L) 51 Occipital Mid (R) 52 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (R) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (R) 72 Caudate (R) 73 Putamen (R) 75 Pallidum (R) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschi (L) <th>45</th> <th>Cuneus (L)</th> | 45 | Cuneus (L) |
| 48 Lingual (R) 49 Occipital Sup (R) 50 Occipital Mid (L) 51 Occipital Mid (R) 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 46 | Cuneus (R) |
| 49 Occipital Sup (R) 51 Occipital Mid (L) 52 Occipital Mid (R) 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (L) 61 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (R) 75 Pallidum (R) 75 Pallidum (R) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 47 | Lingual (L) |
| 50 Occipital Sup (R) 51 Occipital Mid (L) 52 Occipital Mid (R) 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (L) 61 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (R) 67 Precuncus (L) 68 Precuncus (R) 69 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 48 | Lingual (R) |
| 51 Occipital Mid (L) 52 Occipital Mid (R) 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 49 | Occipital Sup (L) |
| 52 Occipital Mid (R) 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 50 | Occipital Sup (R) |
| 53 Occipital Inf (L) 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (L) 61 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 51 | Occipital Mid (L) |
| 54 Occipital Inf (R) 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (R) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 52 | Occipital Mid (R) |
| 55 Fusiform (L) 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Sup (R) 61 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (R) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 53 | Occipital Inf (L) |
| 56 Fusiform (R) 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Sup (R) 61 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 54 | Occipital Inf (R) |
| 57 Postcentral (L) 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Sup (R) 61 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 55 | Fusiform (L) |
| 58 Postcentral (R) 59 Parietal Sup (L) 60 Parietal Sup (R) 61 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 56 | Fusiform (R) |
| 59 Parietal Sup (L) 60 Parietal Sup (R) 61 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 57 | Postcentral (L) |
| 60 Parietal Sup (R) 61 Parietal Inf (L) 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 58 | Postcentral (R) |
| 61 | 59 | Parietal Sup (L) |
| 62 Parietal Inf (R) 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 60 | Parietal Sup (R) |
| 63 SupraMarginal (L) 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 61 | Parietal Inf (L) |
| 64 SupraMarginal (R) 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 62 | Parietal Inf (R) |
| 65 Angular (L) 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 63 | SupraMarginal (L) |
| 66 Angular (R) 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 64 | SupraMarginal (R) |
| 67 Precuneus (L) 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 65 | Angular (L) |
| 68 Precuneus (R) 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 66 | Angular (R) |
| 69 Paracentral Lobule (L) 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 67 | Precuneus (L) |
| 70 Paracentral Lobule (R) 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 68 | Precuneus (R) |
| 71 Caudate (L) 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 69 | Paracentral Lobule (L) |
| 72 Caudate (R) 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 70 | Paracentral Lobule (R) |
| 73 Putamen (L) 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 71 | Caudate (L) |
| 74 Putamen (R) 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 72 | Caudate (R) |
| 75 Pallidum (L) 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 73 | Putamen (L) |
| 76 Pallidum (R) 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 74 | Putamen (R) |
| 77 Thalamus (L) 78 Thalamus (R) 79 Heschl (L) | 75 | Pallidum (L) |
| 78 Thalamus (R) 79 Heschl (L) | 76 | Pallidum (R) |
| 79 Heschl (L) | 77 | Thalamus (L) |
| | 78 | Thalamus (R) |
| 80 Heschl (R) | 79 | Heschl (L) |
| | 80 | Heschl (R) |

| 81 | Temporal Sup (L) |
|-----|-----------------------|
| 82 | Temporal Sup (R) |
| 83 | Temporal Pole Sup (L) |
| 84 | Temporal Pole Sup (R) |
| 85 | Temporal Mid (L) |
| 86 | Temporal Mid (R) |
| 87 | Temporal Pole Mid (L) |
| 88 | Temporal Pole Mid (R) |
| 89 | Temporal Inf (L) |
| 90 | Temporal Inf (R) |
| 91 | Cerebelum Crus1 (L) |
| 92 | Cerebelum Crus1 (R) |
| 93 | Cerebelum Crus2 (L) |
| 94 | Cerebelum Crus2 (R) |
| 95 | Cerebelum 3 (L) |
| 96 | Cerebelum 3 (R) |
| 97 | Cerebelum 4 5 (L) |
| 98 | Cerebelum 4 5 (R) |
| 99 | Cerebelum 6 (L) |
| 100 | Cerebelum 6 (R) |
| 101 | Cerebelum 7b (L) |
| 102 | Cerebelum 7b (R) |
| 103 | Cerebelum 8 (L) |
| 104 | Cerebelum 8 (R) |
| 105 | Cerebelum 9 (L) |
| 106 | Cerebelum 9 (R) |
| 107 | Cerebelum 10 (L) |
| 108 | Cerebelum 10 (R) |
| 109 | Vermis 1 2 |
| 110 | Vermis 3 |
| 111 | Vermis 4 5 |
| 112 | Vermis 6 |
| 113 | Vermis 7 |
| 114 | Vermis 8 |
| 115 | Vermis 9 |
| 116 | Vermis 10 |

5. Task design

The procedure "TaskDesign" in the folder VDB (D: \\VDB\\TaskDesign) is an example of designing task. The integrated development environment of this procedure is Microsoft Visual studio 2008 or above. Several functions have been built in the class "TaskConstruction", and these functions are responsible for constructing tasks or reading constructed signals. The testing functions have been built in the classes "CTaskDesignDoc" and "CTaskDesignView", and these functions are responsible for displaying the waveforms of the design matrix and constructed task signals (Figure 12). Users can add member functions in these classes to achieve task design and waveform display. The software "VDB1.0" can be used to analysis brain region activations based on these constructed task signals.



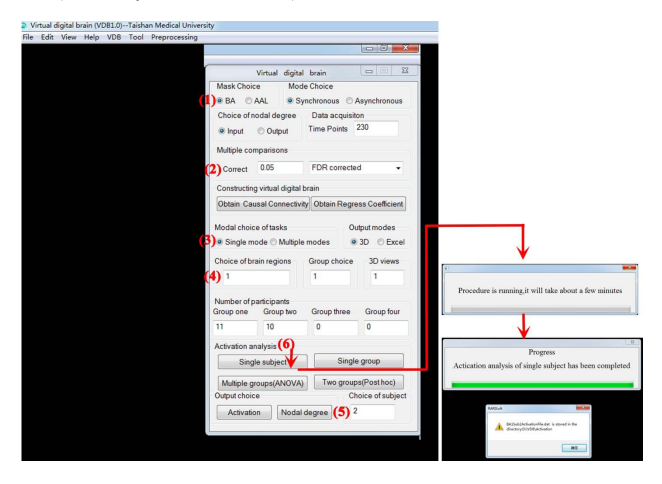
Figure 12. Task design and waveform display

6. Task-based activation analysis

6.1. Activation analysis of single subject

1. Activation of single subject. Select "BA" or "AAL" in the mask choice, corrected parameter in the multiple comparisons (correction is for the results of activations), "Single mode or Multiple modes" in the modal choice of tasks, and then fill the index of brain region (if need to fill multiple brain regions, the format is as follows: brain region A, B, C, D,…. For example, 1, 2, 3. The string "1, 2, 3" indicates that the task stimulating signal will be exerted to 3 brain regions, and the indexes of these brain regions are 1, 2 and 3. When select "Multiple modes" in the modal choice of tasks, the format is as follows: brain region A, B; C, D; E, F. For example, 61, 62; 1, 2; 9, 10. The string "61, 62" indicates that the first task stimulating signal will be exerted to brain regions

61 and 62; the string "1, 2" indicates that the second task stimulating signal will be exerted to brain regions 1 and 2; the string "9, 10" indicates that the third task stimulating signal will be exerted to brain regions 9 and 10) in the editor control "Choice of brain regions". In addition, the number of participants must be filled in these editor controls (Group one to four). Fill the index of subject in the editor control "Choice of subject". Finally, click on the button "Single subject" and the procedure starts to run (Figure 13). The result of analysis is named as "BAxSubxActivationFile.dat or AALxSubxActivationFile.dat" and is automatically stored in the folder "Activation" (the directory: D: \\VDB\\Activation).



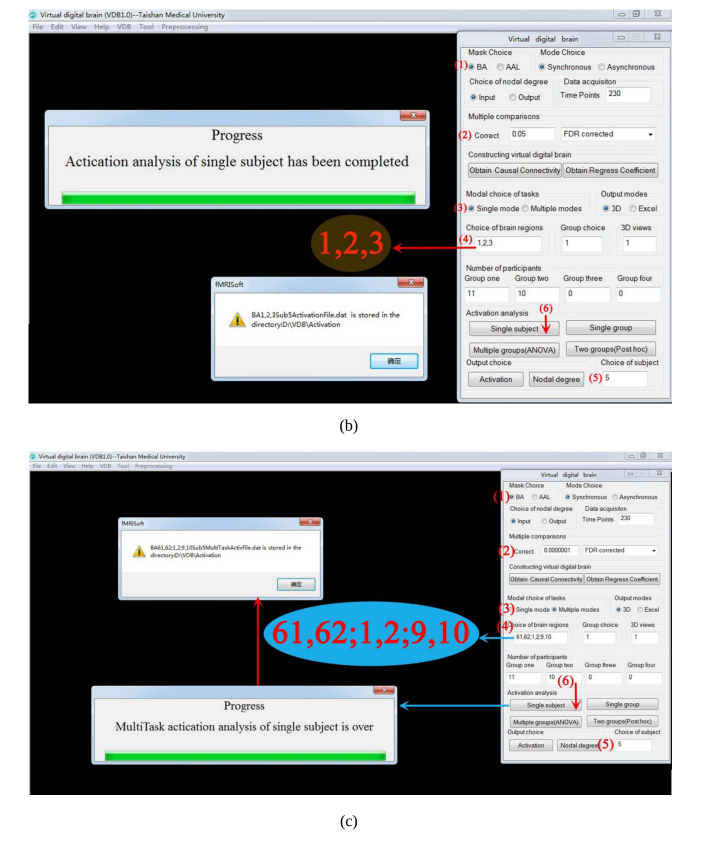


Figure 13. Activation analysis of single subject

generated in previous step. Click the button "Open" and then the result is displayed in an excel table (Figure 14). In this table, the numbers in the column "CH" indicate the index of activated brain regions, and the column "CG" is the strength of activation. Positive real numbers indicate positive strengths of activation. On the contrary, negative real numbers indicate negative strengths of activation. Positive real numbers in every row indicate the strengths of synchronous causal connectivity among activated brain regions, and negative real numbers indicate the strengths of asynchronous causal connectivity. The real numbers of every row indicate the strengths of output causal connectivity corresponding to every node, and the real numbers of every column indicate the strengths of input causal connectivity corresponding to every node. It is worth noting that the values in the excel table are actual values of activation strengths and causal connectivity.

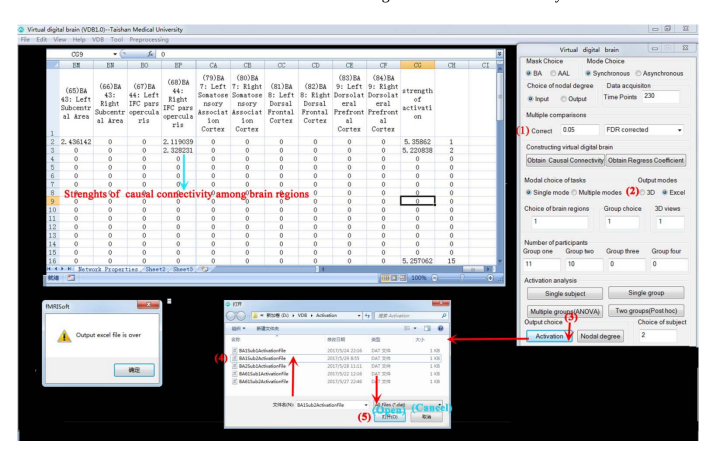


Figure 14. Activation result of single subject

(2) Output the activation results through 3D visualization. Select "BA" or "AAL" in the mask choice, "Synchronous" or "Asynchronous" in the mode choice, corrected parameter in the multiple comparisons (correction is for the strengths of causal connectivity among activated brain regions), and "3D" in the output modes, and then fill the index of 3D view in the editor control "3D views" (the index "1" indicates the superior

view, "2" indicates the inferior view, "3" indicates the left view, and "4" indicates the right view. In these views, every color sphere indicates an activated brain region. The size of the sphere indicates the strength of the brain region activation (It is worth noting that the sizes are not responding to actual strengths of brain region activations, these sizes are responding to standardized strengths of activations. The actual strengths of activations can be displayed through the excel table), every bar among spheres indicates the casual connectivity among these brain regions, and the diameter of the bar denotes the strength of the interregional causality connectivity (It is worth noting that the diameters are not responding to actual strengths of interregional causality connectivity, these diameters are responding to standardized strengths of interregional causality connectivity. The actual strengths of causality connectivity can be obtained through the excel table). The gold bar denotes the synchronous causality connectivity, and the light blue bar denotes the asynchronous causality connectivity. The direction of the arrow denotes the direction of causality connectivity. Especially, when the index of 3D view is bigger than 4, we display activated brain regions by using color areas. Different colors indicate distinct strengths of brain region activations. Blue is corresponding to weaker activated strength and yellow indicates stronger activated strength. Color changes of the color bar are corresponding to changes of activated strengths of brain regions. The index "5" of 3D view indicates that the activated brain regions are projected to this view from superior to inferior; "6" indicates that the activated brain regions are projected to this view from left to right; "7" indicates that the activated brain regions in the superior cerebral hemisphere are projected to this view from inferior to superior; "8" indicates that the activated brain regions in the inferior cerebral hemisphere are projected to this view from superior to inferior; "9" indicates that the activated brain regions in the left cerebral hemisphere are projected to this view from right to left; "10" indicates that the activated brain regions in the right cerebral hemisphere are projected to this view from left to right). Click on an opened dialog box. These files have been generated in previous step. Click on the button "Open" and then the result is showed in the left of client area (Figure 15).

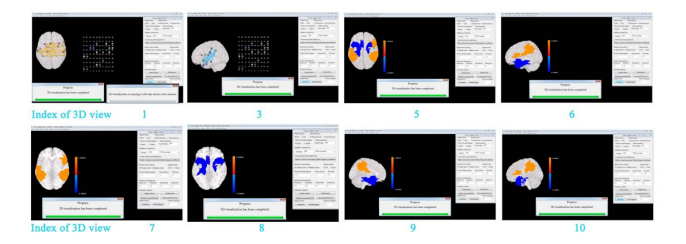


Figure 15. 3D visualization of brain region activations of single subject

6.2. Activation analysis of single group

1. Activation analysis of single group. Select "BA" or "AAL" in the mask choice, corrected parameter in the multiple comparisons (correction is for the results of activations), "Single mode or Multiple modes" in the modal choice of tasks, and then fill the index of brain region (if need to fill multiple brain regions, the format is as follows: brain regions A, B, C, D,… For example, 61, 62. The string "61, 62" indicates that the task stimulating signal will be exerted to 2 brain regions, and the indexes of these brain regions are 61 and 62) in the editor control "Choice of brain regions". In addition, the number of participants must be filled in these editor controls (Group one to four). Fill the index of group in the editor control "Group choice". Finally, click on the button "Single group" and the procedure starts to run (Figure 16). The result of analysis is named as "BAxGrpxActivationFile.dat" and is automatically stored in the folder "Activation" (the directory: D: \\VDB\\Activation).

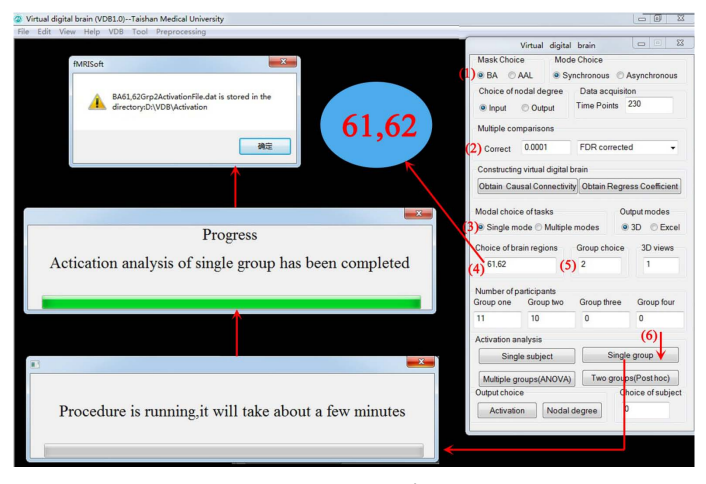


Figure 16. Activation analysis of single group

2. Activation results of single group. (1) Output the activation results through the excel table. Select "BA" or "AAL" in the mask choice, corrected parameter in the multiple comparisons (correction is for the strengths of causal connectivity among activated brain regions), "Excel" in the "Output modes", and then click on the button "Activation". Select one file in the folder "Activation" (the directory: D: \\VDB\\Activation\) through an opened dialog box. These files have been generated in previous step. Click on the button "Ok" and then the result is displayed in an excel table (Figure 17). In this table, the numbers in the column "CH" indicate the index of activated brain regions, and the column "CG" is the strength of activation. Positive real numbers indicate positive strengths of activation. On the contrary, negative real numbers indicate negative strengths of activation. Positive real numbers in every row indicate the strengths of synchronous causal connectivity among activated brain regions, and negative real numbers indicate the strengths of asynchronous causal connectivity. The real numbers of every row indicate the strengths of input causal connectivity corresponding to every node, and the real numbers of every column indicate the strengths of input causal connectivity corresponding to every

node (see also Figure 14 for details). In addition, the number of participants must be filled in these editor controls (Group one to four).

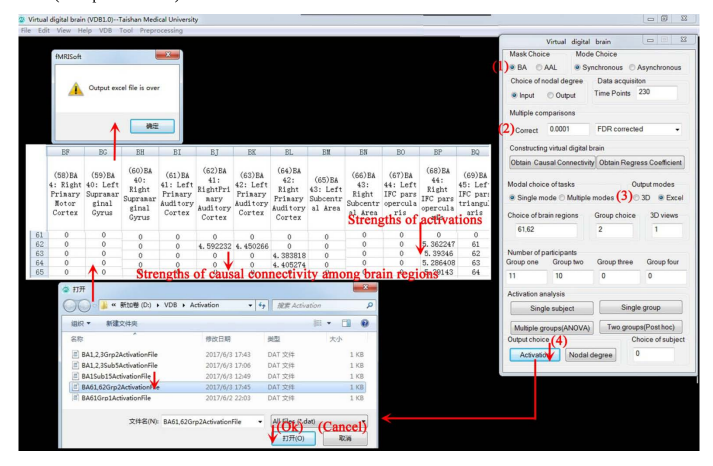


Figure 17. Activation result of single group

(2) Output the activation results through 3D visualization. Select "BA" or "AAL" in the mask choice, "Synchronous" or "Asynchronous" in the mode choice, corrected parameter in the multiple comparisons (correction is for the strengths of causal connectivity among activated brain regions), and "3D" in the output modes, and then fill the index of 3D view in the editor control "3D views" (the index "1" indicates the superior view, "2" indicates the inferior view, "3" indicates the left view, and "4" indicates the right view. In these views, every color sphere indicates an activated brain region. The size of the sphere indicates the strength of the brain region activation, every bar among spheres indicates the casual connectivity among these brain regions, and the diameter of the bar denotes the strength of the interregional causality connectivity. The gold bar denotes the synchronous causality connectivity, and the light blue bar denotes the asynchronous causality connectivity. The direction of the arrow denotes the direction of causality connectivity. Especially, when the index of 3D view is bigger than 4, we display activated brain regions by using color areas. Different colors indicate distinct

strengths of brain region activations. Blue is corresponding to weaker activated strength and yellow indicates stronger activated strength. Color changes of the color bar are corresponding to changes of activated strengths of brain regions. The index "5" of 3D view indicates that the activated brain regions are projected to this view from superior to inferior; "6" indicates that the activated brain regions are projected to this view from left to right; "7" indicates that the activated brain regions in the superior cerebral hemisphere are projected to this view from inferior to superior; "8" indicates that the activated brain regions in the inferior cerebral hemisphere are projected to this view from superior to inferior; "9" indicates that the activated brain regions in the left cerebral hemisphere are projected to this view from right to left; "10" indicates that the activated brain regions in the right cerebral hemisphere are projected to this view from left to right). Click on the button "Activation". Select one file in the folder "Activation" (the directory: D: \\\VDB\\Activation\) through an opened dialog box. These files have been generated in previous step. Click on the button "Open" and then the result is showed in the left of client area (Figure 18). In addition, the number of participants must be filled in these editor controls (Group one to four).

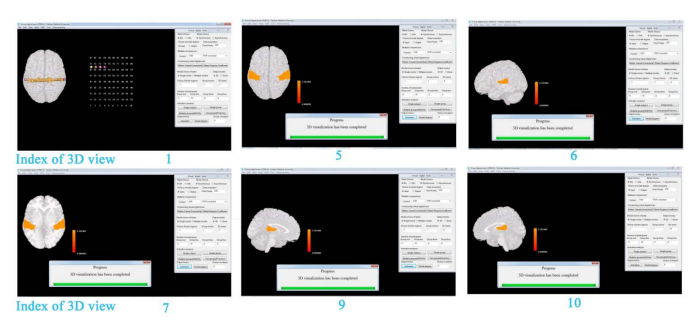


Figure 18. 3D visualization of brain region activations of single group

6.3. Activation analysis of two groups

1. Activation analysis of two groups. Select "BA" or "AAL" in the mask choice, corrected parameter in the multiple comparisons (correction is for the results of activations), "Single mode or Multiple modes" in the

modal choice of tasks, and then fill the index of brain region (if multiple brain regions need to be filled in the editor control, then the format is as follows: brain regions A, B, C, D,... For example, 61, 62. The string "61, 62" indicates that the task stimulating signal will be exerted to 2 brain regions, and the indexes of these brain regions are 61 and 62) in the editor control "Choice of brain regions". In addition, the number of participants must be filled in these editor controls (Group one to four). Fill the indexes of groups in the editor control "Group choice" (The format is as follows: Groups A, B. For example, 1, 2. The string "1, 2" indicates that the groups 1 and 2 participate in the test. Two-sample t-test will be implemented to compare activation results of group 1 to 2.). Finally, click on the button "Two groups" and the procedure starts to run (Figure 19). The result of analysis is named as "BAxTrpxActivationFile.dat or AALxTrpxActivationFile.dat" and is automatically stored in the folder "Activation" (the directory: D: \\\VDB\\Activation\).

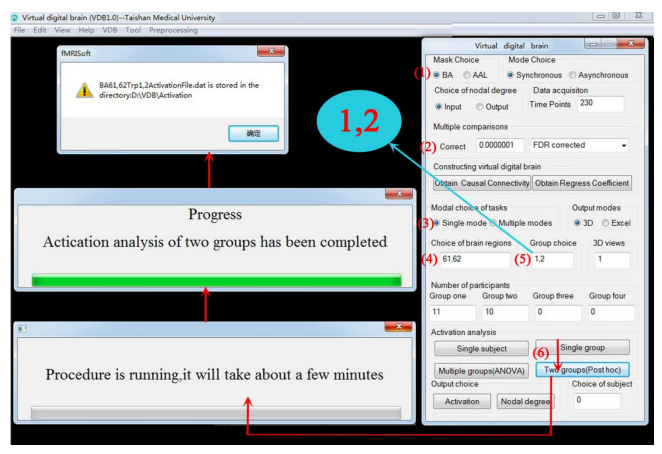


Figure 19. Activation analysis of two groups

2. Activation results of two groups. (1) Output the activation results through the excel table. Select "BA" or "AAL" in the mask choice, corrected parameter in the multiple comparisons (correction is for the strengths of causal connectivity among activated brain regions), "Excel" in the "Output modes", and then click on the

button "Activation". Select one file in the folder "Activation" (the directory: D: \\VDB\\Activation\) through an opened dialog box. These files have been generated in previous step. Click on the button "Ok" and then the result is displayed in an excel table (Figure 20). In this table, the numbers in the column "CH" indicate the index of activated brain regions, and the column "CG" is the strength of activation. Positive real numbers indicate positive strengths of activation. On the contrary, negative real numbers indicate negative strengths of activation. Positive real numbers in every row indicate the strengths of synchronous causal connectivity among activated brain regions, and negative real numbers indicate the strengths of asynchronous causal connectivity. The real numbers of every row indicate the strengths of output causal connectivity corresponding to every node, and the real numbers of every column indicate the strengths of input causal connectivity corresponding to every node (see also Figure 14 and 17for details). In addition, the number of participants must be filled in these editor controls (Group one to four).

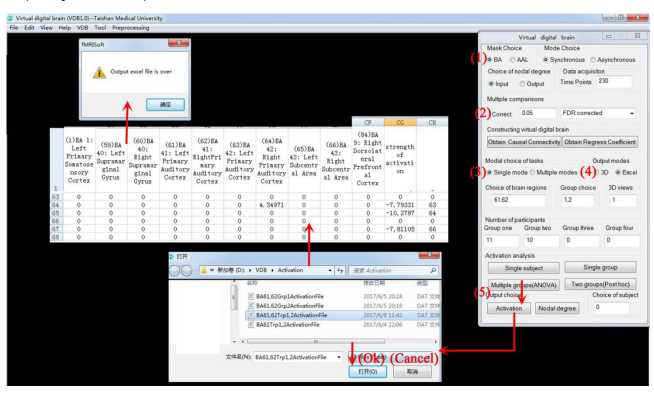


Figure 20. Activation result of two groups

(2) Output the activation results through 3D visualization. Select "BA" or "AAL" in the mask choice, "Synchronous" or "Asynchronous" in the mode choice, corrected parameter in the multiple comparisons (correction is for the strengths of causal connectivity among activated brain regions), and "3D" in the output modes, and then fill the index of 3D view in the editor control "3D views" (the index "1" indicates the superior

view, "2" indicates the inferior view, "3" indicates the left view, and "4" indicates the right view. In these views, every color sphere indicates an activated brain region. The size of the sphere indicates the strength of the brain region activation, every bar among spheres indicates the casual connectivity among these brain regions, and the diameter of the bar denotes the strength of the interregional causality connectivity. The gold bar denotes the synchronous causality connectivity, and the light blue bar denotes the asynchronous causality connectivity. The direction of the arrow denotes the direction of causality connectivity. Especially, when the index of 3D view is bigger than 4, we display activated brain regions by using color areas. Different colors indicate distinct strengths of brain region activations. Blue is corresponding to weaker activation strength and yellow indicates stronger activation strength. Color changes of the color bar are corresponding to changes of activated strengths of brain regions. The index "5" of 3D view indicates that the activated brain regions are projected to this view from superior to inferior; "6" indicates that the activated brain regions are projected to this view from left to right; "7" indicates that the activated brain regions in the superior cerebral hemisphere are projected to this view from inferior to superior; "8" indicates that the activated brain regions in the inferior cerebral hemisphere are projected to this view from superior to inferior; "9" indicates that the activated brain regions in the left cerebral hemisphere are projected to this view from right to left; "10" indicates that the activated brain regions in the right cerebral hemisphere are projected to this view from left to right). Click on the button "Activation". Select one file in the folder "Activation" (the directory: D: \\VDB\\Activation) through an opened dialog box. These files have been generated in previous step. Click on the button "Open" and then the result is showed in the left of client area (Figure 21). In addition, the number of participants must be filled in these editor controls (Group one to four).

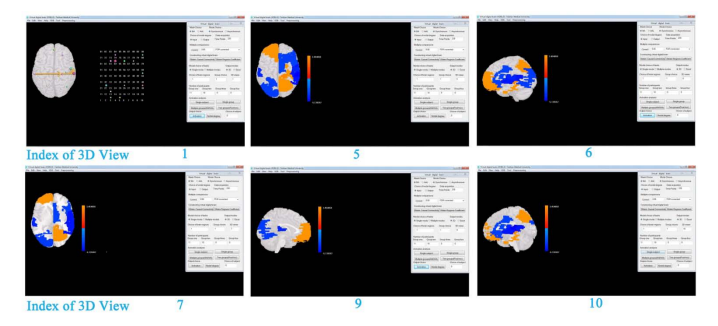


Figure 21. 3D visualization of brain region activations of two groups

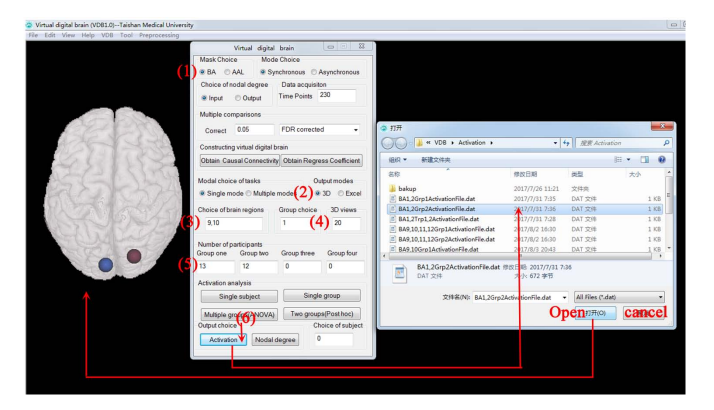


Figure 22. 3D visualization of selected brain regions

6.4. Activation visualization

1. Firstly, click on the menu VDB, and then select the "Visualization" in the drop-down menu. As shown in Figure 23.

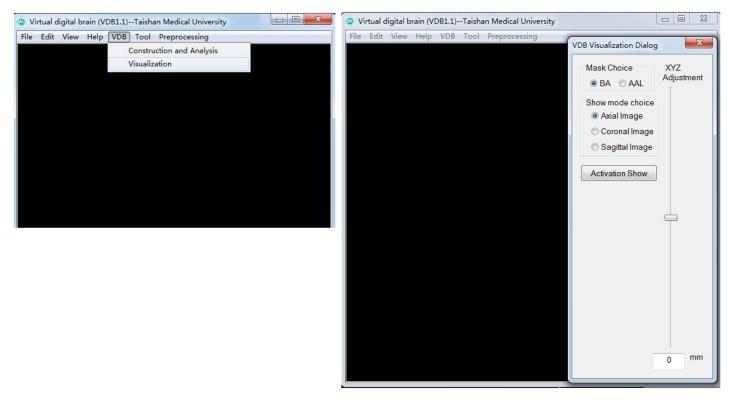


Figure 23. Diagram of opening visualization dialog.

2. Select "BA" or "AAL" in the mask choice, "Axial Image" or "Coronal Image" or "Sagittal Image" in the show mode choice, and then click on the button "Activation Show" and open the destination folder "Activation". Select a file in the opened folder and click on the button "Open", and then the axial image is displayed in the client area, as shown in figure 24. The color bar indicates the strengths of activation, the negative value denotes the strength of negative activation, and the positive value denotes the strength of positive activation. The numbers beside the activated regions indicate the index of Brodmann Area (BA), the "L" denotes the left, and the "R" denotes the ight.

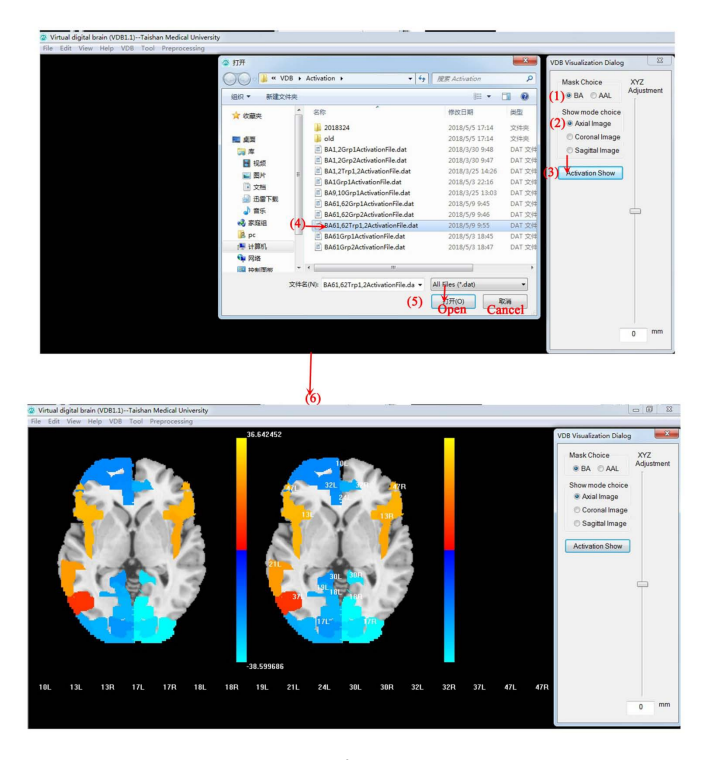
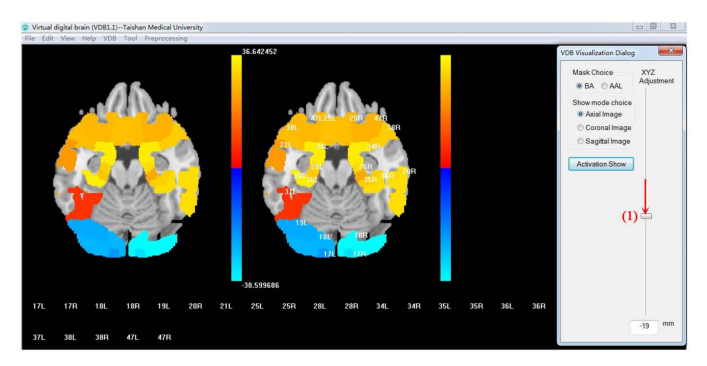


Figure 24. Diagram of the axial image display

3. Remove the slider and alter the location of the activated images (for example, the axial image), as shown in figure 25.



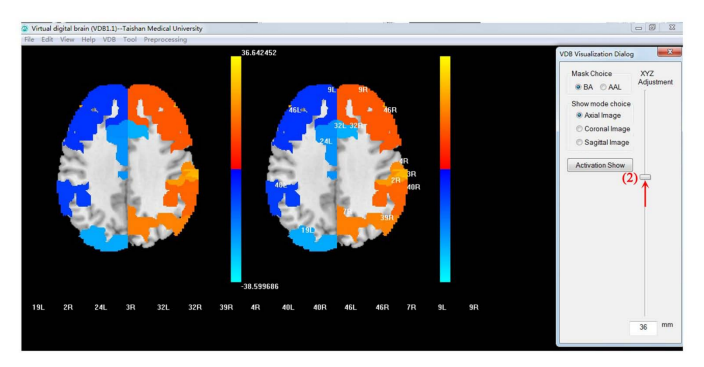


Figure 25. Diagram of the axial image change

7. The shortest path

1. The shortest path between brain regions. Select "BA" or "AAL" in the mask choice, corrected parameter in the multiple comparisons (correction is for the strengths of causal connectivity among activated brain regions), "3D" in the output modes, and then fill "A,B" in the editor control "Brain regions of SP"("A,B" denotes the direction of the shortest path is from the brain region A to B). Click on the button "Shortest Path (SP)" and

select the activation file in the folder "Activation" through the opened dialog. Click on the button "Open" and obtain the 3D view of the shortest path between brain regions A and B. The result is showed in figure 26.

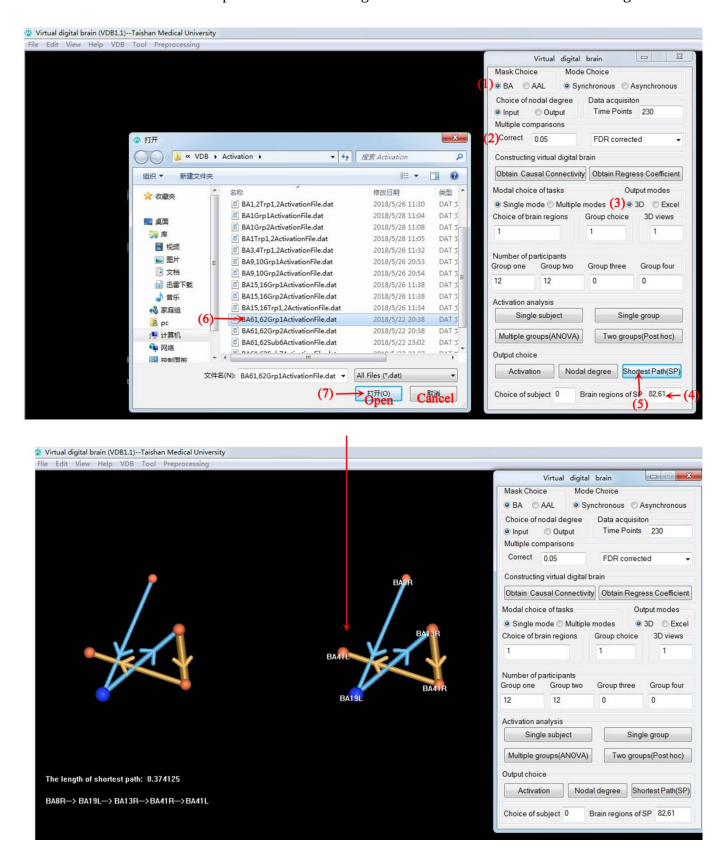


Figure 26. 3D view of the shortest path

In figure 26, the red sphere denotes positive activation, and the blue sphere denotes negative activation.

causality connectivity, and the light blue bar denotes the asynchronous causality connectivity. The direction of the arrow denotes the direction of causality connectivity, and the diameter of the bar denotes the strength of the interregional causality connectivity.

2. The shortest path lengths of all subjects. Select "BA" or "AAL" in the mask choice, corrected parameter in the multiple comparisons, "Single mode" or "Multiple modes" in the modal choice of tasks, "Excel" in the output modes, and then fill the codes of brain regions in the editor controls "Choice of brain regions" and "Brain regions of SP"(such as "61,62", "1,61"). In addition, the number of participants must be filled in these editor controls (Group one to four). Click on the button "Shortest path(SP)" and obtain the shortest path lengths of all subjects. The result is showed in an excel file (Figure 27). Here, the codes of brain regions in the editor controls "Choice of brain regions" denote those brain regions which the virtual task signal will be exerted to.

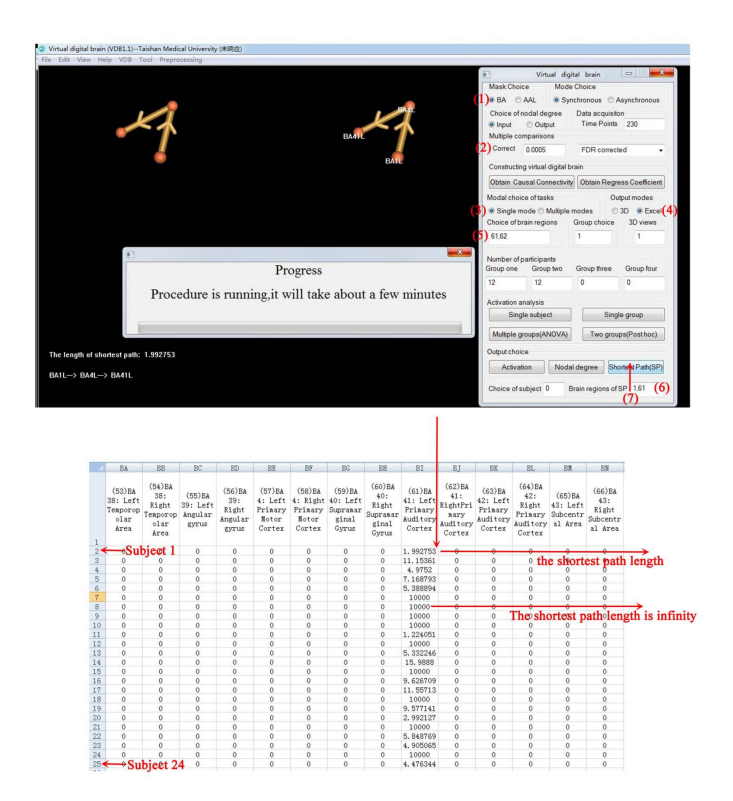


Figure 27. The shortest path lengths of all subjects