```
clear all;
close all;
cd ('/Users/Aurina/Documents/Genetics_connectome/Gen_Cog/Data/
Connectomes/WithoutACT/'); %choose the folder
Parcellation = {'aparcaseg'}; % choose parcellation
Tract = {'iFOD2_'}; % choose tract method
Name = strcat(Tract{1}, Parcellation{1});
load (sprintf('%s.mat', Name));
Density = zeros(size(density,2),1);
for i=1:size(density,2)
    figure; imagesc(log(density{i})); title(sprintf('Subject %d',
 sub_id{i}));
    Strength = sum(density{i});
    figure; histogram(Strength, 20); title(sprintf('Strength Subject
 %d', sub_id{i}));
end
% clear all;
% close all;
% cd ('/Users/Aurina/Documents/Genetics_connectome/Gen_Cog/Data/
Connectomes/WithACT/'); %choose the folder
% Parcellation = {'aparcaseg'}; % choose parcellation
% Tract = {'iFOD2_'}; % choose tract method
% Name = strcat(Tract{1}, Parcellation{1});
% load (sprintf('%s.mat', Name));
% Density = zeros(size(density,2),1);
% for i=1:size(density,2)
      figure; imagesc(log(density{i})); title(sprintf('Subject %d',
sub_id{i}));
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      Strength = sum(density{i});
0
      figure; histogram(Strength, 20); title(sprintf('Strength Subject
%d', sub id{i}));
% end
```

















































































