Protein Structure Networks with Bio3D

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Overview

The Bio3D package (version 2.1 and above) contains functionality for the creation and analysis of protein structure networks. Here we will introduce this functionality by building and analyzing networks obtained from atomic correlation data derived from normal mode analysis (NMA) and molecular dynamics (MD) simulation data (**Figure 1**).

Basic network generation and visualization

Bio3D contains extensive functionality for NMA and MD analysis. An introduction to each of these areas is detailed in separate NMA and MD vignettes available online. In this section we will focus on correlation network analysis derived from NMA, and in subsequent sections MD.

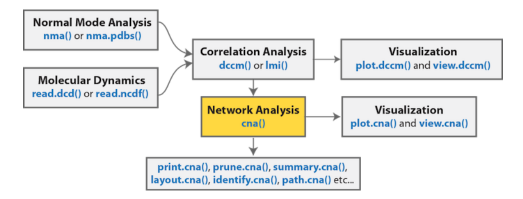


Figure 1: Overview of a typical Bio3D network analysis protocol with key functions detailed in blue. Normal mode and/or molecular dynamics input first undergoes correlation analysis (with the functions dccm() or lmi()). The output from these functions is typically a matrix of residue-by-residue cross-correlations. Correlation network analysis can then be performed (with the cna() function) to generate a correlation network with residues as nodes that are linked by weighted edges that are proportional to their degree of correlated motion. The cna() function also performs a community clustering analysis to identify communities of highly correlated residues. The full residue network and coarse-grained community network can be visualized and analyzed further. This methodology can provide valuable insight not readily available from the measures of correlation alone.

Network generation from normal mode analysis data

The code snippet below first loads the Bio3D package, then reads an example PDB structure from the RCSB online database. We then follow the flow in Figure 1 by performing NMA, then dynamic cross-correlation analysis, and finally network analysis with the **cna()** function.

```
library(bio3d)
pdbfile <- system.file("examples/hivp.pdb", package = "bio3d")
pdb <- read.pdb( pdbfile )
modes <- nma(pdb)
cij <- dccm(modes)
net <- cna(cij, cutoff.cij=0.3)</pre>
```

To enable quick examination of the network analysis results we provide **cna** specific variants of the **print()**, **summary()** and **plot()** functions, as demonstrated below:

```
print(net)
```

```
##
## Call:
     cna.dccm(cij = cij, cutoff.cij = 0.3)
##
##
## Structure:
   - NETWORK NODES#:
##
                         198
                                 EDGES#: 752
##
     COMMUNITY NODES#: 8
                             EDGES#: 11
##
    + attr: network, communities, community.network,
##
##
           community.cij, cij, call
```

```
# Summary information as a table
x <- summary(net)</pre>
##
    id size
                                  members
##
                  c(1:8, 89:107, 190:198)
     1
         36
                    c(9:11, 22:32, 83:88)
##
     2
         20
                          c(12:21, 60:73)
##
     3
         24
##
     4
         28
                  c(33:46, 55:59, 74:82)
##
     5
         16
                        c(47:54, 146:153)
##
         17
                c(108, 122:130, 183:189)
     6
                      c(109:121, 159:173)
     7
##
         28
     8
         29 c(131:145, 154:158, 174:182)
##
# Other useful things are returned from **summary.cna()**
attributes(x)
## $names
## [1] "id"
                            "members" "tbl"
                  "size"
# List the residues belonging to the 5th community
x$members[5]
## $\5\
                                 54 146 147 148 149 150 151 152 153
##
    47
        48
            49
                50
                    51
                         52
                             53
                                 54 146 147 148 149 150 151 152 153
                    51
                         52 53
# Plot both the 'full' all-residue network and simplified community network
par(mfcol = c(1, 2), mar = c(0, 0, 0, 0))
plot(net, pdb, full = TRUE)
plot(net, pdb)
```

Tip: The raw correlation matrix obtained from normal modes (or indeed molecular dynamics analysis) very likely has a non-zero value for each element. We need to exclude some of the weakest values (close to zero) to avoid building a network with $N^2 - N$ edges, which would considerably slow down further analysis. The *cutoff.cij** argument in the **cna()** function specifies the lower boundary for the correlation values (default is 0.4), all the values lower than this cutoff value will be ignored when defining network edges.

Note that the above code plots both a *full* all-residue network as well as a more coarse grained community network - see **Figure 2**. In this example, and by default, the Girvan-Newman clustering method was used to map the full network into communities of highly intra-connected but loosely inter-connected nodes that were in turn used to derive the simplified network displayed in **Figure 2**. We will discuss community detection in further detail below and simply highlight here that a comparison of the distribution and connectivity of communities between similar biomolecular systems (e.g. in the presence and absence of a binding partner or mutation) has the potential to highlight differences in coupled motions that may be indicative of potential allosteric pathways. This approach has previously been applied to investigate allostery in tRNA-protein complexes, kinesin motor domains, G-proteins, thrombin, and other systems [1-4].

Another useful function for the inspection of community structure is the **view.cna()** function, which permits interactive network visualization in the VMD molecular graphics program (see **Figure 3**).

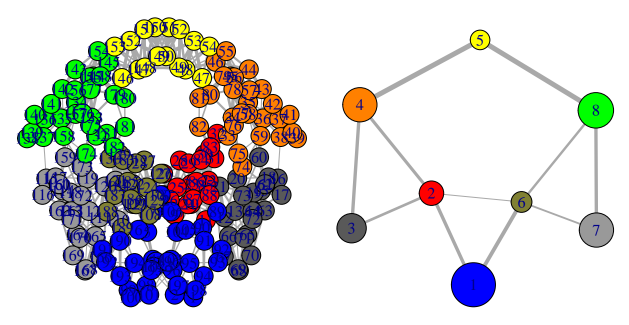


Figure 2: Full all-residue network and simplified community network

```
# View the network in VMD (see Figure 3)
view.cna(net, pdb, launch = TRUE)
```

One of the main advantages of correlation network analysis is that it often facilitates the interpretation of correlated motions that can be challenging to rationalize from the output of a dynamic correlation map alone (e.g. the output of the dccm() function). For example, the identified communities of highly intra-connected residues can be tied into the visualization of the raw correlation data (**Figure 4**) and cross referenced to the network visualizations presented in **Figures 2** and **3**:

```
plot.dccm(cij, margin.segments = net$communities$membership)
```

To more fully appreciate the results of network analysis one should become somewhat familiar with the structure of the object returned from the **cna()** function. This will allow you to generate customized visualizations beyond those in **Figures 2** to **4** and also enable you to more fully examine the effect of different parameters on network structure and topology.

The structure of cna() network objects

The output of the **cna()** function is a list object of class cna with both network and community structure attributes. These are summarized at the bottom of the **print(net)** results above and can be explicitly listed with the **attributes(cna)** command:

```
attributes(net)
```

```
## $names
## [1] "network" "communities" "community.network"
## [4] "community.cij" "cij" "call"
##
## $class
## [1] "cna"
```

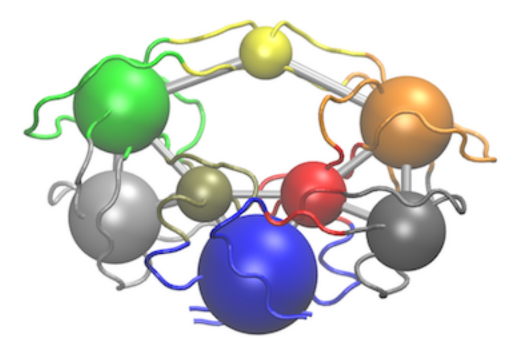


Figure 3: Example of **view.cna()** output. Here within VMD we have set display style to 'Tube' and color to 'Color by Chain' to see corresponding regions of structure colored by community along with the community protein structure network

Residue Cross Correlation

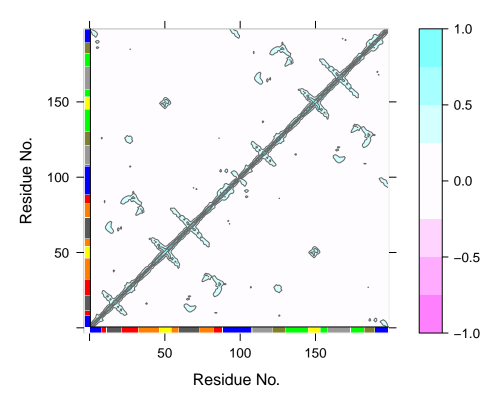


Figure 4: DCCM plot. Note that unique colors are assigned to communities and that these correspond to those annotated in all previous Figures and those used by VMD.

Here we introduce each of these components in turn:

- **\$network**: The *full* protein structure network typically with a node per residue and connecting edges weighted by their corresponding correlation values (i.e. input cij values optionally filtered by the user defined 'cutoff.cij'). This filtered cij matrix is also returned in the output as **\$cij**, as listed below.
- **\$communities**: A community clustering object detailing the results of clustering the full **\$network**. This is itself a list object with a number of attributes (see attributes(net\$communities) for details). Notable attributes include **\$communities\$membership**, **\$communities\$modularity** and **\$communities\$algorithm**. These components detail the nodes (typically residues) belonging to each community; the modularity (which represents the difference in probability of intra- and intercommunity connections for a given network division); and the clustering algorithm used respectively.
- \$cij: The filtered correlation matrix used to build the full \$network.
- **\$community.network**: A coarse-grained community network object with the number of nodes equal to the number of communities present in **\$communities** (i.e. the community clustering of the full network) and edges based on the inter-community coupling as determined by the input 'collapse.method' (by default this is the maximum coupling between the original nodes of the respective communities).
- **\$community.cij**: A cij matrix obtained by applying a "collapse.method" on \$cij. The rows and columns match the number of communities in \$communities. The values are based on the cij couplings of inter-communities residue. (collapse.method available are: max, mean, trimmed and median).

Additional network annotation

Both the network and network attributes are igraph package network objects [5] and contain additional node (or vertex) and edge annotations that can be accessed with the V() and E() functions in the following way:

```
# Basic data access follows the igraph package scheme
head(V(net$network)$color)

## [1] "#0000FF" "#0000FF" "#0000FF" "#0000FF" "#0000FF"

V(net$community.network)$size

## [1] 36 20 24 28 16 17 28 29

E(net$community.network)$weight

## [1] 0.4960 0.4580 0.5620 0.5190 0.7274 0.4989 0.3649 0.3649 0.6414 0.5868

## [11] 0.4990

V(net$community.network)$name
```

```
## [1] "1" "2" "3" "4" "5" "6" "7" "8"
```

In addition to various Bio3D functions, the full set of extensive igraph package features can be used to further analyze these network objects.

The community clustering procedure

The **\$communities** attribute returned by the **cna()** function provides details of the community clustering procedure. This procedure aims to split the full network into highly correlated local substructures (referred to as communities) within which the network connections are dense but between which they are sparser. A number of community detection algorithms can be used to solve the graph-partitioning problem. The default Girvan-Newman edge-betweenness approach is a divisive algorithm that is based on the use of the edge betweenness as a partitioning criterion. "Betweenness" is calculated by finding the shortest path(s) between a pair of vertexes and scoring each of the edges on this/these path(s) with the inverse value of the number of shortest paths. (So if there was only one path of the shortest length, each edge on it would score 1 and if there were 10 paths of that length, each edge would score 1/10.) This is done for every pair of vertexes. In this way each edge accumulates a "betweenness" score for the whole network. The network is separated into clusters by removing the edge with the highest "betweenness", then recalculating betweenness and repeating. The method is fully described in (Girvan and Newman, 2002 PNAS) [6].

A useful quantity used to measure the quality of a given community partition is the **modularity** (detailed in **\$communities\$modularity** component of **cna()** output). The modularity represents the difference in probability of intra- and inter-community connections for a given network division. Modularity values fall in the range from 0 to 1, with higher values indicating higher quality of the community structure. The optimum community structures obtained for typical correlation networks fall in the 0.4 to 0.7 range.

```
head(net$communities$modularity)

## [1] -5.770e-03 -4.431e-03 -3.617e-03 -8.542e-04 2.043e-05 2.077e-03

max(net$communities$modularity)
```

```
## [1] 0.7381
```

The **community.tree()** function can be used to reconstruct the community membership vector for each clustering step. Here we plot modularity vs number of communities.

```
tree <- community.tree(net, rescale=TRUE)
plot( tree$num.of.comms, tree$modularity )</pre>
```

Below we inspect the maximum modularity value partitioning and the number of communities (k) at the point of maximum modularity. Based on a quick inspection of the clustering dendrogram we explore a new membership partitioning (at k=3). Again this example is purely for demonstrating some of the available functionality within the package and we encourage motivated readers to explore the true structure within their data by examining multiple structures, input parameters and clustering procedures.

```
max.mod.ind <- which.max(tree$modularity)
tree$num.of.comms[ max.mod.ind ]</pre>
```

[1] 8

```
# Membership vector at this partition point
tree$tree[max.mod.ind,]
```

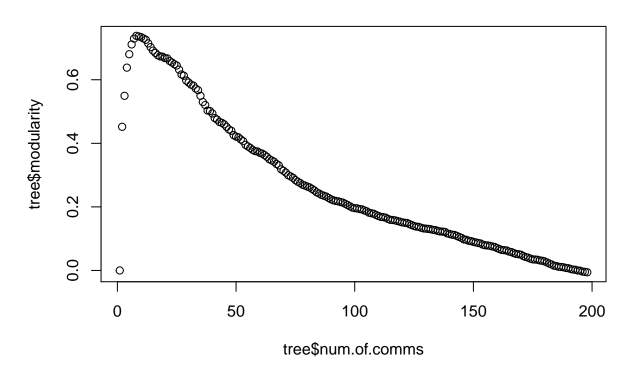


Figure 5: Network modularity upon removing edges during the betweenness clustering procedure for community annotation

```
 \hbox{\tt \#\#} \hbox{\tt [141]} \hbox{\tt 888885555555558888877777777777777777788} 
# Should be the same as that contained in the original CNA network object
all( net$communities$membership == tree$tree[max.mod.ind,] )
## [1] TRUE
# Inspect the clustering dendrogram, to match colors use 'colors=vmd.colors()[memb.k3]'
h <- as.hclust(net$communities)</pre>
hclustplot(h, k=3)
# Produce a new k=3 community network
net.3 <- network.amendment(net, memb.k3)</pre>
plot(net.3, pdb)
## Obtaining layout from PDB structure
# View the network in VMD (see Figure 8)
view.cna(net.3, trim.pdb(pdb, atom.select(pdb,"calpha")), launch=TRUE )
```

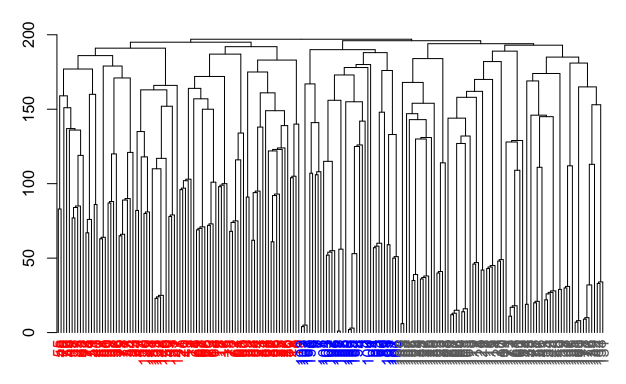


Figure 6: Viewing community partitioning progress as a dendrogram with colored communities

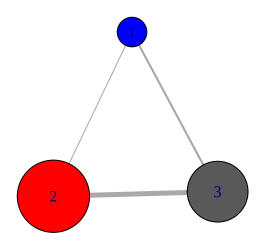


Figure 7: Example k=3 community network

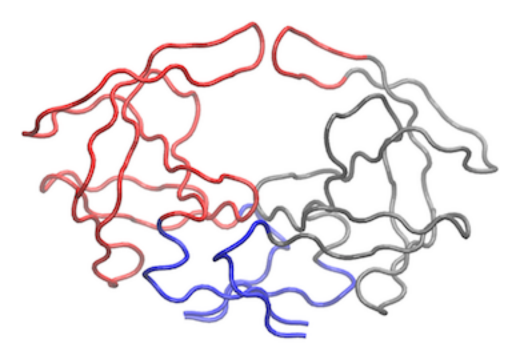


Figure 8: Example of **view.cna()** output for modified k=3 community network. Here within VMD we have set display style to 'Tube' and color to 'Color by Chain' as well as removing community spheres by deleting graphical objects.

Network generation from molecular dynamics data

For this example section we apply correlation network analysis to a short molecular dynamics trajectory of Human Immunodeficiency Virus aspartic protease (HIVpr). This trajectory is included with the Bio3D package and stored in CHARMM/NAMD DCD format. To reduce file size and speed up example execution, non C-alpha atoms have been previously removed and the frames superposed. Note that typically an input trajectory would require superposition with the fit.xyz() function prior to correlation analysis with the dccm() function (see the Trajectory Analysis vignette for full details).

The code snippet below first sets the file paths for the example HIVpr starting structure (pdbfile) and trajectory data (dcdfile), then reads these files (producing the objects trj and pdb).

Once we have the superposed trajectory frames we can asses the extent to which the atomic fluctuations of individual residues (in this very short example simulation) are correlated with one another and build a network from this data:

```
cij <- dccm(trj)
net <- cna(cij)
plot(net, pdb)</pre>
```

Obtaining layout from PDB structure

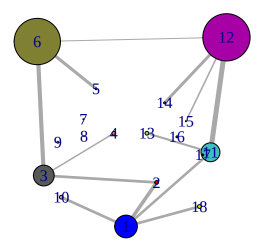


Figure 9: Network analysis of MD data

Tip: Due to the intrinsic noise of the correlations calculated on atomic fluctuations from short MD simulations, we suggest running longer simulations and/or multiple replica simulations from which a consensus correlation matrix and consensus network can be generated and compared to results obtained from individual simulations. For example, please read the following works: Scarabelli G and Grant BJ, Kinesin-5 allosteric inhibitors uncouple the dynamics of nucleotide, microtubule and neck-linker binding sites, Biophys J. in press and Yao X and Grant BJ XXXX submitted.

Note that the dccm() function can be slow to run on very large trajectories. In such cases you may want to use multiple CPU cores for the calculation by setting the 'ncore' option to an appropriate value for your computer - see help(dccm) for details.

```
# View the correlations in pymol
view.dccm(cij, pdb, launch = TRUE)

# View the structure mapped network
view.cna(net)
```

Using a contact map filter:

In the original Luthy-Shulten and co-workers approach to correlation network analysis [1] edges were only included between "in contact" nodes. Contacting nodes were defined as those having any heavy atom within 4.5 Å for greater than 75% of the simulation. The weight of these contact edges was then taken from the

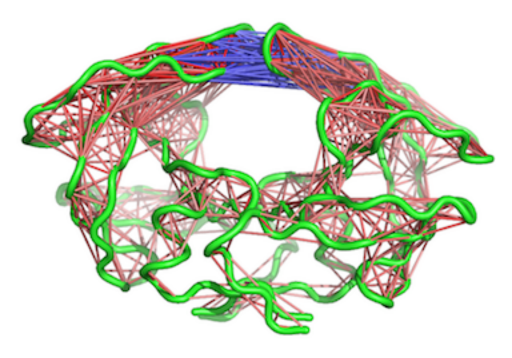


Figure 10: Example of view.dccm() output for an MD trajectory

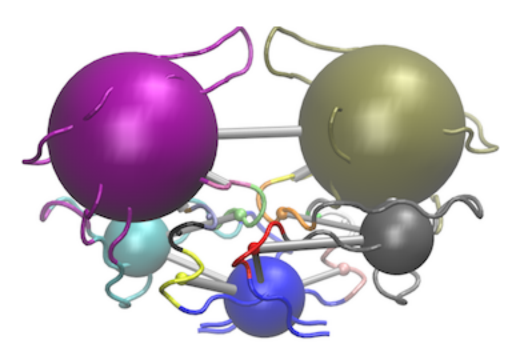


Figure 11: The view.cna() output for the HIVpr example MD trajectory

cross-correlation data. This approach effectively ignores long range correlations between residues that are not in physical contact for the majority of the trajectory. To replicate this approach one can simply define a contact map with the **cmap()** function and provide this to the **cna()** function along with the required cross-correlation matrix, e.g.:

```
cm <- cmap(trj, dcut = 5, scut = 0, pcut = 0.75, mask.lower = FALSE)
net.cut <- cna(cij, cm = cm)</pre>
```

One could also simply multiple the correlation matrix by the contact map to zero out these long-range correlations. However, we currently prefer not to use this filtering step as we have found that this approach can remove potentially interesting correlations that might prove to be important for the types of long-range coupling we are most interested in exploring. For example, compare the plot of correlation values below to those in **Figure X**.

```
par(mfcol = c(1, 2), mar = c(0, 0, 0, 0))
plot.dccm((cij * cm), margin.segments = net.cut$communities$membership)
```

Residue Cross Correlation

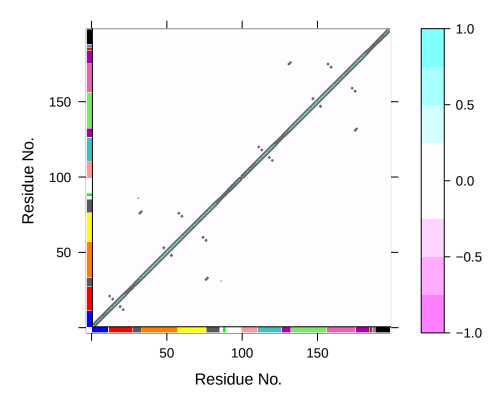


Figure 12: DCCM plot with annotated communities

```
plot.dccm(cij, margin.segments = net$communities$membership)
```

NOTE: Need to fix potential error message if mask.lower=TRUE is used in the cmap() call. It is also possible to use a consensus of contact map and dynamical correlation matrices from replica simulations with the functions cmap.filter() and filter.dccm() - see their respective help pages for details.

Residue Cross Correlation

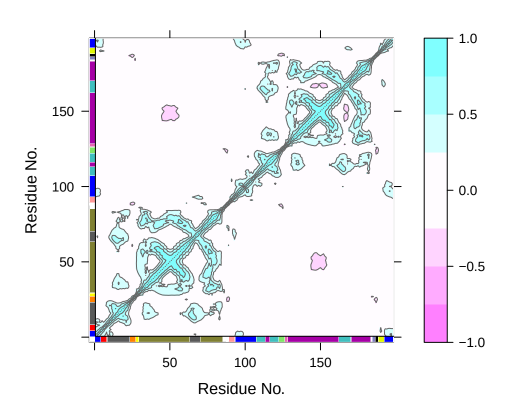


Figure 13: DCCM plot with annotated communities

Network visualization

The layout.cna() function can be used to determine the *geometric center* of protein residues and communities and thus provide 3D and 2D network and community graph layouts that most closely match the protein systems 3D coords.

```
layout3D <- layout.cna(net, pdb, k=3)
layout2D <- layout.cna(net, pdb, k=2)</pre>
```

Other network layout options are available via the igraph package, see ?igraph::layout in R for details.

Identifying nodes/communities and their composite residues

The identify.cna() function allows one to click on the nodes of a plot.cna() network plot to identify the residues within a give set of nodes. This can also be a useful first step to determine the placement of text labels for a plot -see *help(identify.cna)* for details.

```
## plotCNA, eval=FALSE
## # Click with the right mouse to label, left button to exit.
## xy <- plot.cna(net)
## identify.cna(xy, cna = net)</pre>
```

Color settings and node/edge labels

The **vmd.colors()** function is used by default to define node color. This can be changed to any input color vector as demonstrated below. Note that the **vmd.colors()** function outputs a vector of colors that is ordered as they are in the VMD molecular graphics program and changing the colors will likely break this correspondence.

```
grp.col <- bwr.colors(20)[net$communities$membership]</pre>
```

Useful options in **plot.cna()** to be aware of include the optional flags 'mark.groups' and 'mark.col', which let you to draw and color areas around groups of residues corresponding to the community clustering or any other residue partitioning of interest:

```
grp.col <- list()
for (i in 1:max(net$communities$membership)) {
  grp.tmp <- which(net$communities$membership == i)
  grp.col[[i]] <- grp.tmp
}</pre>
```

```
colbar.full <- vmd.colors()[net$communities$membership]
colbar.comms <- vmd.colors(max(net$communities$membership), alpha = 0.5)
plot(net, pdb, full = TRUE, mark.groups = grp.col, mark.col = colbar.comms)</pre>
```

Obtaining layout from PDB structure

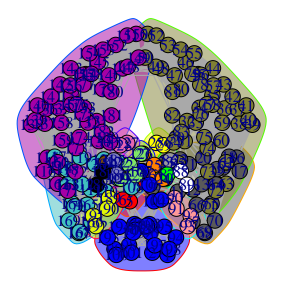


Figure 14: \$network colored according to communities grouping

Network manipulation

The **prune.cna()** function allows one to remove communities under a certain size or with less less than a certain number of edges to other communities prior to visualization or further analysis. For example, to remove communities composed of only 2 residues:

```
net.pruned <- prune.cna(net, size.min = 3)</pre>
```

```
##
    id size
                                     {\tt members}
                   c(1:4, 94:107, 193:198)
##
     1
          24
##
     2
           4
                                         5:8
##
     3
          22
                             c(9:23, 64:70)
     4
##
           4
                                       24:27
##
     5
           2
                                       28:29
                            c(30:63, 71:85)
##
     6
          49
##
     7
           1
##
     8
           1
                                          87
##
     9
           2
                                       88:89
##
    10
                                       90:93
##
    11
          20 c(108:113, 117:122, 163:170)
##
    12
          50 c(114:116, 129:162, 171:183)
##
    13
           4
                                     123:126
    14
           2
                                     127:128
##
##
    15
           1
                                         184
           2
                                     185:186
##
    16
           2
##
    17
                                     187:188
##
                                     189:192
## Removing Nodes: 7, 8, 9, 16, 17, 5, 14, 15
## id
         size
                  edges
                          members
                 28:29
## 5
         2
             1
## 7
         1
             0
                 86
## 8
             0
         1
                 87
## 9
         2
             0
                 88:89
         2
## 14
                 127:128
             1
```

Obtaining estimated layout with fruchterman.reingold

plot.cna(net)

```
plot.cna(net.pruned)
```

Obtaining estimated layout with fruchterman.reingold

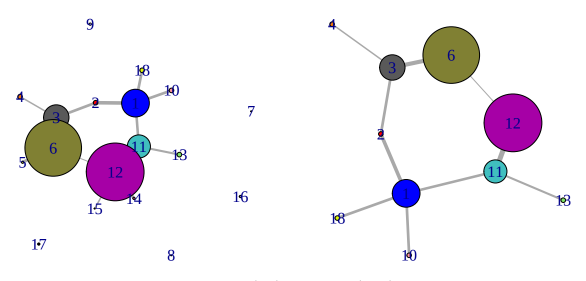


Figure 15: Original (left) and pruned (right) networks

Applying this command to our network example, you can see how the communities composed by less than 3 residues are deleted from the graph.

Calculating network node centrality and suboptimal paths.

Node centrality describes the distribution of the edges in the network. In protein-networks, identifying the nodes with many edges (hubs) as well as the segments with a high number of connections allows to get insights on the internal coordination of the protein regions. This measure can also be employed to characterize the protein sections showing conformational-dependent differences in the protein dynamics.

There are different measures of node centrality, from the simplest one that counts the number of edges of each node (node degree) to the more complex such as the betweenness centrality. The betweenness centrality of a node is defined as the number of unique-shortest paths crossing that node and has the advantage of considering in the calculation the whole network topology and not only the closest neighbors as the node degree.

The igraph package provides different functions to perform such calculations as **degree**, **betweenness** and **closeness**. Below we report the output of the betweenness centrality on the HIVP example case:

plot(node.betweenness, xlab="Residue No", ylab="Centrality", type="h")

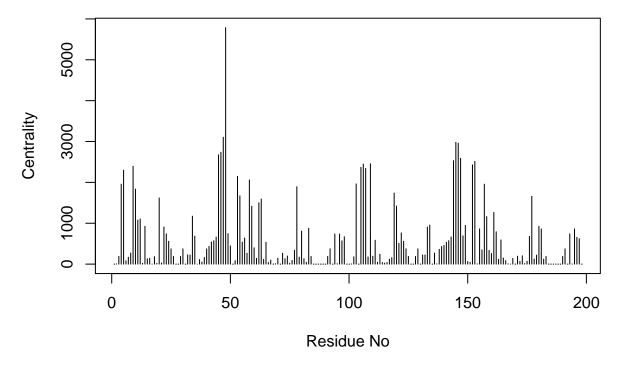


Figure 16: Betweenness centrality

Interestingly, in **FigureXX**, the regions with the highest centrality values are the two flaps of the HIVP structure, whose dynamics and interaction is fundamental for the protein enzymatic function.

Suboptimal paths calculation. Identifying residues potentially involved in the dynamic coupling of distal protein regions can be facilitated by calculating possible linking paths through the correlation network objects. To run such an analysis, we can use the WISP program [7]. It is easy to interface the protein network built using Bio3d with WISP to perform a search of X shortest paths between two selected network nodes. WISP requires in input the adjacency matrix of the network, which can be saved in a file using the following command:

```
write.table(net$cij, quote=FALSE, row.names=FALSE, col.names=FALSE, file="adj.matrix.txt")
```

To view an example of the output obtained and how key residues involved in the allosteric signaling were identified, please read the following work: Scarabelli G and Grant BJ, Kinesin-5 allosteric inhibitors uncouple the dynamics of nucleotide, microtubule and neck-linker binding sites, Biophys J. 2014, in press. and Yao X and Grant BJ Characterizing Nucleotide Dependent Allostery in G-Proteins with Molecular Dynamics and Normal Mode Analysis, submitted.

Summary

In this vignette we have confined ourselves to introducing relatively simple correlation network generation protocols. However, the individual Bio3D functions for network analysis permit extensive customization of the network generation procedure. For example, one can chose different node representations (including all heavy

atoms, residue center of mass, Calpha atoms for each residue, or collections of user defined atoms), different correlation methods (including Pearson and mutual information), different community detection methods (including Girvan-Newman betweenness, Random walk, and Greedy modularity optimization approaches), as well as customized network layouts and result visualizations. We encourage users to explore these options and provide feedback as appropriate.

Although the protocols detailed herein can provide compelling results in a number of cases, defining an optimal procedure for network generation is an area of active research. For example rearrangement of some residues in the communities and changes in the total number of communities may be observed among different community structures based on different independent MD simulations of the same system. These variations of community structures are partially due to the small changes in the calculated cij values that reflect statistical fluctuations among independent simulations. However, changes in mutual information or correlation values may still be relatively small and differences among independent community structures may be largely due to limitations of the modularity metric, a methodology that has been developed for binary adjacency matrices, where edges are counted without taking into account their lengths. We are currently developing a modularity methodology based on natural familial networks, where the strength of a community is defined by the number of close connections between members. We are also developing integrated 3D viewing functions (so VMD and PyMol will no longer be required), as well as functionality to facilitate analysis of the interactions that generate changes in the community network.

References

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Document Details

This document is shipped with the Bio3D package in both R and PDF formats. All code can be extracted and automatically executed to generate Figures and/or the PDF with the following commands:

```
library(rmarkdown)
render("cna_vignette.R")
```

Information About the Current Bio3D Session

```
print(sessionInfo(), FALSE)

## R version 3.0.2 (2013-09-25)

## Platform: x86_64-apple-darwin10.8.0 (64-bit)

##

## attached base packages:
```

```
## [1] grid
                parallel stats
                                    graphics grDevices utils
                                                                  datasets
## [8] methods
                base
##
## other attached packages:
## [1] lattice_0.20-29 igraph_0.7.1
                                      bio3d_2.1-0
                                                      rmarkdown_0.3.3
##
## loaded via a namespace (and not attached):
## [1] digest_0.6.4
                      evaluate_0.5.5 formatR_0.10
                                                      htmltools_0.2.4
## [5] knitr_1.6
                      stringr_0.6.2
                                      tools_3.0.2
                                                      yaml_2.1.13
```

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