# Vignette: Grammar of graphics of genealogy (ggenealogy)

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This LATEX vignette document is created using the R function Sweave on the R package ggenealogy. It is automatically downloaded with the package and can be accessed with the R command vignette("ggenealogy").

## Summary

#### Description

The ggenealogy package provides tools to examine genealogical data, generating basic statistics on their graphical structures using parent and child connections, and displaying the results. The genealogy can be drawn in relation to additional variables, such as development year, and the shortest path distances between genetic lines can be determined and displayed. Production of pairwise distance matrices and phylogenetic diagrams constrained by generation count are also available in the visualization toolkit. This vignette is intended to walk readers through the different methods available in the ggenealogy package.

#### Caution

igraph must be used with version >= 0.7.1

#### Introduction

#### Installation

 $\mathsf{R}$  is a open source software project for statistical computing, and can be freely downloaded from the Comprehensive R Archive Network (CRAN) website. The link to contributed documentation on the CRAN website offers practical resources for an introduction to R , in several languages. After downloading and installing R , the installation of additional packages is straightforward. To install the <code>ggenealogy</code> package from R , use the command:

```
> install.packages("ggenealogy")
```

The ggenealogy package should now be successfully installed. Next, to render it accessible to the current R session, simply type:

```
> library(ggenealogy)
```

To access help pages with example syntax and documentation for the available functions of the ggenealogy package, please type:

```
> help(package="ggenealogy")
```

To access more detailed information about a specific function in the ggenealogy package, use the following help command on that function, such as:

```
> help(getChild)
```

The above command will return the help file for the getChild function. The help file often includes freestanding example syntax to illustrate how function commands are executed. In the case of the getChild function, the example syntax is the following three lines, which can be pasted directly into an R session.

```
> data(sbGeneal)
> getChild("Tokyo", sbGeneal)
> getChild("Essex", sbGeneal)
```

#### Preprocessing pipeline

> data(sbGeneal)

> str(sbGeneal)

In the ggenealogy package, there is an example dataset containing genealogical information on soybean varieties called sbGeneal.rda. It may be helpful to load that example file so that you can follow along with the commands and options introduced in this vignette. To ensure that you have uploaded the correct, raw sbGeneal.rda file, you can observe the first six lines of the file, and determine its dimension and structure:

```
> head(sbGeneal)
          child year yield year.imputed
                                             parent
1
          5601T 1981
                         NA
                                     TRUE Hutcheson
2
          Adams 1948
                       2734
                                    FALSE Dunfield
3
           A.K. 1910
                                     TRUE
                                                <NA>
4 A.K. (Harrow) 1912
                                    FALSE
                                                A.K.
                       2665
5
         Altona 1968
                         NA
                                    FALSE
                                           Flambeau
6
          Amcor 1979
                       2981
                                    FALSE
                                           Amsoy 71
> dim(sbGeneal)
[1] 390
          5
```

```
'data.frame': 390 obs. of 5 variables:

$ child : chr "5601T" "Adams" "A.K." "A.K. (Harrow)" ...

$ year : num 1981 1948 1910 1912 1968 ...

$ yield : int NA 2734 NA 2665 NA 2981 2887 2817 NA NA ...
```

\$ year.imputed: logi TRUE FALSE TRUE FALSE FALSE FALSE ...
\$ parent : chr "Hutcheson" "Dunfield" NA "A.K." ...

We see that the sbGeneal data file is a data frame structure with 390 rows (observations) and 5 columns (variables). Each row contains a child node character label and parent node character label. Each row also contains a numeric value corresponding to the year the child node was introduced, an

integer value of the protein yield of the child node, and a logical value year.imputed, which indicates whether or not the year of introduction of the child node was imputed.

Now that the sbGeneal file has been loaded as a data frame, it must next be converted into a graph object using the dfToIG() function. The dfToIG() function requires a data frame as input, and that data frame should be structured such that each row represents an edge with a child and parent relationship. For more information, try using the help command on the function:

#### > help(dfToIG)

We see that the function takes optional parameter arguments, such as vertexinfo (a list of columns of the data frame which provide information for the starting "child" vertex, or a separate data frame containing information for each vertex with the first column as the vertex name), edgeweights (a column that contains edge values, with a default value of unity), and isDirected (a boolean value that describes whether the graph is directed (true) or undirected (false); the default is false).

In this example, we want to produce an undirected graph object that contains all edge weight values of one, because our goal is to set an edge value of unity for every pair of vertices (individuals) that are related as parent and child. The dfToIG() function uses the software igraph to convert the data frame into a graph object. For clarity, we will assign the outputted graph object the name ig (for igraph object), and then examine its class type:

```
> ig <- dfToIG(sbGeneal)
> class(ig)
[1] "igraph"
```

Above, we confirmed that the ig object is of class type igraph. The ig object is required as input in many ggenealogy functions, which will be demonstrated below.

# General (non-plotting) methods of genealogical data

The ggenealogy package offers several functions that result in useful information beside plots. Below is a brief introduction to some of the available non-plotting functions.

#### Functions for individual vertices

The ggenealogy package offers several functions that you can use to obtain information for individual vertices. First, the function <code>isParent()</code> can return a logical variable to indicate whether or not the second variety is a parent of the first variety.

```
> isParent("Young","Essex",sbGeneal)
[1] TRUE
```

```
> isParent("Essex", "Young", sbGeneal)
[1] FALSE
We see that "Essex" is a parent of "Young", and not vice-versa. Similarly, the function isChild() can
return a logical variable to indicate whether or not the first variety is a child of the second variety.
> isChild("Young", "Essex", sbGeneal)
[1] TRUE
> isChild("Essex", "Young", sbGeneal)
[1] FALSE
We see that, as expected, "Young" is a child of "Essex", and not vice-versa. It is also possible to derive
the year of a given variety using the getYear() function:
> getYear("Young",sbGeneal)
[1] 1968
> getYear("Essex",sbGeneal)
[1] 1962
Fortunately, the returned year values are consistent, as the "Young" variety (1968) is a child to the
"Essex" variety (1962) by an age difference of 6 years. In some cases, you may wish to obtain a complete
list of all the parents of a given variety. This can be achieved using the getParent() function:
> getParent("Young",sbGeneal)
[1] "Davis" "Essex"
> getParent("Tokyo",sbGeneal)
character(0)
```

> getYear("Tokyo", sbGeneal)

#### [1] 1907

We learn from this that "Essex" is not the only parent of "Young"; "Young" also has a parent "Davis". We also see that "Tokyo" does not have any documented parents in this dataset, and has an older year of introduction (1907) than other varieties we have examined thusfar. Likewise, in other cases, you may wish to obtain a complete list of all the children of a given variety. This can be achieved using the getChild() function:

```
[1] "Ogden" "Volstate"

> getChild("Ogden",sbGeneal)

[1] "C1069" "C1079" "D51-2427" "D55-4090" "D55-4159"
[6] "D55-4168" "Kent" "N44-92" "N45-745" "N48-1101"
[11] "Ogden x CNS" "Ralsoy x Ogden"
```

We find that even though the "Tokyo" variety is a grandparent of the dataset, it only has two children, "Ogden" and "Volstate". However, one of its children, "Ogden", produced 12 children.

If we want to obtain a list that contains more than just one generation past or previous to a given variety, then we can use the getAncestors() and getDescendants() functions, where we specify the number of generations we wish to view. This will return a data frame to us with the labels of each ancestor or descendant, along with the number of generations each one is from the given variety.

If we only look at one generation of ancestors of the "Young" variety, we should see the same information we did earlier when we used the getParent() function of the Young variety:

```
> getAncestors("Young",sbGeneal,1)
```

> getChild("Tokyo",sbGeneal)

```
label gen
2 Davis 1
1 Essex 1
```

Indeed, we consistently see that the "Young" variety has only 2 ancestors within one generation, "Davis" and "Essex". However, if we view the first five generations of ancestors of the "Young" variety, we can view four more generations of ancestors past simply the parents:

#### > getAncestors("Young",sbGeneal,5)

	label	gen
27	Davis	1
26	Essex	1

```
25
             Ralsoy x Ogden
24 Roanoke x (Ogden x CNS)
                                 2
                                 2
23
                          Lee
22
                    S55-7075
                                 2
                                 3
21
                        Ogden
                                 3
20
                      Ralsoy
19
                 Ogden x CNS
                                 3
                                 3
17
                          CNS
18
                     Roanoke
                                 3
                                 3
                        S 100
16
                    N48-1248
                                 3
15
                                 3
14
                        Perry
10
                        Ogden
                                 4
                    PI 54610
                                 4
13
12
                        Tokyo
                                 4
11
                          CNS
9
                     Clemson
                                 4
6
                     Roanoke
                                 4
8
                      Illini
                                 4
7
   N45-745 x (Ogden x CNS)
4
                    PI 54610
                                 5
3
                                 5
                        Tokyo
                 Ogden x CNS
                                 5
1
5
                     Clemson
                                 5
2
                         A.K.
                                 5
```

> nrow(getAncestors("Young",sbGeneal,5))

#### [1] 27

In the second line of code above, we determined the dimensions of the returned data frame, and see that there are 27 ancestors within the first five ancestral generations of the "Young" variety.

Similarly, if we only look at the first generation of descendants of the "Ogden" variety, we should see the same information as we did earlier when we used the getChild() function on the "Ogden" variety:

#### > getDescendants("Ogden",sbGeneal,1)

	label	gen
12	C1069	1
11	C1079	1
10	D51-2427	1
9	D55-4090	1
8	D55-4159	1
7	D55-4168	1
6	Kent	1
5	N44-92	1
4	N45-745	1

```
3 N48-1101 1
2 Ogden x CNS 1
1 Ralsoy x Ogden 1
```

Indeed, we see again that "Ogden" has 12 children. Additionally, if we want to view not only the children, but also the grandchildren, of the "Ogden" variety, then we can use this function, only now specifying two generations of descendants:

#### > getDescendants("Ogden",sbGeneal,2)

				label	-
28	C1069				1 1
27	C1079				
26	D51-2427				
25	D55-4090				
24	D55-4159				
23	D55-4168				
22	. Kent				
21	N44-92				
20					
19			N4	18-1101	1
18			Ogder	n x CNS	1
17		F	Ralsoy 2	c Ogden	1
16	• •				2
15	Cutler				2
14	C1266R				2
13	Semmes				2
11	D60-7965				2
12	D60-7965			2	
10	D59-9289				2
9	Beeson				
8	Calland				2 2
7	Hood				
6	N48-1867				2 2
5	D52-810				
4	N45-745	х	(Ogden	x CNS)	2 2
3			•	R54-168	2
2	Roanoke	x	(Ogden	x CNS)	2
1			-	Davis	2

We see that variety "Ogden" has 16 grandchildren from its 12 children.

#### Functions for pairs of vertices

Say you have a pair of vertices, and you wish to determine the degree of the shortest path between them, where edges represent parent-child relationships. You can accomplish that with the getDegree() function.

```
> getDegree("Tokyo", "Ogden", ig, sbGeneal)
[1] 1
> getDegree("Tokyo", "Holladay", ig, sbGeneal)
[1] 7
```

As expected, the shortest path between the "Tokyo" and "Ogden" varieties has a value of 1, as we already determined that they have a direct parent-child relationship. However, the shortest path between "Tokyo" and one of its descendants, "Holladay", has a much higher degree of 7.

Note that degree calculations in this case are not limited to one linear string of parent-child relationships; cousins and siblings and products thereof will also have computable degrees via nonlinear strings of parent-child relationships.

#### Functions for the full genealogical structure

There are many parameters about the full genealogical structure that you may wish to know that cannot easily be obtained through images and tables. The function getBasicStatistics() will return graph theoretical measurements of the full genealogy. For instance, is the full genealogy connected? If not, how many separated components does it contain? In addition to these parameters, the getBasicStatistics() function will also return the number of nodes, the number of edges, the average path length, the graph diameter, among others:

```
> getBasicStatistics(ig)

$isConnected
[1] FALSE

$numComponents
[1] 11

$avePathLength
[1] 5.333746

$graphDiameter
[1] 13

$numNodes
[1] 230

$numEdges
[1] 340

$logN
[1] 5.438079
```

In this case, we learn that our full genealogical structure is not all connected by parent-child edges. Instead, it is composed of 11 separate components. The average path length of the full genealogy is 5.333746, that the graph diameter is 13, and that the logN value is 5.438079. We also see that the number of nodes in the full genealogy is 230, and the number of edges in the full genealogy is 340.

But can we view a list of these nodes and edges? To do so, we can call the getNodes() and getEdges() commands to obtain lists of all the unique nodes and edges in the full genealogical structure. Here, we obtain a list of the 340 edges (with each row containing the names of the two connected vertices, and an edge weight, if existent). We will simply view the first six rows of the object, and determine the number of edges by counting the number of rows (340):

```
> eList = getEdges(ig, sbGeneal)
> head(eList)
     child
                      parent
[1,] "5601T"
                      "Hutcheson"
[2,] "Adams"
                      "Dunfield"
[3,] "A.K. (Harrow)" "A.K."
[4,] "Altona"
                      "Flambeau"
[5,] "Amcor"
                      "Amsoy 71"
[6,] "Amsoy"
                      "Adams"
> nrow(eList)
[1] 340
```

We then obtain a list of the 230 nodes. Again, we only view the first six rows of the object, and determine the number of nodes by counting the number of indices (230).

# Plotting methods of genealogical data

Until this point, the vignette has introduced functions that return lists, data frames, and statistics about the genealogical dataset. However, the ggenealogy package also contains visualization tools for genealogical datasets. Access to various types of visual plots and diagrams of the lineage can allow genealogical researchers to more efficiently and accurately explore an otherwise complicated data structure. Below, we introduce functions in ggenealogy that produce visual outputs of the dataset.

#### Plotting the ancestors and descendants of a vertex

One visualization tool, plotAncDes(), allows the user to view the ancestors and descendants of a given variety. The inputted variety is highlighted in the center of the plot, ancestors are displayed to the left of the center, and descendants are displayed to the right of the center. The further left or right from the center, the larger the number of generations that particular ancestor/descendant is from the inputted and centered variety.

As such, this plotting command does not provide visual information about specific years associated with each related variety (as is done in some of the visualization tools introduced later), but it does group all varieties from each generation group onto the same position of the horizontal axis. Here, we specify that we want to plot 5 ancestor generations and 4 descendant generations of the variety "Lee":

#### > plotAncDes("Lee", sbGeneal,5,4)

We immediately see in Figure 1 that this visual representation of the ancestors and descendants of a given variety can often provide enhanced readability compared to the list output provided in the previous functions, getAncestors() and getDescendants(). We notice that even though we specified for 5 generations of ancestors, the extent of documented ancestors of "Lee" includes only 3 generations.

We also see now that some node labels are repeated. For instance, the "5601T" variety appears twice, once as a great-grandchild (third generation descendant) of "Lee", and once as a great-granchild (fourth generation descendant) of "Lee". This is because there are two separate parent-child pathways between "Lee" and "5601T", one pathway with only two nodes ("Essex" and "Hutchson") between them, and one pathway with three nodes ("Essex", "T80-69", and "TN89-39") between them.

Why does this happen? In this visual tool, we are constraining the horizontal axis to generation count. Without allowing nodes to repeat, this data information cannot be clearly and succinctly presented. Most graph visualization software that genealogists might use to view their datasets do not allow for repeated nodes, as per the definition of a graph. Hence, the plotAncDes() function is one of the more unique visual tools of the ggenealogy package.

It should be noted that the plotAncDes() function, by default, highlights the centered variety label in pink. However, the user can alter this color, as we will show next. Furthermore, the user can specify additional grammar of graphics plotting tools (from the ggplot2 package) to tailor the output of the plotAncDes() function, which we also show below.

For example, we will now change the color of the center variety label vColor to be highlighted in blue. Also, we will add a horizontal axis label called "Generation index", using the ggplot2 syntax. Note that this time we do not specify the generational count for ancestors and descendants, and so the default value of three generations is applied to both cases. Remember, to determine such default values, as well as all function parameters, simply run the help command on the function of interest.

```
> plotAncDes("Tokyo", sbGeneal, vColor = "blue") + ggplot2::labs(x="Generation index",y="")
```

We verify immediately from Figure 2 that the "Tokyo" variety does not have any ancestors in this dataset, an observation consistent with what we discovered earlier. We also see the "Tokyo" variety only has two children, but has many more grandchildren, and great-grand children.

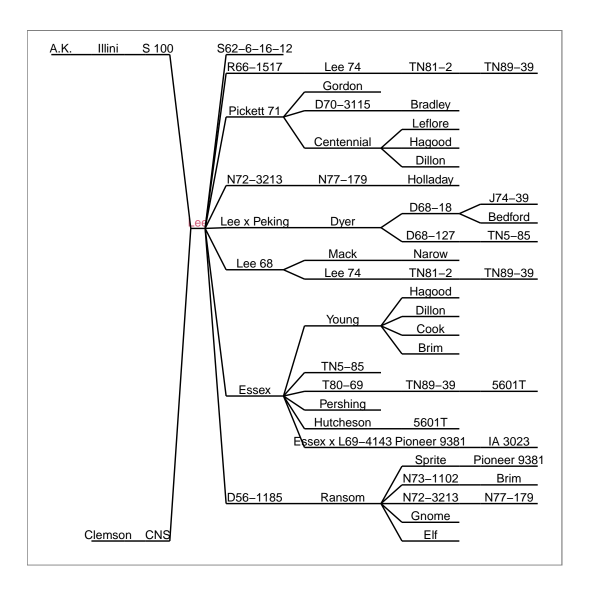


Figure 1: Ancestors and descendants of the "Lee" variety, constrained on the horizontal axis by generational separation from "Lee".

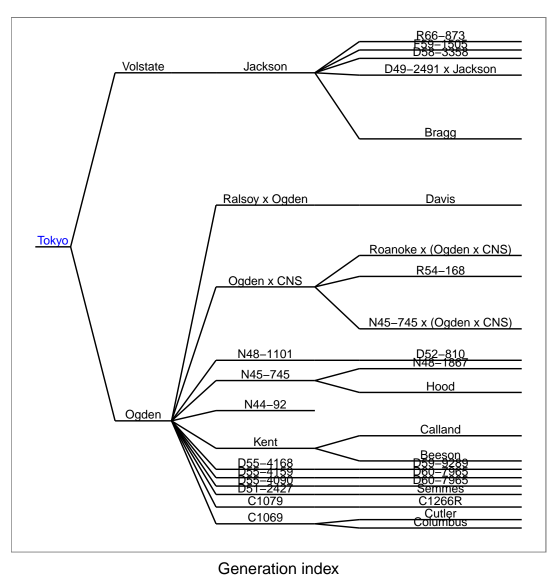


Figure 2: Ancestors and descendants of the "Tokyo" variety, constrained on the horizontal axis by generational separation from "Tokyo".

#### Plotting the shortest path between two vertices

As this data set deals with soybean lineages, it may be useful for agronomists to track how two varieties are related to each other via parent-child relationships. Then, any dramatic changes in protein yield, SNP varieties, and other measures of interest between the two varieties can be tracked across their genetic timeline, and pinpointed to certain paths within their historical lineage.

The ggenealogy software allows users to select two varieties of interest, and determine the shortest pathway of parent-child relationships between them, using the getPath() function. This will return a list path object that contains the variety names and their years in the path. The returned path object can then be plotted using the plotPath() function, which we now demonstrate.

The getPath() function determines the shortest path between the two inputted vertices, and takes into account whether or not the graph is directed with the parameter isDirected, which defaults to false. The getPath() function will check both directions and return the path if it exists:

```
> getPath("Brim", "Bedford", ig, sbGeneal, isDirected=FALSE)

$pathVertices
[1] "Brim" "Young" "Essex" "T80-69" "J74-40" "Forrest" "Bedford"

$yearVertices
[1] "1977" "1968" "1962" "1975" "1975" "1973" "1978"
```

We see that there is a path between "Brim" and "Bedford" varieties, with 5 varieties separating them. We are not considering direction, however, because the ig object is undirected. However, to demonstrate the importance of direction, we will recompute the path where the direction matters. We first produce a directed igraph object dirIG, and then try to determine the path between the same two vertices, "Brim" and "Bedford".

```
> dirIG = dfToIG(sbGeneal, isDirected = TRUE)
> getPath("Brim", "Bedford", dirIG, sbGeneal, isDirected = TRUE)
```

Now that we are considering the direction, we are only considering paths where each edge represents a parent-child relationship in the same direction as the one before it. We now would receive an error warning that we cannot compute a directed path on an undirected graph. We next try to reverse the input order of the vertices, as shown below, but we will receive the same error message:

```
> getPath("Bedford", "Brim", dirIG, sbGeneal, isDirected=TRUE)
```

We can derive from the errors returned in the last two commands that the varieties "Brim" and "Bedford" are not connected by a linear sequence of parent-child relationships. Rather, the path between them branches at some point, involving siblings and/or cousins.

Hence, unless you are working with a dataset that must be analyzed as a directed graph, it is best to use the getPath() function with the default third parameter indicating lack of direction, and to use an igraph object without direction, such as our original ig object. We do just that, and save the path between these two varieties to a variable called path:

#### > pathBB = getPath("Bedford", "Brim", ig, sbGeneal, isDirected=FALSE)

Now that we have a non-empty pathBB object that consists of two lists (for variety names and years), we can plot the relationship between the two using the plotPath() function.

#### > plotPath(pathBB)

This produces a neat visual (see Figure 3) that informs us of all the varieties involved in the shortest path between "Brim" and "Bedford". In this plot, the years of all varieties involved in the path are indicated on the horizontal axis, while the vertical axis has no meaning other than to simply to display the labels evenly spaced vertically.

Although a call to the ggenealogy function getYear() indicates that "Bedford" was developed in 1978 and "Brim" in 1977, we quickly determine from the plot that "Brim" is not a parent, grandparent, nor

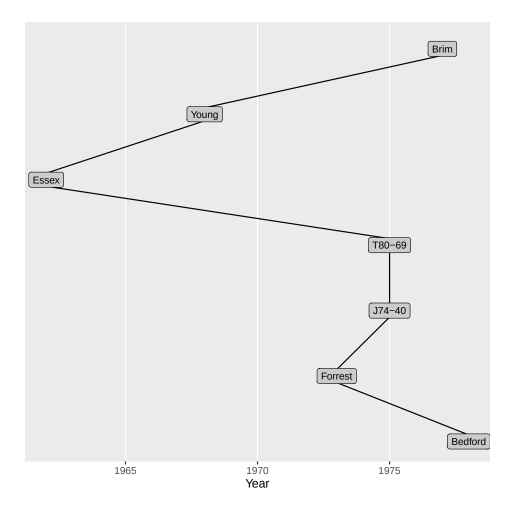


Figure 3: The shortest path between varieties "Brim" and "Bedford" is not strictly composed of unidirectional parent-child relationships, but instead, includes cousin-like relationships.

any great-grandparent of "Bedford". Instead, we see that these two varieties are not related through a unidirectional parent-child lineage, but have a cousin-like relationship. The oldest common ancestor between "Bedford" and "Brim" is the variety "Essex", which was developed in 1962.

However, there are other cases of pairs of varieties that are connected by a linear, unidirectional combination of parent-child relationships, as we see below:

```
> pathNT = getPath("Narow", "Tokyo", ig, sbGeneal, isDirected=FALSE)
> plotPath(pathNT)
```

From the output, shown in Figure 4, we see that the variety "Tokyo" is an ancestor of "Narow" via four linear parent-child relationships. Because of this, we can still view the pathway, even when we use an igraph object dirIG that is directed, and set the boolean isDirected variable to true.

Either ordering of the two varieties will produce the exact same result. In other words, the following

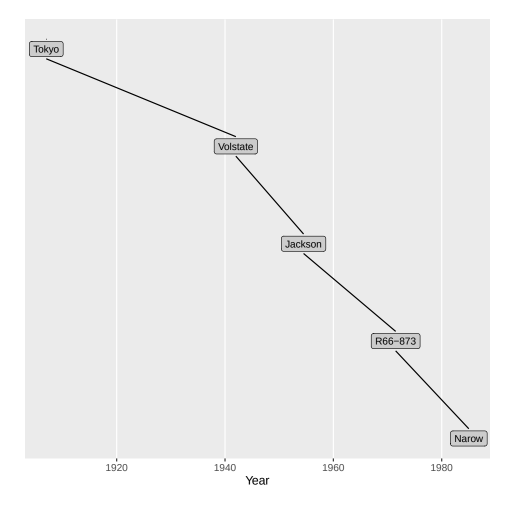


Figure 4: The shortest path between varieties "Narow" and "Tokyo" is strictly composed of a unidirectional sequence of parent-child relationships.

two code blocks would produce the same figure we saw in Figure 4:

```
> pathNT = getPath("Narow", "Tokyo", dirIG, sbGeneal, isDirected=TRUE)
> plotPath(pathNT)

> pathTN = getPath("Tokyo", "Narow", dirIG, sbGeneal, isDirected=TRUE)
> plotPath(pathTN)
```

#### Plotting shortest paths superimposed on full genealogical structure

Now that we can create and plot path objects, we may wish to know how those paths are positioned in comparison to the genealogical lineage of the entire data structure. For instance, of the documented soybean cultivar lineage varieties, where does the shortest path between two varieties of interest exist? Are these two varieties comparatively older compared to the overall data structure? Are they newer? Or, do they span the entire structure, and represent two extreme ends of documented time points?

There is a function available in the <code>ggenealogy</code> package, <code>plotPathOnAll()</code>, that allows users to quickly visualize their path of interest superimposed over all varieties and edges present in the whole data structure. Here we will produce a plot of the previously-determined shortest path between varieties "Tokyo" and "Narow" across the entire dataset (in this particular dataset, some edges are not plotted, as they contain NA values):

```
> plotPathOnAll(pathNT, sbGeneal, ig, binVector = 1:3)
```

The resulting plot is shown in Figure 5.

While the first three explicit parameters to the function plotPathOnAll() have been introduced earlier, the fourth parameter (binVector) requires some explanation. The motivation of the plotPathOnAll() function is to write variety text labels on a plot, with the center of each variety label constricted on the horizontal axis to its developmental year. As is the case for the plots before, the vertical axis has no specific meaning. Unfortunately, for large datasets, this motivation can be a difficult task because the text labels of the varieties can overlap if they are assigned a similar y coordinate, have a similar year (x coordinate), and have labels with large numbers of characters (width of x coordinate).

For each variety, the x coordinate (year) and width of the x coordinate (text label width) cannot be altered, as they provide useful information. However, the vertical coordinate is arbitrary. Hence, in an attempt to mitigate text overlap, the plotPathOnAll() function does not randomly assign the vertical coordinate. Instead, it allows users to specify the number of bins (binVector), which partially controls the vertical positions.

If the user determines to produce a plot using three bins, as in the example code above, then the varieties are all grouped into three bins based on their years of development. In other words, there will be bin 1 (the "oldest bin") which includes the one-third of all varieties with the oldest developmental years, bin 2 (the "middle bin"), and bin 3 (the "youngest bin").

Then, in order to decrease text overlap, consecutively increasing vertical positions are alternatively assigned to the three bins (For example: bin 1, then bin 2, then bin 3, etc.) repeatedly until all

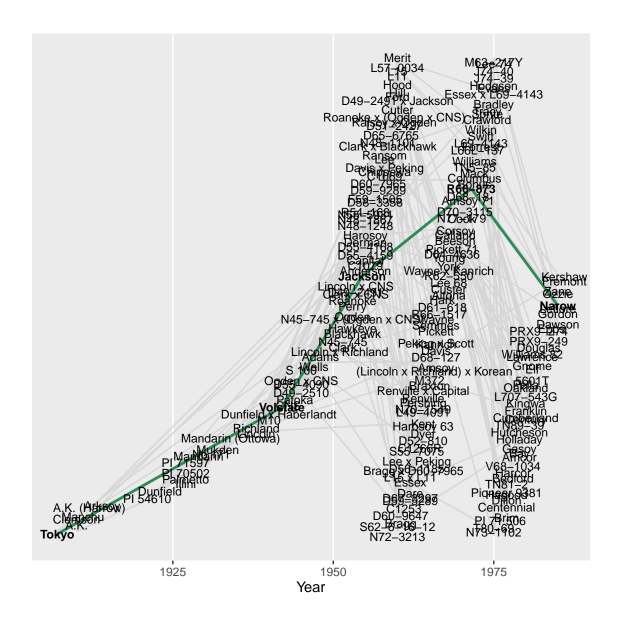


Figure 5: Plot of the shortest path, highlighted in green, between the varieties "Tokyo" and "Narow" superimposed on the full genealogical structure, using a binVector size of 3.

varieties are accounted. This algorithm means that there are at least two vertical positions separating any pair of varieties from the same bin.

In this plot, edges not on the path of interest are thin and gray, whereas edges on the path of interest are bolded and green, by default. Also, variety labels in the path of interest are boldfaced, by default.

Using the plot, we immediately recognize that the path spans most of the years in the full data structure: "Tokyo" appears to be the oldest variety in the data, and "Narow" appears to be among the youngest. We note that many varieties have development years between 1950 and 1970.

However, this plot has significant empty spaces between the distinct bins, and almost all text labels are overlapping, causing decreased readability. To force variety text labels into these spaces, the user may consider choosing a larger number of bins. Hence, we next examine a binVector size of six:

> plotPathOnAll(pathNT, sbGeneal, ig, binVector = 1:6)

Figure 6 shows that the binVector size of six successfully mitigated text overlap compared to Figure 5, which had a binVector size of three. Most of the remaining textual overlap is confined to the range

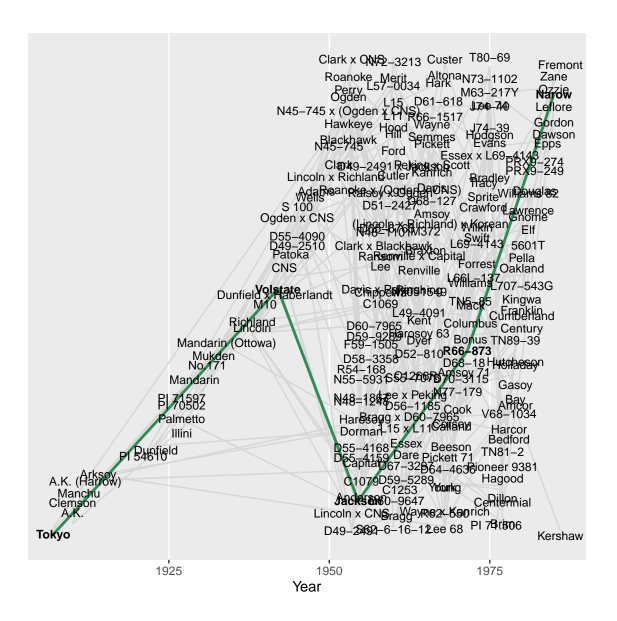


Figure 6: Plot of the shortest path, highlighted in green, between the varieties "Tokyo" and "Narow" superimposed on the full genealogical structure, using a binVector size of 6.

of years (1950-1970) of which the most varieties had development years.

#### Plotting pairwise distance matrices between a set of vertices

It may also be of interest to generate matrices where the cell colors indicate the magnitude of a variable (such as the degree of the shortest path) between all pairwise combinations of inputted varieties. The package ggenealogy also provides a function plotDegMatrix() for that purpose.

Here, we plot a distance matrix for a set of 8 varieties, defining both the x- and y- axes titles as "Soybean label", and the legend label as "Degree". Syntax from the ggplot2 package can be appended to tailor the output of the plotDegMatrix() function. In this case, we denote pairs with small degrees to be colored white, and pairs with large degrees to be colored dark green, using scale\_fill\_continuous:

```
> varieties=c("Brim", "Bedford", "Calland", "Narow", "Pella", "Tokyo", "Young", "Zane")
> p = plotDegMatrix(varieties, ig, sbGeneal, "Soybean label", "Soybean label", "Degree")
> p + ggplot2::scale_fill_continuous(low="white", high="darkgreen")
```

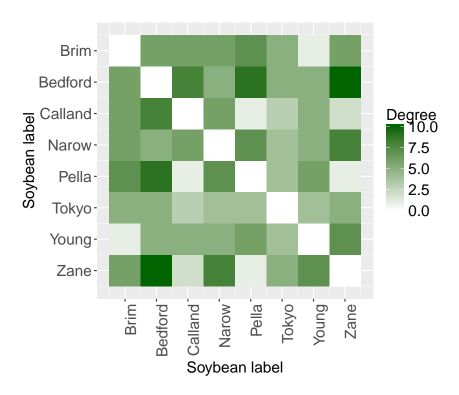


Figure 7: Colored matrix plot showing the degrees of the shortest paths between all pair combinations from a set of eight varieties of interest.

Figure 7 shows that the degree of the shortest path between varieties "Bedford" and "Zane" seems to be the largest in the dataset, which should be around 10. We can verify this simply with:

> getDegree("Bedford", "Zane", ig, sbGeneal)

#### [1] 10

Indeed, the degree of the shortest path between "Bedford" and "Zane" is 10. The distance matrix plot provides us additional information: The degree of 10 may be a comparatively large degree within the given soybean dataset sbGeneal, seeing that the degrees of the shortest paths for the other 27 pairwise combinations of the eight varieties that we explored here are less than 10.

In a similar function plotYearMatrix(), the difference in years between all pairwise combinations of vertices can be constructed and viewed:

> varieties=c("Brim", "Bedford", "Calland", "Narow", "Pella", "Tokyo", "Young", "Zane")
> plotYearMatrix(varieties,sbGeneal)

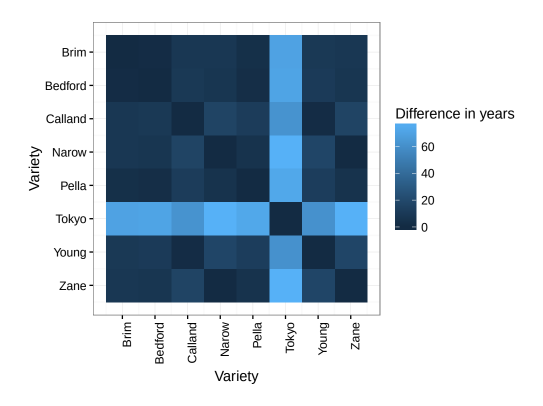


Figure 8: Colored matrix plot showing the year differences between all pair combinations from a set of eight varieties of interest.

Here, we did not change any defaults. As such, the resulting plot in Figure 8 contains the default values of "Variety" for the x-and y-axis labels, and "Difference in years" for the legend label. It also uses the default colors of dark blue for small year difference and light blue for large year difference.

Running this function on this particular set of eight vertices suggests that most combinations of varieties are only one or two decades apart in year introduction, with the exception of the "Tokyo" variety, which appears to be separated from each of the other seven varieties by about six decades. This is not surprising, because we have seen througout the tutorial that the "Tokyo" variety is the oldest variety in the dataset.

### Conclusions

The ggenealogy package offers various plotting tools that can assist those studying genealogical lineages in the data exploration phases. As each plot comes with its advantages and disadvantages, we recommend for users to explore several of the available visualization tools.

This vignette briefly introduced some of the capabilities of the ggenealogy package. Inevitably, new approaches will necessitate new features in subsequent versions and might reveal unforeseen bugs. Please send comments, suggestions, questions, and bug reports to lrutter@iastate.edu.