

# Analysis of high-throughput sequencing of T and B cell receptors with LymphoSeq

David G. Coffey, MD

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Application of high-throughput sequencing of T and B lymphocyte antigen receptors has great potential for improving the monitoring of lymphoid malignancies, assessing immune reconstitution after hematopoietic stem cell transplantation, and characterizing the composition of lymphocyte repertoires.<sup>1</sup> LymphoSeq is an R package designed to import, analyze, and visualize antigen receptor sequencing from Adaptive Biotechnologies' ImmunoSEQ assay.<sup>2</sup> The package is also adaptable to the analysis of T and B cell receptor sequencing processed using other platforms such as MiXCR<sup>3</sup> or IMGT/HighV-QUEST.<sup>4</sup> This vignette has been written to highlight some of the features of LymphoSeq and guide the user through a typical workflow.

## Importing data

The LymphoSeq function `readImmunoSeq` imports tab-separated value (.tsv) files exported by Adaptive Biotechnologies ImmunoSEQ analyzer where each row represents a unique sequence and each column is a variable with information about that sequence such as read count, frequency, or variable gene name. Only files with the extension .tsv are imported while all other are disregarded. It is possible to import files processed using other platforms as long as the files are tab-delimited with the extension .tsv and have identical column names as the ImmunoSEQs files (see `readImmunoSeq` manual for a list of column names).<sup>5</sup>

In the example below, `system.file("extdata", "TCRB_sequencing", package = "LymphoSeq")` refers to a path to a directory embedded within the LymphoSeq package containing an example data set of T cell receptor beta (TCRB) sequencing from the peripheral blood of a patient who underwent a bone marrow transplant. The files are named according to the day posttransplant and the immunophenotype of the T cells if sorted prior to sequencing. The function `readImmunoSeq` is used to import the files in as a list object where each file becomes a data frame. You can import all columns from each file by setting the `columns` parameter to "all" or list just those columns you are interested in as shown below. Be aware that Adaptive Biotechnologies has changed the column names of their files over time and if the headings of your files are not all the same, you will need to specify "all" or provide all variations of the column header as done for "count" and "frequencyCount" in the example below. If the directory you are pointing to contains subdirectories with additional sequencing files, you can choose to import those by setting the parameter `recursive` to TRUE, otherwise only the files in the top directory will be imported.

```
library(LymphoSeq)
```

```
## Loading required package: LymphoSeqDB
```

```
file.path <- system.file("extdata", "TCRB_sequencing", package = "LymphoSeq")
```

```
file.list <- readImmunoSeq(path = file.path,
                           columns = c("aminoAcid", "nucleotide", "count", "count (templates)",
                                         "count (reads)", "frequencyCount", "frequencyCount (%)",
                                         "estimatedNumberGenomes", "vFamilyName", "dFamilyName",
                                         "jFamilyName"),
                           recursive = FALSE)
```

Notice that each data frame listed in the `file.list` object is named according the ImmunoSEQ file name files. If different names are desired, you may rename the original .tsv files or assign `names(file.list)` to a new character vector of desired names within R.

```
names(file.list)
```

```
[1] "TCRB_Day0_Unsorted"      "TCRB_Day1320_CD8_CMV"
[3] "TCRB_Day1320_Unsorted"   "TCRB_Day32_Unsorted"
```

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<sup>1</sup>Warren, E. H. *et al. Blood* 2013;122:19–22.

<sup>2</sup><http://www.adaptivebiotech.com/immunoseq>

<sup>3</sup><http://mixcr.milaboratory.com>

<sup>4</sup><http://www.imgt.org/HighV-QUEST>

<sup>5</sup>Some functionality may be lost. Extensive testing with other platforms has not been performed.

```
[5] "TCRB_Day369_CD8_CMV"      "TCRB_Day369_Unsorted"
[7] "TCRB_Day83_CD8_CMV"       "TCRB_Day83_Unsorted"
[9] "TCRB_Day949_CD4"          "TCRB_Day949_CD8"
[11] "TCRB_Day949_Unsorted"
```

Having the data in the form of a list makes it easy to apply a function over that list using the base functions `sapply` (outputs a matrix) or `lapply` (outputs a list). For example, you may use the function `dim` to report the dimensions of each data frame as shown below. Notice that in the example data set, each data frame has less than 1000 rows (e.g. sequences) since it has been truncated from its original size for demonstration purposes. There are 8 columns to each data frame as specified by the `readImmunoSeq` function. In place of `dim`, you may also use `colnames`, `nrow`, `ncol`, or other more complex functions that perform operations on subsetting columns.

```
sapply(file.list, dim)
```

```
TCRB_Day0_Unsorted TCRB_Day1320_CD8_CMV TCRB_Day1320_Unsorted
[1,]                999                  40                  999
[2,]                 8                   8                   8
TCRB_Day32_Unsorted TCRB_Day369_CD8_CMV TCRB_Day369_Unsorted
[1,]                920                  414                  999
[2,]                 8                   8                   8
TCRB_Day83_CD8_CMV TCRB_Day83_Unsorted TCRB_Day949_CD4
[1,]                201                  999                  999
[2,]                 8                   8                   8
TCRB_Day949_CD8 TCRB_Day949_Unsorted
[1,]                999                  999
[2,]                 8                   8
```

## Subsetting data

If you imported all of the files from your project but just want to perform an analysis on a subset, use standard R methods to subset the list. Remember that a single bracket `[` returns a list and a double bracket `[[` returns a single data frame.

```
CMV <- file.list[grep("CMV", names(file.list))]
names(CMV)
```

```
[1] "TCRB_Day1320_CD8_CMV" "TCRB_Day369_CD8_CMV" "TCRB_Day83_CD8_CMV"
```

```
TCRB_Day0_Unsorted <- file.list[["TCRB_Day0_Unsorted"]]
head(TCRB_Day0_Unsorted)
```

```

nucleotide
1 TCAATTCCTGGAGCTTGGTGACTCTGCTGTGTATTTCTGTGCCAGCAGCCATCGGGACAGAGAACACTGAAGCTTTCTTTGGACAA
2 CTGATTCTGGAGTCCGCCAGCACCAACCAGACATCTATGTACCTCTGTGCCAGCAGCCCCGTGAGCAATGAGCAGTTCTTCGGGCCA
3 ATCAATTCCTGGAGCTTGGTGACTCTGCTGTGTATTTCTGTGCCAGCAGCCAAGAAGTTCCGCCTTACCAAGCTTTCTTTGGACAA
4 TGCCATCCCCAACAGACAGCTCTTTACTTCTGTGCCACCAAGTGTCCACAAACAGGGGGCAGGACCGGGGAGCTGTTTTTTGGAGAA
5 CACACCCTGCAGCCAGAAGACTCGGCCCTGTATCTCTGCGCCAGCAGCCAAGAGGCTAGCGGGGAGACAGACCCAGTACTTCGGGCCA
6 GCCAGCACCAACCAGACATCTATGTACCTCTGTGCCAGCAGTTTGGAGCACACGGGTGCAACTAATGAAAAACTGTTTTTTGGCAGT

aminoAcid count frequencyCount vFamilyName dFamilyName
1          89285          6.606637      TCRBV03      TCRBD01
2      CASSPVSNEQFF 50511          3.737558      TCRBV28      TCRBD02
3      CASSQEVPPYQAF 49129          3.635297      TCRBV03
4          43293          3.203462      TCRBV24      TCRBD01
5      CASSQEASGRQTQYF 43264          3.201317      TCRBV04      TCRBD02
6 CASSLEHTGATNEKLFF 40119          2.968602      TCRBV28      TCRBD02
jFamilyName estimatedNumberGenomes
1      TCRBJ01          1450
2      TCRBJ02           822
3      TCRBJ01           797
4      TCRBJ02           702
5      TCRBJ02           704
6      TCRBJ01           653
```

For more complex subsetting, you can use a metadata file where one column contains the file names and the other columns have additional information about the sample files (e.g pretreatment and posttreatment). You can then subset the metadata file using criteria from the other columns to give you just a character vector of file names that you can use to subset file.list.

```
metadata <- read.csv(system.file("extdata", "metadata.csv", package = "LymphoSeq"))
metadata
```

	samples	day	timePoint	phenotype
1	TCRB_Day0_Unsorted	0	Pretransplant	Unsorted
2	TCRB_Day32_Unsorted	32	1 Month	Unsorted
3	TCRB_Day83_Unsorted	83	3 Months	Unsorted
4	TCRB_Day83_CD8_CMV	83	3 Months	CD8+CMV+
5	TCRB_Day369_Unsorted	369	1 Year	Unsorted
6	TCRB_Day369_CD8_CMV	369	1 Year	CD8+CMV+
7	TCRB_Day949_Unsorted	949	2 Years	Unsorted
8	TCRB_Day949_CD4	949	2 Years	CD4+
9	TCRB_Day949_CD8	949	2 Years	CD8+
10	TCRB_Day1320_Unsorted	1320	3 Years	Unsorted
11	TCRB_Day1320_CD8_CMV	1320	3 Years	CD8+CMV+

```
selected <- as.character(metadata[metadata$phenotype == "Unsorted" &
                               metadata$day > 300, "samples"])
file.list.selected <- file.list[selected]
names(file.list.selected)
```

```
[1] "TCRB_Day369_Unsorted" "TCRB_Day949_Unsorted" "TCRB_Day1320_Unsorted"
```

## Extracting productive sequences

A productive sequence is defined as a sequences that is in frame and does not have an early stop codon. If you sequenced genomic DNA as opposed to complimentary DNA made from RNA, then you will have unproductive and productive sequences in your data files. Use the function `productiveSeq` to remove unproductive sequences and recompute the frequencyCount for each of your samples.

If you are interested in just the complementarity determining region 3 (CDR3) amino acid sequences, then set aggregate to "aminoAcid" and the count and estimated number of genomes for duplicate amino acid sequences will be summed. Note that the resulting list of data frames will have columns corresponding to "aminoAcid", "count", "frequencyCount", and "estimatedNumberGenomes" (if this column is available) only. All other columns, such as those corresponding to the V, D, and J gene names, will be removed if they were included in your original file list. The reason for this is to avoid confusion since a single amino acid CDR3 sequence may be encoded by multiple different nucleotide sequences with differing V, D, and J genes.

```
productive.aa <- productiveSeq(file.list = file.list, aggregate = "aminoAcid",
                               prevalence = FALSE)
```

Alternatively, you may set aggregate to "nucleotide" and the resulting list of data frames will all have the same columns as your original file list. Take note that some LymphoSeq functions require a productive sequence list aggregated by amino acid or nucleotide.

```
productive.nt <- productiveSeq(file.list = file.list, aggregate = "nucleotide",
                               prevalence = FALSE)
```

If the parameter `prevalence` is set to TRUE, then a new column is added to each of the data frames giving the prevalence (%) of each CDR3 amino acid sequence in 55 healthy donor peripheral blood samples. Values range from 0 to 100% where 100% means the sequence appeared in the blood of all 55 individuals. The data for this operation resides in a separate package that is automatically loaded called LymphoSeqDB. Please refer to that package manual for more details.

Notice in the example below that there are no amino acid sequences given in the first and fourth row of the file.list data frame for sample "TCRB\_Day949\_Unsorted". This is because the nucleotide sequence is out of frame and does not produce a productively transcribed amino acid sequence. If an asterisk (\*) appears in the amino acid sequences, this would indicate an early stop codon.

```
head(file.list[["TCRB_Day0_Unsorted"]])
```

nucleotide

```

1 TCAATTCCTGGAGCTTGGTGACTCTGCTGTGTATTTCTGTGCCAGCAGCCATCGGGACAGAGAACACTGAAGCTTTCTTTGGACAA
2 CTGATTCTGGAGTCCGCCAGCACCAACCAGACATCTATGTACCTCTGTGCCAGCAGCCCCGTGAGCAATGAGCAGTTCTTCGGGCCA
3 ATCAATTCCTGGAGCTTGGTGACTCTGCTGTGTATTTCTGTGCCAGCAGCCAAGAAGTTCCGCCTTACCAAGCTTTCTTTGGACAA
4 TGCCATCCCCAACAGACAGCTCTTTACTTCTGTGCCACCAGTGTCCACAAACAGGGGGCAGGACCGGGGAGCTGTTTTTTGGAGAA
5 CACACCCTGCAGCCAGAAGACTCGGCCCTGTATCTCTGCGCCAGCAGCCAAGAGGCTAGCGGGAGACAGACCCAGTACTTCGGGCCA
6 GCCAGCACCAACCAGACATCTATGTACCTCTGTGCCAGCAGTTTGGAGCACACGGGTGCAACTAATGAAAAACTGTTTTTTGGCAGT

```

	aminoAcid	count	frequencyCount	vFamilyName	dFamilyName
1		89285	6.606637	TCRBV03	TCRBD01
2	CASSPVSNEQFF	50511	3.737558	TCRBV28	TCRBD02
3	CASSQEVPPYQAFF	49129	3.635297	TCRBV03	
4		43293	3.203462	TCRBV24	TCRBD01
5	CASSQEASGRQTQYF	43264	3.201317	TCRBV04	TCRBD02
6	CASSLEHTGATNEKLFF	40119	2.968602	TCRBV28	TCRBD02

	jFamilyName	estimatedNumberGenomes
1	TCRBJ01	1450
2	TCRBJ02	822
3	TCRBJ01	797
4	TCRBJ02	702
5	TCRBJ02	704
6	TCRBJ01	653

After `productiveSeq` is run, the unproductive sequences are removed and the `frequencyCount` is recalculated for each sequence. If there were two identical amino acid sequences that differed in their nucleotide sequence, they would be combined and their counts added together.

```
head(productive.aa[["TCRB_Day0_Unsorted"]])
```

	aminoAcid	count	frequencyCount	estimatedNumberGenomes
1	CASSPVSNEQFF	50511	5.543769	822
2	CASSQEVPPYQAFF	49129	5.392090	797
3	CASSQEASGRQTQYF	43264	4.748384	704
4	CASSLEHTGATNEKLFF	40119	4.403209	653
5	CASSPGDEQYF	38100	4.181616	619
6	CSARSPSTGTLAEAFF	26377	2.894973	429

Finally, notice that the `productive.nt` data frame for sample "TCRB\_Day949\_Unsorted" below has additional columns not present in `productive.aa` but are in `file.list`. This is because the data frame was aggregated by nucleotide sequence and all of the original columns from `file.list` were carried over.

```
head(productive.nt[["TCRB_Day0_Unsorted"]])
```

						nucleotide
1	CTGATTCTGGAGTCCGCCAGCACCAACCAGACATCTATGTACCTCTGTGCCAGCAGCCCCGTGAGCAATGAGCAGTTCTTCGGGCCA					
2	ATCAATTCCTGGAGCTTGGTGACTCTGCTGTGTATTTCTGTGCCAGCAGCCAAGAAGTTCCGCCTTACCAAGCTTTCTTTGGACAA					
3	CACACCCTGCAGCCAGAAGACTCGGCCCTGTATCTCTGCGCCAGCAGCCAAGAGGCTAGCGGGAGACAGACCCAGTACTTCGGGCCA					
4	GCCAGCACCAACCAGACATCTATGTACCTCTGTGCCAGCAGTTTGGAGCACACGGGTGCAACTAATGAAAAACTGTTTTTTGGCAGT					
5	CCCCTGACCCTGGAGTCTGCCAGGCCCTCACATACCTCTCAGTACCTCTGTGCCAGCAGTCCGGGGGACGAGCAGTACTTCGGGGCCG					
6	AGTGCCCATCCTGAAGACAGCAGCTTCTACATCTGCAGTGCTAGATCACCCAGTACAGGGACCCCTCGCTGAAGCTTTCTTTGGACAA					

	aminoAcid	count	frequencyCount	vFamilyName	dFamilyName
1	CASSPVSNEQFF	50511	5.543769	TCRBV28	TCRBD02
2	CASSQEVPPYQAFF	49129	5.392090	TCRBV03	
3	CASSQEASGRQTQYF	43264	4.748384	TCRBV04	TCRBD02
4	CASSLEHTGATNEKLFF	40119	4.403209	TCRBV28	TCRBD02
5	CASSPGDEQYF	38100	4.181616	TCRBV25	TCRBD02
6	CSARSPSTGTLAEAFF	26377	2.894973	TCRBV20	TCRBD01

	jFamilyName	estimatedNumberGenomes
1	TCRBJ02	822
2	TCRBJ01	797
3	TCRBJ02	704
4	TCRBJ01	653
5	TCRBJ02	619
6	TCRBJ01	429

## Create a table of summary statistics

To create a table summarizing the total number of sequences, number of unique productive sequences, number of genomes, entropy, clonality, Gini coefficient, and the frequency (%) of the top productive sequence in each imported file, use the function `clonality`.

```
clonality(file.list = file.list)
```

	samples	totalSequences	uniqueProductiveSequences		
1	TCRB_Day949_CD4	999	845		
2	TCRB_Day949_CD8	999	796		
3	TCRB_Day0_Unsorted	999	837		
4	TCRB_Day83_CD8_CMV	201	122		
5	TCRB_Day32_Unsorted	920	767		
6	TCRB_Day369_CD8_CMV	414	281		
7	TCRB_Day83_Unsorted	999	830		
8	TCRB_Day1320_CD8_CMV	40	25		
9	TCRB_Day369_Unsorted	999	828		
10	TCRB_Day949_Unsorted	999	831		
11	TCRB_Day1320_Unsorted	999	833		
	totalGenomes	totalCount	entropy	clonality	giniCoefficient
1	25767	1795561	5.597259	0.42431656	0.8486265
2	26236	2161314	5.535828	0.42554289	0.9018374
3	18215	1158510	7.144318	0.26416150	0.7956718
4	254	4553	5.891883	0.14989093	0.6133629
5	NA	31078	8.296630	0.13424209	0.6007820
6	1794	52480	5.011083	0.38396606	0.8677666
7	NA	427427	7.285945	0.24863671	0.7100983
8	53	53	4.486348	0.03391748	0.2313514
9	NA	725668	6.037979	0.37710971	0.8043414
10	6547	1486480	6.774799	0.30147383	0.7739322
11	180079	180079	5.708193	0.41165830	0.8845652
	topProductiveSequence				
1	29.232143				
2	19.131268				
3	5.543769				
4	8.917656				
5	4.865016				
6	17.718550				
7	13.821673				
8	10.810811				
9	17.568430				
10	13.510800				
11	14.422142				

The clonality score is derived from the Shannon entropy, which is calculated from the frequencies of all productive sequences divided by the logarithm of the total number of unique productive sequences. This normalized entropy value is then inverted ( $1 - \text{normalized entropy}$ ) to produce the clonality metric.

The Gini coefficient<sup>6</sup> is an alternative metric used to calculate repertoire diversity and is derived from the Lorenz curve. The Lorenz curve is drawn such that x-axis represents the cumulative percentage of unique sequences and the y-axis represents the cumulative percentage of reads.<sup>7</sup> A line passing through the origin with a slope of 1 reflects equal frequencies of all clones. The Gini coefficient is the ratio of the area between the line of equality and the observed Lorenz curve over the total area under the line of equality.

Both Gini coefficient and clonality are reported on a scale from 0 to 1 where 0 indicates all sequences have the same frequency and 1 indicates the repertoire is dominated by a single sequence.

<sup>6</sup>[https://en.wikipedia.org/wiki/Gini\\_coefficient](https://en.wikipedia.org/wiki/Gini_coefficient)

<sup>7</sup>[https://en.wikipedia.org/wiki/Lorenz\\_curve](https://en.wikipedia.org/wiki/Lorenz_curve)

## Searching for sequences

To search for one or more amino acid or nucleotide CDR3 sequences in a list of data frames, use the function `searchSeq`. You may specify to search in either a list of productive or unproductive data frames.

```
searchSeq(list = productive.aa, sequence = "CASSPVSNEQFF", type = "aminoAcid",
          match = "global", editDistance = 0)
```

	sample	aminoAcid	count	frequencyCount
1	TCRB_Day0_Unsorted	CASSPVSNEQFF	50511	5.54376923
260	TCRB_Day369_Unsorted	CASSPVSNEQFF	211	0.03315421
64	TCRB_Day83_CD8_CMV	CASSPVSNEQFF	10	0.35958288
334	TCRB_Day949_CD8	CASSPVSNEQFF	214	0.01159812
	estimatedNumberGenomes			
1		822		
260		0		
64		1		
334		2		

If you have only a partial sequence, set the parameter `match` to `"partial"`. If you are looking for related sequences that differ by one or more nucleotides or amino acids, then increase the `editDistance` value. Edit distance is a way of quantifying how dissimilar two sequences are to one another by counting the minimum number of operations required to transform one sequence into the other. For example, an edit distance of 0 means the sequences are identical and an edit distance of 1 indicates that the sequences differ by a single amino acid or nucleotide.

## Searching for published T cell sequences with known antigen specificity

To search your entire list of data frames for a published amino acid CDR3 TCRB sequence with known antigen specificity, use the function `searchPublished`.

```
published <- searchPublished(list = productive.aa)
head(published)
```

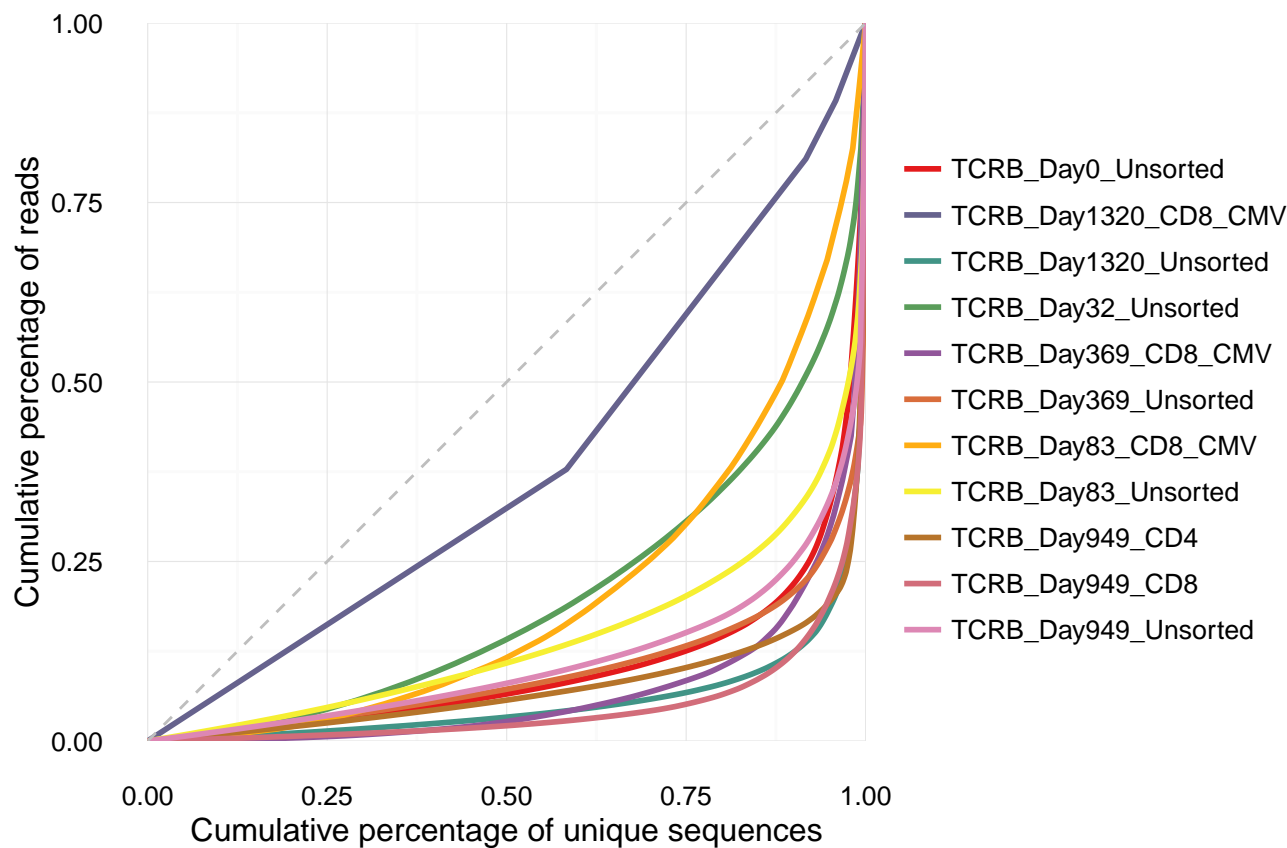
	sample	aminoAcid	count	frequencyCount			
1	TCRB_Day32_Unsorted	CASASSGTDQYF	33	0.131594688			
2	TCRB_Day949_Unsorted	CASSFSTDTQYF	279	0.023538126			
3	TCRB_Day1320_Unsorted	CASSIRSAYEQYF	15	0.010202902			
4	TCRB_Day949_CD8	CASSIRSAYEQYF	593	0.032138727			
5	TCRB_Day32_Unsorted	CASSLAPSYEQYF	17	0.067791203			
6	TCRB_Day1320_Unsorted	CASSLGEQPQHF	10	0.006801934			
	estimatedNumberGenomes	PMID	HLA	antigen	epitope		
1	0	20647322	HLA-A*24:02	Leukemia	<NA>		
2	2	23267020	HLA-A*02	EBV	BMFL1-GLCTLVAML		
3	15	21048112	HLA-A*02	EBV	BMLF1-GLCTLVAML		
4	7	21048112	HLA-A*02	EBV	BMLF1-GLCTLVAML		
5	0	23267020	HLA-B*08	EBV	BZLF1-RAKFKQLL		
6	10	23521884	HLA-B*27:05	HIV	KK10-KRWIILGLNK		
	prevalence						
1	7.3						
2	85.5						
3	18.2						
4	18.2						
5	69.1						
6	29.1						

For each found sequence, a table is provided listing the antigen, epitope, HLA type, PubMed ID (PMID), and prevalence (%) of the sequence among 55 healthy donor blood samples. The data for this function resides in the separate `LymphoSeqDB` package that is automatically loaded when the function is called. Please refer to that package manual for more details.

## Visualizing repertoire diversity

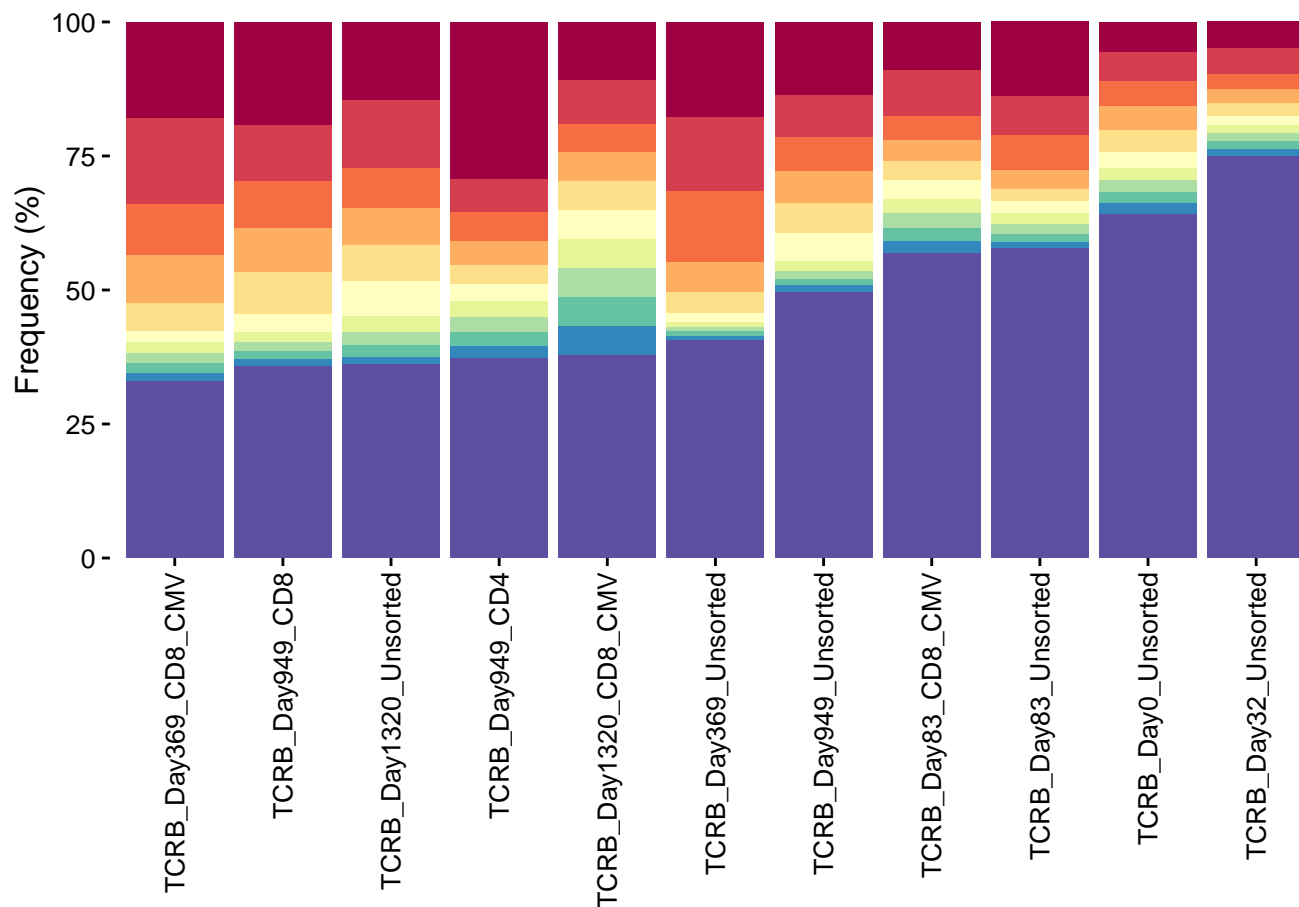
Antigen receptor repertoire diversity can be characterized by a number such as clonality or Gini coefficient calculated by the `clonality` function. Alternatively, you can visualize the repertoire diversity by plotting the Lorenz curve for each sample as defined above. In this plot, the more diverse samples will appear near the dotted diagonal line (the line of equality) whereas the more clonal samples will appear to have a more bowed shape.

```
lorenzCurve(samples = names(productive.aa), list = productive.aa)
```



Alternatively, you can get a feel for the repertoire diversity by plotting the cumulative frequency of a selected number of the top most frequent clones using the function `topSeqsPlot`. In this case, each of the top sequences are represented by a different color and all less frequent clones will be assigned a single color (violet).

```
topSeqsPlot(list = productive.aa, top = 10)
```



Both of these functions are built using the `ggplot2` package<sup>8</sup>. You can reformat the plot using `ggplot2` functions. Please refer to the `lorenzCurve` and `topSeqsPlot` manual for specific examples.

## Comparing samples

To compare the T or B cell repertoires of all samples in a pairwise fashion, use the `bhattacharyyaMatrix` or `similarityMatrix` functions. Both the Bhattacharyya coefficient and similarity score are measures of the amount of overlap between two samples. The value for each ranges from 0 to 1 where 1 indicates the sequence frequencies are identical in the two samples and 0 indicates no shared frequencies. The Bhattacharyya coefficient differs from the similarity score in that it involves weighting each shared sequence in the two distributions by the arithmetic mean of the frequency of each sequence, while calculating the similarity scores involves weighting each shared sequence in the two distributions by the geometric mean of the frequency of each sequence in the two distributions.

```
bhattacharyya.matrix <- bhattacharyyaMatrix(productive.seqs = productive.aa)
bhattacharyya.matrix[,1:2]
```

	TCRB_Day1320_Unsorted	TCRB_Day949_Unsorted
TCRB_Day1320_Unsorted	1.00000000	0.88013589
TCRB_Day949_Unsorted	0.88013589	1.00000000
TCRB_Day369_Unsorted	0.84210238	0.77225070
TCRB_Day1320_CD8_CMV	0.46204244	0.39421126
TCRB_Day83_Unsorted	0.62582845	0.58334619
TCRB_Day369_CD8_CMV	0.79605192	0.72972153
TCRB_Day32_Unsorted	0.32118959	0.27850843
TCRB_Day83_CD8_CMV	0.42168018	0.40854427
TCRB_Day0_Unsorted	0.01700697	0.01416233
TCRB_Day949_CD8	0.81836252	0.81470456
TCRB_Day949_CD4	0.44279429	0.42687836

<sup>8</sup><http://www.cookbook-r.com/Graphs/>

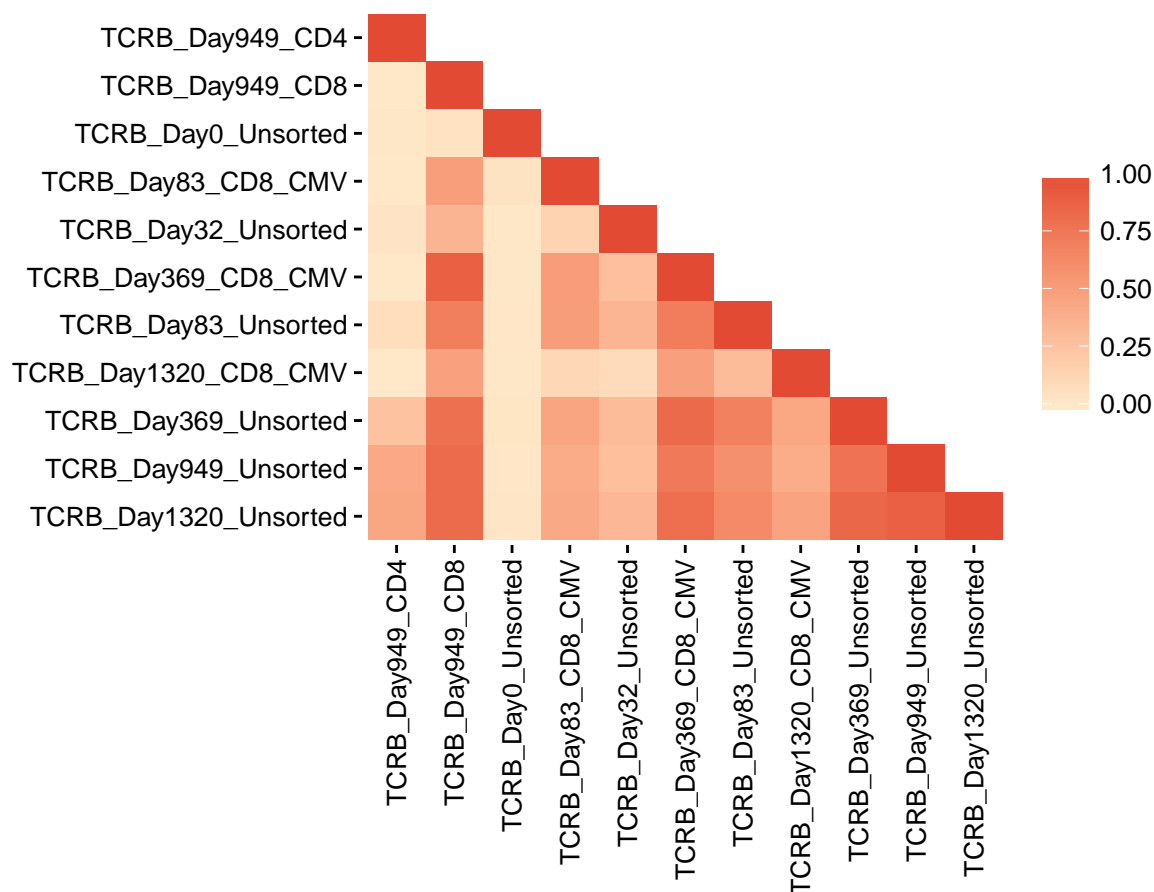


```
similarity.matrix <- similarityMatrix(productive.seqs = productive.aa)
similarity.matrix[,1:2]
```

	TCRB_Day1320_Unsorted	TCRB_Day949_Unsorted
TCRB_Day1320_Unsorted	1.00000000	0.85741424
TCRB_Day949_Unsorted	0.85741424	1.00000000
TCRB_Day369_Unsorted	0.84852643	0.78458455
TCRB_Day1320_CD8_CMV	0.48448189	0.37158455
TCRB_Day83_Unsorted	0.62977574	0.58472269
TCRB_Day369_CD8_CMV	0.75230736	0.59370996
TCRB_Day32_Unsorted	0.40554581	0.37243182
TCRB_Day83_CD8_CMV	0.43445840	0.39263374
TCRB_Day0_Unsorted	0.01231397	0.02166242
TCRB_Day949_CD8	0.91435304	0.85504038
TCRB_Day949_CD4	0.80667431	0.54676809

The results of either function can be visualized by the `pairwisePlot` function.

```
pairwisePlot(matrix = bhattacharyya.matrix)
```



To view sequences shared between two or more samples, use the function `commonSeqs`. This function requires that a productive amino acid list be specified.

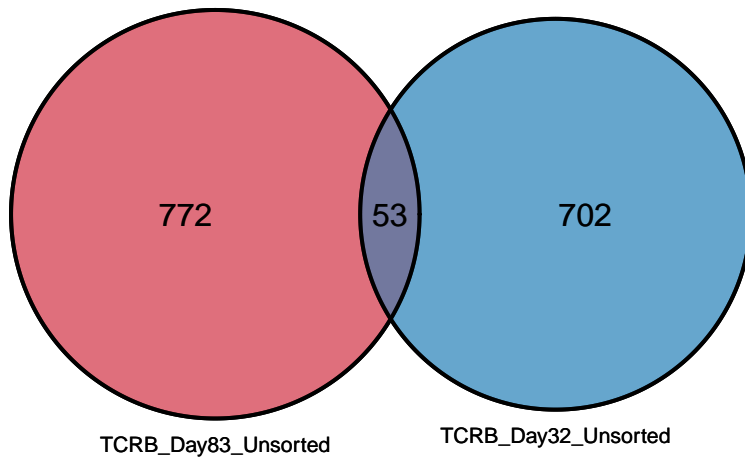
```
common <- commonSeqs(samples = c("TCRB_Day0_Unsorted", "TCRB_Day32_Unsorted"),
                      productive.aa = productive.aa)
head(common)
```

	aminoAcid	TCRB_Day0_Unsorted	TCRB_Day32_Unsorted
1	CASSQDRIGQYGYTF	0.47380673	0.80551900
2	CAWTGGTTEAFF	0.10887567	0.15153328
3	CAISEGNYGYTF	0.03566995	0.18742274

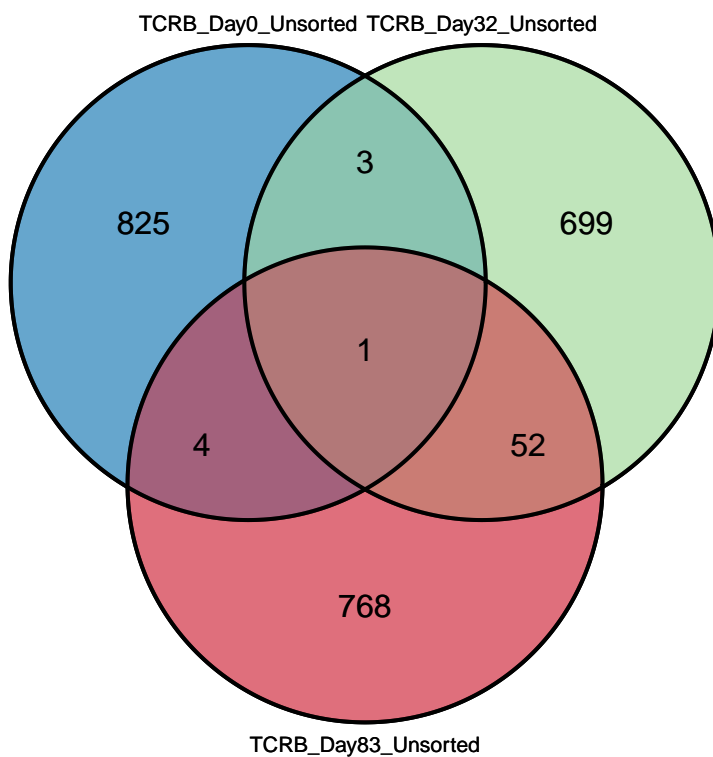
4 CASSFGIQETQYF 0.01393872 0.09570523

To visualize the number of overlapping sequences between two or three samples in the form of a Venn diagram, use the function `commonSeqVenn`.

```
commonSeqsVenn(samples = c("TCRB_Day32_Unsorted", "TCRB_Day83_Unsorted"),  
               productive.seqs = productive.aa)
```

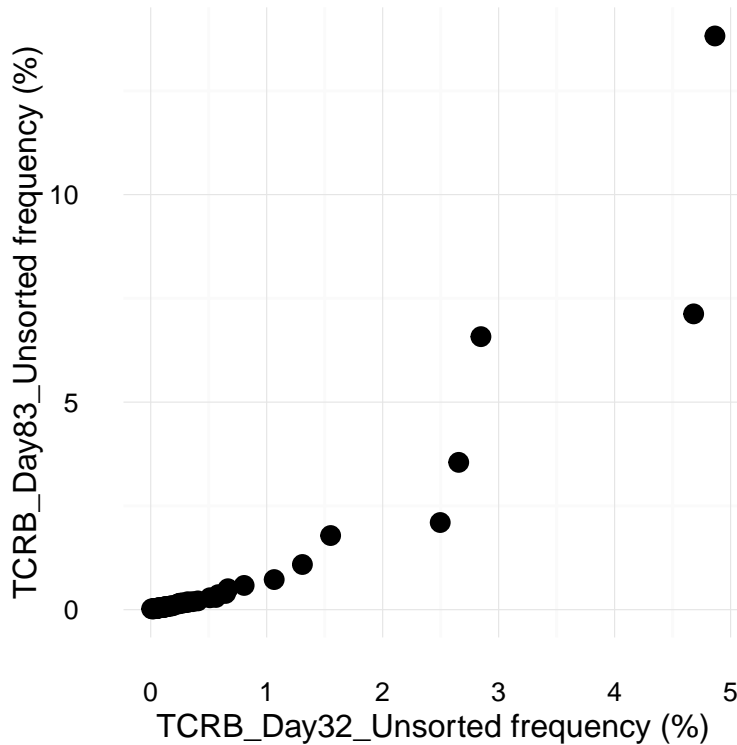


```
commonSeqsVenn(samples = c("TCRB_Day0_Unsorted", "TCRB_Day32_Unsorted", "TCRB_Day83_Unsorted"),  
               productive.seqs = productive.aa)
```



To compare the frequency of sequences between two samples as a scatter plot, use the function `commonSeqsPlot`.

```
commonSeqsPlot("TCRB_Day32_Unsorted", "TCRB_Day83_Unsorted",
  productive.aa = productive.aa)
```



## Revealing sequences that appear in multiple samples

To create a data frame of unique, productive amino acid sequences as rows and sample names as headers use the `seqMatrix` function. Each value in the data frame represents the frequency that each sequence appears in the sample. You can specify your own list of sequences or all unique sequences in the list using the output of the function `uniqueSeqs`. The `uniqueSeqs` function creates a data frame of all unique, productive sequences and reports the total count in all samples.

```
unique.seqs <- uniqueSeqs(productive.aa = productive.aa)
head(unique.seqs)
```

	aminoAcid	count
2465	CASSPAGAYYNEQFF	666188
1070	CASSESAGSTGELFF	430262
2699	CASSPPTGERDTQYF	358432
2998	CASSQDLMTVDSL FAGANVLTF	357006
2154	CASSLQGREKLFF	320305
3103	CASSQDWERLGEQFF	307744

```
sequence.matrix <- seqMatrix(productive.aa = productive.aa, sequences = unique.seqs$aminoAcid)
head(sequence.matrix)[1:6]
```

	aminoAcid	numberSamples	TCRB_Day0_Unsorted
1	CASSPAGAYYNEQFF	9	0.0000000
2	CASSPPTGERDTQYF	9	0.0000000
3	CASSQDLMTVDSL FAGANVLTF	9	0.0000000
4	CASSQDRTGQYGYTF	9	0.4738067
5	CASSLQGREKLFF	8	0.0000000
6	CASSQDSSDTEAFF	8	0.0000000

	TCRB_Day1320_CD8_CMV	TCRB_Day1320_Unsorted	TCRB_Day32_Unsorted
1	8.108108	14.4459484	0.03987718
2	5.405405	12.8039614	0.40674722
3	2.702703	3.0853575	1.55122224

4	0.000000	0.4278417	0.80551900
5	0.000000	7.2937143	4.86501575
6	0.000000	0.1285566	0.66594888

If just the top clones with a frequency greater than a specified amount are of interest to you, then use the `topFreq` function. This creates a data frame of the top productive amino acid sequences having a minimum specified frequency and reports the minimum, maximum, and mean frequency that the sequence appears in a list of samples. For TCRB sequences, the prevalence (%) and the published antigen specificity of that sequence are also provided.

```
top.freq <- topFreq(productive.aa = productive.aa, percent = 0.1)
head(top.freq)
```

	aminoAcid	minFrequency	maxFrequency	meanFrequency
373	CASSPPTGERDTQYF	0.4067472	17.798977	8.0931668
412	CASSQDLMTVDSLAFGANVLTF	1.5512222	10.542803	5.6504197
418	CASSQDRTGQYGYTF	0.4278417	1.088690	0.6955508
342	CASSPAGAYYNEQFF	0.3955412	19.139614	11.6126951
296	CASSLQGREKLFF	4.0790673	13.821673	7.3429132
556	CASSWPGLASFNEQFF	0.2489508	5.405405	1.0181321

	numberSamples	prevalence	antigen
373	9	1.8	
412	9	1.8	
418	9	3.6	
342	8	5.5	
296	8	30.9	
556	8	1.8	

One very useful thing to do is merge the output of `seqMatrix` and `topFreq`.

```
top.freq <- topFreq(productive.aa = productive.aa, percent = 0)
top.freq.matrix <- merge(top.freq, sequence.matrix)
head(top.freq.matrix)[1:12]
```

	aminoAcid	numberSamples	minFrequency	maxFrequency	meanFrequency
1	CAAGDTTLYEQYF	1	0.01799961	0.01799961	0.01799961
2	CAAGRDLNIQYF	1	0.03950662	0.03950662	0.03950662
3	CAAGTSGDTQYF	1	0.01830743	0.01830743	0.01830743
4	CAARGGESYEYF	1	0.03654798	0.03654798	0.03654798
5	CAATRRQGDVMNTEAFF	1	0.03771095	0.03771095	0.03771095
6	CACSRDRGSDTQYF	1	0.01957292	0.01957292	0.01957292

	prevalence	antigen	TCRB_Day0_Unsorted	TCRB_Day1320_CD8_CMV
1	0.0		0.01799961	0
2	0.0		0.00000000	0
3	1.8		0.00000000	0
4	0.0		0.03654798	0
5	0.0		0.00000000	0
6	0.0		0.00000000	0

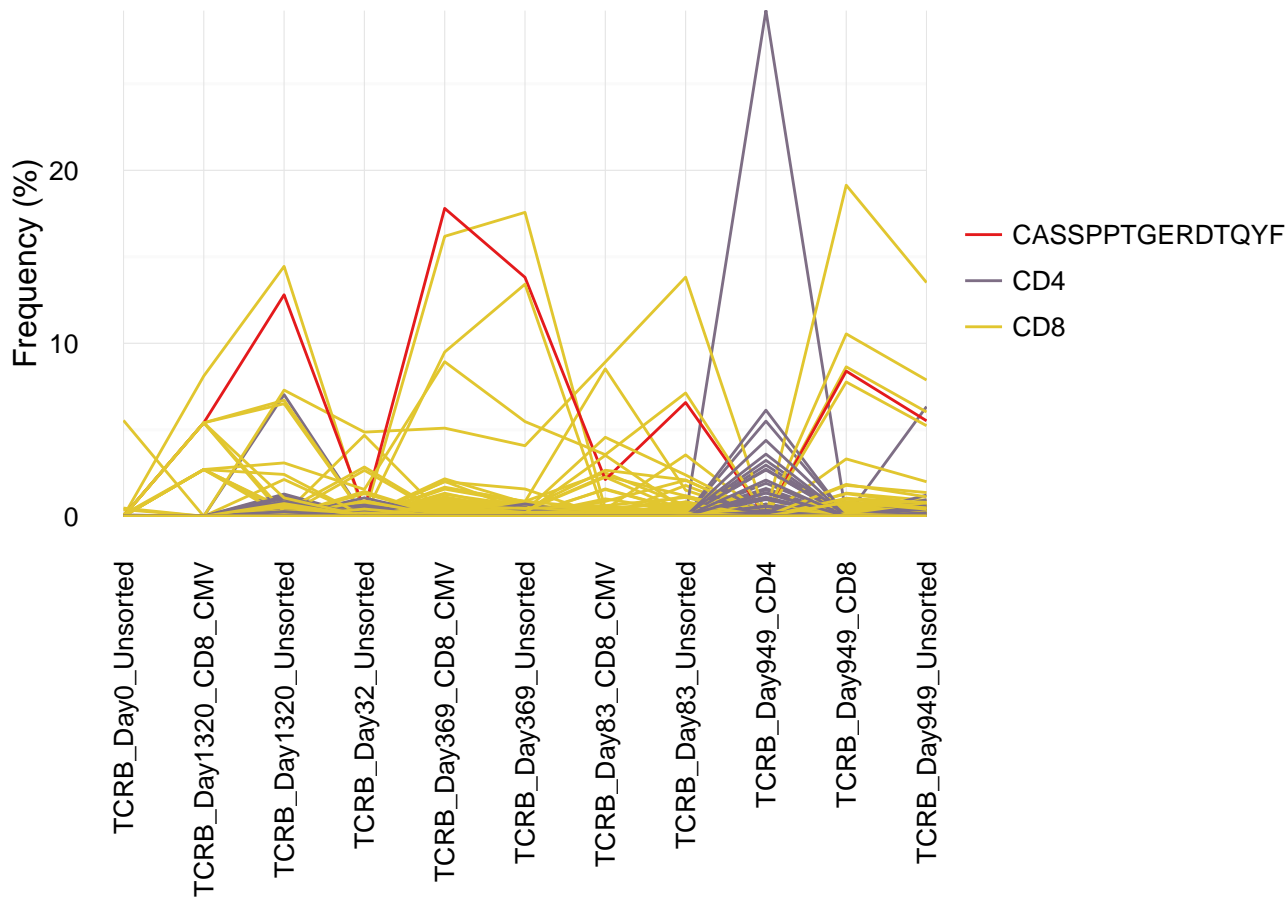
	TCRB_Day1320_Unsorted	TCRB_Day32_Unsorted	TCRB_Day369_CD8_CMV
1	0	0	0
2	0	0	0
3	0	0	0
4	0	0	0
5	0	0	0
6	0	0	0

## Tracking sequences across samples

To visually track the frequency of sequences across multiple samples, use the function `cloneTrack`. This function takes the output from the `seqMatrix` function. You can specify a character vector of amino acid sequences using the parameter `track` to highlight those sequences with a different color. Alternatively, you can highlight all of the sequences from a given sample using the parameter `map`. If the mapping feature is use, then you must specify a productive amino acid list and a character

vector of labels to title the mapped samples. To hide sequences that are not being tracked or mapped, set `unassigned` to `FALSE`.

```
cloneTrack(sequence.matrix = sequence.matrix,
  productive.aa = productive.aa,
  map = c("TCRB_Day949_CD4", "TCRB_Day949_CD8"),
  label = c("CD4", "CD8"),
  track = "CASSPPTGERDTQYF",
  unassigned = FALSE)
```



Refer to the `cloneTrack` manual for examples on how to reformat the chart using `ggplot2` function.

## Comparing V(D)J gene usage

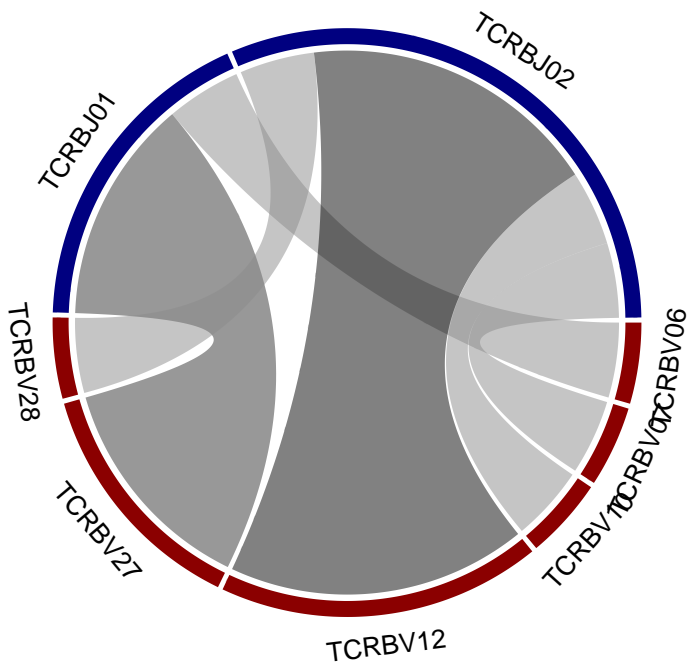
To compare the V, D, and J gene usage across samples, start by creating a data frame of V, D, and J gene counts and frequencies using the function `geneFreq`. You can specify if you are interested in the “VDJ”, “DJ”, “VJ”, “DJ”, “V”, “D”, or “J” loci using the `locus` parameter. Set `family` to `TRUE` if you prefer the family names instead of the gene names as reported by ImmunoSeq.

```
vGenes <- geneFreq(productive.nt = productive.nt, locus = "V", family = TRUE)
head(vGenes)
```

	samples	familyName	count	frequencyGene
1	TCRB_Day949_Unsorted	TCRBV02	16707	1.409503
2	TCRB_Day949_Unsorted	TCRBV03	44147	3.724508
3	TCRB_Day949_Unsorted	TCRBV04	176436	14.885207
4	TCRB_Day949_Unsorted	TCRBV05	89025	7.510687
5	TCRB_Day949_Unsorted	TCRBV06	97462	8.222483
6	TCRB_Day949_Unsorted	TCRBV07	106095	8.950815

To create a chord diagram showing VJ or DJ gene associations from one or more more samples, combine the output of `geneFreq` with the function `chordDiagramVDJ`. This function works well the `topSeqs` function that creates a data frame of a selected number of top productive sequences. In the example below, a chord diagram is made showing the association between V and J genes of just the single dominant clones in each sample. The size of the ribbons connecting VJ genes correspond to the number of samples that have that recombination event. The thicker the ribbon, the higher the frequency of the recombination.

```
top.seqs <- topSeqs(productive.seqs = productive.nt, top = 1)
chordDiagramVDJ(sample = top.seqs,
  association = "VJ",
  colors = c("darkred", "navyblue"))
```

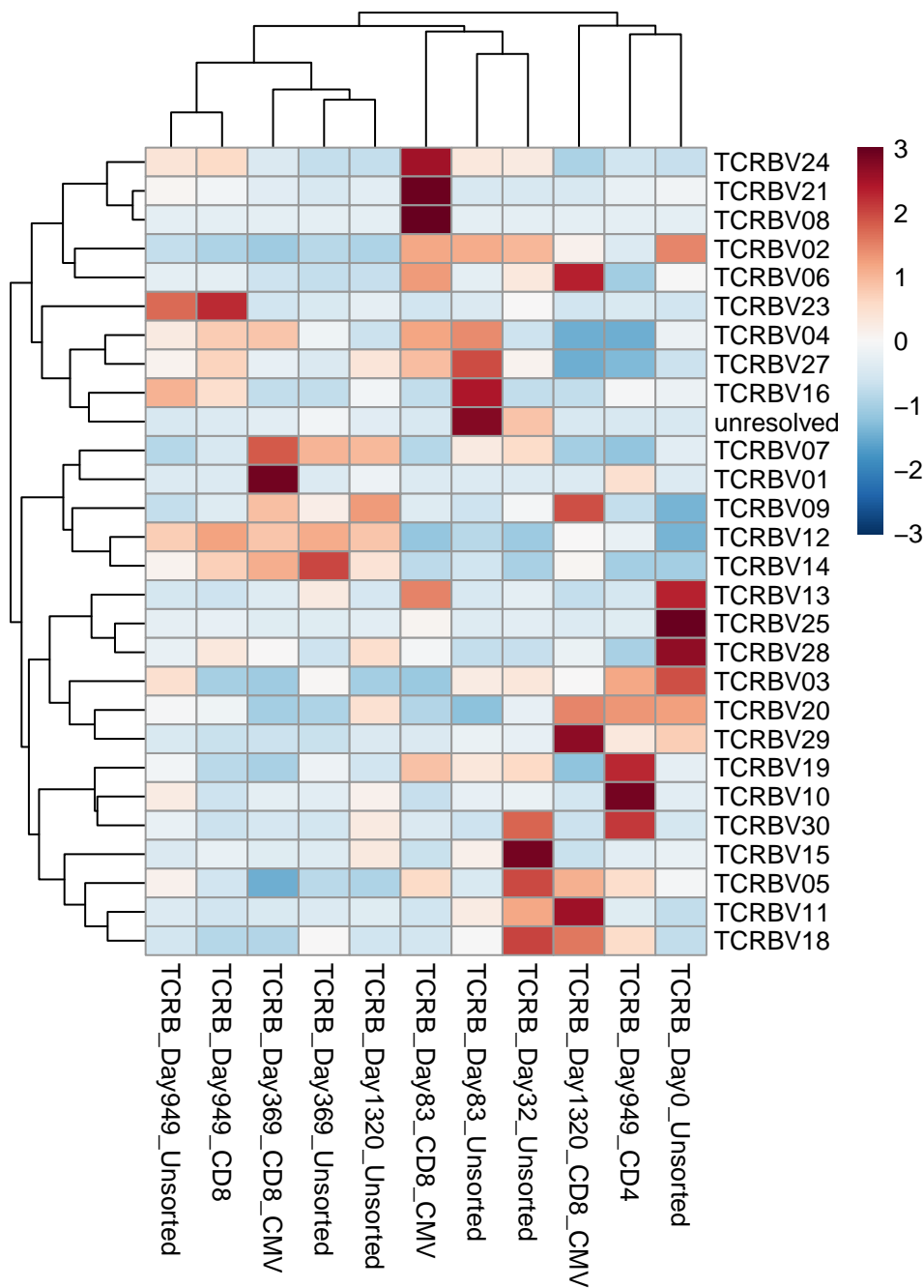


You can also visualize the results of `geneFreq` as a heat map, word cloud, or cumulative frequency bar plot with the support of additional R packages as shown below.

```
vGenes <- geneFreq(productive.nt = productive.nt, locus = "V", family = TRUE)
library(RColorBrewer)
library(grDevices)
RedBlue <- grDevices::colorRampPalette(rev(RColorBrewer::brewer.pal(11, "RdBu")))(256)
library(wordcloud)
wordcloud::wordcloud(words = vGenes[vGenes$samples == "TCRB_Day83_Unsorted", "familyName"],
  freq = vGenes[vGenes$samples == "TCRB_Day83_Unsorted", "frequencyGene"],
  colors = RedBlue)
```

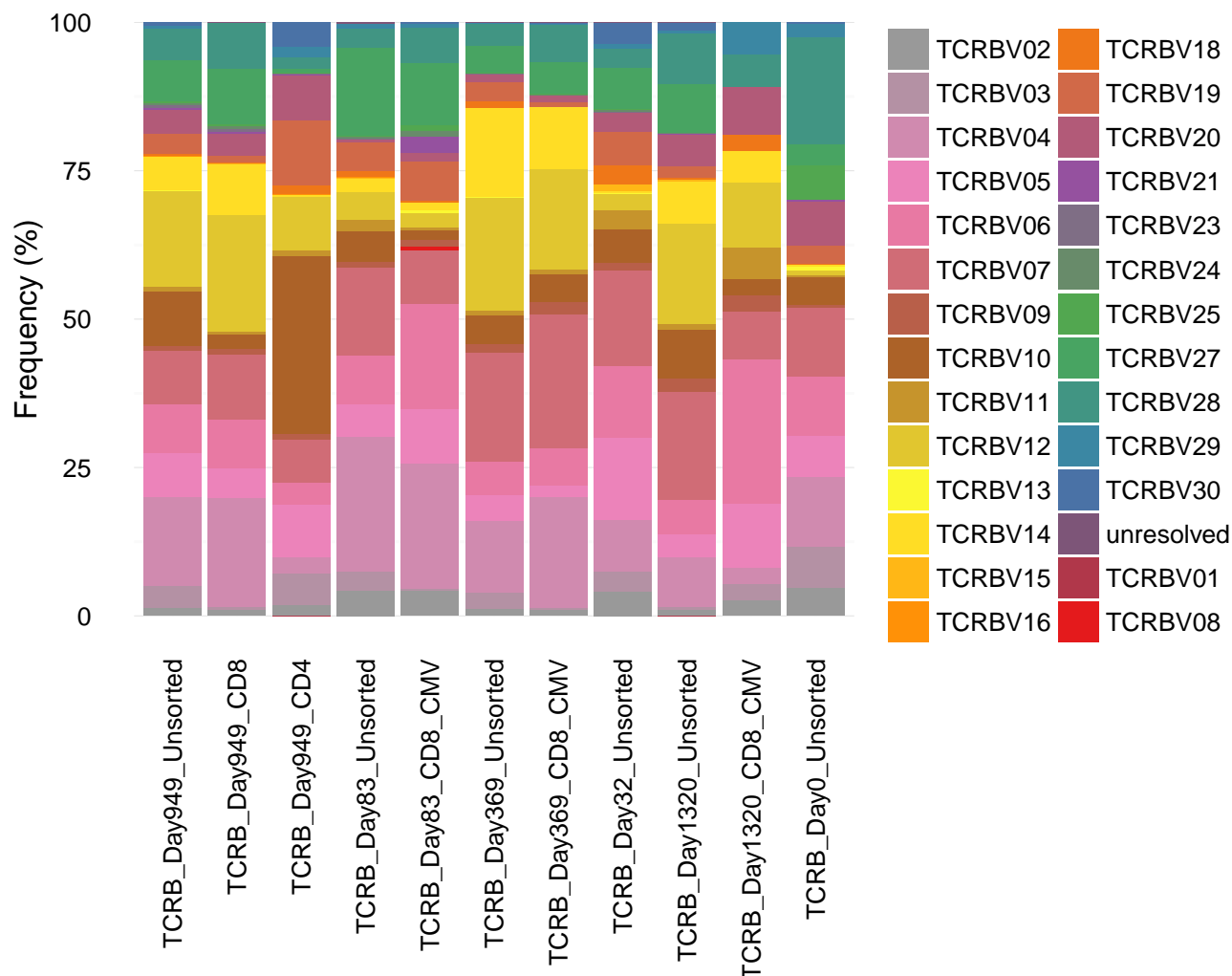


```
library(reshape)
vGenes <- reshape::cast(vGenes, familyName ~ samples, value = "frequencyGene", sum)
rownames(vGenes) = as.character(vGenes$familyName)
vGenes$familyName = NULL
library(pheatmap)
pheatmap::pheatmap(vGenes, color = RedBlue, scale = "row")
```



```
vGenes <- geneFreq(productive.nt = productive.nt, locus = "V", family = TRUE)
library(ggplot2)
multicolors <- grDevices::colorRampPalette(rev(RColorBrewer::brewer.pal(9, "Set1")))(28)
ggplot2::ggplot(vGenes, aes(x = samples, y = frequencyGene, fill = familyName)) +
  geom_bar(stat = "identity") +
  theme_minimal() +
  scale_y_continuous(expand = c(0, 0)) +
  guides(fill = guide_legend(ncol = 2)) +
  scale_fill_manual(values = multicolors) +
  labs(y = "Frequency (%)", x = "", fill = "") +
  theme(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust = 1))
```





## Removing sequences from the dataset

Occasionally you may identify one or more sequences in your data set that appear to be contamination. You can remove an amino acid sequence from all data frames using the function `removeSeq` and recompute `frequencyCount` for all remaining sequences.

```
searchSeq(list = productive.aa, sequence = "CASSDLIGNGKLFF")
```

	sample	aminoAcid	count	frequencyCount
49	TCRB_Day0_Unsorted	CASSDLIGNGKLFF	3494	0.383479434
668	TCRB_Day949_CD8	CASSDLIGNGKLFF	82	0.004444141
	estimatedNumberGenomes			
49		55		
668		1		

```
cleansed <- removeSeq(file.list = productive.aa, sequence = "CASSDLIGNGKLFF")
searchSeq(list = cleansed, sequence = "CASSDLIGNGKLFF")
```

No sequences found.

## Merging samples

If you need to combine multiple samples into one, use the `mergeFiles` function. It merges two or more sample data frames into a single data frame and aggregates count, frequencyCount, and estimatedNumberGenomes.

```
TCRB_Day949_Merged <- mergeFiles(samples = c("TCRB_Day949_CD4", "TCRB_Day949_CD8"),
                                   file.list = file.list)
```

## Conclusion

Advances in high-throughput sequencing have enabled characterizing T and B lymphocyte repertoires with unprecedented depth. LymphoSeq was developed as a tool to assist in the analysis of targeted next generation sequencing of the hypervariable CDR3 region of T and B cell receptors. The three key features of this R package are to characterize lymphocyte repertoire diversity, compare two or more lymphocyte repertoires, and track the frequency of CDR3 sequences across multiple samples. LymphoSeq also provides the unique ability to search for sequences in a curated database of published TCRB sequences with known antigen specificity. Finally, LymphoSeq can assign the percent prevalence that any given TCRB sequence appears in a the peripheral blood in healthy population of donors.

## Session info

```
sessionInfo()
```

```
## R version 3.3.1 (2016-06-21)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
## Running under: Windows Server 2012 R2 x64 (build 9600)
##
## locale:
## [1] LC_COLLATE=C
## [2] LC_CTYPE=English_United States.1252
## [3] LC_MONETARY=English_United States.1252
## [4] LC_NUMERIC=C
## [5] LC_TIME=English_United States.1252
##
## attached base packages:
## [1] stats      graphics  grDevices  utils      datasets  methods    base
##
## other attached packages:
## [1] ggplot2_2.1.0      pheatmap_1.0.8      reshape_0.8.5
## [4] wordcloud_2.5      RColorBrewer_1.1-2  LymphoSeq_1.2.0
## [7] LymphoSeqDB_0.99.2
##
## loaded via a namespace (and not attached):
## [1] Rcpp_0.12.7          knitr_1.14           magrittr_1.5
## [4] munsell_0.4.3        colorspace_1.2-7     R6_2.2.0
## [7] ineq_0.2-13          dplyr_0.5.0          stringr_1.1.0
## [10] plyr_1.8.4           tools_3.3.1          grid_3.3.1
## [13] data.table_1.9.6     gtable_0.2.0         circlize_0.3.9
## [16] DBI_0.5-1            lambda.r_1.1.9       futile.logger_1.4.3
## [19] htmltools_0.3.5      lazyeval_0.2.0       yaml_2.1.13
## [22] assertthat_0.1       digest_0.6.10        tibble_1.2
## [25] formatR_1.4          GlobalOptions_0.0.10 futile.options_1.0.0
## [28] slam_0.1-38          shape_1.4.2          VennDiagram_1.6.17
## [31] evaluate_0.10        rmarkdown_1.1        labeling_0.3
## [34] stringi_1.1.2        scales_0.4.0         chron_2.3-47
```