Chi Squared Tests for Heterogeneity

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Set Up

This analysis requires data on the counts of each 3mer, 5mer, and 7mer polymorphism type private to each of the 1,000 genomes nonadmixed continental groups (AFR, EUR, EAS, and SAS). All of this data has been preprocessed using the process_chrom_counts function in "code/data_wrangling/process_chrom_counts.R". I call these "count dataframes." An example is shown here:

##		Context	X3mer	X1mer	Count	Rate	<pre>context_in_genome</pre>	chr1	chr2
##	1	AAAAA->C	AAA->C	A->C	13218	1.886982e-09	16034992	1110	618
##	2	AAAAA->G	AAA->G	A->G	21558	3.077588e-09	16034992	1824	970
##	3	AAAAA->T	AAA->T	A->T	8686	1.240000e-09	16034992	723	359
##	4	AAAAC->C	AAA->C	A->C	3453	1.523554e-09	5188115	277	154
##	5	AAAAC->G	AAA->G	A->G	4190	1.848737e-09	5188115	317	204

Pairwise Chi Squared Tests

This section details how to perform the pairwise chi squared tests from Harris 2015. These steps were mostly used for replication.

Methodology

Two R functions that I use in this analysis:

- pairwise.chi Given two count dataframes, output a dataframe of chi-squared test results for each context. The arguement 'filter' (set by default to be true), will output "NA" as the p-value for any test for which the chi squared assumptions may not be correct.
- volcano.plot Given two count dataframes, plus the output from pairwise.chi, construct a volcano plot as in Harris 2015. Also takes the arguement lab.lim, which determines the lower p-value limit for which polymorphisms types should be labeled.

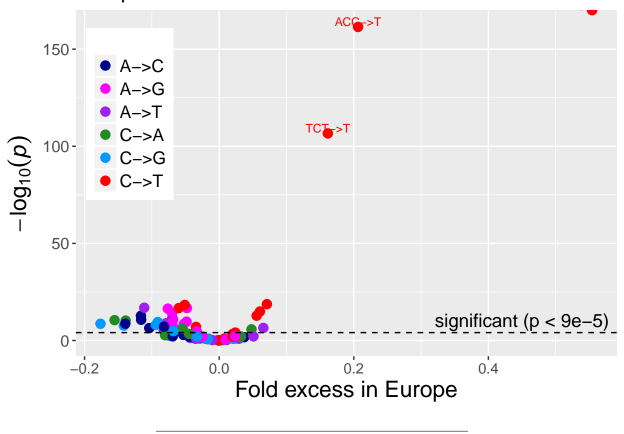
The code used to defint pairwise.chi is shown below:

```
# calculates homogeneity test p values for pairwise comparisons of two dfs of counts
pairwise.chi <- function(counts.1, counts.2, filter = T){</pre>
  n.contexts = length(counts.1$Context)
  result <- data.frame(matrix(ncol=2,nrow=n.contexts))</pre>
  colnames(result) <- c("Context", "p")</pre>
  result$Context <- counts.1$Context
  sum.1 <- sum(counts.1$Count)</pre>
  sum.2 <- sum(counts.2$Count)</pre>
  for (i in 1:n.contexts){
    c.a <- c(counts.1$Count[i], counts.2$Count[i])</pre>
    c.b \leftarrow c(sum.1, sum.2) - c.a
    data <- cbind(c.a, c.b)
    warning <- is(tryCatch(chisq.test(data), warning = function(w) w), "warning")</pre>
    if (filter == T & warning){
      result$p[i] <- NA
    else result$p[i] <- chisq.test(data)$p.value</pre>
  return(result)
```

Examples

As previously mentioned, these methods are not central to my analysis and mostly important for replication. It's worthwhile to note that these functions and data replicate volcano plots like the ones from Kelley Harris's leading figure in PNAS 2015:

Europe v. Africa



Fourway Tests for Homogeneity

In order to lighten the multiple hypothesis testing burden of running $\binom{4}{2} = 6$ pairwise comparisons for each possible polymorphism type, we switched to a homogeneity testing framework, which helps us rank polymorphism types based on how much they vary between populations. This is the dominant analysis technique we use to identify polymorphisms which are heterogeneous across continental groups.

Methodology

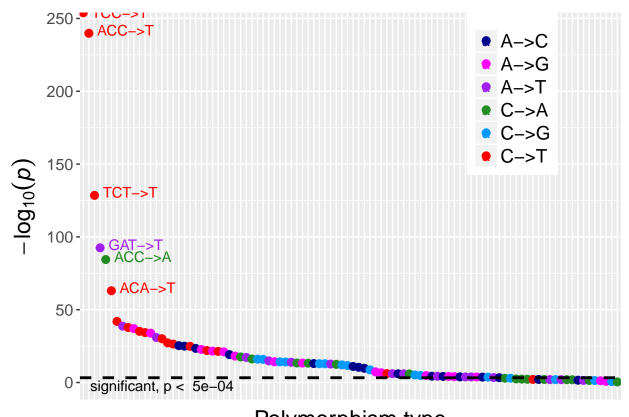
This section defines one function for calculation and two for visualization. Again, I'm hiding the code for the plotting functions in the compiled report because it's not essential to understanding.

- fourway.chi Given four count dataframes, output a dataframe of chi-squared test results for each context.
- hom.test.plot Given the output from fourway.chi, construct a volcano plot as in Harris 2015. Also takes the arguement lab.lim, which determines the lower p-value limit for which polymorphisms types should be labeled, and the boolean, NoTCC, which, when True, leaves out labels for any polymorphism with the 3mer subcontext TCC->T.
- sigs.plot Given the same arguments as hom.test.plot, make a plot of just the significant results.

The r code used to define fourway.chi is shown below:

```
# calculates homogeneity test p values for Fourway comparisons of counts dfs
fourway.chi <- function(AFR, EUR, EAS, SAS, filter = T){</pre>
  n.contexts = length(AFR$Context)
  # make dataframe for results
  result <- data.frame(matrix(ncol=9,nrow=n.contexts))</pre>
  colnames(result) <- c("Context", "X5mer", "X3mer", "X1mer",</pre>
                         "AFR.Count", "EUR.Count", "EAS.Count", "SAS.Count", "p")
  result$Context <- AFR$Context
  result$X5mer <- AFR$X5mer # for smaller contexts, X3mer and X5mer columns do not exist,
  result$X3mer <- AFR$X3mer # and will disappear at this step
  result$X1mer <- AFR$X1mer
  result$AFR.Count <- AFR$Count; result$EUR.Count <- EUR$Count
  result$EAS.Count <- EAS$Count; result$SAS.Count <- SAS$Count
  # start setting up tables
  sums <- c(sum(AFR$Count), sum(EUR$Count), sum(EAS$Count), sum(SAS$Count))</pre>
  # set up table and run test for each context
  for (i in 1:n.contexts){
    c.a <- c(AFR$Count[i], EUR$Count[i], EAS$Count[i], SAS$Count[i])</pre>
    c.b <- sums - c.a
    data <- cbind(c.a, c.b)
    warning <- is(tryCatch(chisq.test(data), warning = function(w) w), "warning")</pre>
    if (filter == T & warning){
      result$p[i] <- NA}
    else result$p[i] <- chisq.test(data)$p.value</pre>
  return(result)
```

Now using these functions, we can run the following tests for 3mer polymorphism types which are heterogeneous across ancestral groups. We can begin with the 3mer context paradigm, which is most commonly used in the literature in this area. The table below shows all 3mer polymorphism types, ranked according to their p value in a fourway.chi test for heterogeneity across populations.



Polymorphism type

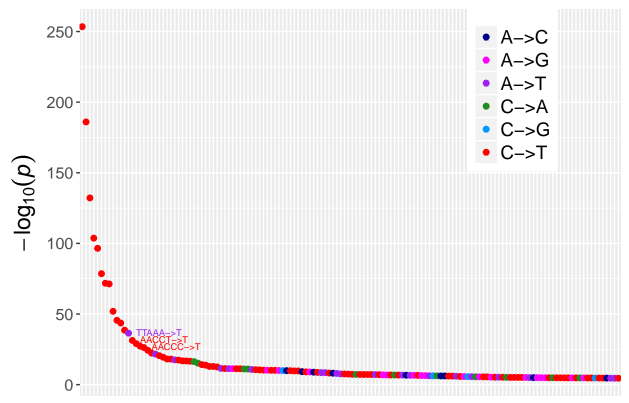
Table 1: 10 most significant 3mer polymorphisms

Context	X1mer	AFR.Count	EUR.Count	EAS.Count	SAS.Count	p
TCC->T	C->T	116126	30482	29565	37953	0.000000e+00
ACC->T	C->T	120823	24618	28466	35222	1.504434e-240
TCT->T	C->T	128789	25265	32178	37226	3.058550 e-129
GAT->T	A->T	42007	7561	12866	12948	3.229764 e-93
ACC->A	C->A	68375	12103	19959	20513	3.698720e-85
ACA->T	C->T	202388	32435	47413	53182	9.412389e-64
ACG->T	C->T	295109	50212	77743	83636	1.123409e-42
TAA->T	A->T	39856	5982	9412	9503	1.914092e-39
CCC->T	C->T	119810	21461	28885	33443	1.784362e-38
CAC->G	A->G	83359	13090	20315	20634	9.071360e-38

This plot highlights the top six contexts, which are significant at p<1e-60. They include TCC->T, ACC->T, and TCT->T, which have been previously reported as part of a European signal of C->T elevation. The next three contexts have not been noted by any previous analyses of mutation rate heterogeneity. There are 75 significant polymorphisms falling out from this analysis after Bonferroni correction.

Now we move to higher levels of sequence context, which may capture more detail in how mutation rates vary. In this section, we run the same analysis as above for 5mers, identifying variable polymorphism types which may not have been highlighted at the 3mer level.

The plot below shows the homogeneity test p values for just the 140 5mers which are significant after bonferroni correction.



Polymorphism type

It is clear from this plot (and the one for 7mers) that many of the significant polymorphisms at the 5mer and 7mer level are a part of the signal of C->T elevation that we observe at the 3mer level. This begs the question: how many significant 5mer signals are there outside of the 3mer subcontexts we have already idenified? To answer this question, I removed from the significant 5mer set all mutations whose 3mer subcontexts correspond to the European C->T elevation (TCC->T, ACC->T, TCT->T, and CCC->T), or otherwise highlighted by the heatmap signals 1-3 we identify in another analysis (see heatmaps report) identified in the previous section. This leaves a total of 71 new significant polymorphisms. The following table shows the most highly significant 5mers outside of these 3mer signals that have already been noted:

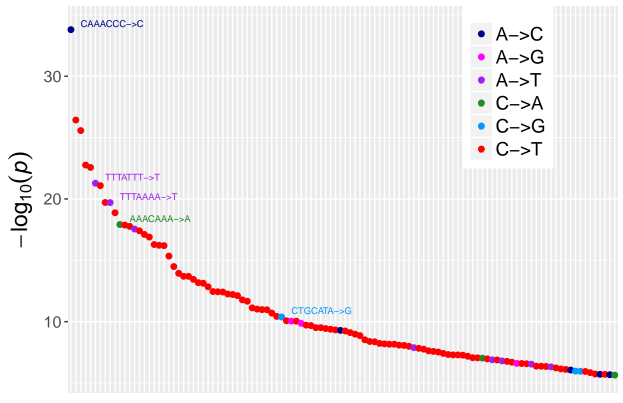
Table 2: 10 most significant 5mers not noted on a 3mer level

Context	AFR.Count	EUR.Count	EAS.Count	SAS.Count	p
TTAAA->T	17863	2569	3960	4035	3.654909e-37
TTATT->T	7641	1194	2062	1668	2.874258e-18
GGCTT->T	8807	1462	2710	2429	4.331903e-18
GGCCC->T	6516	1172	2059	1971	1.464061e-17
GGCTG->T	9312	1536	2790	2681	1.630722e-14
TACAT->T	18249	2892	4094	4615	3.012244e-13

Context	AFR.Count	EUR.Count	EAS.Count	SAS.Count	p
AAAAA->T	8686	1300	2029	1999	4.894078e-12
ACACC->G	4800	707	1005	1081	5.120720e-12
GGCTC->T	8438	1461	2509	2457	2.611590e-11
GGCTA->T	5397	921	1669	1518	3.710670e-11

Note that the most highly significant new 5mer is TTAAA->T, which corresponds to the 8th most significant 3mer, TAA->T. As we will see, the 7mer TTTAAAA->T is also one of the top significantly variable 7mers.

Now we move to the same analysis at the 7mer level, beginning with a homogeneity test plot:



Polymorphism type

The plot above shows heterogeneity test p values for the 112 7mers significant after bonferroni correction. We can ask the same question about these results as we did with the 5mers: which of these 7mers are results that we have not previously picked out from our 3mer analysis? Filtering these signals leaves 24 significant results, the top ten of which are shown below:

Table 3. 1	10 most significant	7mers not noted	on a 3mer level
Table o. 1	to most significant	THIELS HOL HOLED	on a omer iever

Context	AFR.Count	EUR.Count	EAS.Count	SAS.Count	p
CAAACCC->C	118	16	107	10	1.614606e-34
TTTATTT->T	2494	344	652	410	5.250459 e-22
TTTAAAA->T	10654	1565	2350	2424	1.994090e-20
AAACAAA->A	2773	377	620	489	1.223094e-18
ATTAAAA->T	3355	402	699	691	2.845707e-18
CTGCATA->G	65	13	51	12	4.167821e-11
TATATAT->G	6250	943	1349	1461	9.034974e-11
ACTAAAA->G	1956	410	662	590	1.382442e-10
AGTACAG->C	47	12	41	9	5.157272e-10
TATATTT->T	949	123	170	171	1.323000 e-08

Summary

The following table summarizes the numbers of significant results from this section.

Context Model	Number Significant	Number New
3mer	75	_
5mer	140	71
7mer	112	24

False Discovery Rate Corrections

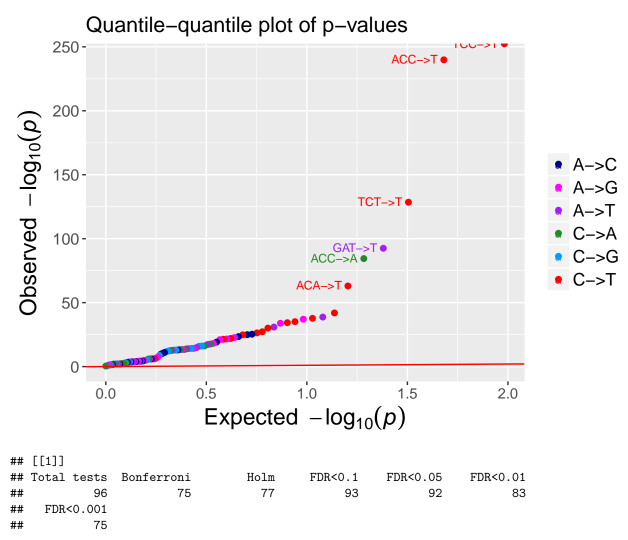
All of the tests in the above section use the Bonferroni Correction, which is conservative even when hypothesis tests are positively correlated (as is most-likely the case here.) However, the Bonferroni correction is often criticized as being *too* conservative. For these reasons, it may be useful to apply other significance thresholds which account for the multiple testing burden.

Methodology

Initially, I tried to use the qualue package to perform false discovery rate analysis. However, this package proved difficult to use, since our p-values from our homogeneity tests don't follow a uniform [0,1] distribution (they range from 0-0.45). Instead, I decided to use the built-in R function, p.adjust(), which uses Benjamini-Hoochberg-Yekutieli. These methods should be acceptable even when the p-values are positively correlated. The following function, fdr, performs simple fdr analysis on an output dataframe from a chi-squared function.

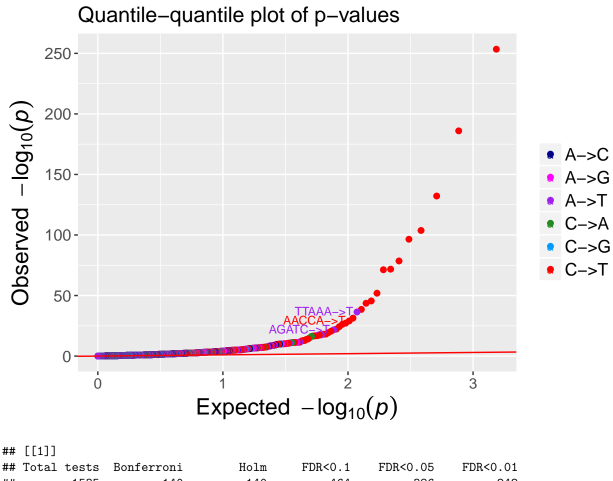
```
fdr <- function(p.data){</pre>
  p.data <- p.data[complete.cases(p.data),]</pre>
  # This uses Benjamini-Hochberg-Yekutieli fdr
  p.data$fdr <- p.adjust(p.data$p, method = "fdr")</pre>
  # multiple hypothesis correction by holm
  p.data$holm <- p.adjust(p.data$p, method = "holm")</pre>
  alpha = 0.05/length(p.data$p)
  p.data <- p.data[complete.cases(p.data), ]</pre>
  n.sig <- c(length(p.data$p), sum(p.data$p < alpha), sum(p.data$holm< 0.05),
              sum(p.data$fdr< 0.1), sum(p.data$fdr< 0.05),</pre>
              sum(p.data$fdr< 0.01), sum(p.data$fdr< 0.001))</pre>
  names(n.sig) <- c("Total tests", "Bonferroni", "Holm",</pre>
                      "FDR<0.1", "FDR<0.05",
                     "FDR<0.01", "FDR<0.001")
  return(list(n.sig, p.data))
}
```

I am additionally defining the funtion **qq.labels**, which takes in a p-value dataframe, a lab.lim, a title (default = "Quantile-quantile plot of p-values"), and the NoTCC arguement and returns a qq plot of all contexts, color-coded and labeled. In the following section, I will construct qq plots and run fdr analysis for each of the 3mer, 5mer, and 7mer models.



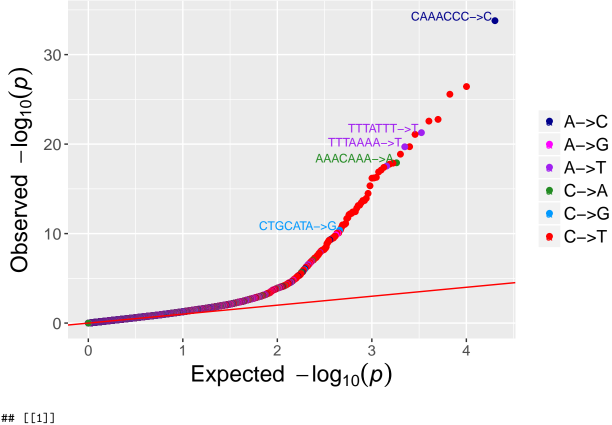
The qq plots shown above display relatively the same information as the p-value plots by context. However, it is worth noting that the observed p values, even at the lower end, are above expected p-value quantiles. This may suggest that in fact, every context is significant so that the null distribution of p values does not hold. More realistically, this appears to be an artifact of the fact that hypothesis tests set up as above are actually positively correlated (that is, a small p-value in one test probably increases the likelihood of a small p-value in another test).

One possible solution to this problem would be to simulate null-distributed datasets to approximate an emprical distribution for expected p-value.



##	[[1]]					
##	Total tests	Bonferroni	Holm	FDR<0.1	FDR<0.05	FDR<0.01
##	1535	140	140	464	386	242
##	FDR<0.001					
##	159					





##	[[1]]					
##	Total tests	Bonferroni	Holm	FDR<0.1	FDR<0.05	FDR<0.01
##	20076	112	112	372	272	184
##	FDR<0.001					
##	123					

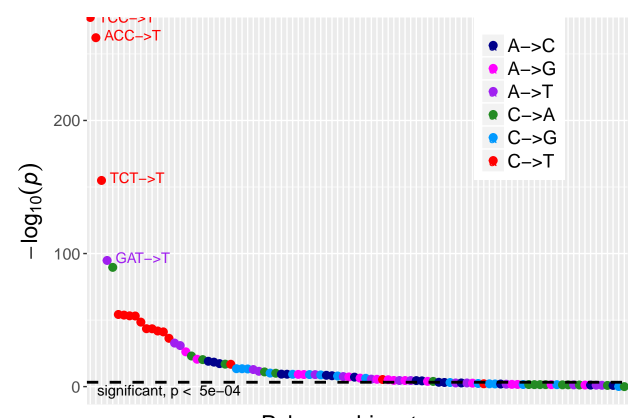
Ordered p-values

Methodology

The following function, **ordered.p**, returns a p-value dataframe with p calculated based on the methods from Harris and Pritchard, 2017. This method is proven to give less-significant results, but helps partially combat the problem of positive correlation between p values using our original methods.

```
ordered.p <- function(pdata){</pre>
  #preprocess data to order and remove nas
  pdata <- pdata[complete.cases(pdata$p),]</pre>
  myorder <- order(pdata$p)</pre>
  n.muts <- length(pdata$p)</pre>
  p.ordered <- rep(0, n.muts)</pre>
  #set largest p-value
  j <- myorder[n.muts]</pre>
  p.ordered[j] <- pdata$p[j]</pre>
  #initialize not mutated counts based on this lowest p-value mutation
  not.mut <- c(pdata$AFR.Count[j], pdata$EUR.Count[j],</pre>
                pdata$EAS.Count[j], pdata$SAS.Count[j])
  for (i in n.muts:1){
    j <- myorder[i]</pre>
    mut <-c(pdata$AFR.Count[j], pdata$EUR.Count[j],</pre>
             pdata$EAS.Count[j], pdata$SAS.Count[j])
    data <- cbind(mut, not.mut)</pre>
    p.ordered[j] <- chisq.test(data)$p.value</pre>
    #add these mutations to the not.mutated counts for future tests
    not.mut <- not.mut + mut</pre>
  pdata$p <- p.ordered
  return(pdata)
}
```

In the following sections, I will repeat all of the above analyses for fourway homogeneity test in terms of ordered p value.



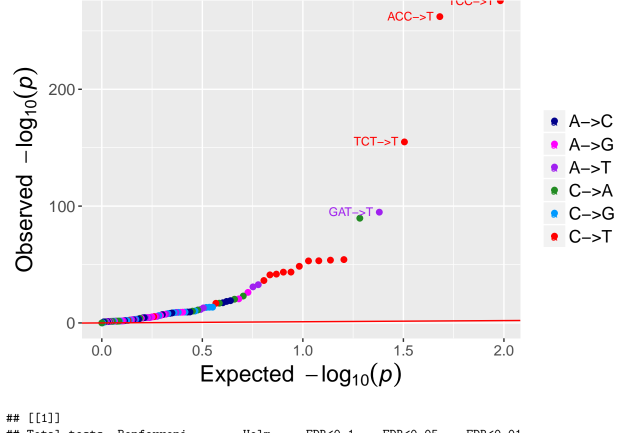
Polymorphism type

Table 5: 10 most significant 3mers using ordered p value correction

Context	X1mer	AFR.Count	EUR.Count	EAS.Count	SAS.Count	p
$\overline{\text{TCC->T}}$	C->T	116126	30482	29565	37953	0.000000e+00
ACC->T	C->T	120823	24618	28466	35222	6.419786e-263
TCT->T	C->T	128789	25265	32178	37226	1.175837e-155
GAT->T	A->T	42007	7561	12866	12948	1.553628e-95
ACC->A	C->A	68375	12103	19959	20513	2.377759e-90
CCC->T	C->T	119810	21461	28885	33443	6.127063e-55
TCG->T	C->T	181089	31320	47553	51985	1.833727e-54
ACG->T	C->T	295109	50212	77743	83636	6.391329e-54
ACA->T	C->T	202388	32435	47413	53182	8.420241e-54
GCG->T	C->T	206633	35598	54787	58748	3.410460e-49

Notice that, as before, GAT->T and ACC->A are the 4th and 5th most significant results. Meanwhile, ACA->T, the third highly significant signal from earlier, is moved from the 6th to the 9th place in terms of significance, and no longer sticks out from the remaining mutation types as it once did. Certain mutations (for example TAA->T) have dropped in significance notably (from 8th to 17th), while C->T mutations seem to be featured much more prominently among the most significant polymorphism types.

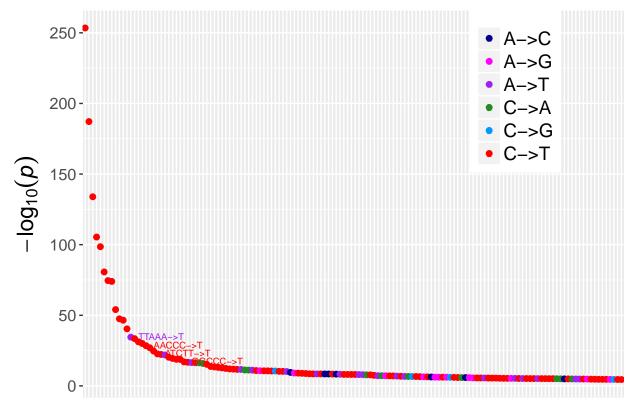




##	[[1]]					
##	Total tests	Bonferroni	Holm	FDR<0.1	FDR<0.05	FDR<0.01
##	96	62	65	92	85	71
##	FDR<0.001					
##	63					

[1] 142

[1] 65



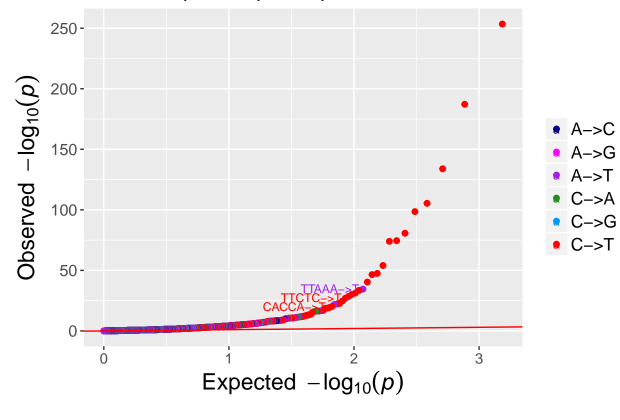
Polymorphism type

Table 6: 10 most significant new 5mers using ordered p value correction

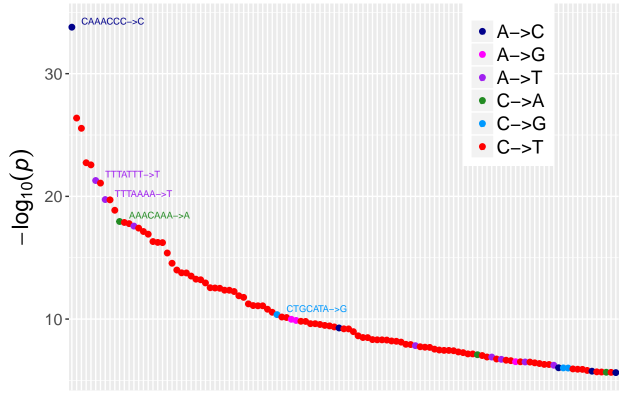
Context	AFR.Count	EUR.Count	EAS.Count	SAS.Count	p
$\overline{\text{TTAAA->T}}$	17863	2569	3960	4035	3.010762e-35
GGCCC->T	6516	1172	2059	1971	1.071839e-17
GGCTT->T	8807	1462	2710	2429	1.925726e-17
TTATT->T	7641	1194	2062	1668	3.152808e-17
GGCTG->T	9312	1536	2790	2681	3.959126e-14
TACAT->T	18249	2892	4094	4615	2.114546e-12
ACACC->G	4800	707	1005	1081	1.941371e-11
GGCTC->T	8438	1461	2509	2457	2.151745e-11
GGCCT->T	9977	1787	2933	2791	2.228892e-11
CCCCT->G	4877	904	1472	1530	2.973484e-11

Notice that, as before, TTAAA->T is the most significant new 5mer, with a p-value several orders of magnitude smaller than the other new significant 5mers. Again, many of the top results are present in a different order than in the original test.





##	[[1]]					
##	Total tests	Bonferroni	Holm	FDR<0.1	FDR<0.05	FDR<0.01
##	1535	142	143	427	361	250
##	FDR<0.001					
##	162					



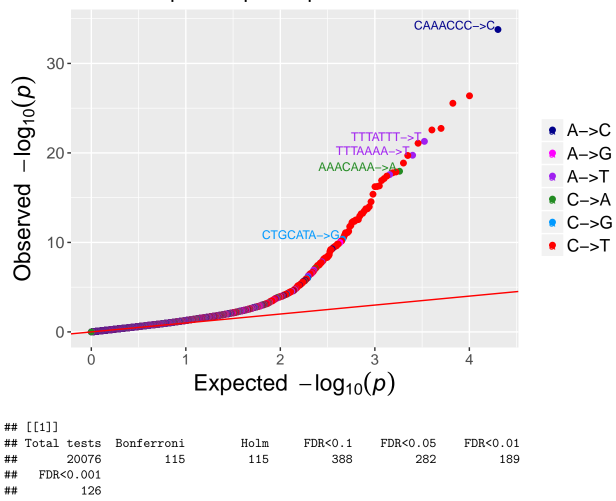
Polymorphism type

Table 7: 10 most significant new 7mers using ordered p value correction

Context	AFR.Count	EUR.Count	EAS.Count	SAS.Count	p
CAAACCC->C	118	16	107	10	1.628163e-34
TTTATTT->T	2494	344	652	410	5.112185e-22
TTTAAAA->T	10654	1565	2350	2424	1.816997e-20
AAACAAA->A	2773	377	620	489	1.116270e-18
ATTAAAA->T	3355	402	699	691	2.639385e-18
CTGCATA->G	65	13	51	12	4.348405e- 11
TATATAT->G	6250	943	1349	1461	1.040820e-10
ACTAAAA->G	1956	410	662	590	1.294195e-10
AGTACAG->C	47	12	41	9	5.319299e-10
TATATTT->T	949	123	170	171	1.469412e-08

Notice that, for 7mers, the ordering of the top ten most significant results is entirely unchanged.

Quantile-quantile plot of p-values



Summary

The following table summarizes the numbers of significant results from this section.

Context Model	Number Significant	Number New
3mer	62	_
5mer	142	65
$7 \mathrm{mer}$	115	24

Notice that using the ordered p value calculation causes us to pick up far fewer significant 3mers, but slightly more 5mers and 7mers. Moreover, it seems that using p ordered on 3mers has a much greater effect than on 5mers and 7mers. This might be expected because, for the most significant 7mers or 5mers, our p-value calculations still include most of the data, however, for our highly significant 3mers, a much larger portion of the data is excluded.