

# Supplementary material for UniqTag: Content-derived unique and stable identifiers for gene annotation

*Shaun D. Jackman*

## Supplementary material

The following supplementary material of the UniqTag paper present the code, shown in Listings A and B, and the data, shown in supplementary Table A, used to generate Figures 1 and 2 of the main manuscript.

### Load libraries

```
library(ggplot2)
library(knitr) # for kable
library(reshape2)
library(scales) # for alpha
```

### Read the data

```
data.orig <- read.delim('UniqTag-supp.tsv',
  colClasses = c(A = 'factor', B = 'factor'))
x <- do.call(rbind, strsplit(as.character(data.orig$Table), '.', fixed = TRUE))
colnames(x) <- c('Data', 'Transform', 'Identifier')
data <- cbind(data.orig, x)
rm(x)
data$k <- as.integer(gsub('^[a-z]*', '', data$Identifier))

build.wide <- with(data,
  data.frame(Build.A = A, Build.B = B,
  Num.A = Only.A + Both, Num.B = Only.B + Both))
build.tall <- melt(build.wide, id.vars = c('Build.A', 'Build.B'),
  variable.name = 'Build', value.name = 'Count')
```

### Figure 1. Plot the number of common identifiers vs. other build

The number of common UniqTag identifiers between build 75 of the Ensembl human genome and nine other builds, the number of common gene and protein identifiers between builds, and the number of genes with peptide sequences that are identical between builds.

```
data.subset <- subset(data, data$k == 9 | is.na(data$k))
aes.data <- aes(x = A, y = Both,
  group = Table, colour = Identifier)
aes.build <- aes(x = Build.A, y = Count,
```

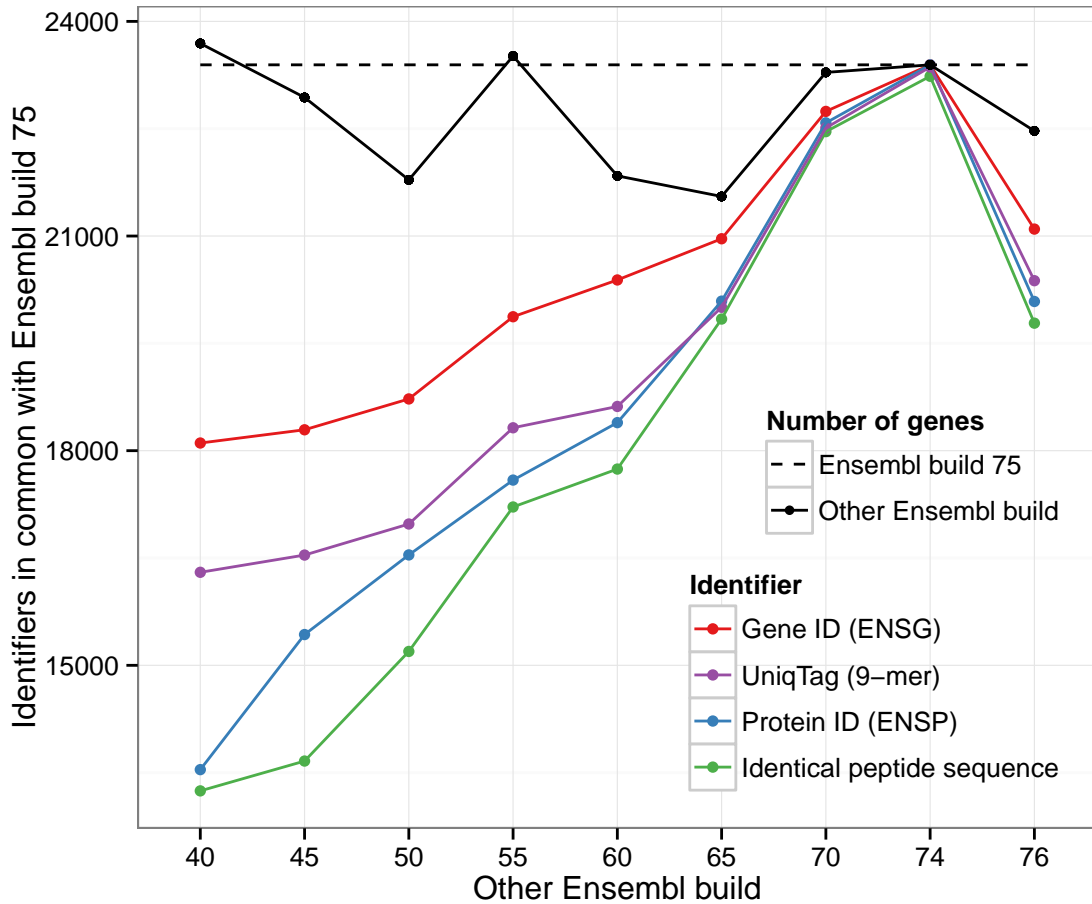
```

group = Build, linetype = Build, shape = Build)
ggplot() +
  geom_point(aes.data, data.subset) +
  geom_line(aes.data, data.subset) +
  scale_colour_brewer(palette = 'Set1',
    breaks = c('gene', 'uniqtag9', 'id', 'seq'),
    labels = c('Gene ID (ENSG)', 'UniqTag (9-mer)',
      'Protein ID (ENSP)', 'Identical peptide sequence')) +

  geom_point(aes.build, build.tall) +
  geom_line(aes.build, build.tall) +
  scale_linetype_manual(name = 'Number of genes',
    breaks = c('Num.B', 'Num.A'),
    labels = c('Ensembl build 75', 'Other Ensembl build'),
    values = c('solid', 'dashed')) +
  scale_shape_manual(name = 'Number of genes',
    breaks = c('Num.B', 'Num.A'),
    labels = c('Ensembl build 75', 'Other Ensembl build'),
    values = c(20, 32)) +

  theme_bw() +
  theme(legend.position = c(1.0, 0),
    legend.justification = c(1, 0),
    legend.box.just = 'right',
    legend.background = element_rect(fill = alpha('white', 0))) +
  xlab('Other Ensembl build') +
  ylab('Identifiers in common with Ensembl build 75')

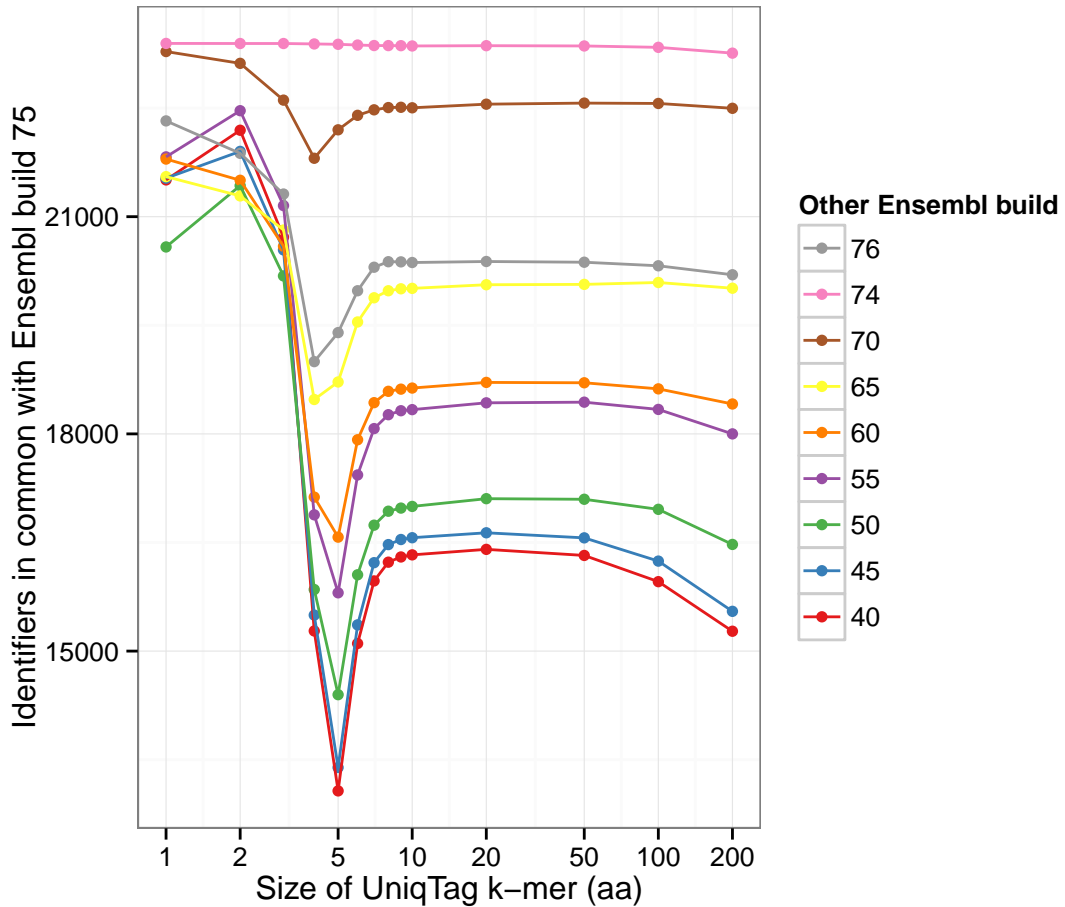
```



**Figure 2. Plot the number of common identifiers vs.  $k$**

The number of common UniqTag identifiers between build 75 of the Ensembl human genome and nine other builds for different values of  $k$ .

```
ggplot(na.omit(data), aes(x = k, y = Both, group = A, colour = A)) +
  geom_point() +
  geom_line() +
  scale_x_continuous(trans = log_trans(),
                     breaks = c(1, 2, 5, 10, 20, 50, 100, 200)) +
  scale_colour_brewer(name = 'Other Ensembl build', palette = 'Set1') +
  guides(colour = guide_legend(reverse = TRUE)) +
  theme_bw() +
  xlab('Size of UniqTag k-mer (aa)') +
  ylab('Identifiers in common with Ensembl build 75')
```



## Listing A. UniqTag 1.0

This listing shows the source of [UniqTag 1.0](#), implemented in Ruby.

```
#!/usr/bin/env ruby
# Determine a unique substring (k-mer) of each string
# Copyright 2014 Shaun Jackman

require 'optparse'

class String
  # Iterate over each k-mer
  def each_kmer k
    return enum_for(:each_kmer, k) unless block_given?
    (0 .. length - k).each { |i|
      kmer = self[i, k]
      yield kmer unless kmer =~ /~/
    }
  end
end

class Array
  # Append a serial number to distinguish duplicate strings
```

```

def dedup
  each_with_object(Hash.new(0)).map { |x, count|
    "#{$x}-#{count[x]} += 1"
  }
end
end

# Count the k-mers in a set of strings
def count_kmer seqs, k
  seqs.each_with_object(Hash.new(0)) { |seq, counts|
    seq.each_kmer(k).to_a.uniq.each { |kmer|
      counts[kmer] += 1
    }
  }
end

# Return the unique tag of the specified string
def get_tag seq, kmer_counts, k
  _, tag = seq.each_kmer(k).map { |kmer|
    [kmer_counts[kmer], kmer]
  }.min
  tag || seq.split('`').min
end

# Parse command line options
k = 9
OptionParser.new do |opts|
  opts.banner = "Usage: uniqtag [-k N] [FILE]..."
  opts.version = "0.1.0"
  opts.release = nil

  opts.on("-k", "--kmer N", OptionParser::DecimalInteger,
         "Size of the unique tag (default 9)") do |n|
    k = n
  end
end.parse!

# Read strings and write unique tags
seqs = ARGF.each_line.reject { |s|
  s =~ />/
}.map { |s|
  s.chomp.upcase
}
kmer_counts = count_kmer seqs, k
puts seqs.map { |seq| get_tag(seq, kmer_counts, k) }.dedup

```

## Listing B. Calculate the number of common identifiers

This [Makefile](#) script calculates the data used to plot the above figures.

```

# The supplementary material for the UniqTag paper
# UniqTag: Content-derived unique and stable identifiers for gene annotation
# Copyright 2014 Shaun Jackman

# Download the data and compute the results
all: UniqTag.tsv

# Remove all computed files
clean:
    rm -f *.comm *.gene *.id *.seq *.sort *.tsv *.unihtag *.venn

# Install dependencies
install-deps: /usr/local/bin/brew
    brew install coreutils seqtk untag wget

# Check for Homebrew
/usr/local/bin/brew:
    @if brew --version >/dev/null 2>/dev/null; then \
        echo Install Homebrew http://brew.sh/ or Linuxbrew http://brew.sh/linuxbrew/; \
    fi

.PHONY: all clean install-deps
.DELETE_ON_ERROR:
.SECONDARY:

# Download Ensembl Human genome NCBI36 build 40
Homo_sapiens.NCBI36.40.pep.all.fa.gz:
    wget ftp://ftp.ensembl.org/pub/release-40/homo_sapiens_40_36b/data/fasta/pep/Homo_sapiens.NCBI36.40

# Download Ensembl Human genome NCBI36 build 45
Homo_sapiens.NCBI36.45.pep.all.fa.gz:
    wget ftp://ftp.ensembl.org/pub/release-45/homo_sapiens_45_36g/data/fasta/pep/Homo_sapiens.NCBI36.45

# Download Ensembl Human genome NCBI36
Homo_sapiens.NCBI36.%_.pep.all.fa.gz:
    wget ftp://ftp.ensembl.org/pub/release-$*/fasta/homo_sapiens/pep/Homo_sapiens.NCBI36.$*.pep.all.fa.gz

# Download Ensembl Human genome GRCh37
Homo_sapiens.GRCh37.%_.pep.all.fa.gz:
    wget ftp://ftp.ensembl.org/pub/release-$*/fasta/homo_sapiens/pep/Homo_sapiens.GRCh37.$*.pep.all.fa.gz

# Download Ensembl Human genome GRCh38
Homo_sapiens.GRCh38.%_.pep.all.fa.gz:
    wget -O $@ ftp://ftp.ensembl.org/pub/release-$*/fasta/homo_sapiens/pep/Homo_sapiens.GRCh38.pep.all.fa.gz

# Uncompress FASTA and remove line breaks
%.fa: %.fa.gz
    seqtk seq $< >$@

# Remove the headers from a FASTA file
%.seq: %.fa
    grep -v '^>' $< >$@
```

```

# Convert a FASTA file to sorted TSV of ID, gene name and sequence
%.all.fa.tsv: %.all.fa
    awk -vORS=' ' '{print $$1 "\t" $$4; getline; print "\t" $$0 "\n" }' $< |sort -k2,2 -k1 >$@

# Keep the first protein isoform in the FASTA file
%.uniqgene.fa: %.fa.tsv
    awk '$x[$$2]++ == 0 { print $$1 " " $$2 "\n" $$3 }' $< >$@

# Join all protein isoforms separated by tilde
%.allgene.fa: %.fa.tsv
    awk '$x[$$2]++ == 0 { print $$1 " " $$2 "\n" $$3; next } \
        { print "~" $$3 }' $< |seqtk seq - >$@

# Extract the gene name from the FASTA header
%.gene: %.fa
    sed -En 's/^>.*gene:([^ ]*) .* /1/p' $< >$@

# Extract the ID from the FASTA header
%.id: %.fa
    sed -En 's/^>([^\n]*).*/\1/p' $< >$@

# Compute the UniqTag for each sequence in the FASTA file
ks=1 2 3 4 5 6 7 8 9 10 20 50 100 200
$(foreach k, $(ks), $(eval %.unihtag$k: %.fa; unihtag -k$k $$< >$$@))

# Join the gene name, ID and UniqTag into a TSV file
%.tsv: %.gene %.id %.unihtag7
    (printf "gene\tid\tunihtag\n" && paste $^) >$@

# Join the TSV of identifiers of two builds on the gene name
Homo_sapiens.GRCh37.70.75%.tsv: Homo_sapiens.GRCh37.70%.tsv Homo_sapiens.GRCh37.75%.tsv
join $^ |tr ' ' '\t' >$@

# Sort the file
%.sort: %
    sort $< >$@

# Compare an older Ensembl build to build 75
# Note: BSD comm has a bug possibly related to long lines and so GNU comm is
# used instead.
Homo_sapiens.Ensembl.40.75%.comm: Homo_sapiens.NCBI36.40%.sort Homo_sapiens.GRCh37.75%.sort
gcomm $^ >$@

Homo_sapiens.Ensembl.45.75%.comm: Homo_sapiens.NCBI36.45%.sort Homo_sapiens.GRCh37.75%.sort
gcomm $^ >$@

Homo_sapiens.Ensembl.50.75%.comm: Homo_sapiens.NCBI36.50%.sort Homo_sapiens.GRCh37.75%.sort
gcomm $^ >$@

Homo_sapiens.GRCh37.55.75%.comm: Homo_sapiens.GRCh37.55%.sort Homo_sapiens.GRCh37.75%.sort
gcomm $^ >$@

Homo_sapiens.GRCh37.60.75%.comm: Homo_sapiens.GRCh37.60%.sort Homo_sapiens.GRCh37.75%.sort

```

```

gcomm $^ >$@

Homo_sapiens.GRCh37.65.75%.comm: Homo_sapiens.GRCh37.65%.sort Homo_sapiens.GRCh37.75%.sort
gcomm $^ >$@

Homo_sapiens.GRCh37.70.75%.comm: Homo_sapiens.GRCh37.70%.sort Homo_sapiens.GRCh37.75%.sort
gcomm $^ >$@

Homo_sapiens.GRCh37.74.75%.comm: Homo_sapiens.GRCh37.74%.sort Homo_sapiens.GRCh37.75%.sort
gcomm $^ >$@

Homo_sapiens.GRCh38.76.75%.comm: Homo_sapiens.GRCh38.76%.sort Homo_sapiens.GRCh37.75%.sort
gcomm $^ >$@

# Count the overlap of two sets
%.venn: %.comm
printf "%u\t%u\t%u\n" `grep -c $$'`[^t]' $<` \
`grep -c $$'`t\t' $<` \
`grep -c $$'`t[^t]' $<` >$@

# Create the experimental design table
%-design.tsv:
printf "%s\t%s\t%s\n" >$@ \
Table A B \
$* 40 75 \
$* 45 75 \
$* 50 75 \
$* 55 75 \
$* 60 75 \
$* 65 75 \
$* 70 75 \
$* 74 75 \
$* 76 75

# Compute the experimental data table
%-data.tsv: \
Homo_sapiens.Ensembl.40.75.pep%.venn \
Homo_sapiens.Ensembl.45.75.pep%.venn \
Homo_sapiens.Ensembl.50.75.pep%.venn \
Homo_sapiens.GRCh37.55.75.pep%.venn \
Homo_sapiens.GRCh37.60.75.pep%.venn \
Homo_sapiens.GRCh37.65.75.pep%.venn \
Homo_sapiens.GRCh37.70.75.pep%.venn \
Homo_sapiens.GRCh37.74.75.pep%.venn \
Homo_sapiens.GRCh38.76.75.pep%.venn
	printf 'Only.A\tBoth\tOnly.B\n' && cat $^) >$@

# Join the experimental design and data tables
%.tsv: %-design.tsv %-data.tsv
paste $^ >$@

# Compute the data table
UniqTag.tsv: \

```

```

all.uniqgene.gene.tsv \
all.uniqgene.id.tsv \
all.uniqgene.seq.tsv \
all.uniqgene.uniqtag1.tsv \
all.uniqgene.uniqtag2.tsv \
all.uniqgene.uniqtag3.tsv \
all.uniqgene.uniqtag4.tsv \
all.uniqgene.uniqtag5.tsv \
all.uniqgene.uniqtag6.tsv \
all.uniqgene.uniqtag7.tsv \
all.uniqgene.uniqtag8.tsv \
all.uniqgene.uniqtag9.tsv \
all.uniqgene.uniqtag10.tsv \
all.uniqgene.uniqtag20.tsv \
all.uniqgene.uniqtag50.tsv \
all.uniqgene.uniqtag100.tsv \
all.uniqgene.uniqtag200.tsv
(head -n1 $< && tail -qn+2 $^) >$@

```

**Table A. The number of common identifiers**

These data are used to plot the above figures. They are also available in [tab-separated values \(TSV\) format](#).

`kable(data)`

| Table             | A  | B  | Only.A | Both  | Only.B | Data | Transform | Identifier | k  |
|-------------------|----|----|--------|-------|--------|------|-----------|------------|----|
| all.uniqgene.gene | 40 | 75 | 5585   | 18107 | 5286   | all  | uniqgene  | gene       | NA |
| all.uniqgene.gene | 45 | 75 | 4645   | 18292 | 5101   | all  | uniqgene  | gene       | NA |
| all.uniqgene.gene | 50 | 75 | 3062   | 18723 | 4670   | all  | uniqgene  | gene       | NA |
| all.uniqgene.gene | 55 | 75 | 3644   | 19872 | 3521   | all  | uniqgene  | gene       | NA |
| all.uniqgene.gene | 60 | 75 | 1455   | 20386 | 3007   | all  | uniqgene  | gene       | NA |
| all.uniqgene.gene | 65 | 75 | 591    | 20962 | 2431   | all  | uniqgene  | gene       | NA |
| all.uniqgene.gene | 70 | 75 | 545    | 22742 | 651    | all  | uniqgene  | gene       | NA |
| all.uniqgene.gene | 74 | 75 | 0      | 23393 | 0      | all  | uniqgene  | gene       | NA |
| all.uniqgene.gene | 76 | 75 | 1372   | 21097 | 2296   | all  | uniqgene  | gene       | NA |
| all.uniqgene.id   | 40 | 75 | 10150  | 13542 | 9851   | all  | uniqgene  | id         | NA |
| all.uniqgene.id   | 45 | 75 | 7507   | 15430 | 7963   | all  | uniqgene  | id         | NA |
| all.uniqgene.id   | 50 | 75 | 5242   | 16543 | 6850   | all  | uniqgene  | id         | NA |
| all.uniqgene.id   | 55 | 75 | 5927   | 17589 | 5804   | all  | uniqgene  | id         | NA |
| all.uniqgene.id   | 60 | 75 | 3449   | 18392 | 5001   | all  | uniqgene  | id         | NA |
| all.uniqgene.id   | 65 | 75 | 1463   | 20090 | 3303   | all  | uniqgene  | id         | NA |
| all.uniqgene.id   | 70 | 75 | 705    | 22582 | 811    | all  | uniqgene  | id         | NA |
| all.uniqgene.id   | 74 | 75 | 0      | 23393 | 0      | all  | uniqgene  | id         | NA |
| all.uniqgene.id   | 76 | 75 | 2384   | 20085 | 3308   | all  | uniqgene  | id         | NA |
| all.uniqgene.seq  | 40 | 75 | 10447  | 13245 | 10148  | all  | uniqgene  | seq        | NA |
| all.uniqgene.seq  | 45 | 75 | 9275   | 13662 | 9731   | all  | uniqgene  | seq        | NA |
| all.uniqgene.seq  | 50 | 75 | 6591   | 15194 | 8199   | all  | uniqgene  | seq        | NA |
| all.uniqgene.seq  | 55 | 75 | 6303   | 17213 | 6180   | all  | uniqgene  | seq        | NA |
| all.uniqgene.seq  | 60 | 75 | 4098   | 17743 | 5650   | all  | uniqgene  | seq        | NA |
| all.uniqgene.seq  | 65 | 75 | 1713   | 19840 | 3553   | all  | uniqgene  | seq        | NA |
| all.uniqgene.seq  | 70 | 75 | 828    | 22459 | 934    | all  | uniqgene  | seq        | NA |

| Table                 | A  | B  | Only.A | Both  | Only.B | Data | Transform | Identifier | k  |
|-----------------------|----|----|--------|-------|--------|------|-----------|------------|----|
| all.uniqgene.seq      | 74 | 75 | 160    | 23233 | 160    | all  | uniqgene  | seq        | NA |
| all.uniqgene.seq      | 76 | 75 | 2687   | 19782 | 3611   | all  | uniqgene  | seq        | NA |
| all.uniqgene.unihtag1 | 40 | 75 | 2184   | 21508 | 1885   | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag1 | 45 | 75 | 1405   | 21532 | 1861   | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag1 | 50 | 75 | 1203   | 20582 | 2811   | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag1 | 55 | 75 | 1690   | 21826 | 1567   | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag1 | 60 | 75 | 45     | 21796 | 1597   | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag1 | 65 | 75 | 0      | 21553 | 1840   | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag1 | 70 | 75 | 6      | 23281 | 112    | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag1 | 74 | 75 | 0      | 23393 | 0      | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag1 | 76 | 75 | 146    | 22323 | 1070   | all  | uniqgene  | unihtag1   | 1  |
| all.uniqgene.unihtag2 | 40 | 75 | 1498   | 22194 | 1199   | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag2 | 45 | 75 | 1035   | 21902 | 1491   | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag2 | 50 | 75 | 356    | 21429 | 1964   | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag2 | 55 | 75 | 1052   | 22464 | 929    | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag2 | 60 | 75 | 338    | 21503 | 1890   | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag2 | 65 | 75 | 266    | 21287 | 2106   | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag2 | 70 | 75 | 169    | 23118 | 275    | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag2 | 74 | 75 | 1      | 23392 | 1      | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag2 | 76 | 75 | 594    | 21875 | 1518   | all  | uniqgene  | unihtag2   | 2  |
| all.uniqgene.unihtag3 | 40 | 75 | 2975   | 20717 | 2676   | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag3 | 45 | 75 | 2396   | 20541 | 2852   | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag3 | 50 | 75 | 1603   | 20182 | 3211   | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag3 | 55 | 75 | 2363   | 21153 | 2240   | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag3 | 60 | 75 | 1249   | 20592 | 2801   | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag3 | 65 | 75 | 737    | 20816 | 2577   | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag3 | 70 | 75 | 677    | 22610 | 783    | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag3 | 74 | 75 | 1      | 23392 | 1      | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag3 | 76 | 75 | 1156   | 21313 | 2080   | all  | uniqgene  | unihtag3   | 3  |
| all.uniqgene.unihtag4 | 40 | 75 | 8414   | 15278 | 8115   | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag4 | 45 | 75 | 7440   | 15497 | 7896   | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag4 | 50 | 75 | 5935   | 15850 | 7543   | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag4 | 55 | 75 | 6634   | 16882 | 6511   | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag4 | 60 | 75 | 4714   | 17127 | 6266   | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag4 | 65 | 75 | 3078   | 18475 | 4918   | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag4 | 70 | 75 | 1480   | 21807 | 1586   | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag4 | 74 | 75 | 7      | 23386 | 7      | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag4 | 76 | 75 | 3471   | 18998 | 4395   | all  | uniqgene  | unihtag4   | 4  |
| all.uniqgene.unihtag5 | 40 | 75 | 10623  | 13069 | 10324  | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag5 | 45 | 75 | 9545   | 13392 | 10001  | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag5 | 50 | 75 | 7387   | 14398 | 8995   | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag5 | 55 | 75 | 7711   | 15805 | 7588   | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag5 | 60 | 75 | 5267   | 16574 | 6819   | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag5 | 65 | 75 | 2836   | 18717 | 4676   | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag5 | 70 | 75 | 1087   | 22200 | 1193   | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag5 | 74 | 75 | 12     | 23381 | 12     | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag5 | 76 | 75 | 3070   | 19399 | 3994   | all  | uniqgene  | unihtag5   | 5  |
| all.uniqgene.unihtag6 | 40 | 75 | 8587   | 15105 | 8288   | all  | uniqgene  | unihtag6   | 6  |
| all.uniqgene.unihtag6 | 45 | 75 | 7575   | 15362 | 8031   | all  | uniqgene  | unihtag6   | 6  |
| all.uniqgene.unihtag6 | 50 | 75 | 5731   | 16054 | 7339   | all  | uniqgene  | unihtag6   | 6  |
| all.uniqgene.unihtag6 | 55 | 75 | 6083   | 17433 | 5960   | all  | uniqgene  | unihtag6   | 6  |
| all.uniqgene.unihtag6 | 60 | 75 | 3922   | 17919 | 5474   | all  | uniqgene  | unihtag6   | 6  |

| Table                  | A  | B  | Only.A | Both  | Only.B | Data | Transform | Identifier | k  |
|------------------------|----|----|--------|-------|--------|------|-----------|------------|----|
| all.uniqgene.uniqtag6  | 65 | 75 | 2007   | 19546 | 3847   | all  | uniqgene  | uniqtag6   | 6  |
| all.uniqgene.uniqtag6  | 70 | 75 | 887    | 22400 | 993    | all  | uniqgene  | uniqtag6   | 6  |
| all.uniqgene.uniqtag6  | 74 | 75 | 22     | 23371 | 22     | all  | uniqgene  | uniqtag6   | 6  |
| all.uniqgene.uniqtag6  | 76 | 75 | 2492   | 19977 | 3416   | all  | uniqgene  | uniqtag6   | 6  |
| all.uniqgene.uniqtag7  | 40 | 75 | 7723   | 15969 | 7424   | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag7  | 45 | 75 | 6716   | 16221 | 7172   | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag7  | 50 | 75 | 5046   | 16739 | 6654   | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag7  | 55 | 75 | 5443   | 18073 | 5320   | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag7  | 60 | 75 | 3410   | 18431 | 4962   | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag7  | 65 | 75 | 1673   | 19880 | 3513   | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag7  | 70 | 75 | 811    | 22476 | 917    | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag7  | 74 | 75 | 29     | 23364 | 29     | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag7  | 76 | 75 | 2167   | 20302 | 3091   | all  | uniqgene  | uniqtag7   | 7  |
| all.uniqgene.uniqtag8  | 40 | 75 | 7464   | 16228 | 7165   | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag8  | 45 | 75 | 6466   | 16471 | 6922   | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag8  | 50 | 75 | 4853   | 16932 | 6461   | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag8  | 55 | 75 | 5251   | 18265 | 5128   | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag8  | 60 | 75 | 3253   | 18588 | 4805   | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag8  | 65 | 75 | 1576   | 19977 | 3416   | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag8  | 70 | 75 | 780    | 22507 | 886    | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag8  | 74 | 75 | 30     | 23363 | 30     | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag8  | 76 | 75 | 2091   | 20378 | 3015   | all  | uniqgene  | uniqtag8   | 8  |
| all.uniqgene.uniqtag9  | 40 | 75 | 7392   | 16300 | 7093   | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag9  | 45 | 75 | 6396   | 16541 | 6852   | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag9  | 50 | 75 | 4810   | 16975 | 6418   | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag9  | 55 | 75 | 5196   | 18320 | 5073   | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag9  | 60 | 75 | 3223   | 18618 | 4775   | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag9  | 65 | 75 | 1549   | 20004 | 3389   | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag9  | 70 | 75 | 776    | 22511 | 882    | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag9  | 74 | 75 | 31     | 23362 | 31     | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag9  | 76 | 75 | 2093   | 20376 | 3017   | all  | uniqgene  | uniqtag9   | 9  |
| all.uniqgene.uniqtag10 | 40 | 75 | 7363   | 16329 | 7064   | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag10 | 45 | 75 | 6371   | 16566 | 6827   | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag10 | 50 | 75 | 4787   | 16998 | 6395   | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag10 | 55 | 75 | 5181   | 18335 | 5058   | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag10 | 60 | 75 | 3208   | 18633 | 4760   | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag10 | 65 | 75 | 1543   | 20010 | 3383   | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag10 | 70 | 75 | 783    | 22504 | 889    | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag10 | 74 | 75 | 35     | 23358 | 35     | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag10 | 76 | 75 | 2102   | 20367 | 3026   | all  | uniqgene  | uniqtag10  | 10 |
| all.uniqgene.uniqtag20 | 40 | 75 | 7287   | 16405 | 6988   | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag20 | 45 | 75 | 6303   | 16634 | 6759   | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag20 | 50 | 75 | 4680   | 17105 | 6288   | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag20 | 55 | 75 | 5087   | 18429 | 4964   | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag20 | 60 | 75 | 3130   | 18711 | 4682   | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag20 | 65 | 75 | 1493   | 20060 | 3333   | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag20 | 70 | 75 | 733    | 22554 | 839    | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag20 | 74 | 75 | 31     | 23362 | 31     | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag20 | 76 | 75 | 2088   | 20381 | 3012   | all  | uniqgene  | uniqtag20  | 20 |
| all.uniqgene.uniqtag50 | 40 | 75 | 7371   | 16321 | 7072   | all  | uniqgene  | uniqtag50  | 50 |
| all.uniqgene.uniqtag50 | 45 | 75 | 6373   | 16564 | 6829   | all  | uniqgene  | uniqtag50  | 50 |
| all.uniqgene.uniqtag50 | 50 | 75 | 4688   | 17097 | 6296   | all  | uniqgene  | uniqtag50  | 50 |

| Table                   | A  | B  | Only.A | Both  | Only.B | Data | Transform | Identifier | k   |
|-------------------------|----|----|--------|-------|--------|------|-----------|------------|-----|
| all.uniqgene.unihtag50  | 55 | 75 | 5078   | 18438 | 4955   | all  | uniqgene  | unihtag50  | 50  |
| all.uniqgene.unihtag50  | 60 | 75 | 3135   | 18706 | 4687   | all  | uniqgene  | unihtag50  | 50  |
| all.uniqgene.unihtag50  | 65 | 75 | 1488   | 20065 | 3328   | all  | uniqgene  | unihtag50  | 50  |
| all.uniqgene.unihtag50  | 70 | 75 | 718    | 22569 | 824    | all  | uniqgene  | unihtag50  | 50  |
| all.uniqgene.unihtag50  | 74 | 75 | 35     | 23358 | 35     | all  | uniqgene  | unihtag50  | 50  |
| all.uniqgene.unihtag50  | 76 | 75 | 2098   | 20371 | 3022   | all  | uniqgene  | unihtag50  | 50  |
| all.uniqgene.unihtag100 | 40 | 75 | 7733   | 15959 | 7434   | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag100 | 45 | 75 | 6694   | 16243 | 7150   | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag100 | 50 | 75 | 4827   | 16958 | 6435   | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag100 | 55 | 75 | 5178   | 18338 | 5055   | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag100 | 60 | 75 | 3219   | 18622 | 4771   | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag100 | 65 | 75 | 1462   | 20091 | 3302   | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag100 | 70 | 75 | 723    | 22564 | 829    | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag100 | 74 | 75 | 54     | 23339 | 54     | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag100 | 76 | 75 | 2147   | 20322 | 3071   | all  | uniqgene  | unihtag100 | 100 |
| all.uniqgene.unihtag200 | 40 | 75 | 8418   | 15274 | 8119   | all  | uniqgene  | unihtag200 | 200 |
| all.uniqgene.unihtag200 | 45 | 75 | 7388   | 15549 | 7844   | all  | uniqgene  | unihtag200 | 200 |
| all.uniqgene.unihtag200 | 50 | 75 | 5312   | 16473 | 6920   | all  | uniqgene  | unihtag200 | 200 |
| all.uniqgene.unihtag200 | 55 | 75 | 5516   | 18000 | 5393   | all  | uniqgene  | unihtag200 | 200 |
| all.uniqgene.unihtag200 | 60 | 75 | 3428   | 18413 | 4980   | all  | uniqgene  | unihtag200 | 200 |
| all.uniqgene.unihtag200 | 65 | 75 | 1541   | 20012 | 3381   | all  | uniqgene  | unihtag200 | 200 |
| all.uniqgene.unihtag200 | 70 | 75 | 790    | 22497 | 896    | all  | uniqgene  | unihtag200 | 200 |
| all.uniqgene.unihtag200 | 74 | 75 | 134    | 23259 | 134    | all  | uniqgene  | unihtag200 | 200 |
| all.uniqgene.unihtag200 | 76 | 75 | 2271   | 20198 | 3195   | all  | uniqgene  | unihtag200 | 200 |