# The annmap Cookbook; Examples and Patterns for annmap

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### 1 Introduction

annmap is a Bioconductor package that provides annotation data and cross-mappings between, amongst other things, genes, transcripts, exons, proteins, domains and Affymetrix probesets. This document provides examples of its usage, and recipes to help you get started.

# 2 Initial Steps

annmap makes use of a MySQL database, annmap, to provide its annotation data (downloadable from the Annmap website<sup>1</sup>). A variety of different species are supported - each with their own separate database.

This document assumes that you have downloaded and installed one of the available MySQL databases (as described in the INSTALL vignette that came with this package).

It is also possible to connect to the annmap webservice instead of installing your own database – with some caveats. Please see the section 2.3 for details.

#### 2.1 Preamble

This cookbook is designed to walk you through the annmap functions from first principles to you being able to generate plots such as Figure 1 for your Next-Generation Sequencing data.

```
> gene = symbolToGene( 'SHH' )
> strand( gene ) = '*'
> ngsBridgePlot( gene, bamfileData )
```

<sup>1</sup>http://annmap.cruk.manchester.ac.uk/download

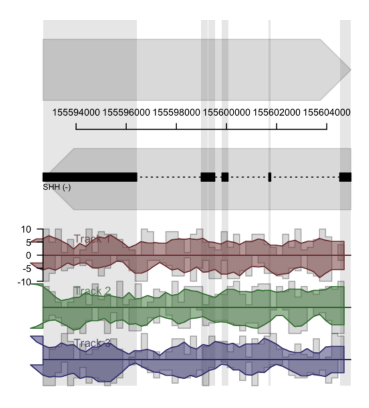


Figure 1: Dummy (in this example) NGS data shown alongside SHH. Grey boxes represent individual values. Coloured regions represent smoothed data.

### 2.2 Loading the annmap package and connecting to the database

Assuming everything is installed correctly, you should be able to load the package as normal:

```
> library( annmap )
```

If this is your first time running the package, you might need to create a datasource. These are stored by default in a folder called .annmap in your home directory, but this location can be customised by setting an environment variable ANNMAP\_HOME to point to a folder of your choosing. To add a datasource, we will use the annmapAddConnection method.

Assuming you have installed the annmap\_homo\_sapiens\_66 database into a MySQL instance running on your local machine accessed by the username 'test' and with a blank password, you can run:

```
> annmapAddConnection( 'human66', 'homo_sapiens', '66', username='test' )
```

You then need to tell annuap to connect to this annotation database:

```
annmapConnect( 'human66' )
Connected to annmap_homo_sapiens_66 (localhost)
Selected array 'HuEx-1_0' as a default.
```

A typical installation will support a variety of species, each with their own version of the annmap database. You can specify which database to use in the function call. Here, for example, human66 is the name we gave to our new connection. If you enter annmapConnect() without any parameters, it will show you a list of possible databases to choose from<sup>2</sup>.

<sup>&</sup>lt;sup>2</sup>unless you only have one database defined, in which case it will simply connect you to that

### 2.3 Using the annmap webservice

It's also possible to connect to the annmap webservice to get your data.

This is slower than using a locally installed database, less reliable as it relies on the annmap website which is occasionally taken down for updating.

Also, the annmap website operates a 'one in one out' rule, so when a new species version is added, the oldest one is removed to save disk space.

Those caveats aside, it's a great way of testing things out without the need to install MySQL and the database.

To connect to the webservice, simply call:

```
> annmapConnect( use.webservice=TRUE )
```

and you will be presented with a list of available connections to use.

As with the non-webservice connection, you can also do:

```
> annmapConnect( 'homo_sapiens.72', use.webservice=TRUE )
```

To connect to a known named webservice connection (so long as it exists).

### 2.4 Disconnecting from the database

To disconnect from the database, use:

```
annmapDisconnect()
Disconnecting from annmap_homo_sapiens_66 (localhost)
```

Note that you do not have to disconnect before connecting to another database; annmap will do this automatically for you when annmapConnect is called.

### 2.5 Overview

The majority of the calls in annmap fall into one of four categories: 'all' queries that fetch all known instances of an object type, 'details' queries that provide more detailed annotation for a list of IDs, 'to' queries that provide mappings between things (e.g. from genes to exons), and 'range' queries that find the things that lie between a pair of coordinates. These are summarised in table 1.

								Т	0							]		
		Gene	Transcript	Exon	EST Gene	EST Transcript	EST Exon	Prediction Transcript	Prediction Exon	Probeset	Probe	Hit	Protein	Domain	Symbol		InRange	Details
	Gene	N	Υ	Υ	N	N	N	N	N	Υ	N	N	Υ	Υ	Υ		Υ	Υ
	Transcript	Υ	N	Υ	N	N	N	N	N	Υ	N	N	Υ	Υ	N		Υ	Υ
	Exon	Υ	Υ	N	N	N	N	N	N	Υ	N	N	N	N	N		Υ	Y
	EST Gene	N	N	N	N	Υ	Υ	N	N	Υ	N	N	N	N	N		Υ	Υ
	EST Transcript	N	N	N	Υ	N	Υ	N	N	Υ	N	N	N	Ν	N		Υ	Y
	EST Exon	N	N	N	Υ	Υ	N	N	N	Υ	N	N	N	Ν	N		Υ	Υ
From	Prediction Transcript	N	N	N	N	N	N	N	Υ	Υ	N	N	N	Ν	N		Υ	Υ
ᅸ	Prediction Exon	N	N	N	N	N	N	N	N	N	N	N	N	Ν	N		N	N
	Probeset	Υ	Υ	Υ	Υ	Υ	Υ	Υ	N	N	Υ	Υ	Υ	Υ	N		Υ	Υ
	Probe	N	N	N	N	N	N	N	N	Υ	N	Υ	N	Ν	N		Υ	Υ
	Hit	N	N	N	N	N	N	N	N	N	N	N	N	Ν	N		N	N
	Protein	Υ	Υ	N	N	N	N	N	N	Υ	N	N	N	Υ	N		Υ	Υ
	Domain	Υ	Υ	N	N	N	N	N	N	Υ	N	N	Υ	Ν	N		Υ	Υ
	Symbol	Υ	Υ	N	Υ	Υ	N	N	N	N	N	N	N	Ν	N		N	N

Table 1: The mappings and functions available in annmap database. Mapping queries are of the form XXXXToYYY(), searches using genome coordinates, XXXInRange(), and detailed annotation XXXDetails()

# 3 Fetching Data

### 3.1 Retrieving all instances of a given type.

It is often useful to get a list of, for example, all genes or transcripts in the database. For this we would use functions such as allGenes or allTranscripts. All the functions of this type are defined in the documentation object ?annmap.all.

For example, to get a list of all the chromosomes in the human database, we can simply type:

```
annmapConnect( 'human66' )
Connected to annmap_homo_sapiens_66 (localhost)
Selected array 'HuEx-1_0' as a default.
> allChromosomes()
GRanges with 25 ranges and 0 metadata columns:
      seqnames
                   ranges strand
         <Rle>
                    <IRanges> <Rle>
             1 [1, 249250621]
   [1]
   [2]
             10 [1, 135534747]
   [3]
             11 [1, 135006516]
   [4]
            12 [1, 133851895]
   [5]
            13 [1, 115169878]
  [21]
             8 [1, 146364022]
             9 [1, 141213431]
  [22]
```

By default these functions all return a GRanges object (if the concept of genomic coordinates is appropriate), or a data.frame (if a genomic region doesn't exist for the object). It is also possible to just get the names of the chromosomes by passing as.vector=TRUE as a parameter:

```
> allChromosomes( as.vector=TRUE )
[1] "11" "21" "7" "Y" "2" "17" "22" "1" "18" "13" "16" "6" "X" "3" "9"
[16] "12" "14" "15" "20" "8" "4" "10" "19" "5" "MT"
```

#### 3.1.1 Possible values for as.vector and as.data.frame

Almost all of the functions in annmap accept an as.vector parameter, and will return a vector of identifiers rather than the full table of available information. Setting as.vector='data.frame' will return a data.frame, rather than a GRanges object:

```
as.vector=TRUE => a character vector of ids as.vector=FALSE => A GRanges object (default) as.vector='data.frame' => A data.frame
```

All of the XXXDetails() functions and functions that map to Hits or coding regions take a parameter as.data.frame instead (where a vector would not be possible):

```
as.data.frame=FALSE => A GRanges object (default) as.data.frame=TRUE => A data.frame
```

### 3.2 Getting detailed annotation based on database identifiers.

Sometimes, you will have a list of IDs for which you require more detailed annotation. You can use the ?annmapDetails methods for this.

For example, given a list of chromosome names, we can find their length:

```
> chr.list = c( '1', '2', '3' )
> chromosomeDetails( chr.list )
GRanges with 3 ranges and 0 metadata columns:
      segnames
                        ranges strand
         \langle R.1e \rangle
                     <IRanges> <Rle>
              1 [1, 249250621]
  [1]
  [2]
              2 [1, 243199373]
             3 [1, 198022430]
  [3]
  ___
  seqlengths:
    1 2 3
   NA NA NA
```

Or we can look for the details of a few microarray probes by their sequence:

Note that Probe sequences inside the annmap database are not the exact physical sequences to be found on the corresponding array. Instead, we always return the sequence of the genomic target.

#### 3.2.1 The order of things



When calling the following transformation queries (XXXTOYYY, and XXXDetails), you should not rely on the order of the output being the same as the input order.

If you need to look things up based on the input id you gave them, you should either subset based on the identifier column (ie: d[ elementMetadata( d )\$stable\_id == 'PTEN', ]), or you can split the results based on a column (see section 3.3.3)

### 3.3 One thing to another

In this section, we look at the available functions for mapping between annotation items. The majority of mappings are available, and all take the form XXXToYYY<sup>3</sup> (Table 1)

#### 3.3.1 Finding a gene by its symbol

For example, we can fetch a gene by its symbol:

```
> symbolToGene( 'PTEN' )
GRanges with 1 range and 9 metadata columns:
                           ranges strand | IN1
     seqnames
                                                             stable_id
                         <IRanges> <Rle> | <character>
                                                           <character>
  [1]
           10 [89622870, 89731687]
                                       + |
                                             PTEN ENSG00000171862
            biotype
                     status
        <character> <character>
  [1] protein_coding
                         KNOWN
                                                     description
                                                     <character>
  [1] phosphatase and tensin homolog [Source: HGNC Symbol; Acc: 9588]
     db_display_name
                      symbol
                                            symbol_description
         <character> <character>
                                                   <character>
  [1]
         HGNC Symbol
                           PTEN phosphatase and tensin homolog
           analysis_name
             <character>
  [1] ensembl_havana_gene
 seqlengths:
  10
```

<sup>&</sup>lt;sup>3</sup>A full list of functions can be found by entering ?annmapTo in an R console.

```
Or by vector of symbols:
```

```
> symbolToGene( c( 'PTEN', 'SHH' ) )
   GRanges with 2 ranges and 9 metadata columns:
          seqnames
                                   ranges strand |
                                                            IN1
                                                                      stable_id
             <Rle>
                                <IRanges> <Rle> | <character>
                                                                    <character>
      [1]
                10 [ 89622870, 89731687]
                                               + |
                                                      PTEN ENSG00000171862
      [2]
                 7 [155592680, 155604967]
                                                - |
                                                           SHH ENSG00000164690
                 biotype
                              status
             <character> <character>
      [1] protein_coding
                               KNOWN
      [2] protein_coding
                               KNOWN
                                                            description
                                                            <character>
      [1] phosphatase and tensin homolog [Source: HGNC Symbol; Acc: 9588]
      [2]
                         sonic hedgehog [Source: HGNC Symbol; Acc: 10848]
          db_display_name
                               symbol
                                                   symbol_description
              <character> <character>
                                                          <character>
      [1]
              HGNC Symbol
                                 PTEN phosphatase and tensin homolog
      [2]
              HGNC Symbol
                                  SHH
                                                       sonic hedgehog
                analysis_name
                  <character>
      [1] ensembl_havana_gene
      [2] ensembl_havana_gene
      seqlengths:
      10 7
      NA NA
   Again, as.vector works as before:
   > symbolToGene( c( 'PTEN', 'SHH' ), as.vector=TRUE )
    [1] "ENSG00000171862" "ENSG00000164690"
      Mapping from a gene to its transcripts
3.3.2
    > gene = symbolToGene( c( 'PTEN', 'SHH' ) )
    > geneToTranscript( gene )
   GRanges with 10 ranges and 13 metadata columns:
           seqnames
                                    ranges strand |
                                                                 IN1
                                                                            stable_id
              <Rle>
                                 <IRanges> <Rle> |
                                                         <character>
                                                                          <character>
       [1]
                 10 [ 89622870, 89731687]
                                                 + | ENSG00000171862 ENST00000371953
       [2]
                 10 [ 89624206, 89626806]
                                                 + | ENSG00000171862 ENST00000487939
                 10 [ 89624225, 89654083]
                                                 + | ENSG00000171862 ENST00000462694
       [3]
                                                 + | ENSG00000171862 ENST00000498703
       [4]
                 10 [ 89685280, 89693244]
       [5]
                 10 [ 89711956, 89721066]
                                                 + | ENSG00000171862 ENST00000472832
                  7 [155592680, 155604967]
       [6]
                                                 - | ENSG00000164690 ENST00000297261
                  7 [155592733, 155601766]
       [7]
                                                 - | ENSG00000164690 ENST00000441114
                  7 [155592734, 155601766]
                                                 - | ENSG00000164690 ENST00000430104
       [8]
       [9]
                  7 [155592744, 155601766]
                                                 - | ENSG00000164690 ENST00000435425
      [10]
                  7 [155599276, 155600064]
                                                 - | ENSG00000164690 ENST00000472308
                                        status description
                                                                 db_display_name
                           biotype
```

```
<character> <character> <character>
                                                                 <character>
 [1]
                                    KNOWN
                                                  <NA> HGNC transcript name
              protein_coding
                                                  <NA> HGNC transcript name
 [2]
                                    KNOWN
        processed_transcript
 [3]
        processed_transcript
                                    KNOWN
                                                  <NA> HGNC transcript name
 [4]
        processed_transcript
                                    KNOWN
                                                  <NA> HGNC transcript name
 [5]
        processed_transcript
                                    KNOWN
                                                  <NA> HGNC transcript name
 [6]
                                                  <NA> HGNC transcript name
              protein_coding
                                    KNOWN
                                                  <NA> HGNC transcript name
 [7] nonsense_mediated_decay
                                    KNOWN
              protein_coding
                                 PUTATIVE
                                                  <NA> HGNC transcript name
 [9] nonsense_mediated_decay
                                    KNOWN
                                                  <NA> HGNC transcript name
[10]
        processed_transcript
                                    KNOWN
                                                  <NA> HGNC transcript name
          symbol
                              symbol_description translation_start
     <character>
                                      <character>
                                                           <integer>
 [1]
        PTEN-001 phosphatase and tensin homolog
                                                                1358
 [2]
        PTEN-002 phosphatase and tensin homolog
                                                                <NA>
        PTEN-003 phosphatase and tensin homolog
 [3]
                                                                <NA>
 [4]
        PTEN-004 phosphatase and tensin homolog
                                                                <NA>
 [5]
        PTEN-005 phosphatase and tensin homolog
                                                                <NA>
 [6]
         SHH-001
                                                                 152
                                  sonic hedgehog
 [7]
         SHH-005
                                  sonic hedgehog
                                                                  91
 [8]
         SHH-003
                                  sonic hedgehog
                                                                  91
 [9]
         SHH-004
                                  sonic hedgehog
                                                                  91
[10]
         SHH-002
                                  sonic hedgehog
                                                                <NA>
     translation_start_exon translation_end translation_end_exon
                  <integer>
                                   <integer>
                                                         <integer>
 [1]
                     7035840
                                                            7036264
                                          186
 [2]
                        <NA>
                                         <NA>
                                                               <NA>
 [3]
                        <NA>
                                         <NA>
                                                               <NA>
 [4]
                                         <NA>
                        <NA>
                                                               <NA>
 [5]
                        <NA>
                                                               <NA>
                                         <NA>
 [6]
                     6965401
                                          827
                                                            6965465
 [7]
                     6965509
                                          80
                                                            6965518
 [8]
                                          206
                     6965509
                                                            6965536
 [9]
                                          71
                                                            6965551
                     6965509
[10]
                        <NA>
                                         <NA>
                                                               <NA>
                  analysis_name
                    <character>
 [1] ensembl_havana_transcript
 [2]
                         havana
 [3]
                         havana
 [4]
                         havana
 [5]
                         havana
 [6] ensembl_havana_transcript
 [7]
                         havana
 [8]
                         havana
 [9]
                         havana
[10]
                         havana
seqlengths:
 10 7
NA NA
```

And with as.vector:

```
> transcripts = geneToTranscript( gene, as.vector=TRUE )
> transcripts
[1] "ENST00000371953" "ENST00000487939" "ENST00000462694" "ENST00000498703"
[5] "ENST00000472832" "ENST00000297261" "ENST00000441114" "ENST00000430104"
[9] "ENST00000435425" "ENST00000472308"
```

We have tried to keep all of the methods in annmap "type agnostic" - so this will work passing a character vector as the parameter in place of a GRanges object:

```
> gene = symbolToGene( 'PTEN', as.vector=TRUE )
> class( gene )
[1] "character"
> geneToTranscript( gene )
GRanges with 5 ranges and 13 metadata columns:
                            ranges strand |
                                                         IN1
     seqnames
                                                                    stable_id
         <Rle>
                          <IRanges> <Rle> |
                                                 <character>
                                                                 <character>
  [1]
            10 [89622870, 89731687]
                                      + | ENSG00000171862 ENST00000371953
            10 [89624206, 89626806]
  [2]
                                        + | ENSG00000171862 ENST00000487939
            10 [89624225, 89654083]
  [3]
                                         + | ENSG00000171862 ENST00000462694
  [4]
            10 [89685280, 89693244]
                                         + | ENSG00000171862 ENST00000498703
            10 [89711956, 89721066]
  [5]
                                         + | ENSG00000171862 ENST00000472832
                   biotype
                               status description db_display_name
               <character> <character> <character>
                                                            <character>
                                           <NA> HGNC transcript name
  [1]
           protein_coding
                               KNOWN
  [2] processed_transcript
                                KNOWN
                                             <NA> HGNC transcript name
                                KNOWN
                                             <NA> HGNC transcript name
  [3] processed_transcript
  [4] processed_transcript
                                KNOWN
                                              <NA> HGNC transcript name
                                KNOWN
                                              <NA> HGNC transcript name
  [5] processed_transcript
           symbol
                              symbol_description translation_start
      <character>
                                     <character>
                                                         <integer>
  [1]
         PTEN-001 phosphatase and tensin homolog
                                                              1358
  [2]
                                                               <NA>
         PTEN-002 phosphatase and tensin homolog
  [3]
         PTEN-003 phosphatase and tensin homolog
                                                               <NA>
  [4]
         PTEN-004 phosphatase and tensin homolog
                                                               <NA>
  [5]
         PTEN-005 phosphatase and tensin homolog
     {\tt translation\_start\_exon} \ {\tt translation\_end} \ {\tt translation\_end\_exon}
                   <integer> <integer>
                                                       <integer>
  [1]
                     7035840
                                        186
                                                          7036264
  [2]
                        <NA>
                                        <NA>
                                                             <NA>
  [3]
                        <NA>
                                                             <NA>
                                        <NA>
  [4]
                                        <NA>
                                                             <NA>
                        <NA>
  [5]
                        <NA>
                                        <NA>
                                                              <NA>
                  analysis_name
                    <character>
  [1] ensembl_havana_transcript
  [2]
                         havana
  [3]
                         havana
  [4]
                         havana
  [5]
                         havana
  seqlengths:
  10
  NA
```

#### 3.3.3 The IN1 field in 'to' results

The GRanges and data.frame returned by 'to' queries, contain a column IN1. This corresponds to the initial query that led to the row being generated. If we extract the IN1 column, we get the original 2 genes that we sent to the query:

```
> elementMetadata( geneToTranscript( symbolToGene( c( 'PTEN', 'SHH' ) ) ) )$IN1
[1] "ENSG00000171862" "ENSG00000171862" "ENSG00000171862" "ENSG00000171862"
[5] "ENSG00000171862" "ENSG00000164690" "ENSG00000164690"
[9] "ENSG00000164690" "ENSG00000164690"
```

This can be very useful when we wish to use, for example, merge, to combine a table of expression data with the annotation data supplied by annuap.

It can also be used with the split method to get the returned data into a list, for example, to generate a list, one element per gene, each containing the exons found in that gene:

```
> exons = geneToExon( symbolToGene( c( 'PTEN', 'SHH' ) ) )
> split( elementMetadata( exons )[['stable_id']], elementMetadata( exons )[['IN1']] )
$ENSG00000164690
[1] "ENSE00001086614" "ENSE00001086617" "ENSE00001149618" "ENSE00001795984"
[5] "ENSE00001760492" "ENSE00001612491" "ENSE00001676036" "ENSE00001758320"
[9] "ENSE00001655961" "ENSE00001800592" "ENSE00001908904" "ENSE00001889423"

$ENSG00000171862
[1] "ENSE00001456562" "ENSE00001156351" "ENSE00001156344" "ENSE00003595610"
[5] "ENSE00001156330" "ENSE00001156327" "ENSE00003601394" "ENSE00001156315"
[9] "ENSE00001456541" "ENSE00001813529" "ENSE00001881230" "ENSE00001926457"
[13] "ENSE00001920439" "ENSE00001837210" "ENSE000003597859" "ENSE00001893700"
[17] "ENSE00001876418" "ENSE00003657126" "ENSE00001939988"
```

#### 3.3.4 Microarray probeset mappings

annmap also provides probe mappings for Affymetrix microarrays. When you connect to a database, annmap will pick an array by default (if a mapping exists for that species).

To select your array of choice, you can use the arrayType() method:

```
> arrayType( 'HuEx-1_0' )
Using array 'HuEx-1_0'.
```

As with annmapConnect() calling this function with no parameters will give you a menu of available arrays to choose from.

Probesets are treated like any other feature, so:

```
> exonToProbeset(geneToExon(symbolToGene('MPI')),as.vector=TRUE)

[1] "3601956" "3601960" "3601963" "3601964" "3601965" "3601969" "3601971"

[8] "3601959" "3601974" "3601975" "3645143" "3975448" "2746540" "2880800"

[15] "3535951" "2410077" "3655014" "3462599" "2662284" "3638759" "2766811"

[22] "3822407" "2830115" "3585206" "3825799" "3961893" "2735060" "2646089"

[29] "3471746" "3720274" "3017565" "3328152" "3248968" "2713240" "2685561"

[36] "23285899" "3415635" "3869300" "3067623" "3259813" "3707032" "3731058"

[43] "3190116" "3384517" "3235475" "2676567" "2796100" "2923872" "2591241"

[50] "3056411" "3474991" "2756434" "2813774" "2861407" "3601973" "2544232"

[57] "3336154" "2393742" "3318836" "3470332" "3841120" "3553844" "3866317"

[64] "3464331" "3665054" "3109991" "2622458" "3970147" "2396528" "3454391"

[71] "3472536" "3601957" "3601970" "3601958" "3601961" "3601966" "3601972"

[78] "3601968"
```

will find all probesets that map to exons in the gene MPI (as a vector).

Ommitting the as.vector=TRUE parameter in the call will give us the results in a data.frame. Here's the details of the first probeset from our last query:

> exonToProbeset(geneToExon(symbolToGene('MPI')) )[1,]

```
IN1 stable_id array_name probe_count hit_score gene_score

1 ENSE00002608534 3601956 HuEx-1_0 4 1 1
transcript_score exon_score est_gene_score est_transcript_score

1 2 2 0 0 0
est_exon_score prediction_transcript_score prediction_exon_score

1 0 0 0
protein_score domain_score

1 0 0
```

As you can see, a probeset's details, comprises a set of scores. These scores are always 0, 1 or 2;

- 0. One or more of the probes miss the item of interest
- 1. All of the probes hit the item of interest once (and only once)
- 2. One or more of the probes hit more than one item of interest

So a hit\_score of 1 means that each and every probe in the probeset hit the genome once and only once, but a score of 2 for gene\_score means that one or more of the probes hits more than one gene. Since if it also has a hit score of 1, this means that the match is at a region where 2 genes are overlapping.

#### 3.3.5 Some subtleties with probeset mappings

Probeset mappings in annmap are done in two ways:

- 1. Probes are mapped to the entire genome, and their match locations recorded, and
- 2. they are also mapped directly to cDNA sequences in order to pick up probes and probesets that fall on exon junctions.

 $probe set To C dnatrans cript \ and \ transcript To C dna probe set \ provide \ the \ mappings \ needed \ to \ retrieve \ these \ probesets.$ 

## 4 Searching by genome coordinates

### 4.1 Finding things by their location

7 NA

It is also possible to provide a set of genomic regions and find the features they contain. For example:

```
> geneInRange( '7', 1000000, 1060000, 1)
   GRanges with 1 range and 8 metadata columns:
         seqnames
                               ranges strand |
                                                     stable_id
                                                                       biotype
                           <IRanges> <Rle> |
                                                    <character>
                                                                   <character>
      [1]
                 7 [1022835, 1029276]
                                         + | ENSG00000073067 protein_coding
               status
          <character>
      [1]
               KNOWN
                                                                                     description
                                                                                     <character>
      [1] cytochrome P450, family 2, subfamily W, polypeptide 1 [Source: HGNC Symbol; Acc: 20243]
                               symbol
         db_display_name
              <character> <character>
      [1]
              HGNC Symbol
                           CYP2W1
                                             symbol_description
                                                                       analysis_name
                                                     <character>
                                                                         <character>
     [1] cytochrome P450, family 2, subfamily W, polypeptide 1 ensembl_havana_gene
     seqlengths:
       7
      NΑ
  will find all of the genes on the forward strand of chromosome '7' that lie between positions 1000000
and 1060000. XXXInRange queries also accept data.frame, RangedData and GRanges objects (see
?annmapRange):
   > # Use a data.frame as the initial parameter
   > .df = data.frame( chromosome_name='7', start=1000000, end=1060000, strand=1 )
   > geneInRange( .df, as.vector=TRUE )
   [1] "ENSG00000073067"
   > # Use a RangedData object as the initial parameter
   > .rd = RangedData( space='7', ranges=IRanges( start=1000000, end=1060000 ), strand=1 )
   > geneInRange( .rd, as.vector=TRUE )
   [1] "ENSG00000073067"
   > # Use a GRanges object as the initial parameter
   > .rd = RangedData( space='7', ranges=IRanges( start=1000000, end=1060000 ), strand=1L )
   > .rd = as( .rd, 'GRanges' )
   > .rd
   GRanges with 1 range and 0 metadata columns:
         seqnames
                               ranges strand
             <Rle>
                            <IRanges> <Rle>
               7 [1000000, 1060000]
      [1]
     seqlengths:
```

```
> geneInRange( .rd, as.vector=TRUE )
[1] "ENSG00000073067"
```

This means that it is possible to chain InRange queries as you would most of the other queries, ie;

```
> geneToSymbol(
                                 # Return the gene symbols
   exonToGene(
                                 # get the gene for each exon
     exonInRange(
                                 # get all exons contained in the GRanges
       symbolToGene( 'PTEN' ) ) ) # get a GRanges object for PTEN
ENSG00000171862 ENSG00000171862 ENSG00000171862 ENSG00000171862 ENSG00000224745
                                    "PTEN" "PTEN" "RP11-380G5.2"
        "PTEN"
                      "PTEN"
ENSG00000224745 ENSG00000224745 ENSG00000171862 ENSG00000171862 ENSG00000171862
"RP11-380G5.2" "RP11-380G5.2" "PTEN" "PTEN"
                                                                 "PTFN"
ENSG00000171862 ENSG00000171862 ENSG00000171862 ENSG00000171862 ENSG00000171862
        "PTEN" "PTEN" "PTEN" "PTEN"
ENSG00000213613 ENSG00000171862 ENSG00000171862 ENSG00000171862 ENSG00000171862
"RP11-380G5.3"
                    "PTEN"
                                   "PTEN"
                                             "PTEN"
                                                                "PTEN"
ENSG00000171862 ENSG00000171862 ENSG00000171862
       "PTEN"
                     "PTEN"
                                    "PTEN"
```

For example, to find all probes that target within 1Kb of the 3´ end of each gene in genes, we can first alter the range of these objects so they are just covering our region of interest:

```
> genes = geneInRange( '7', 1000000, 1060000, 1 )
> start(genes) = end( genes ) - 1000
```

Before calling probeInRange on the modified GRanges object:

### > probeInRange( genes )

GRanges with 27 ranges and 2 metadata columns:

```
seqnames
                                 ranges strand
                                                                               sequence
          <Rle>
                             <IRanges> <Rle>
                                                      <character>
           7 [1028280, 1028304] + | TCTGTGTTGGGGAGCGCCTGGCCAG
 [1]
 [2]
              7 [1028310, 1028334]
                                                + | AGCTCTTCCTGCTGTTTTGCCGGCCT
 [3]
                                               + | TCCTGCAGAGGTACCGCCTGCTGCC
             7 [1028334, 1028358]
 [4]
             7 [1028398, 1028422]
                                               + | CGCCCGGGCTTTTACCATGAGGCCG
             7 [1028399, 1028423]
 [5]
                                              + | GCCCGGGCTTTTACCATGAGGCCGA
             7 [1028980, 1029004] + | AGCCACTGGGGCCATGCGTATGACT
7 [1029043, 1029067] + | AGGCACTGGCGCCAGAGGCTTCCTT
7 [1029089, 1029113] + | AGCCCCTGAAGACAAGCAGCACTGC
7 [1029122, 1029146] + | AAATGGAAACACTGACCCGGTGCGG
7 [1029125, 1029149] + | TGGAAACACTGACCCGGTGCGGTGC
[23]
[24]
[25]
[26]
[27]
     probe_hit_count
             <numeric>
 [1]
 [2]
                        1
 [3]
 [4]
                        1
 [5]
                       1
 . . .
                      . . .
[23]
                       1
[24]
                       1
[25]
                       1
```

```
[26]
[27]
---
seqlengths:
7
NA
```

As with other queries, the as.vector parameter can be used to return more concise results:

```
> probeInRange( genes, as.vector=TRUE )

[1] "TCTGTGTTGGGGAGCGCCTGGCCAG" "AGCTCTTCCTGCTGTTTGCCGGCCT"
[3] "TCCTGCAGAGGTACCGCCTGCTGCC" "CGCCCGGGCTTTTACCATGAGGCCG"
[5] "GCCCGGGCTTTTACCATGAGGCCGA" "CCGGGCTTTTACCATGAGGCCGAGG"
[7] "GGGCTTTTACCATGAGGCCGAGGGC" "CCCACAGCTCGGACTGCTCTGGGAG"
[9] "GTCAGCAACTGCTTCCGGTTACACC" "TGCTTCCGGTTACACCCAGGACTAC"
[11] "CTGAAGCTGCACTCCCACCCACCTA" "CTGCAGGGAGACAACGGGTGGCTGC"
[13] "GAGACAACGGGTGGCTGCATCCAGC" "GACAACGGGTGGCTGCATCCAGCCA"
[15] "CTGCATCCAGCCAGAGACAGGCGCA" "TGGGTGTCCTCAGCGTGCGAGCCCT"
[17] "GGTGTCCTCAGCGTGCGAGCCCTGC" "ACTCCATCCCGCTCCTGGAACACT"
[21] "TCCATTCCCGCTCCTGGAACACTTC" "TGTGCCTGGAGGCCTTGCAC"
[23] "AGCCACTGGGGCCATGCGTATGACT" "AGGCACTGGCGCCAGAGGCTTCCTT"
```

For example, it is possible to find estGenes which overlap known genes as follows:

[25] "AGCCCCTGAAGACAAGCAGCACTGC" "AAATGGAAACACTGACCCGGTGCGG"

```
> estGeneInRange( symbolToGene( c( 'lama3', 'tp53', 'shh' ) ), as.vector=TRUE )

[1] "ENSESTG00000004861" "ENSESTG00000004918" "ENSESTG00000032897"

[4] "ENSESTG00000032910" "ENSESTG00000032912" "ENSESTG00000032918"

[7] "ENSESTG00000032924" "ENSESTG00000032930" "ENSESTG00000032952"

[10] "ENSESTG00000033006" "ENSESTG00000010677"
```

### 4.2 Adventures with annmapRangeApply

[27] "TGGAAACACTGACCCGGTGCGGTGG"

A common task is to iterate down a GRanges object and perform a function per row. This functionality is provided by the method annmapRangeApply().

To take a contrived example, lets write a function that takes a GRanges object, and returns a character vector of the form chr:location (to provide labels for a graph, say):

```
> contrived.function = function( chromosome, location ) {
+  paste( chromosome, ':', location, sep='' )
+ }
```

In a moment, we're going to use annmapRangeApply to map this down a set of transcripts, which we'll retrieve now:

```
> transcripts = geneToTranscript( symbolToGene( c( 'shh', 'tp53' ) ) )
```

Before we can do this, though, we need to tell the function which columns in the GRanges object to map onto 'chromosome' and 'location', and also to tell it what datatype each of these parameters should be.

This is done using two additional parameters in annmapRangeApply: filter and coerce. The first defines the column names in the GRanges object we are interested in, the second, the functions we need to coerce these to the expected types<sup>4</sup>:

```
> contrived.filter = c( chromosome='space', location='start' )
> contrived.coerce = c( as.character, as.numeric )
```

Once these are defined, we can then fire off our apply statement, as follows:

<sup>&</sup>lt;sup>4</sup>The coerce is needed because we can't always rely on R to correctly guess the data type we're expecting ('space' in a GRanges object will often be converted to a factor, not a character vector as required by paste in our contrived. function)

# 5 Affymetrix<sup>™</sup>Array Annotation

As well as providing Ensembl annotation, annmap also provides mappings for Affymetrix ™arrays, as described in *An annotation infrastructure for the analysis and interpretation of Affymetrix exon array data* Genome Biology 2007, 8:R79, doi:10.1186/gb-2007-8-5-r79.

### 5.1 Exonic, intronic, intergenic and unreliable

Probesets in annuap are defined as being either exonic, intronic, intergenic or unreliable. These terms are best described by:

- Exonic: Probesets are classed as exonic if all of their probes map to the genome only once, and every one of these mappings falls within an exon boundary.
- Intronic: Probesets are classed as intronic if all of their probes map to the genome only once, but at least one probe misses an exon region, but still falls within the boundary of a gene.
- Intergenic: Probesets are classed as intergenic if all of their probes map to the genome only once, but at least one probe misses all the known genes.
- Unreliable: Probesets with one or more multi-targetting probes, or with one or more probes that do not map to the genome, are classed as unreliable.

These categories are a very broad filter. An unreliable probeset may have 3 probes which all hit in a single location and all hit an exon, but the single probe which misses mapping to the genome will result in the entire probeset being classed as to unreliable.

### 5.2 The filtering methods

To filter a collection of probesets by their relevant category, the obvious methods exist. For example, we can look at the probesets that are hitting around the gene for TP53:

```
> g = symbolToGene('TP53')
> ps = geneToProbeset( g, as.vector=TRUE )
> length(ps)
[1] 237
```

We can then see numbers of probesets that match each category:

```
> length( exonic( ps ) )
Building probeset specificity cache.....done
[1] 18
> length( intronic( ps ) )
[1] 14
> length( intergenic( ps ) )
[1] 1
> length( unreliable( ps ) )
[1] 204
```

So as you can see, hitting TP53, we have 18 exonic, 14 intronic, 1 intergenic<sup>5</sup> and 204 unreliable probesets.

All of these functions, will take an exclude parameter that results in an inverted list. So to get the number of all of the probesets that are not unreliable, we can do:

<sup>&</sup>lt;sup>5</sup>This may seem odd, but note that as only a single probe in a probeset needs to be intergenic for the entire probeset to be categorised as such. So it is possible for a probeset that mostly maps to a gene to be flagged as intergenic.

```
> length( unreliable( ps, exclude=T ) )
[1] 33
```

Since anything that is not unreliable must hit the genome somewhere, the following should evaluate to TRUE:

```
> u.ps = sort( unreliable( ps, exclude=T ) )
> c.ps = sort( c( intronic( ps ), exonic( ps ), intergenic( ps ) ) )
> all( u.ps == c.ps )
[1] TRUE
```

### 5.3 UTR filtering

The function utrProbesets can be used to find those probesets that hit UTRs:

```
> gene = symbolToGene( 'TP53' )
> probesets = geneToProbeset( gene )
> utrProbesets( probesets )
  [1] "3743926" "3743927" "3743928" "3743929" "3743930" "3743931" "3743932"
  [8] "3743933" "3743938" "3743939" "3743940" "3743908" "3743926" "3743927"
 [15] "3743928" "3743929" "3743930" "3743931" "3743932" "3743938" "3743939"
 [22] "3743940" "3743908" "3743909" "3743910" "3743911" "3743912" "3743935"
 [29] "3743926" "3743927" "3743928" "3743929" "3743930" "3743931" "3743932"
 [36] "3743938" "3743939" "3743940" "3743908" "3743909" "3743910" "3743911"
 [43] "3743912" "3743935" "3743926" "3743927" "3743928" "3743929" "3743930"
 [50] "3743931" "3743932" "3743938" "3743939" "3743940" "3743908" "3743921"
 [57] "3743922" "3743923" "3743924" "3743925" "3743926" "3743927" "3743928"
 [64] "3743929" "3743930" "3743931" "3743932" "3743938" "3743939" "3743940"
 [71] "3743926" "3743927" "3743928" "3743929" "3743930" "3743931" "3743932"
 [78] "3743938" "3743939" "3743940" "3743926" "3743927" "3743928" "3743929"
 [85] "3743930" "3743931" "3743932" "3743938" "3743939" "3743940" "3743926"
 [92] "3743927" "3743928" "3743929" "3743930" "3743931" "3743932" "3743938"
 [99] "3743939" "3743940" "3743909" "3743926" "3743927" "3743928"
```

In this form, the function will attempt to match probesets to all possible transcripts and then filter by UTR, so any UTR targeting probeset will be found.

You can also pass a vector of transcripts in, in which case the search is limited to these:

```
> transcripts = geneToTranscript( gene, as.vector=TRUE )[1:6]
> transcripts
[1] "ENST00000413465" "ENST00000359597" "ENST00000504290" "ENST00000510385"
[5] "ENST00000504937" "ENST00000269305"
> utrProbesets( probesets, transcripts )
  [1] "3743926" "3743927" "3743928" "3743929" "3743930" "3743931" "3743932"
  [8] "3743933" "3743938" "3743939" "3743940" "3743908"
```

So these probesets have at least one probe in the UTR regions of the first six transcripts of TP53. It is also possible to limit the search to only the 3' or 5' UTR:

```
> utrProbesets( probesets, transcripts, end='3' )
[1] "3743908"
> utrProbesets( probesets, transcripts, end='5' )
```

```
[1] "3743926" "3743927" "3743928" "3743929" "3743930" "3743931" "3743932" [8] "3743933" "3743938" "3743939" "3743940"
```

It is also possible to omit the list of probesets and the utr.probeset function will generate one for you:

```
> utrProbesets( NULL, transcripts )

[1] "3743908" "3743909" "3743910" "3743911" "3743912" "3743913" "3743914"
[8] "3743915" "3743916" "3743917" "3743918" "3743919" "3743920" "3743935"
[15] "3743936" "3743908" "3743909" "3743910" "3743911" "3743912" "3743913"
[22] "3743914" "3743915" "3743916" "3743917" "3743918" "3743919" "3743920"
[29] "3743935" "3743936" "3743908" "3743909" "3743910" "3743911" "3743912"
[36] "3743913" "3743914" "3743915" "3743916" "3743917" "3743918" "3743919"
[43] "3743920" "3743935" "3743936" "3743926" "3743927" "3743928" "3743929"
[50] "3743930" "3743931" "3743932" "3743933" "3743938" "3743939" "3743940"
[57] "3743908"
```

Or you can get a list of probesets which fall inside the coding region of a transcript by using the complementary codingProbesets function:

```
> codingProbesets( NULL, transcripts )
[1] "3743905" "3743907" "3743908" "3743909" "3743910" "3743911" "3743912"
[8] "3743913" "3743914" "3743915" "3743916" "3743917" "3743918" "3743919"
[15] "3743920" "3743921" "3743922" "3743923" "3743924" "3743925" "3743935"
[22] "3743936" "3743907" "3743908" "3743909" "3743910" "3743911" "3743912"
[29] "3743913" "3743914" "3743915" "3743916" "3743917" "3743918" "3743919"
[36] "3743920" "3743921" "3743922" "3743923" "3743924" "3743925" "3743935"
[43] "3743915" "3743916" "3743911" "3743912" "3743912" "3743913" "3743914"
[50] "3743915" "3743916" "3743917" "3743918" "3743919" "3743920" "3743921"
[57] "3743922" "3743923" "3743924" "3743925" "3743936"
```

### 5.4 Coding regions and UTRs

It is possible to manipulate a list of transcripts so that you find the genomic location of their coding region or UTR. This is done using the two complementary methods transcriptToUtrRange and transcriptToCodingRange. Here for example is the normal details for a given transcript ENST00000417324 which falls on the reverse strand of Chromosome 1 in human:

```
> transcriptDetails( 'ENST00000417324' )
GRanges with 1 range and 12 metadata columns:
     seqnames
                      ranges strand |
                                           stable_id
                                                           biotype
                                                                         status
                   <IRanges> <Rle> |
                                         <character> <character> <character>
                                  - | ENST00000417324
  [1]
            1 [34554, 36081]
                                                           lincRNA
                                                                         KNOWN
     description
                      db_display_name
                           <character> <character>
      <character>
  [1]
             <NA> HGNC transcript name FAM138A-001
                                 symbol_description translation_start
                                        <character>
                                                            <integer>
  [1] family with sequence similarity 138, member A
     {\tt translation\_start\_exon\ translation\_end\ translation\_end\_exon}
                   <integer>
                                   <integer>
                                                       <integer>
  [1]
                        <NA>
                                        <NA>
                                                             <NA>
                  analysis_name
```

```
<character>
     [1] ensembl_havana_transcript
     seqlengths:
       1
      NA
  We can also get the genomic coordinates of its coding region:
   > transcriptToCodingRange( 'ENST00000417324' )
   GRanges with 0 ranges and 20 metadata columns:
                 ranges strand | IN1 transcript_id stable_id
         <Rle> <IRanges> <Rle> | <character>
                                              <numeric> <character> <numeric>
      chromosome_id
                     biotype
                                  status description synonym_id external_db_id
          <numeric> <character> <character> <character> <numeric>
                                                                   <numeric>
      db_display_name symbol_description translation_start
          <character> <character> <character>
      translation_start_exon translation_end translation_end_exon analysis_name
                  <integer>
                                 <integer>
                                                     <integer>
                                                                <character>
          phase end.phase
      <logical> <logical>
     seqlengths:
  Or, we can get the regions covered by the UTR of this same transcript:
   > transcriptToUtrRange( 'ENST00000417324' )
   GRanges with 1 range and 4 metadata columns:
         seqnames
                        ranges strand |
                                                   IN1
                                                         prime
                      1 [34554, 36081]
                                 - | ENST00000417324
                                                                5
     [1]
         translated
          <logical>
             FALSE
     [1]
     seqlengths:
       1
      NA
  Again, these two methods take an optional 'end' parameter to specify which end of the transcript
you want to work with<sup>6</sup>:
   > transcriptToCodingRange( 'ENST00000417324', end='3' )
   GRanges with 1 range and 20 metadata columns:
         seqnames
                        ranges strand |
                                                   IN1 transcript_id
                      <IRanges> <Rle> | <character>
                                                           <numeric>
     [1]
               1 [34554, 36081]
                                 - | ENST00000417324
                                                             1949354
              stable_id gene_id chromosome_id
                                                  biotype
                                                             status
             <character> <numeric> <numeric> <character> <character>
     [1] ENST00000417324
                         615192
                                         27511 lincRNA
                                                          KNOWN
```

<numeric>

db\_display\_name

<character> <character>

symbol

description synonym\_id external\_db\_id

<character> <numeric>

<sup>&</sup>lt;sup>6</sup>if you pass an 'end' to transcriptToCodingRange, you are choosing which UTR to omit from the range, i.e. passing end='3' returns you the range that covers both the coding range and the 5' end, but not the 3' end.

```
[1]
           <NA>
                   3545894
                                  50609 HGNC transcript name FAM138A-001
                              symbol_description translation_start
                                    <character>
                                                       <integer>
 [1] family with sequence similarity 138, member A
                                                           <NA>
     translation_start_exon translation_end translation_end_exon
                               <integer>
                 <integer>
                                                   <integer>
                      <NA>
 [1]
                                    <NA>
                                                        <NA>
                analysis_name
                                phase end.phase
                  <character> <logical> <logical>
 [1] ensembl_havana_transcript
                                <NA>
 seqlengths:
   1
  NA
> transcriptToUtrRange( 'ENST00000417324', end='3')
GRanges with 1 range and 4 metadata columns:
     seqnames
                    ranges strand |
                                              IN1
                                                       prime
                  [1]
           1 [34554, 36081]
                             - | ENST00000417324
     translated
      <logical>
 [1]
         FALSE
 seqlengths:
   1
  NA
```

# 6 annmap's local cache

You may have noticed that the first time we called exonic on page 17, annmap told us that it was 'Building probeset specificity cache.....'. This cache allows us to filter large numbers of probesets much quicker than doing each in turn. However, for smaller queries of less than about 1000 probesets (on our system) it is quicker not to suffer the  $\approx$  1s load time for the cache to be loaded into memory from disk.

If you are running lots of these filtering queries on small numbers of probesets (in a loop or apply, for example), it might be worth turning the cache off whilst you run them:

```
> annmapToggleCaching()
[1] FALSE
```

Now, the cache (on by default) has been turned off. (This can be seen by the return value of FALSE). Calling toggle again will turn it back on:

```
> annmapToggleCaching()
[1] TRUE
```

Currently we cache the probeset specificity data used for the filters, described above, and the calls to allXXX. The cache is stored in .annmap/cache.

## 7 The path to the answer matters

### 7.1 The probeset boundary to commutativity

In general, annmap is commutative. For example, if you go from a list of genes to transcripts and back again, you will end up where you started:

However, with probesets, this commutativity is broken. This is because they (even exonic probesets) can hit genes, transcripts or exons other than those in the list that you passed in – for example:

```
> genes = symbolToGene( c( 'pten', 'shh' ), as.vector=TRUE )
> genes
[1] "ENSG00000171862" "ENSG00000164690"
> probesets = exonic( geneToProbeset( genes, as.vector=TRUE ) )
> probesets
```

Indeed, any mapping involving hits, probes or probesets, are not commutative.

## 8 Utility methods

### 8.1 Splicing Index

For this test, we have an eSet object stored locally which contains expression data for an MCF7/MCF10A dataset around the gene 'tp53'.

```
> load( file.path( .path.package('annmap'), 'rdata', 'HuEx-1_0.tp53.expr.RData' ) )
> x.rma

ExpressionSet (storageMode: lockedEnvironment)
assayData: 240 features, 6 samples
    element names: exprs
protocolData: none
phenoData
    rowNames: ex1MCF7_r1.CEL ex1MCF7_r2.CEL ... ex2MCF10A_r3.CEL (6
        total)
    varLabels: sample group
    varMetadata: labelDescription
featureData: none
experimentData: use 'experimentData(object)'
Annotation: huex10stv1
```

So, we can call our spliceIndex method, passing this data, the gene id for 'tp53' and the fact that the first 3 columns are one sample type (mcf7) and the second 3 are the other sample.

```
> spliceIndex( x.rma, symbolToGene( 'TP53' ), gps=list(1:3,4:6) )
$ENSG00000141510
                      p.score t.statistic
                 si
3743908 0.112899062 0.22479731 1.43429922 0.02815739
3743909 -0.157154547 0.51591129 -0.71178196 0.02815739
3743912 -0.044769083 0.70715317 -0.40359763 0.02815739
3743913 -0.186660284 0.33324256 -1.09958985 0.02815739
3743915 -0.162931473 0.08576344 -2.26962208 0.02815739
3743917 0.126866306 0.17884429 1.62805226 0.02815739
3743918 0.056591953 0.32816552 1.11278250 0.02815739
3743919  0.062286225  0.46526044  0.80632965  0.02815739
3743920 -0.156910026 0.05743946 -2.64234332 0.02815739
3743922 0.048387256 0.57010143 0.61787261 0.02815739
3743923 -0.102274940 0.36081354 -1.03096096 0.02815739
3743924  0.008979375  0.93770685  0.08317721  0.02815739
3743925 0.006989562 0.95775111 0.05636913 0.02815739
3743936 -0.865412389 0.11450250 -2.01226730 0.02815739
```

As can be seen, this call returns a list, with one data.frame per gene. In this example, we only have passed a single gene for brevity.

### 8.2 Mapping probes into transcript coordinates

To find the location of probes relative to the mature, spliced mRNA sequence (rather than in genome coordinate space), the function transcriptToTranslatedprobes joins the exons in a transcript (from the 5' end to the 3' end), and then converts the probe locations so that they are an offset from the 5' end of the transcript.

```
> transcriptToTranslatedprobes( c( 'ENST00000462694', 'ENST00000329958' ) )
```

#### \$ENST00000462694 GRanges with 11 ranges and 3 metadata columns: seqnames ranges strand | sequence probe\_hit\_count <Rle> <IRanges> <Rle> | <character> <numeric> [1] 10 [19, 43] + | AGAGATCGTTAGCAGAAACAAAAGG 10 [33, 57] + | GAAACAAAAGGAGATATCAAGAGGA 2 [2] [3] 10 [46, 70] + | ATATCAAGAGGATGGATTCGACTTA 2 + | CAAGAGGATGGATTCGACTTAGACT + | GATGGATTCGACTTAGACTTGACCT [50, 74] [4] 10 10 [56, 80] 2 [5] 10 [137, 161] + | TACAGGAACAATATTGATGATGTAG 10 [255, 279] + | AGATACTTTGTGATGTAAACTATTA 10 [290, 314] + | CTATAATCATTTTTTGGCTTACCGT . . . [7] 2 [8] 1 [9] 10 [305, 329] + | GGCTTACCGTACCTAATGGACTTCA 10 [331, 355] + | GGGGATACAGTTCATTTGATAAGAA Γ10] 10 [305, 329] 1 [11] TN1 <character> [1] ENSE00001926457 [2] ENSE00001926457 [3] ENSE00001926457 [4] ENSE00001926457 [5] ENSE00001926457 [7] ENSE00001920439 [8] ENSE00001920439 [9] ENSE00001920439 [10] ENSE00001920439 [11] ENSE00001920439 seqlengths: 10 NA

Please note, that this function does not return probes that span the exon-junctions of the combined transcript sequence.

### 8.3 Converting between Genomic, Transcript and Protein coordinates

\$ENST00000329958

NULL

In annmap, there are two functions that convert between genomic coordinates and the related transcript/protein coordinates.

```
<IRanges> <Rle> | <character>
  [1] ENST00000413465 [1018, 1018]
                                    * | transcript
 seqlengths:
  ENST00000413465
> # With as.vector=TRUE
> genomeToTranscriptCoords( start( gene ), transcripts, as.vector=TRUE )
ENST00000413465 ENST00000359597 ENST00000504290 ENST00000510385 ENST00000504937
                            NA
                                           NA
ENST00000269305 ENST00000455263 ENST00000420246 ENST00000445888 ENST00000576024
            NA
                           NA
                                           NA
                                                           NA
                                                                           NA
ENST00000509690 ENST00000514944 ENST00000574684 ENST00000505014 ENST00000508793
            NA
                           NA
                                           NA
                                                          NA
ENST00000604348 ENST00000503591
            NΑ
                            NΑ
> # Get the coding range for the transcripts
> coding = transcriptToCodingRange( transcripts )
> # And then to protein coordinates
> genomeToProteinCoords( start( coding ), proteins )
GRanges with 12 ranges and 2 metadata columns:
             seqnames
                         ranges strand
                                               frame coord.space
                <Rle> <IRanges> <Rle>
                                         | <numeric> <character>
   [1] ENSP00000269305 [394, 394]
                                        2
                                                         protein
   [2] ENSP00000352610 [344, 344]
                                                         protein
   [3] ENSP00000391127 [342, 342]
                                                   2
                                         protein
   [4] ENSP00000391478 [394, 394]
                                                   2
                                         protein
   [5] ENSP00000398846 [347, 347]
                                        2
                                                         protein
   [8] ENSP00000424104 [166, 166]
                                        0
                                                        protein
   [9] ENSP00000425104 [199, 199]
                                        2
                                                         protein
  [10] ENSP00000426252 [128, 128]
                                        1
                                                         protein
                                                  2
  [11] ENSP00000458393 [ 32, 32]
                                                         protein
  [12] ENSP00000473895 [143, 143]
                                                         protein
 seqlengths:
  ENSP00000269305 ENSP00000352610 ... ENSP00000458393 ENSP00000473895
               NA
                              NA ...
                                                                  NA
> # With as.vector=TRUE
> genomeToProteinCoords( start( coding ), proteins, as.vector=TRUE )
ENSP00000410739 ENSP00000352610 ENSP00000269305 ENSP00000398846 ENSP00000391127
                          344
                                          394
                                                          347
ENSP00000391478 ENSP00000458393 ENSP00000425104 ENSP00000423862 ENSP00000424104
           394
                           32
                                          199
                                                          156
                                                                          166
ENSP00000473895 ENSP00000426252
```

There are also functions for performing translation in the opposite direction; proteinCoordsToGenome and transcriptCoordsToGenome.

```
> pos.one = transcriptCoordsToGenome( transcripts, 1, as.vector=TRUE )
> pos.one
```

```
ENST00000413465 ENST00000359597 ENST00000504290 ENST00000510385 ENST00000504937
        7579912
                        7579912
                                        7578811
                                                        7578811
                                                                         7578811
ENST00000269305 ENST00000455263 ENST00000420246 ENST00000445888 ENST00000576024
        7590856
                        7590799
                                        7590799
                                                        7590805
                                                                         7576905
ENST00000509690 ENST00000514944 ENST00000574684 ENST00000505014 ENST00000508793
        7590805
                        7590745
                                        7578437
                                                        7590805
                                                                         7580752
ENST00000604348 ENST00000503591
        7590805
                        7590745
```

And as we are on the reverse strand, all these locations should match the end of the transcripts:

```
> pos.one == end( transcripts )
ENST00000413465 ENST00000359597 ENST00000504290 ENST00000510385 ENST00000504937
           TRUE
                          TRUE
                                           TRUE
                                                           TRUE
ENST00000269305 ENST00000455263 ENST00000420246 ENST00000445888 ENST00000576024
           TRUE
                          TRUE
                                           TRUE
                                                           TRUE
                                                                           TRUE
ENST00000509690 ENST00000514944 ENST00000574684 ENST00000505014 ENST00000508793
           TRUE
                          TRUE
                                           TRUE
                                                           TRUE
                                                                           TRUE
ENST00000604348 ENST00000503591
          TRUE
                          TRUE
```

We can also pass cds=TRUE to this method to take the UTRs into account when calculating genomic location:

```
> pos.one = transcriptCoordsToGenome( transcripts, 1, as.vector=TRUE, cds=TRUE )
> pos.one
ENST00000413465 ENST00000359597 ENST00000504290 ENST00000510385 ENST00000504937
       7579912
                     7579912
                                                        NA
                                         NA
ENST00000269305 ENST00000455263 ENST00000420246 ENST00000445888 ENST00000576024
       7579912 7579912
                                    7579912
                                                   7579912
                                                                  7576905
ENST00000509690 ENST00000514944 ENST00000574684 ENST00000505014 ENST00000508793
       7578533 7579912
                                                        NA
                                                                 7579912
ENST00000604348 ENST00000503591
                7579912
       7579912
```

### **8.4 Plotting data with genomicPlot**

genomicPlot will plot the genes and exons found at a given locus.

For example, to plot the genomic region around the gene MPI (Figure 2):

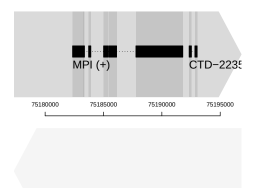
```
> range = symbolToGene( 'MPI' )
> ranges( range ) = ranges( range ) + 5000
> genomicPlot( range )
```

We can also show the opposite strand, in one of two ways. First, passing draw.opposite.strand =TRUE, results in a 'washed out' view of the opposite strand (Figure 3).

```
> genomicPlot( range, draw.opposite.strand=TRUE )
```

Alternatively setting the strand column from the range parameter we are passing in (Figure 4) to '\*' results in both strands being shown.

```
> strand( range ) = '*'
> genomicPlot( range )
```



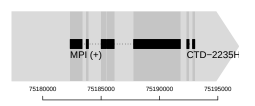
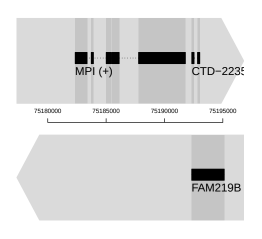


Figure 2: A genomic plot 5000bp either side of MPI.

Figure 3: A genomic plot 5000bp either side of MPI with the opposite strand drawn 'washed out'.



75180000 75185000 75190000 75195000

FAM219B (-)

Figure 4: Both strands 5000bp either side of MPI.

Figure 5: Both strands 5000bp either side of MPI with highlights.

It is also possible to highlight regions of the genome with your own annotation. To do this, you simply need to pass in a data.frame containing columns start, end, strand and name.

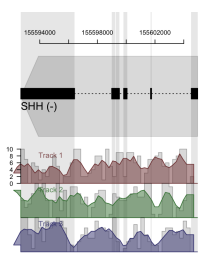
For example, we could add regions to the start and end locations of MPI but on the opposite strand and call them annmap1 and annmap2 (Figure 5);

```
= symbolToGene( 'MPI' )
> a1
> end( a1 )
             = start( a1 ) + 1000
             = symbolToGene( 'MPI' )
> start( a2 ) = end( a2 ) - 1000
             = data.frame( start =c( as.integer( start( a1 ) ), as.integer( start( a2 ) ) ),
> hig
                                 =c( as.integer( end( a1 ) ), as.integer(
                                                                              end( a2 ) ) ),
                           strand=c(
                                       strandAsInteger( a1 ),
                                                                  strandAsInteger( a2 )
                           name = c(
                                       'annmap1',
                                                    'annmap2'))
```

> genomicPlot( range, highlights=hig )

### 8.5 Plotting NGS data with ngsBridgePlot

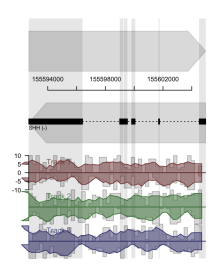
The ngsBridgePlot method allows NGS read data to be plotted alongside a genomicPlot. This was how Figure 1 was generated. There are many options for this plot function; here we will cover the most important ones.



155594000 155598000 155602000 SHH (-)

Figure 6: Dummy (in this example) NGS data shown alongside SHH.

Figure 7: Dummy (in this example) NGS data shown alongside SHH with probes shown.



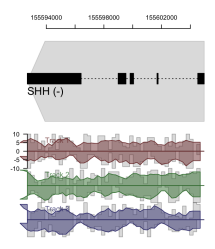


Figure 8: Dummy (in this example) NGS data shown alongside SHH (both strands) with probes shown.

Figure 9: Dummy (in this example) NGS data shown alongside SHH. Both strands of NGS reads are shown. Exon depth plot hidden.

To plot Bamfile data alongside a region (and strand) denoted by a given gene, you can do (Figure 6):

```
> gene = symbolToGene( 'SHH' )
> ngsBridgePlot( gene, bamfileData )
```

If you want to show where the probes for a given array hit this region, you can set the probe.plot parameter to the genomicProbePlot function (Figure 7)

```
> gene = symbolToGene( 'SHH' )
> ngsBridgePlot( gene, bamfileData, probe.plot=genomicProbePlot )
```

And as we saw in the preamble to this document, we can set the strand of the given range to '\*' and both strands of the region will be shown (Figure 8).

```
> gene = symbolToGene( 'SHH' )
> strand( gene ) = '*'
> ngsBridgePlot( gene, bamfileData )
```

Or, we can plot the genomicPlot for a single strand, but show the NGS data for both strands side-by-side using the trace.match.strand parameter and, for clarity, turn off the exon.depth.plot (Figure 9).

```
> gene = symbolToGene( 'SHH' )
> # trace.match.strand=FALSE would have the same effect
> ngsBridgePlot( gene, bamfileData, exon.depth.plot=NULL, trace.match.strand='*' )
```

In all these examples, the bamfileData is a synthetic object we have created for the purposes of this vignette. Its structure is:

```
> str( bamfileData )
List of 3
 $ :List of 3
  ..$ name: chr "Track 1"
  ..$ rle :List of 2
  ....$ +:Formal class 'Rle' [package "IRanges"] with 4 slots
 ..... ... ... @ values : int [1:46] 5 10 5 10 1 6 0 2 10 1 ...
                        : int [1:46] 155592680 245 245 245 245 245 245 245 245 ...
  .. .. .. ..@ lengths
  .. .. .. .. @ elementMetadata: NULL
  ....$ -: Formal class 'Rle' [package "IRanges"] with 4 slots
                         : int [1:42] 3 10 7 0 6 5 1 4 2 5 ...
  .. .. .. ..@ values
  .. .. .. .. @ lengths
                           : int [1:42] 155592680 245 245 245 245 245 245 245 245 ...
 .. .. .. .. @ elementMetadata: NULL
 .. .. .. .. @ metadata
                        : list()
 ..$ col : chr "#804040FF"
 $ :List of 3
 ..$ name: chr "Track 2"
  ..$ rle :List of 2
  ....$ +:Formal class 'Rle' [package "IRanges"] with 4 slots
 .. .. .. ..@ values
                        : int [1:46] 9 8 6 2 0 2 8 5 1 9 ...
                          : int [1:46] 155592680 245 245 245 245 245 245 245 245 ...
  .. .. .. ..@ lengths
  .. .. .. .. @ elementMetadata: NULL
  ....$ -: Formal class 'Rle' [package "IRanges"] with 4 slots
                           : int [1:43] 8 5 3 4 7 10 8 10 5 0 ...
  .. .. .. ..@ values
  ..... @ lengths : int [1:43] 155592680 245 245 245 245 245 245 245 245 ...
  .. .. .. ..@ elementMetadata: NULL
  .. .. .. .. @ metadata
                          : list()
  ..$ col : chr "#408040FF"
 $ :List of 3
  ..$ name: chr "Track 3"
 ..$ rle :List of 2
  ....$ +:Formal class 'Rle' [package "IRanges"] with 4 slots
```

As you can see, it is a list containing a list for each track. Each of these track lists contain the name of the track, col (its colour), and an rle element which contains a list of Rle elements (one for '+', the forward strand and one for the reverse; '-')

To generate this object from actual BAM file data there is a helper function generateBridgeData. Given 3 BAM files you wish to display on the plot — data1.bam, data2.bam and data3.bam:

```
> bamfiles = c( 'data1.bam', 'data2.bam', 'data3.bam' )
> data = generateBridgeData( symbolToGene( 'SHH' ), bamfiles )
```

Will generate the required format with sensible defaults for name and col. These defaults can be overridden by setting the names and colours parameters on the generateBridgeData function.