RSNPper: utilities for SNP data

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1 Introduction

This document describes **RSNPper** version 1.0, added to Bioconductor in October of 2003. This first version focuses on SNP metadata, with functions that retrieve SNP-related data from the Boston Children's Hospital Informatics Program SNPper web service ?.

Earlier non-released versions of this package included considerable code for working with prettybase format and for conducting other tasks in SNP discovery projects. That material has been moved to inst/OLD and may be re-introduced later. Users seeking legacy support should contact the author.

2 How it works

The core of this package is the XML-RPC service at CHIP accessible through the following URL stub:

```
> print(.SNPperBaseURL)
```

[1] "http://snpper.chip.org/bio/rpcserv/dummy?cmd="

The useSNPper function allows you to work directly with the XML-RPC server by packing up appropriate command and argument strings.

```
> dput(useSNPper)
function (cmd, parmstring)
{
    targ <- url(paste(.SNPperBaseURL, cmd, parmstring, sep = ""))</pre>
    open(targ)
    on.exit(close(targ))
    readLines(targ)
}
> print(useSNPper("geneinfo", "&name=CRP")[1:7])
[1] " <SNPPER-RPC SOURCE=\"*RPCSERV-NAME*\" VERSION=\"$Revision: 1.1.1.1 $\" GENOME=\"h
[2] "
       <GENEINFO>"
[3] "
         <GENE ID=\"1735\">"
           <GENEID>1735</GENEID>"
[4] "
[5] "
           <NAME>CRP</NAME>"
[6] "
           <CHROM>chr1</CHROM>"
[7] "
           <STRAND>-</STRAND>"
```

The main functions of *RSNPper* attend to simplifying specification of parameters and parsing and packaging the XML results.

Note on auditability. All functions return textual information coupled with auditing information as a 'toolInfo' attribute, detailing the SNPper supplied information on the human genome sequence build, the dbSNP version, and the SNPper version from which the results are obtained. At present, there is one exception: when itemsInRange is invoked with item='countsnps, no toolInfo data is obtained. This will be corrected once the countsnps command at SNPper returns valid XML element tags.

3 Overview of the functions

The current set of functions intended for investigative use is:

- geneInfo general information about location and nomenclature
- geneLayout information about exon locations
- geneSNPs all SNPs associated with a given gene
- SNPinfo detailed information on a SNP
- itemsInRange supports chromosome scanning for genes, SNPs, or counts of SNPs

An omission: for SNP information, I have not collected information on submitter.

4 Demonstrations

4.1 Obtaining information on genes

The geneInfo function will collect some basic information on a gene. The gene may be specified by HUGO name, mRNA accession number, or SNPper id.

```
> print(geneInfo("CRP"))
```

```
SNPper Gene metadata:
There are 1 entries.
Basic information:
     GENEID NAME CHROM STRAND
                                                             PRODUCT NSNPS
       1735 CRP chr1
GENE
                            - C-reactive protein, pentraxin-related
                                                                       141
      TX.START
                  TX.END CODSEQ.START CODSEQ.END LOCUSLINK
                                                              OMIM UNIGENE
GENE 157948704 157951003
                            157949939
                                       157950899
                                                       1401 123260 Hs.76452
     SWISSPROT
                 MRNAACC
                           PROTACC REFSEQACC
        P02741 NM_000567 NP_000558
GENE
                                        NULL
SNPper info:
     SOURCE
                      VERSION
                                              GENOME DBSNP
[1,] "*RPCSERV-NAME*" "$Revision: 1.1.1.1 $" "hg18" "125"
```

The geneLayout function provides information on exon locations.

> print(geneLayout("546"))

" " "S100PBP" "chr1" "33055763" CODINGSEQ.START TRANSCRIPT.END CODINGSEQ.END exon1.start "33064288" "33097062" "33094226" "33055763" exon1.end exon2.start exon2.end exon3.start "33055877" "33063501" "33063617" "33064286"	ID
"33064288" "33097062" "33094226" "33055763" exon1.end exon2.start exon2.end exon3.start	
exon1.end exon2.start exon2.end exon3.start	CODINGSEQ.START
	"33064288"
"33055877" "33063501" "33063617" "33064286"	exon1.end
	"33055877"
exon3.end exon4.start exon4.end exon5.start	exon3.end
"33065118" "33066181" "33066269" "33068152"	"33065118"
exon5.end exon6.start exon6.end exon7.start	exon5.end
"33068255" "33091267" "33091354" "33094112"	"33068255"
exon7.end	exon7.end
"33097062"	"33097062"
attr(,"toolInfo")	<pre>attr(,"toolInfo")</pre>
SOURCE VERSION GENOME	S
"*RPCSERV-NAME*" "\$Revision: 1.1.1.1 \$" "hg18"	"*RPCSERV-N
DBSNP	
"125"	

Information on all the genes catalogued in a certain chromosomal region can be obtained using *itemsInRange*.

<pre>> print(itemsInRange("genes",</pre>	"chr1", "156400000",	"156500000"))
[[1]]		and on
		CHROM
"CD1D" PRODUCT		"chr1" NSNPS
		NSNPS "114"
"CD1D antigen, d polypeptide"		
[[2]]		
NAME	CHROM	PRODUCT
"CD1A"	"chr1" "(CD1A antigen precursor"
NSNPS		
"53"		
\$CHR		
[1] "chr1"		
\$START		
[1] "156400000"		
\$END		
[1] "156500000"		
\$COUNT		
[1] "2"		
attr(,"toolInfo")		
SOURCE	VERSION	GENOME
"*RPCSERV-NAME*" "\$Revisi	ion: 1.1.1.1 \$"	"hg18"
DBSNP		
"125"		

4.2 Obtaining information on SNPs

Suppose you want information on the SNP with dbSNP id rs25.

```
> print(SNPinfo("25"))
```

```
SNPper SNP metadata:
DBSNPID CHROMOSOME POSITION ALLELES VALIDATED
```

[1,] "rs25" "chr7" "11550667" "A/G" "Y"
There are details on 4 populations
and 1 connections to gene features
SNPper info:
 SOURCE VERSION GENOME DBSNP
[1,] "*RPCSERV-NAME*" "\$Revision: 1.1.1.1 \$" "hg18" "125"

Suppose instead you want information on all the SNPs cataloged in a certain chromosomal region.

```
> ird <- itemsInRange("snps", "chr1", "156400000", "156500000")
> print(length(ird))
```

[1] 557

```
> print(ird[1:3])
```

[[1]]

DBSNPID	TSCID	CHROMOSOME	POSITION	ALLELES	ROLE
"rs16839876"		"chr1"	"156400131"	"A/T"	
RELPOS	AMINO	AMINOPOS			
[[2]]					
DBSNPID	TSCID	CHROMOSOME	POSITION	ALLELES	ROLE
"rs12117055"		"chr1"	"156400300"	"C/T"	
RELPOS	AMINO	AMINOPOS			
н н					
[[3]]					
DBSNPID	TSCID	CHROMOSOME	POSITION	ALLELES	ROLE
"rs17455763"		"chr1"	"156400743"	"A/T"	
RELPOS	AMINO	AMINOPOS			
11 11					

Note that the start and end locations are supplied as strings. This is to avoid coercion to textual scientific notation.

Additional detail on the count of SNPs can be obtained more briefly:

```
> print(itemsInRange("countsnps", "chr1", "156400000", "156500000"))
```

total exonic nonsyn 553 12 2 To see all the SNPs associated with a given gene, use the geneSNPs function. This requires knowledge of the SNPper gene id, which can be obtained using geneInfo.

> gs <- geneSNPs("546") > print(length(gs))						
[1] 150						
> print(gs[1:3])						
[[1]]	DBSNPID "rs11809784" CHROMOSOME "chr1" ALLELES "A/C" RELPOS "-18168" AMINOPOS "" LOCUSLINK "64766" MRNA "NM_022753"	"S100P	binding	protein	Riken	TSCID """ POSITION "33046120" ROLE "Promoter" AMINO "" HUGO "S100PBP" NAME isoform a"
[[2]]	DBSNPID "rs4422972" CHROMOSOME "chr1" ALLELES "G/T" RELPOS "-17788" AMINOPOS "" LOCUSLINK "64766" MRNA "NM_022753"	"S100P	binding	protein	Riken	TSCID "" POSITION "33046500" ROLE "Promoter" AMINO "" HUGO "S100PBP" NAME isoform a"

[[3]]

DBSNPID					TSCID
"rs3845499"					
CHROMOSOME					POSITION
"chr1"					"33047367"
ALLELES					ROLE
"A/G"					"Promoter"
RELPOS					AMINO
"-16921"					
AMINOPOS					HUGO
					"S100PBP"
LOCUSLINK					NAME
"64766"	"S100P	binding	protein	Riken	isoform a"
MRNA					
"NM_022753"					

5 Application: SNP density on chr 1

Human chromosome 1 is approximately 300Mb, and 142,629 SNPs have been recorded as of dbSNP build 106, according to NCBI SNP/maplists/maplist-newmap.html on 13 Sep 03. Let's see if these facilities can recover this sort of information. Counting the number of SNPs on a long chromosomal region seems to take a long time for SNPper, so we will break up the task.

```
> print(itemsInRange("countsnps", "chr1", "1", "100000"))
total exonic nonsyn
   340
           23
                   3
> system("sleep 2")
> print(itemsInRange("countsnps", "chr1", "100001", "200000"))
 total exonic nonsyn
    28
            0
                   0
> system("sleep 2")
> print(itemsInRange("countsnps", "chr1", "200001", "300000"))
total exonic nonsyn
    48
            0
                   0
> system("sleep 2")
```

These runs complete in a reasonable amount of time. Here we will just look at the first 2Mb in intervals of .1Mb.

```
> starts <- as.character(as.integer(seq(1, 2000001, 1e+05)))
> ends <- as.character(as.integer(as.integer(starts) + 99999))
> out <- matrix(NA, nr = 20, nc = 3)
> for (i in 1:20) {
+ cat(i)
+ out[i, ] <- itemsInRange("countsnps", "chr1", starts[i],
+ ends[i])
+ system("sleep 2")
+ }</pre>
```

1234567891011121314151617181920

> print(out)

	[,1]	[,2]	[,3]
[1,]	340	23	3
[2,]	28	0	0
[3,]	48	0	0
[4,]	14	0	0
[5,]	4	0	0
[6,]	288	0	0
[7,]	49	0	0
[8,]	683	12	1
[9,]	454	39	14
[10,]	461	45	12
[11,]	359	13	0
[12,]	442	54	16
[13,]	400	75	25
[14,]	340	36	6
[15,]	431	38	12
[16,]	292	10	2
[17,]	385	37	9
[18,]	201	7	1
[19,]	405	20	4
[20,]	414	9	3