The sceUpstr support in package harbChIP

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| 1 | Introduction | |
| The intent of this package is to allow code like the following: | | |
| > | library(harbChIP) data(sceUpstr) sceUpstr | |
| upstreamSeqs instance, organism sce There are 6674 entries first keys: [1] "YALOO1C" "YALOO2W" "YALOO3W" "YALOO4W" "YALOO5C" | | |
| > | > getUpstream("YAL001C", sceUpstr) | |
| \$Υ | YALOO1C | |

 $\verb|seq: CTGTACCACTATAATAATTTATCTTGATCGTATTAT...AGGACGTTTGGTTGAAGCCAACTAGCCACAAGAAAA|$

2 Building the sceUpstr object

500-letter DNAString object

Upstream sequences of length 500bp were obtained from the SGD website: www.yeastgenome.org -> Download Data -> FTP sequence/genomic_sequence/orf_dna/archive/utr5_sc_500.20040206.fasta.gz

```
> fname = system.file("extdata/utr5_sc_500_20040206.fasta", package="sceUpstr")
> utr5 = readFASTA(fname)
> sceUpstr = buildUpstreamSeqs2(utr5)
> save(sceUpstr, file="sceUpstr.rda")
```

3 Checking a finding of Harbison et al.

It is asserted in Fig 1 B of the paper that GGCGCTA is specifically bound by Snt2. We will examine the frequency of this heptamer in upstream regions and relate to the binding ratio distribution.