

# BSgenome.Cfamiliaris.UCSC.canFam2

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Cfamiliaris

*Canis lupus familiaris full genome (UCSC version canFam2)*

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## Description

Canis lupus familiaris full genome as provided by UCSC (canFam2, May 2005) and stored in Biostrings objects.

## Note

This BSgenome data package was made from the following source data files:

```
sequences: chromFa.tar.gz  
from http://hgdownload.cse.ucsc.edu/goldenPath/canFam2/bigZips/  
AGAPS masks: all the chr*_gap.txt.gz files from ftp://hgdownload.cse.ucsc.edu/go/  
RM masks: http://hgdownload.cse.ucsc.edu/goldenPath/canFam2/bigZips/chromOut.tar  
TRF masks: http://hgdownload.cse.ucsc.edu/goldenPath/canFam2/bigZips/chromTrf.tar
```

See [?BSgenomeForge](#) and the BSgenomeForge vignette (`vignette("BSgenomeForge")`) in the BSgenome software package for how to make a BSgenome data package.

## Author(s)

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## See Also

[BSgenome-class](#), [DNAString-class](#), [available.genomes](#), [BSgenomeForge](#)

## Examples

```
Cfamiliaris  
seqlengths(Cfamiliaris)  
Cfamiliaris$chr1 # same as Cfamiliaris[["chr1"]]  
  
if ("AGAPS" %in% masknames(Cfamiliaris)) {  
  
  ## Check that the assembly gaps contain only Ns:  
  checkOnlyNsInGaps <- function(seq)
```

```
{  
  ## Replace all masks by the inverted AGAPS mask  
  masks(seq) <- gaps(masks(seq) ["AGAPS"] )  
  unique_letters <- uniqueLetters(seq)  
  if (any(unique_letters != "N"))  
    stop("assembly gaps contain more than just Ns")  
}  
  
## A message will be printed each time a sequence is removed  
## from the cache:  
options(verbose=TRUE)  
  
for (seqname in seqnames(Cfamiliaris)) {  
  cat("Checking sequence", seqname, "... ")  
  seq <- Cfamiliaris[[seqname]]  
  checkOnlyNsInGaps(seq)  
  cat("OK\n")  
}  
}  
  
## See the GenomeSearching vignette in the BSgenome software  
## package for some examples of genome-wide motif searching using  
## Biostrings and the BSgenome data packages:  
if (interactive())  
  vignette("GenomeSearching", package="BSgenome")
```

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