

# TBX20 RNA-Seq data subset

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## 1 Introduction

The TBX20 data set [4] provides ChIP-Seq and RNA-Seq data. In here only the RNA-Seq part of the data is utilized. The raw data were downloaded from Gene Expression Omnibus (GEO) [1], accession number GSM767225-GSM767230. TBX20 (T-box 20) in general is a transcriptional regulator essential for cardiac development and maintenance of mouse heart tissue. In this study TBX20 was knocked-out by using a Tamoxifen mediated conditional knock-out system. Transcriptional changes caused by the ablation of the second exon of TBX20 result in rapid onset of heart failures and the subsequent death of the mice. TBX20 knock-out adult heart tissue was compared to wild type adult heart tissue. This package provides a subset of the RNA-Seq data (chromosome 19) for demonstrating the capabilities of the *SpliceGraph* package. The vignette describes how to access the phenotypic data and the raw reads aligned with Bowtie [3] to the mm9 assembly of *Mus musculus* from UCSC Genome Browser [2].

Accessing the experimental design ...

```
> library("TBX20BamSubset")
> fn <- system.file("extdata", "phenoData.txt",
+                     package="TBX20BamSubset")
> pd <- read.table(fn, header=TRUE,
+                     stringsAsFactors=FALSE)
```

Accessing the raw reads ...

```
> library("Rsamtools")
> fls <- getBamFileList()
> bfs <- BamFileList(fls)
```

	SRX	SRR	GSM	condition	replicate
1	SRX085099	SRR316184	GSM767225	normal	1
2	SRX085100	SRR316185	GSM767226	normal	2
3	SRX085101	SRR316186	GSM767227	normal	3
4	SRX085102	SRR316187	GSM767228	Tbx20 knockout	1
5	SRX085103	SRR316188	GSM767229	Tbx20 knockout	2
6	SRX085104	SRR316189	GSM767230	Tbx20 knockout	3

Table 1: Design of the TBX20 experiment

## References

- [1] Tanya Barrett, Dennis~B. Troup, Stephen~E. Wilhite, Pierre Ledoux, Carlos Evangelista, Irene~F. Kim, Maxim Tomashevsky, Kimberly~A. Marshall, Katherine~H. Phillippy, Patti~M. Sherman, Rolf~N. Muertter, Michelle Holko, Oluwabukunmi Ayanbule, Andrey Yefanov, and Alexandra Soboleva. Ncbi geo: archive for functional genomics data sets - 10 years on. *Nucleic Acids Research*, 39(suppl 1):D1005–D1010, 2011.
- [2] Pauline~A. Fujita, Brooke Rhead, Ann~S. Zweig, Angie~S. Hinrichs, Donna Karolchik, Melissa~S. Cline, Mary Goldman, Galt~P. Barber, Hiram Clawson, Antonio Coelho, Mark Diekhans, Timothy~R. Dreszer, Belinda~M. Giardine, Rachel~A. Harte, Jennifer Hillman-Jackson, Fan Hsu, Vanessa Kirkup, Robert~M. Kuhn, Katrina Learned, Chin~H. Li, Laurence~R. Meyer, Andy Pohl, Brian~J. Raney, Kate~R. Rosenbloom, Kayla~E. Smith, David Haussler, and W.~James Kent. The ucsc genome browser database: update 2011. *Nucleic Acids Research*, 2010.
- [3] Ben Langmead, Cole Trapnell, Mihai Pop, and Steven Salzberg. Ultra-fast and memory-efficient alignment of short dna sequences to the human genome. *Genome Biology*, 10(3):R25, 2009.
- [4] Noboru~J. Sakabe, Ivy Aneas, Tao Shen, Leila Shokri, Soo-Young Park, Martha~L. Bulyk, Sylvia~M. Evans, and Marcelo~A. Nobrega. Dual transcriptional activator and repressor roles of tbx20 regulate adult cardiac structure and function. *Human Molecular Genetics*, 2012.

## 2 Session Information

```
R version 3.0.0 (2013-04-03)
Platform: x86_64-unknown-linux-gnu (64-bit)
```

```
locale:  
[1] LC_CTYPE=en_US.UTF-8      LC_NUMERIC=C  
[3] LC_TIME=en_US.UTF-8       LC_COLLATE=C  
[5] LC_MONETARY=en_US.UTF-8   LC_MESSAGES=en_US.UTF-8  
[7] LC_PAPER=C                LC_NAME=C  
[9] LC_ADDRESS=C              LC_TELEPHONE=C  
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C  
  
attached base packages:  
[1] parallel stats      graphics grDevices utils      datasets methods  
[8] base  
  
other attached packages:  
[1] xtable_1.7-1           TBX20BamSubset_0.99.2 Rsamtools_1.12.0  
[4] Biostrings_2.28.0        GenomicRanges_1.12.0  IRanges_1.18.0  
[7] BiocGenerics_0.6.0  
  
loaded via a namespace (and not attached):  
[1] bitops_1.0-5    stats4_3.0.0  tools_3.0.0  zlibbioc_1.6.0
```