

# Gene Expression and Methylation from Lung Genomic Research Consortium (LGRC)

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The data is also available at <https://www.lung-genomics.org/research/>. We provide them here in a processed form to accompany the methods in the package `COPDSexualDimorphism`.

## 1 Clinical Data

Clinical phenotypes of 254 LGRC samples are given as a `data.frame` named `meta`. It has six fields: `tissueid`, `newid`, `GENDER`, `age`, `cigever`, `pkyrs`, and `diagmaj`. `tissueid` identifies the samples and `newid` identifies the subjects. Some subjects might have more than one sample from left/right/upper/lower lung or blood. These are designated by the last two letters of the tissue ID. The information for these samples have been adjudicated as described in Sathirapongsasuti et al (in review).

```
> library(COPDSexualDimorphism.data)
> `%+%` <- function(x,y) paste(x,y,sep="")
> data(lgrc.meta)
> head(meta)

  tissueid newid GENDER age   cigever pkyrs diagmaj
1 LT196199RU LT196199RU 1-Male 82 2-Ever (>100)    60 2-COPD/Emphysema
2 LT073345RU LT073345RU 1-Male 74      3-Never     0     3-Control
3 LT156041LU LT156041LU 2-Female 70 2-Ever (>100)    77 2-COPD/Emphysema
4 LT095342LU LT095342LU 1-Male 60 2-Ever (>100)    19 2-COPD/Emphysema
5 LT155982RU LT155982RU 2-Female 48 2-Ever (>100)    28 2-COPD/Emphysema
6 LT083759RL LT083759RL 1-Male 73 2-Ever (>100)   120 2-COPD/Emphysema
```

## 2 Gene Expression

Gene expression profile for 229 LGRC samples are available in two parts. One is `expr`, a matrix of 14497 Ensembl genes (rows) by 229 samples (columns), and the other is `expr.meta`, a `data.frame` of 229 samples (rows) by the subjects' clinical metadata. The subjects are arranged in the same order in the two objects.

```
> data(lgrc.expr)
> data(lgrc.expr.meta)
> dim(expr)

[1] 14497  229

> head(expr.meta)

  tissueid sample_name newid GENDER age   cigever pkyrs
1 LT001098RU LT001098RU_COPD 161745 2-Female 46 2-Ever (>100)    35
2 LT001796RU LT001796RU_CTRL 212671  1-Male  48 2-Ever (>100)    19
```

```

3 LT005419RU LT005419RU_COPD 291396 1-Male 70 2-Ever (>100) 43
4 LT007392RU LT007392RU_COPD 169067 1-Male 46 2-Ever (>100) 45
5 LT009615LU LT009615LU_CTRL 49801 2-Female 49 2-Ever (>100) 45
6 LT010491LL LT010491LL_COPD 180409 1-Male 78 2-Ever (>100) 51

diagmaj gender
1 2-COPD/Emphysema 2-Female
2 3-Control 1-Male
3 2-COPD/Emphysema 1-Male
4 2-COPD/Emphysema 1-Male
5 3-Control 2-Female
6 2-COPD/Emphysema 1-Male

```

Corresponding to the Ensembl genes in the expression profile is the data frame `genes`. This is a result of a query to BiomaRt database, stored here for convenience.

```

> data(lgrc.genes)
> head(lgrc.genes)

  ensembl_gene_id hgnc_symbol
ENSG000000000003 ENSG000000000003      TSPAN6
ENSG000000000005 ENSG000000000005      TNMD
ENSG000000000419 ENSG000000000419      DPM1
ENSG000000000457 ENSG000000000457      SCYL3
ENSG000000000460 ENSG000000000460      C1orf112
ENSG000000000938 ENSG000000000938      FGR

ENSG000000000003                                     tetraspanin 6 [Source:HGNC Symbol]
ENSG000000000005                                     tenomodulin [Source:HGNC Symbol]
ENSG000000000419 dolichyl-phosphate mannosyltransferase polypeptide 1, catalytic subunit [Source:HGNC Symbol]
ENSG000000000457                                     SCY1-like 3 (S. cerevisiae) [Source:HGNC Symbol]
ENSG000000000460                                     chromosome 1 open reading frame 112 [Source:HGNC Symbol]
ENSG000000000938 Gardner-Rasheed feline sarcoma viral (v-fgr) oncogene homolog [Source:HGNC Symbol]

  chromosome_name band strand start_position end_position
ENSG000000000003        X q22.1    -1    99883667  99894988
ENSG000000000005        X q22.1     1    99839799  99854882
ENSG000000000419        20 q13.13   -1    49551404  49575092
ENSG000000000457        1 q24.2    -1    169821804 169863408
ENSG000000000460        1 q24.2     1    169631245 169823221
ENSG000000000938        1 p36.11   -1    27938575  27961788

  ensembl_gene_id.1 entrezgene
ENSG000000000003      ENSG000000000003      7105
ENSG000000000005      ENSG000000000005      64102
ENSG000000000419      ENSG000000000419      8813
ENSG000000000457      ENSG000000000457      57147
ENSG000000000460      ENSG000000000460      55732
ENSG000000000938      ENSG000000000938      2268

```

### 3 Methylation

Methylation data for 245 LGRC subjects is provided as a data frame `methp` which contains percent methylation for 12094 variably methylated regions (VMRs). Each row provides average median absolute deviation (MAD), length, and the number of probes for a VMR.

```

> data(lgrc.methp)
> methp[1:5, c("name", "ave.mad", "length", "num.probes")]

```

		name	ave.mad	length	num.probes
1	vmr_chr1_932668_932806	0.03778364	139	5	
2	vmr_chr1_939506_939647	0.04619729	142	5	
3	vmr_chr1_966705_966843	0.05257659	139	5	
4	vmr_chr1_989551_989797	0.04155331	247	5	
5	vmr_chr1_1006424_1006565	0.04367978	142	5	

## 4 Session Information

```
> sessionInfo()
```

R version 4.4.0 beta (2024-04-15 r86425)

Platform: x86\_64-pc-linux-gnu

Running under: Ubuntu 22.04.4 LTS

Matrix products: default

BLAS: /home/biocbuild/bbs-3.19-bioc/R/lib/libRblas.so

LAPACK: /usr/lib/x86\_64-linux-gnu/lapack/liblapack.so.3.10.0

locale:

```
[1] LC_CTYPE=en_US.UTF-8      LC_NUMERIC=C
[3] LC_TIME=en_GB            LC_COLLATE=C
[5] LC_MONETARY=en_US.UTF-8   LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=en_US.UTF-8     LC_NAME=C
[9] LC_ADDRESS=C             LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
```

time zone: America/New\_York

tzcode source: system (glibc)

attached base packages:

```
[1] stats      graphics   grDevices utils      datasets  methods    base
```

other attached packages:

```
[1] COPDSexualDimorphism.data_1.40.0
```

loaded via a namespace (and not attached):

```
[1] compiler_4.4.0 tools_4.4.0
```