# The R package bigstatsr: Memory- and ComputationEfficient Tools for Big Matrices

useR!2017 lightning talk

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### About

I'm a PhD Student (2016-2019) in **Predictive Human Genetics** in Grenoble.

 $\overline{\text{Disease} \sim \text{DNA mutations}}$ 



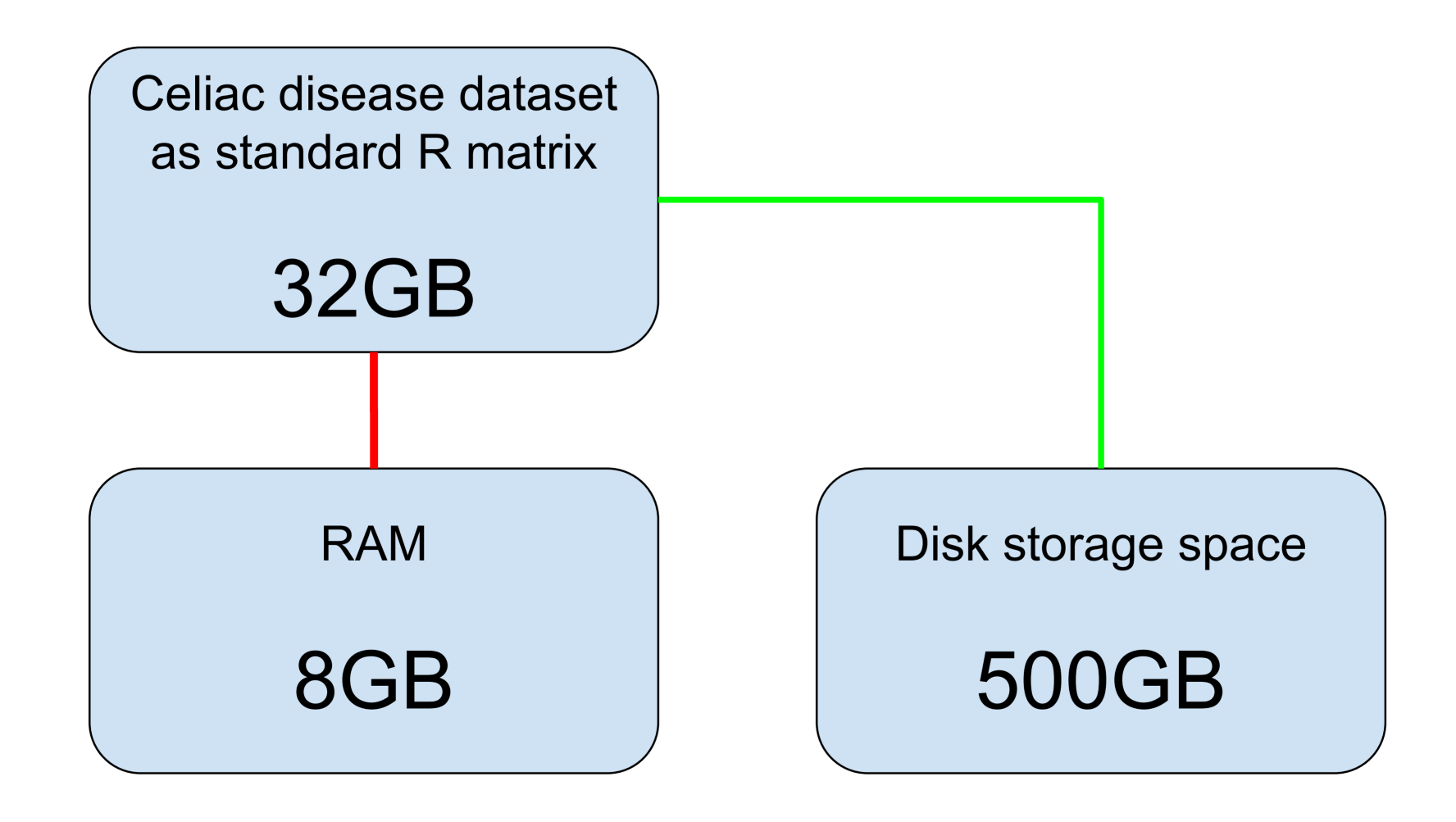
## Very large genotype matrices

• currently: 15K x 300K, celiac disease

• soon: 500K x 800K, UK Biobank

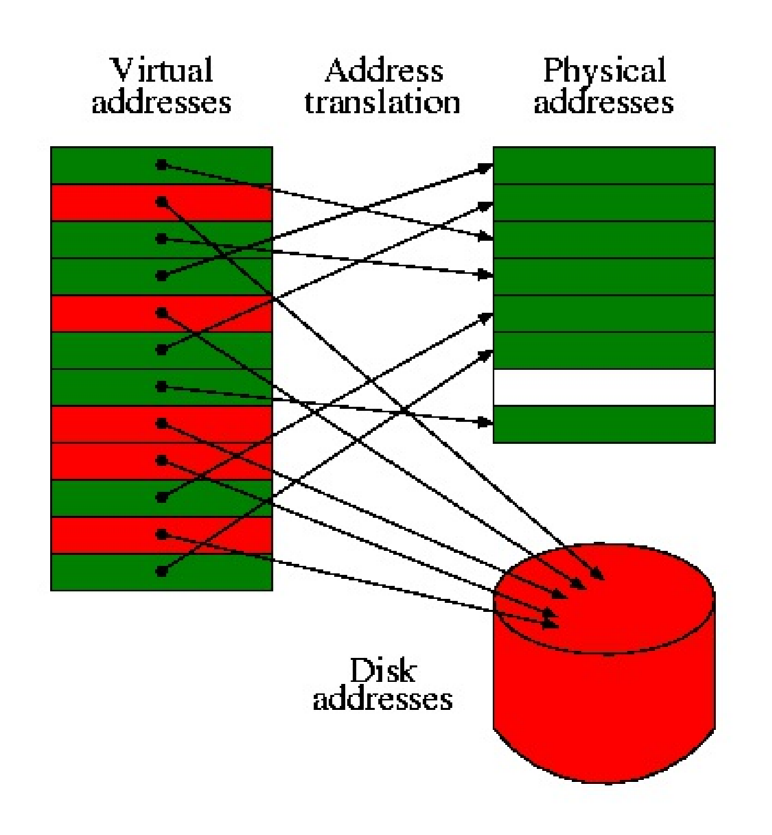


#### Problem I had



## Solution I found: R package bigmemory

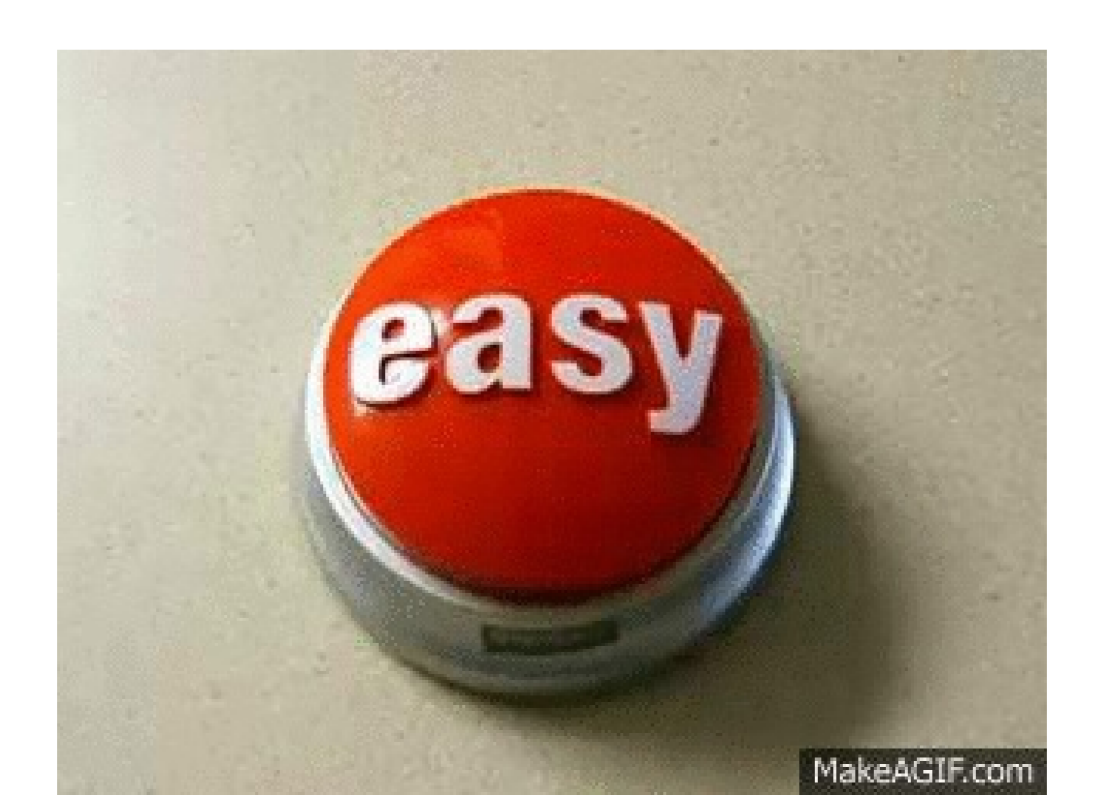
Store matrices on disk and access them from there



Michael J. Kane, John Emerson, Stephen Weston (2013).

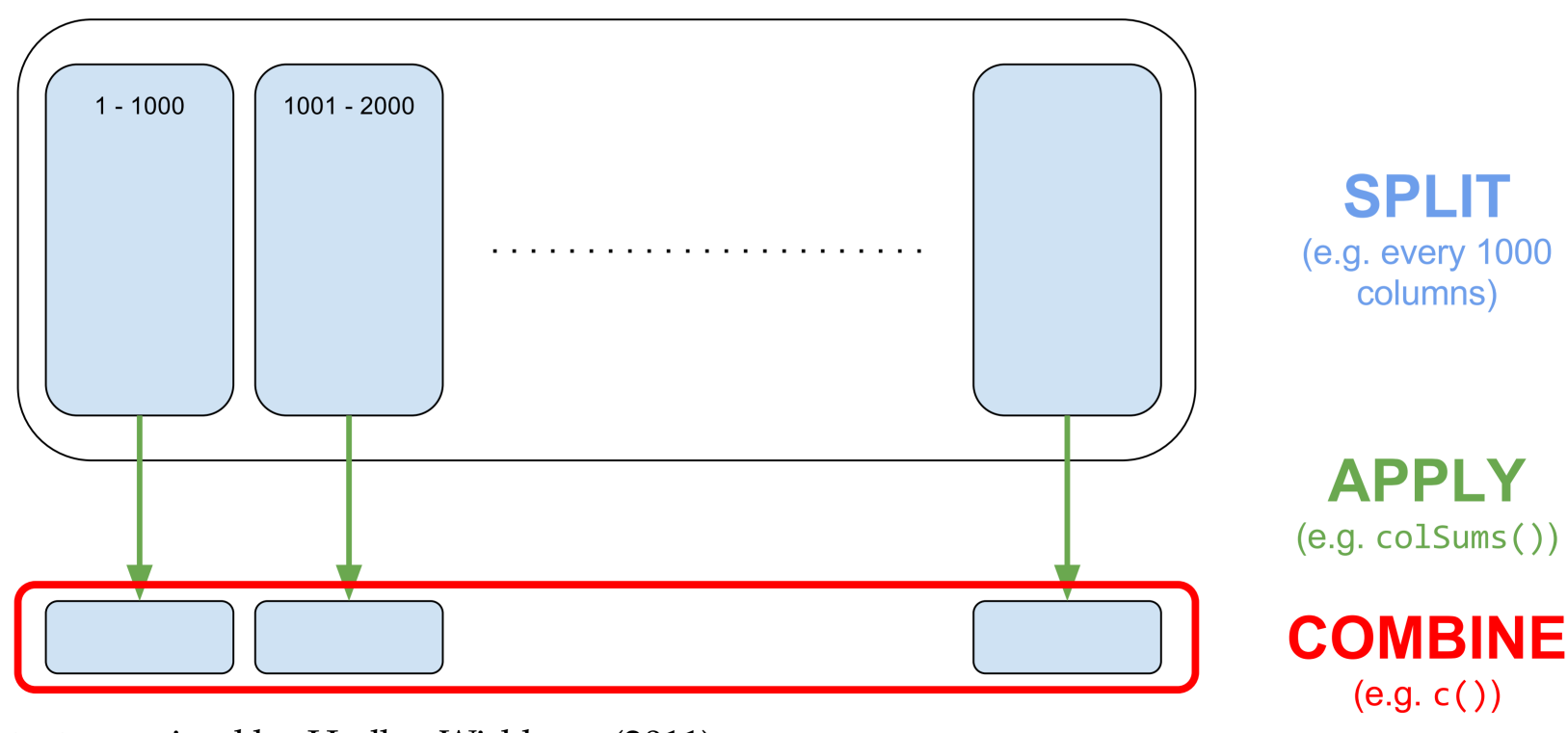
# Accesses almost as if the matrix were in memory

- in **R**: accesses with [, as standard R matrices,
- in Rcpp: accesses single elements with X(i, j), as standard Rcpp matrices.



## Split-(par)Apply-Combine Strategy

Apply standard R functions to big matrices (in parallel)



strategy coined by Hadley Wickham (2011)

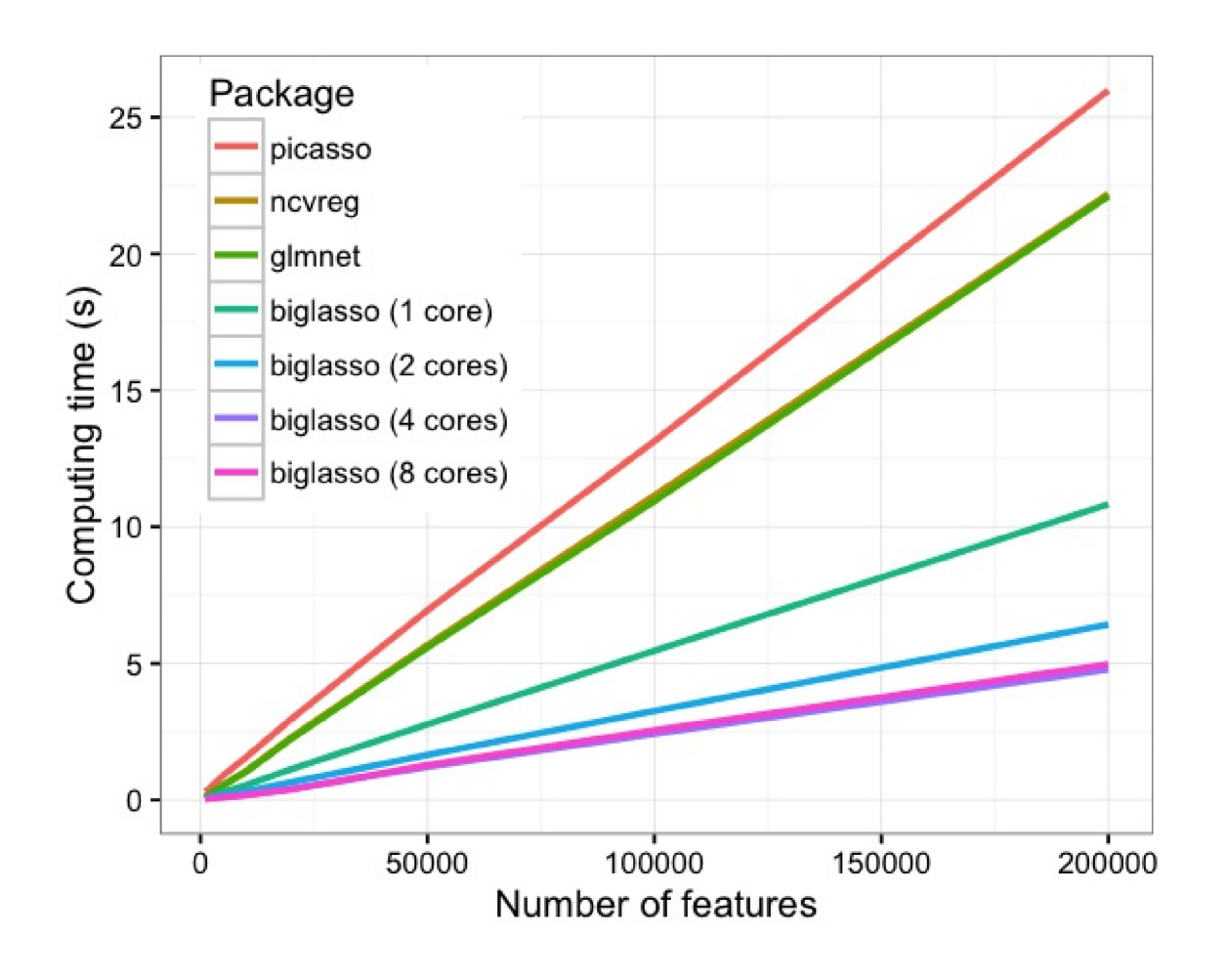
## Matrix operations

- (cross-)products with matrices/vectors
- special tricks for handling scaling (vignette and blog post)

## Example: computation of correlation of a 100,000 x 5000 matrix

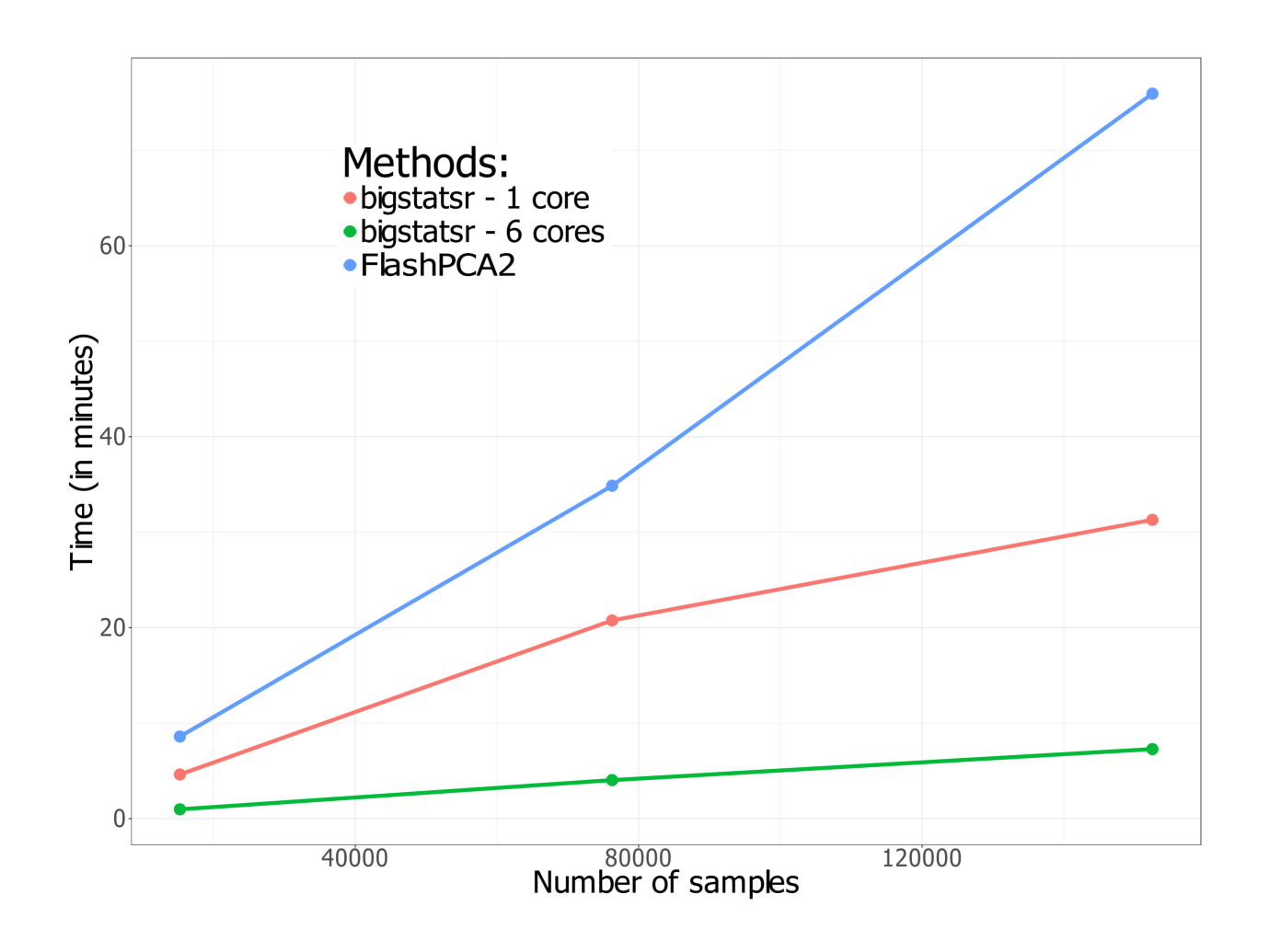
- cor: 22 minutes
- big\_cor: 1 minute

## Sparse linear models: biglasso



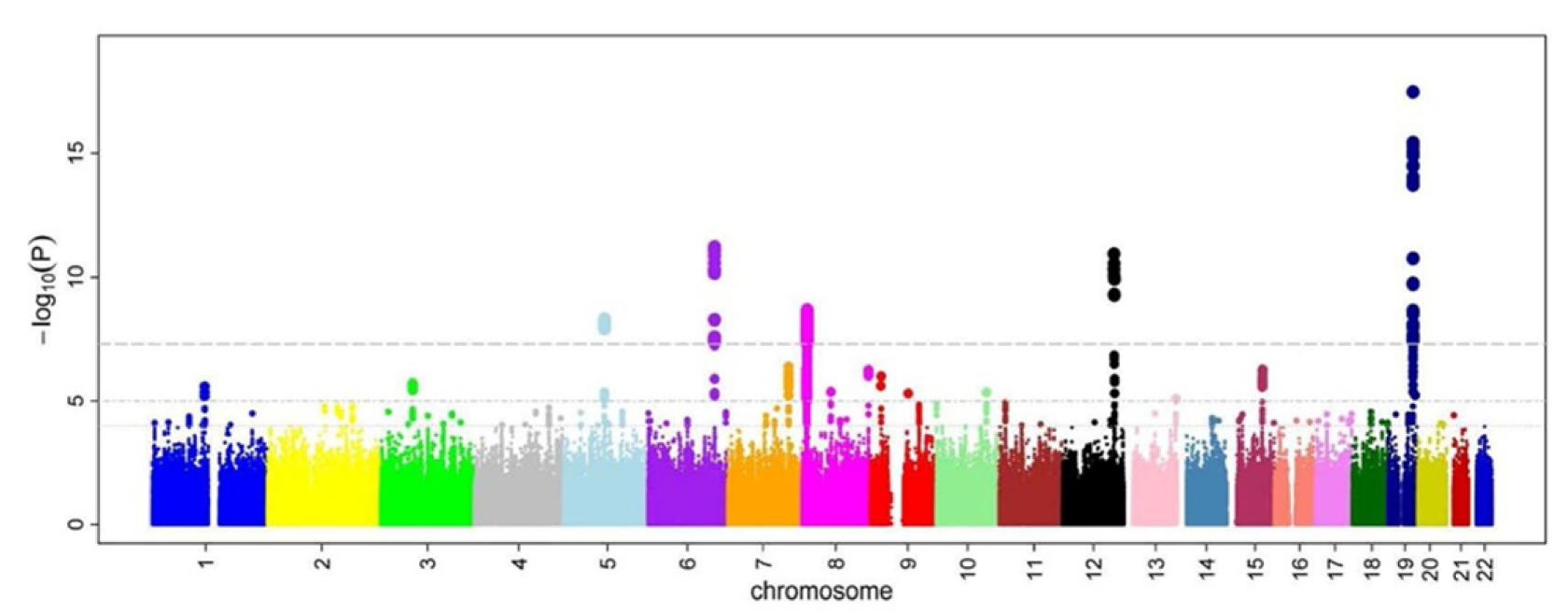
Zeng, Y., and Breheny, P. (2017).

## Partial Singular Value Decomposition



## Test association of each variable with an outcome

In genetics, this is called a Genome-Wide Association Study (GWAS)



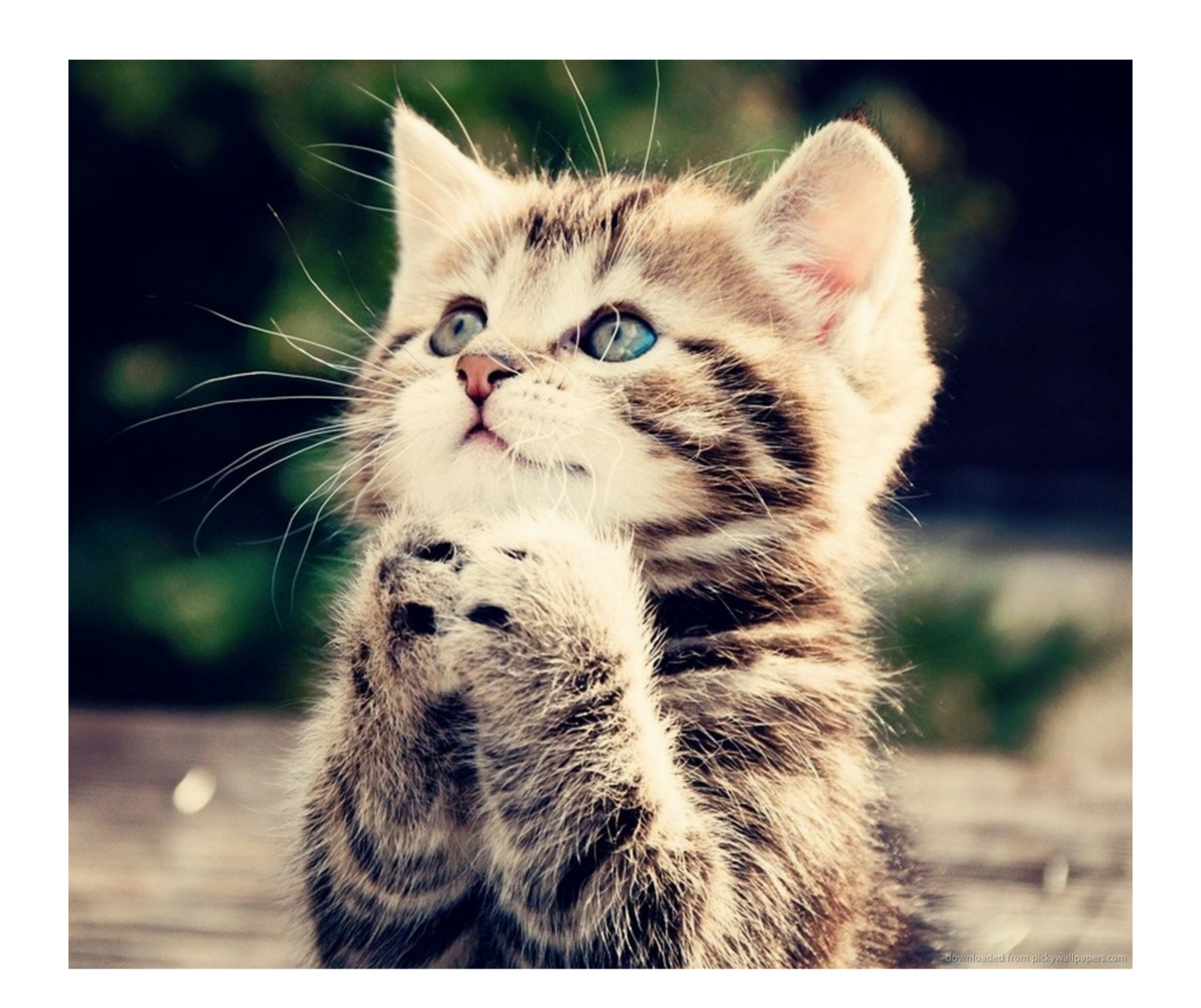
I'm now able to run algorithms on 100GB of data

## R Packages

bigmemory	`big.matrix` object
bigstatsr	Statistical functions for `big.matrix` objects to be used by any field
bigsnpr	Specific functions for SNP arrays

Paper in preparation: "Efficient management and analysis of large-scale genome-wide data with two R packages: bigstatsr and bigsnpr".

## Contributors are welcomed!



## Thanks!

Package's website: https://privefl.github.io/bigstatsr/

Twitter and GitHub: @privefl

Presentation available online: https://goo.gl/nNg0hw

Slides created via the R package xaringan.