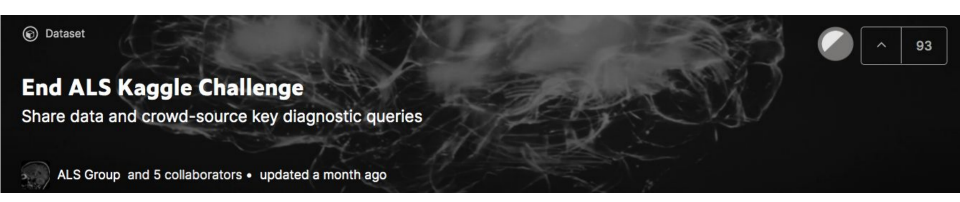
# When the outlier is the signal



Searching new genetic causes of ALS by aberrant gene expression analysis

15<sup>th</sup> May 2021

TUM - UCI team

# Task 1

Does ALS have one mechanism of action?

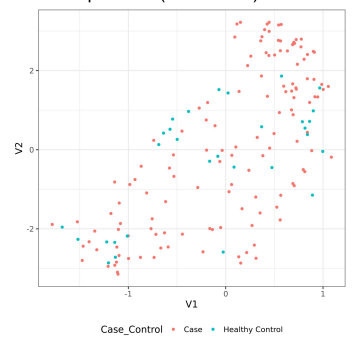
→ one pathway

Or is it caused by multiple independent or different mechanisms of action?

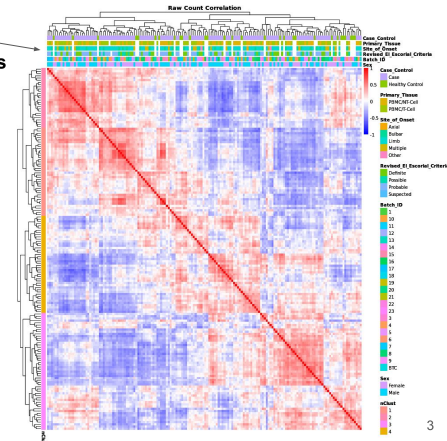
→ multiple pathways

## Gene expression does not naturally cluster donor groups

Cases and controls visualized in first two principal components (V1 and V2)

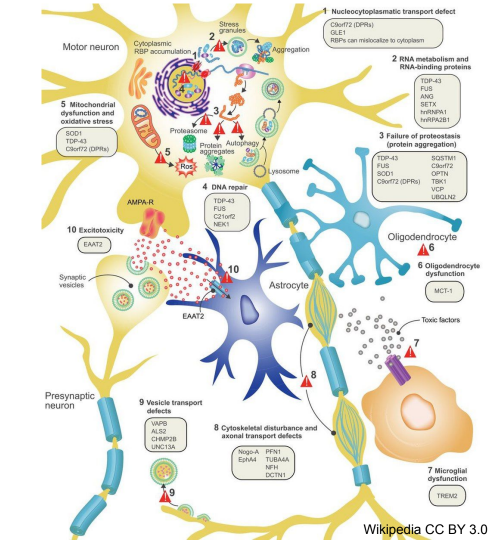


No covariate drives the sample clusters



# In fact, mutations in ca. 40 genes have been implicated in ALS over various molecular pathways<sup>1</sup>

- 1. Gregory et al. Curr Genet Med Rep (2020)
- 2. Hardiman O et al. Nat. Rev. Dis. Primers (2017)
- 3. van Damme et al. Disease Models and Mechanisms (2017)



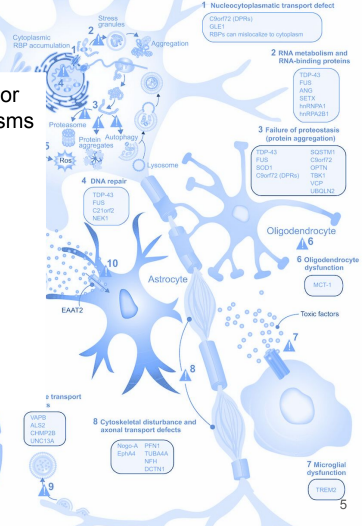
Does ALS have one mechanism of action (one pathway) or is it caused by multiple independent or different mechanisms of action (multiple pathways)?

Motor neuron

#### → Multiple pathways

However:

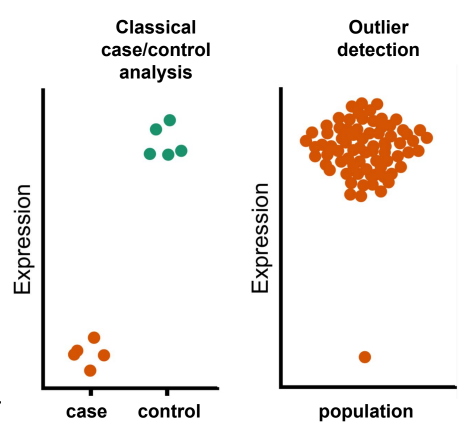
Do we find evidence for new implicated genes? .... and new pathways involved?



## Outlier detection for gene discovery

- No common pattern of gene expression among patients
- Many pathways involved

- → To search for new genes we instead ask: What makes every patient unique?
- → This leads us to focus on expression outlier detection, instead of classical differential case/control expression analysis.

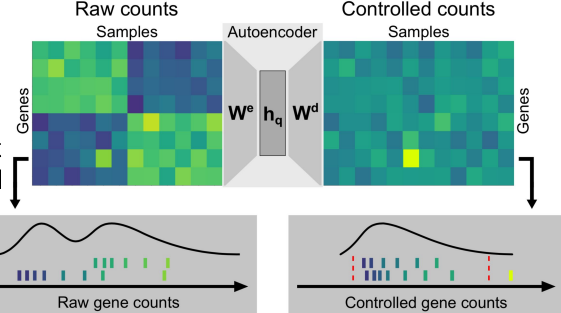


# A denoising autoencoder with a negative binomial loss to control for latent factors in RNA-Seq data

 Use a denoising autoencoder to automatically remove noise and confounding factors

Negative Binomial loss

 Number of latent factors set to maximise precision-recall of artificial outliers





# Selecting optimal latent representation

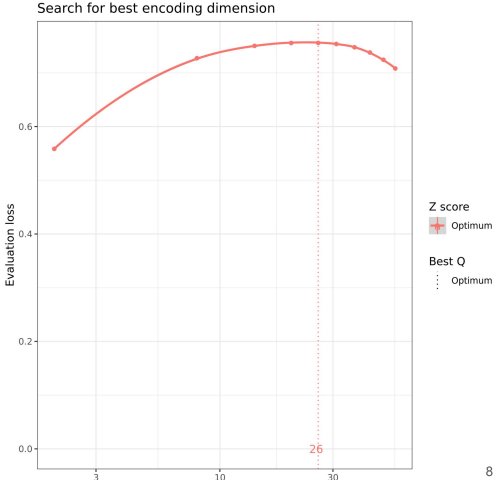
As with image processing denoising autoencoders (AEs), OUTRIDER is optimized to remove artificially injected noise





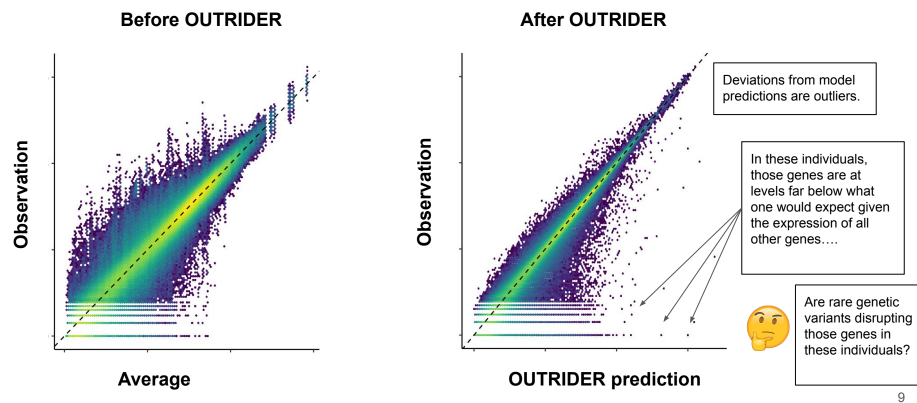
 $f(\mathbf{X^{corrupt.}}, \boldsymbol{\theta})$ 





**Encoding dimensions** 

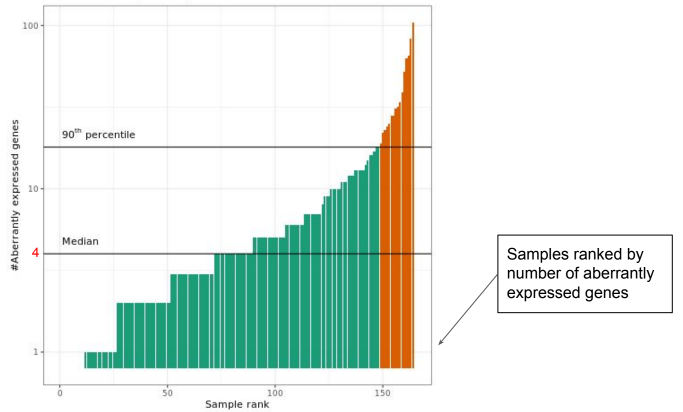
# OUTRIDER accurately predicts expression of each gene per sample and reveals outliers



Scale:  $log_{10}(count +1)$ 

## Individuals typically have 4 outlier genes

We observe a small number of aberrantly expressed genes per sample.

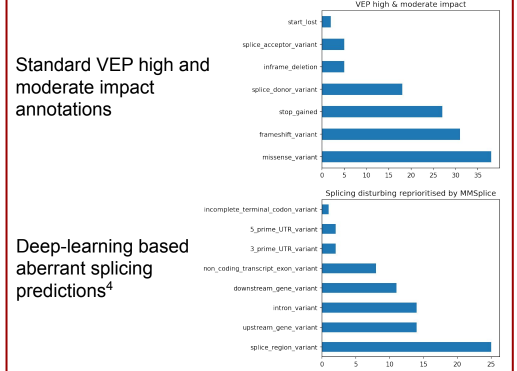


## Genetically supported gene expression outliers

Filtering outliers for having rare and **deleterious** genetic variants (from the same sample with DNA sequencing), impacting coding or splicing.

Frequency in general population < 0.1% (gnomAD<sup>5</sup>)

At most 6 samples in the cohort



<sup>4.</sup> MMSplice, Cheng et al. Genome Biology (2019)

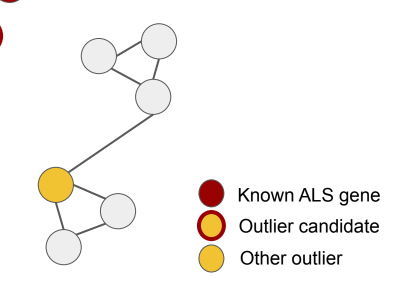
<sup>5.</sup> Karczewski et al. Nature (2020)

# Gene network analysis to study the relationship between newly discovered and known ALS genes

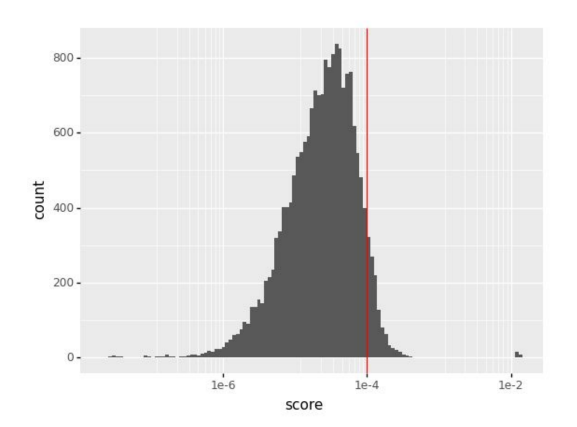
level 1: Outliers in the network vicinity of ALS genes as new

candidates

STRING <a href="https://string-db.org/">https://string-db.org/</a> was used as a gene network.



# Modeling network vicinity with random walks on gene networks

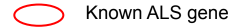


Vicinity to ALS genes modeled as the probability of visiting the gene by random walks starting from an ALS gene.

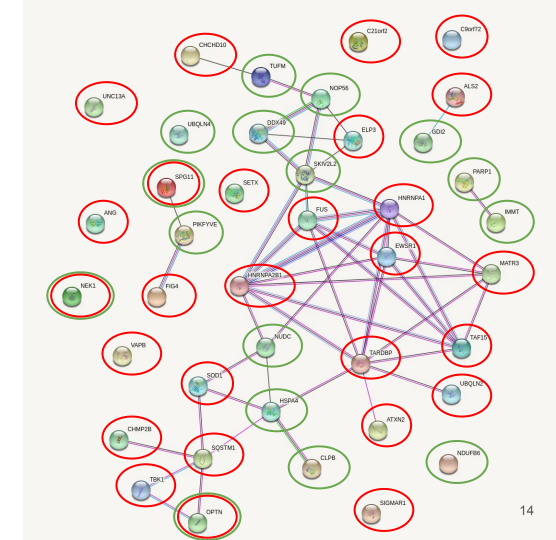
Genes with a prob. larger than 10<sup>-4</sup> (**PPI score**) were considered interesting (right tail after red line).

#### Results

Network of known ALS genes and expression **outlier genes** containing a rare deleterious **variant** and a high **PPI score**.

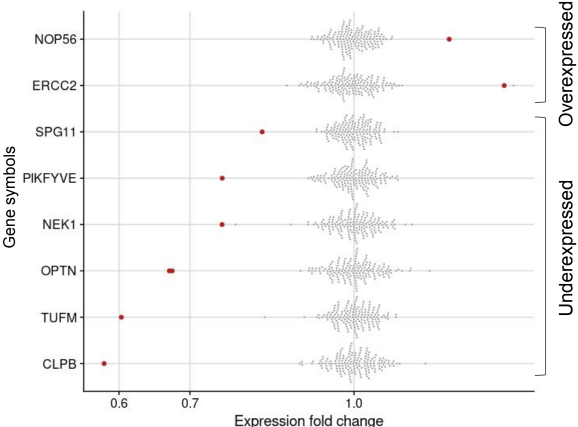


- expression outlier
  - We found 16 expression outliers interacting with known ALS genes.
  - 2. Some of them e.g. PIKFYVE connect known ALS genes



#### Identification of known genes and new interesting candidates

- Gene expression per sample
- Effect size: expression log2
  fold change
- Outliers marked in red



#### Identification of known genes and new interesting candidates

#### List of aberrantly expressed genes (outliers) containing at least one rare high impact variant.

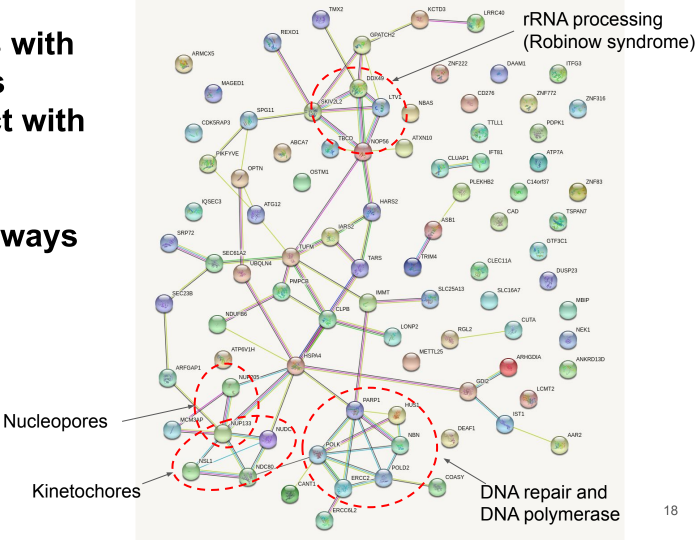
The genes are either **known to cause ALS** (according to ALSoD) or **associated to other relevant diseases**. All genes are close to the established ALS genes in the gene network.

		fold					
Sample	Gene	change	PPI score	Variant	Consequence	ClinVar	Comment
CASE.NEUEK191WYC	NEK1	0.75	1.22E-02	chr4:169424645:G>A	stop		Definitive ALS gene
CASE.NEUBK117YXL	OPTN	0.67	1.23E-02	chr10:13122390:C>A	stop		Definitive ALS gene
CASE.NEUZT557DHF	OPTN	0.67	1.23E-02	chr10:13112464:T>TAG	frameshift		Definitive ALS gene
CASE.NEUVX902YNL	SPG11	0.82	1.23E-02	chr15:44620189:C>A	splice donor	likely pathogenic	Tenuous ALS gene, variant predicted to cause aberrant splicing
CASE.NEULD354RZB	NOP56	1.23	1.61E-04	chr20:2655751:G>A	splice region		Variant predicted to cause aberrant splicing. Gene related to Ataxia.
CASE.NEUTA689LN5	TUFM	0.60	1.06E-04	chr16:28844814:G>A	stop	uncertain significance	Mitochondrial disease gene
CASE.NEUGW326BR V	CLPB	0.58	1.30E-04	chr11:72302312:G>A	stop	pathogenic	Mitochondrial disease gene
CASE.NEUME498PCJ	PIKFYVE	0.75	1.54E-04	chr2:208352730:A>AT	frameshift		Linked to neurodegeneration
CASE.NEURR881FKY	ERCC2	1.38	5.86E-05	chr19:45364832:CCTCA>	splice donor	likely pathogenic	Causes neurological symptoms, e.g. spasticity and reflex abnormalities, and skin manifestations

# Gene network analysis - level 2: Identify clusters of outliers as new candidate pathways

STRING <a href="https://string-db.org/">https://string-db.org/</a> was used as a gene network. Known ALS gene Outlier candidate Other outlier

**Further outliers with** rare deleterious variants interact with each other indicating new implicated pathways



#### **Discussion / outlook**

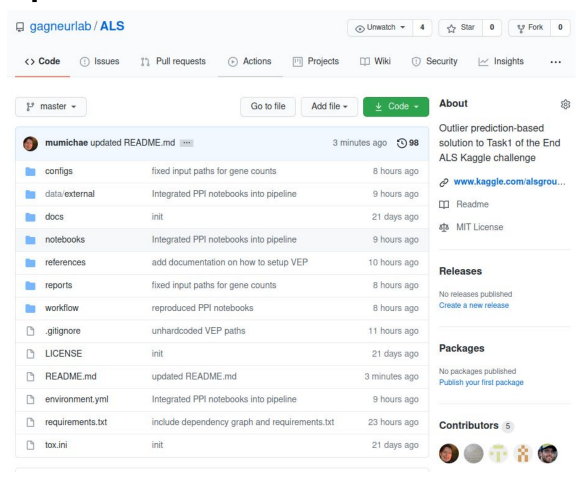
- The outlier analysis provides a new perspective, we believe more informative approach, for studying ALS.
- These new candidate genes could expand the understanding of pathways involved in the etiology of ALS.
- Future analysis would include:
  - Replicating the findings looking at WGS of the entire ALS dataset (other patients with damaging variants in the same genes).
  - Multi-omics outlier analysis: ATAC-seq, splicing, proteomics, to investigate the impact of gene regulatory control, splicing control and protein expression on ALS.
  - Integrative analysis of multi-omics data to obtain a holistic view.
  - Functional follow-ups and collaborations with experimental groups and experts in this area.

#### Conclusion

- We found variants associated with aberrant expression for known ALS genes, potentially characterising those affected patients (n = 4).
- We found new high impact variants in further cases in a gene potentially related to ALS, which would improve our catalogue of pathogenic variants.
- We found new candidate genes in known pathways.
- We found potential new pathways.
- Altogether, this gives a potential genetic explanation to 63 (46%) of the patients and further supports a multi-causal and multi-mechanism view of ALS.

#### Code to reproduce the results

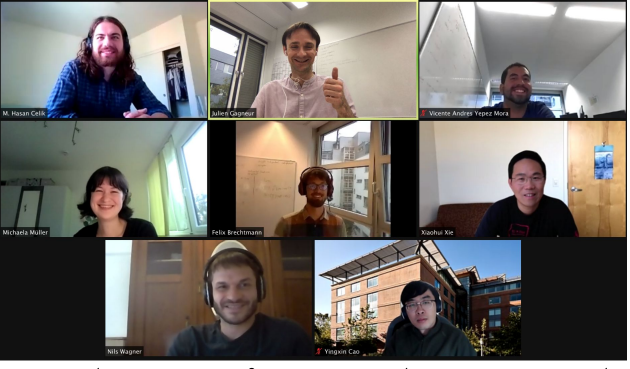
#### https://github.com/gagneurlab/ALS-kaggle





#### The team





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<sup>&</sup>lt;sup>1</sup> Technical University of Munich <sup>2</sup>University of California, Irvine

# **Appendix**

## **Analysis Workflow**

- Reproducible pipeline in Snakemake
- Parallelized and robust
- Main steps:
  - Prepare gene counts
  - OUTRIDER analysis
  - Variant annotation
  - PPI network analysis
  - UMAP on expression space (not shown here)

