

Retinoic Acid Treatment Affects Gene Expression of Immature Neuronal Markers and Up-Regulates SNAP25 Expression in SH-SY5Y cells

Eline Poinsignon-Clavel

INTRODUCTION

- SH-SY5Y cells (neuroblast, immature neuron) ¹
 - Epithelial structure, grow in clumps, not fully differentiated when used for qPCR
 - Neuronal model

What is RA?

- Active form of vit A
- **Increases cell differentiation**
- Decrease cell proliferation
 - Can suppress tumors
- Act as transcription factor in nucleus – gene expression²

RA side effect:

- Oxidative stress
- Less susceptible to get Parkinson's³

- RA Tx: homogeneous differentiated SH-SY5Y neurons
- no RA Tx: high proliferation of undifferentiated SH-SY5Y cells⁴

Research Question/Aim:

How does RA treatment affect gene expression in immature neurons?

Group data

- Collected from 3 Thursday PM classes
- 12 genes -> neuronal markers
- RA-affected because increases differentiation
- binds to receptors in promoting region
- regulates gene transcription in early development⁵ -> (state of SH-SY5Y cells)
- Down-regulated by differentiation -> cell-cell interaction which stops cellular processes, hence decreasing cell proliferation
- Up-regulated by differentiation -> for cell differentiation

Hypothesis:

RA will change gene expression of different neuronal markers

SNAP25

- Collected from 3 Thursday PM classes
- Presynaptic plasma membrane protein part of SNARE complex
 - **Negatively regulates neurotransmitter release**
 - Negatively regulates intracellular Ca^{2+} from V-gated Ca^{2+} channels⁶
- Normal postsynaptic surface expression of NMDARs⁷
- RA Tx -> levels double⁸
 - Increased differentiation & decrease proliferation -> decrease cellular synaptic pathways

Hypothesis:

RA will increase SNAP25 expression (up-regulation)

METHODOLOGY

Independent variable: RA treatment

Dependent variable: Gene expression level

1. SH-SY5Y treated with 10 μ M RA (control = no RA) for 72h
 2. qPCR steps applied as in Figure 1
- Same treatment done on other sample using RNA sequencing (by technicians)

Baselines:

- qPCR normalized w GAPDH
- RNAseq normalized w TPM

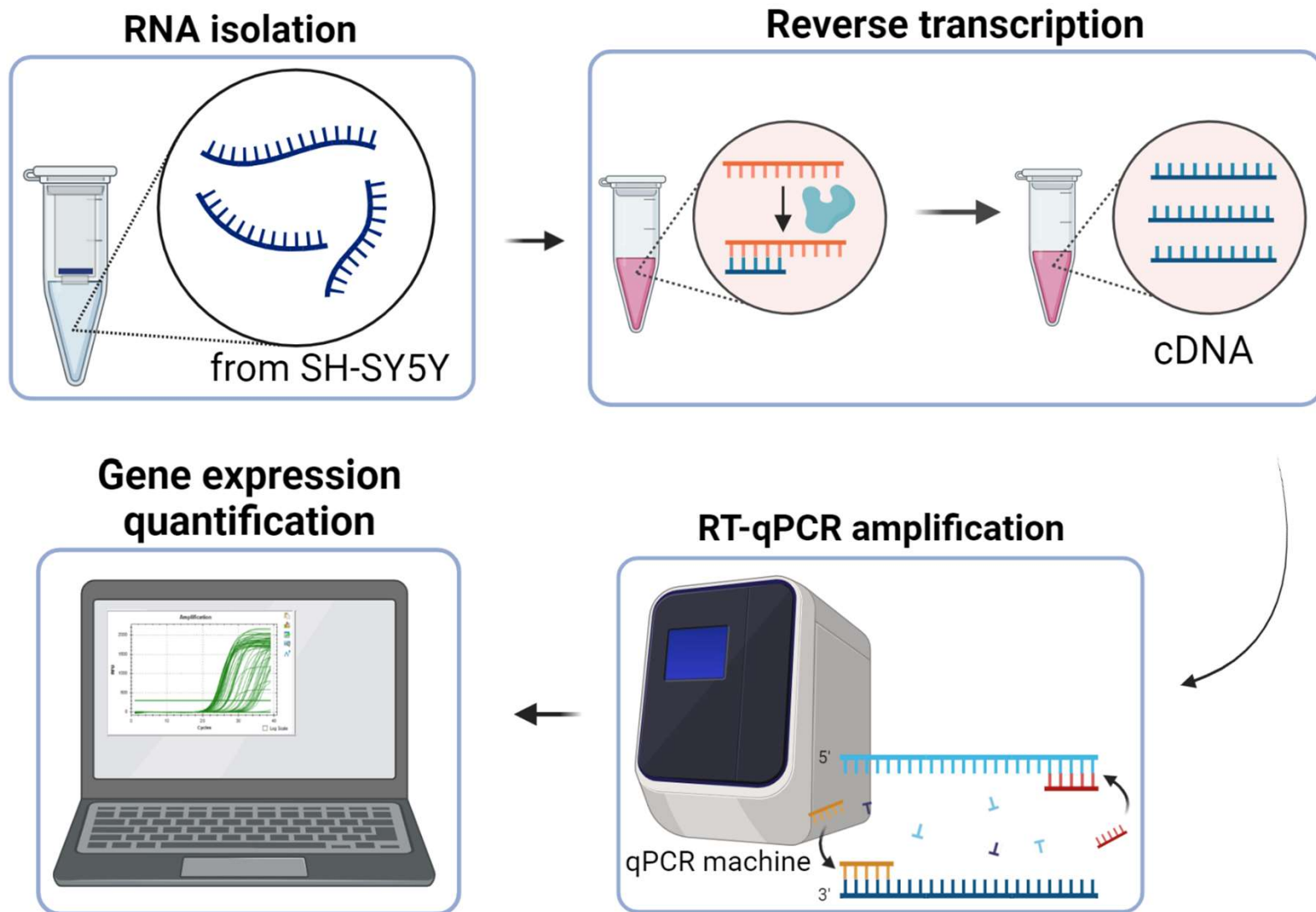
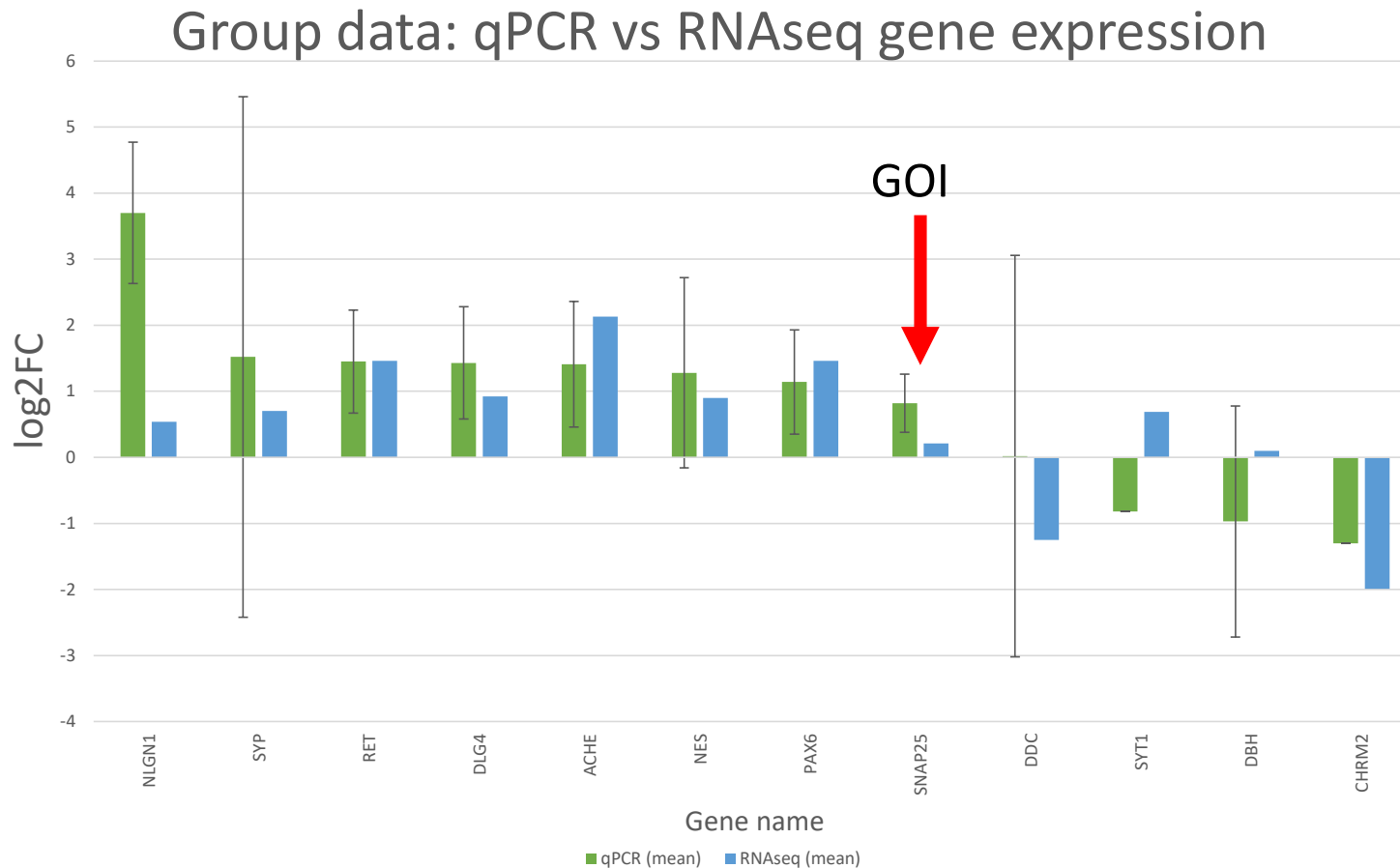


Figure 1: qPCR steps (made with BioRender⁹)

RESULTS

Group Data

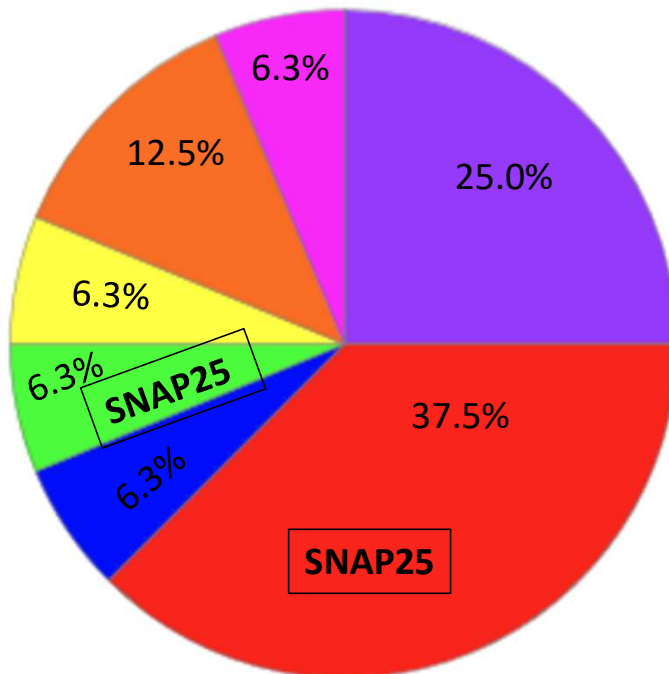


- qPCR data used SD
- DDC, SYT1, DBH had different directionalities (qPCR vs RNAseq)
 - Up-regulated vs down-regulated

Figure 2: Expression level of 12 neuronal markers after RA Tx for qPCR and RNAseq (made with Excel) – data taken from mean of 3 Thursday PM classes

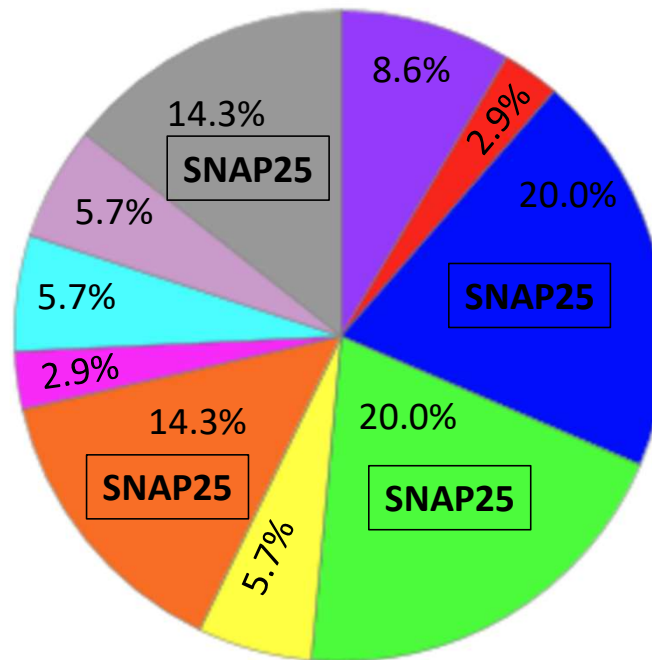
Figure 3: Up-regulated genes (based on qPCR) – Ontological domains (made by PANTHER¹⁰)

Molecular function



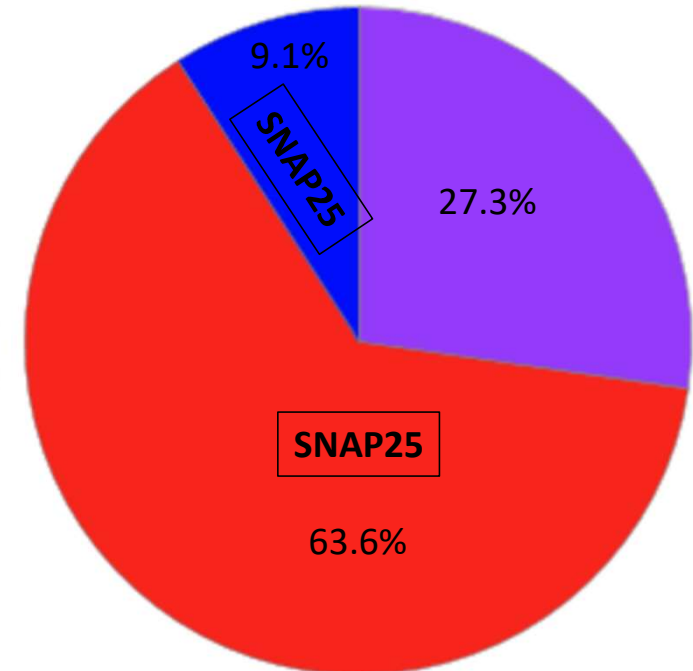
[No PANTHER category is assigned \(UNCLASSIFIED\)](#)
[binding \(GO:0005488\)](#)
[catalytic activity \(GO:0003824\)](#)
[molecular adaptor activity \(GO:0060090\)](#)
[molecular function regulator \(GO:0098772\)](#)
[molecular transducer activity \(GO:0060089\)](#)
[transcription regulator activity \(GO:0140110\)](#)

Biological process



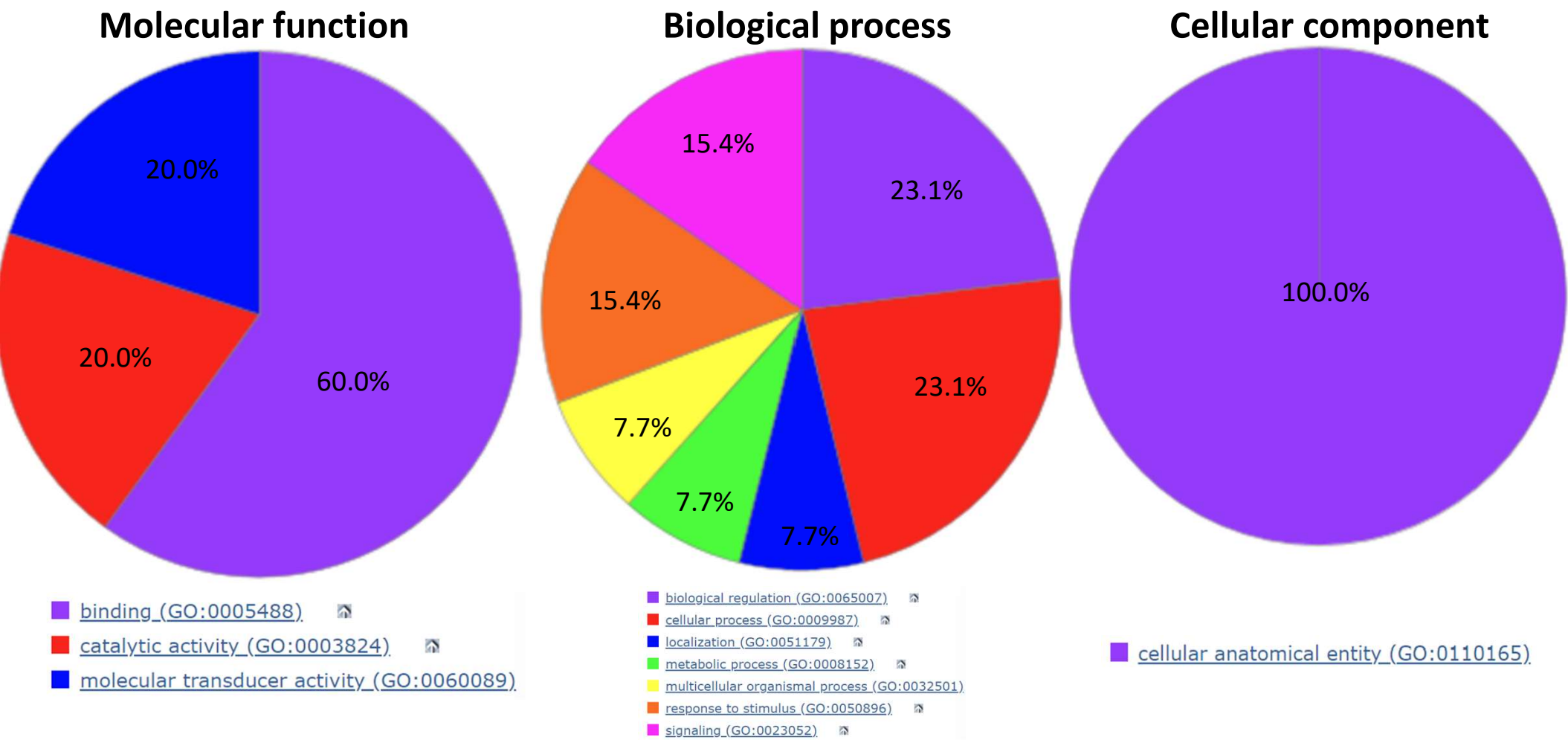
[No PANTHER category is assigned \(UNCLASSIFIED\)](#)
[biological adhesion \(GO:0022610\)](#)
[biological regulation \(GO:0065007\)](#)
[cellular process \(GO:0009987\)](#)
[developmental process \(GO:0032502\)](#)
[localization \(GO:0051179\)](#)
[metabolic process \(GO:0008152\)](#)
[multicellular organismal process \(GO:0032501\)](#)
[response to stimulus \(GO:0050896\)](#)
[signaling \(GO:0023052\)](#)

Cellular component



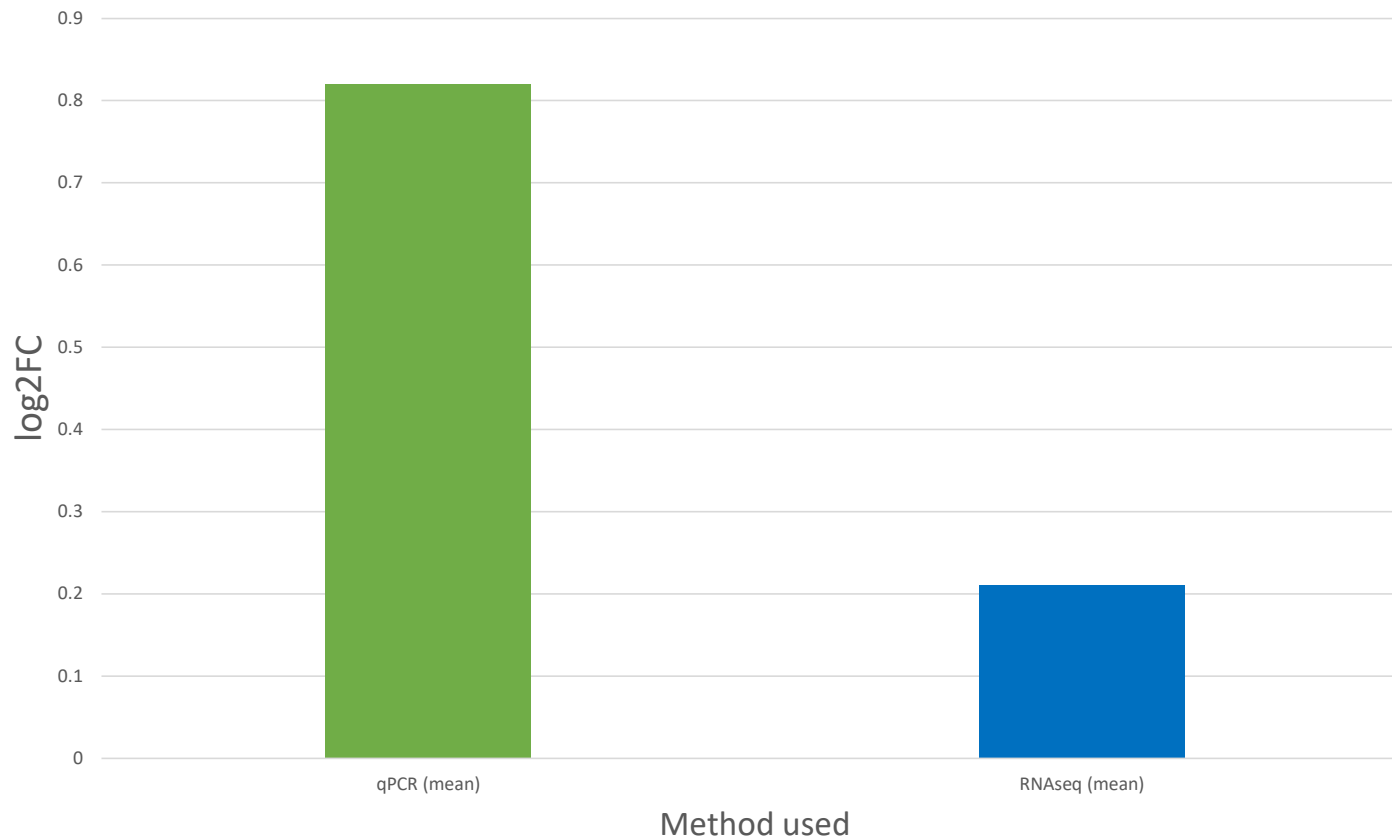
[No PANTHER category is assigned \(UNCLASSIFIED\)](#)
[cellular anatomical entity \(GO:0110165\)](#)
[protein-containing complex \(GO:0032991\)](#)

Figure 4: Down-regulated genes (based on qPCR) – Ontological domains (made by PANTHER¹⁰)



SNAP25 - GOI

SNAP25 expression in qPCR vs RNAseq

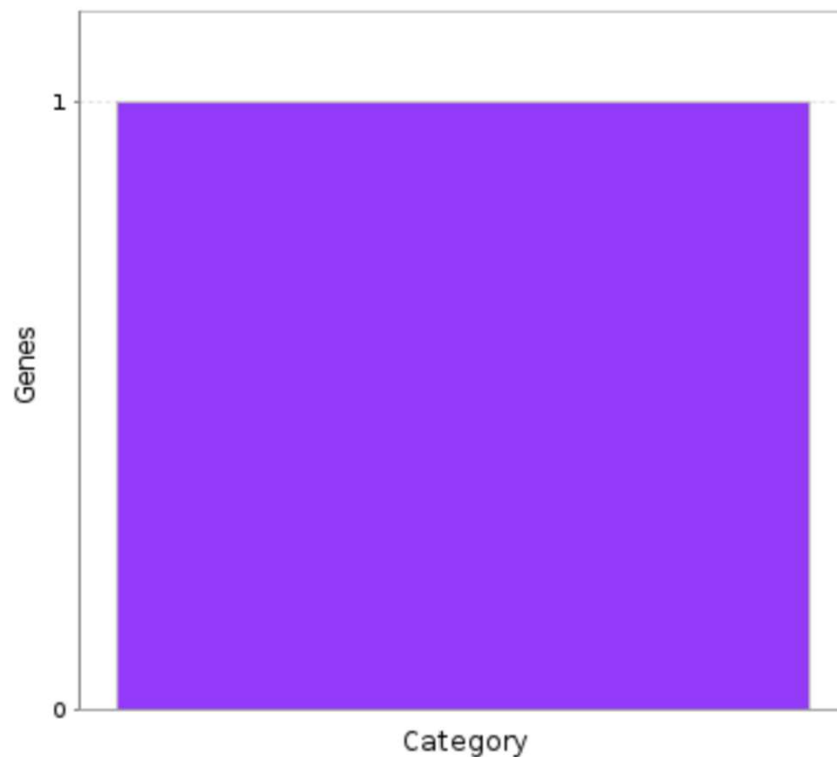


- Same directionality
-> **up-regulated**
- qPCR expression level = 0.82 log2FC
 - increase compared to control

Figure 5: SNAP25 expression level after RA Tx for qPCR and RNAseq (made with Excel) - data taken from mean of 3 Thursday PM classes

Ontological subdomain

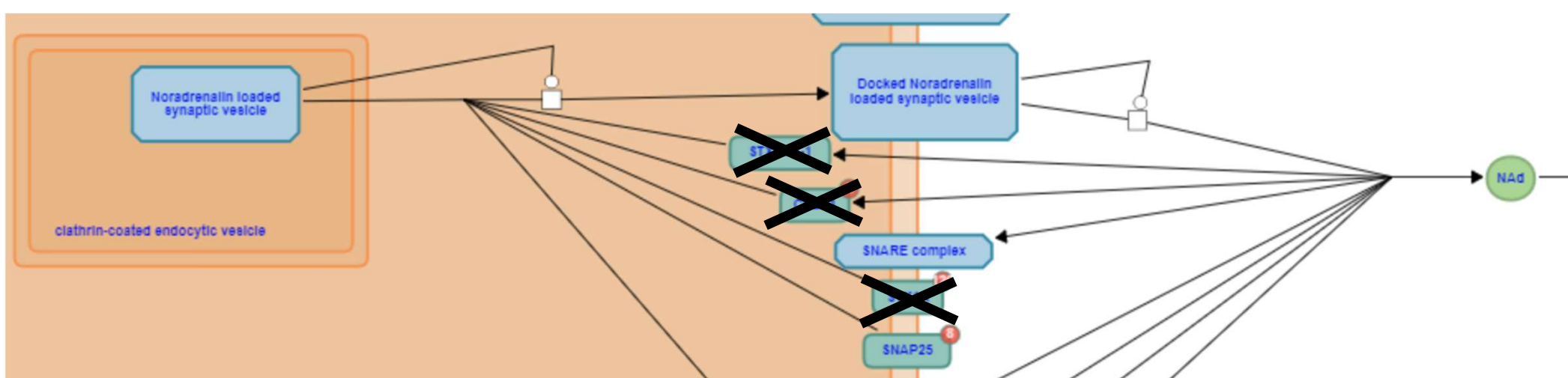
Figure 6: Up-regulated genes - **Protein-containing complex subdomain** of Cellular components (made by PANTHER¹⁰)



- Only SNAP25 (out of up-regulated genes) in protein-containing complex subdomain

■ [membrane protein complex \(GO:0098796\)](#) → SNARE complex

Figure 7: Neurotransmitter release cycle pathway involving SNAP25 (from Reactome¹¹)



- Noradrenaline (NA) vesicle input to SNAP25 (part of SNARE)
- NA gets released and transitioned to extracellular NA
 - Outputs back to SNARE

DISCUSSION - CONCLUSION

- **RA affected gene expression of 12 genes (up/down regulation)**
- **SNAP25 upregulated by RA Tx**
- > **both hypotheses were correct**
- SNAP25 had low variability (qPCR)
- SNAP25 upregulation -> more neurotransmitters released⁸
- BUT it negatively regulates intracellular Ca²⁺, so upregulation should decrease neurotransmitter release
- CHRM2, SYT1, DBH were down-regulated (qPCR) -> involved in cell signalling and synaptic activity
- Different directionality for some genes in qPCR vs RNAseq
 - > qPCR not always homogenous amplification¹²
 - > difference amplification efficiencies¹³

Limitations

- RA treatment not long enough
 - neuroblastoma cells have to be differentiated *in vitro* for at least 7 days for SH-SY5Y cells to be used for experiments¹⁴
- Gene expression change throughout cell differentiation, but was only measured at 72h

Future directions

- Let SH-SY5Y cells differentiate for 1 week
- Samples at different time-points along differentiation process
 - For accurate idea of RA Tx effect on SNAP25

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