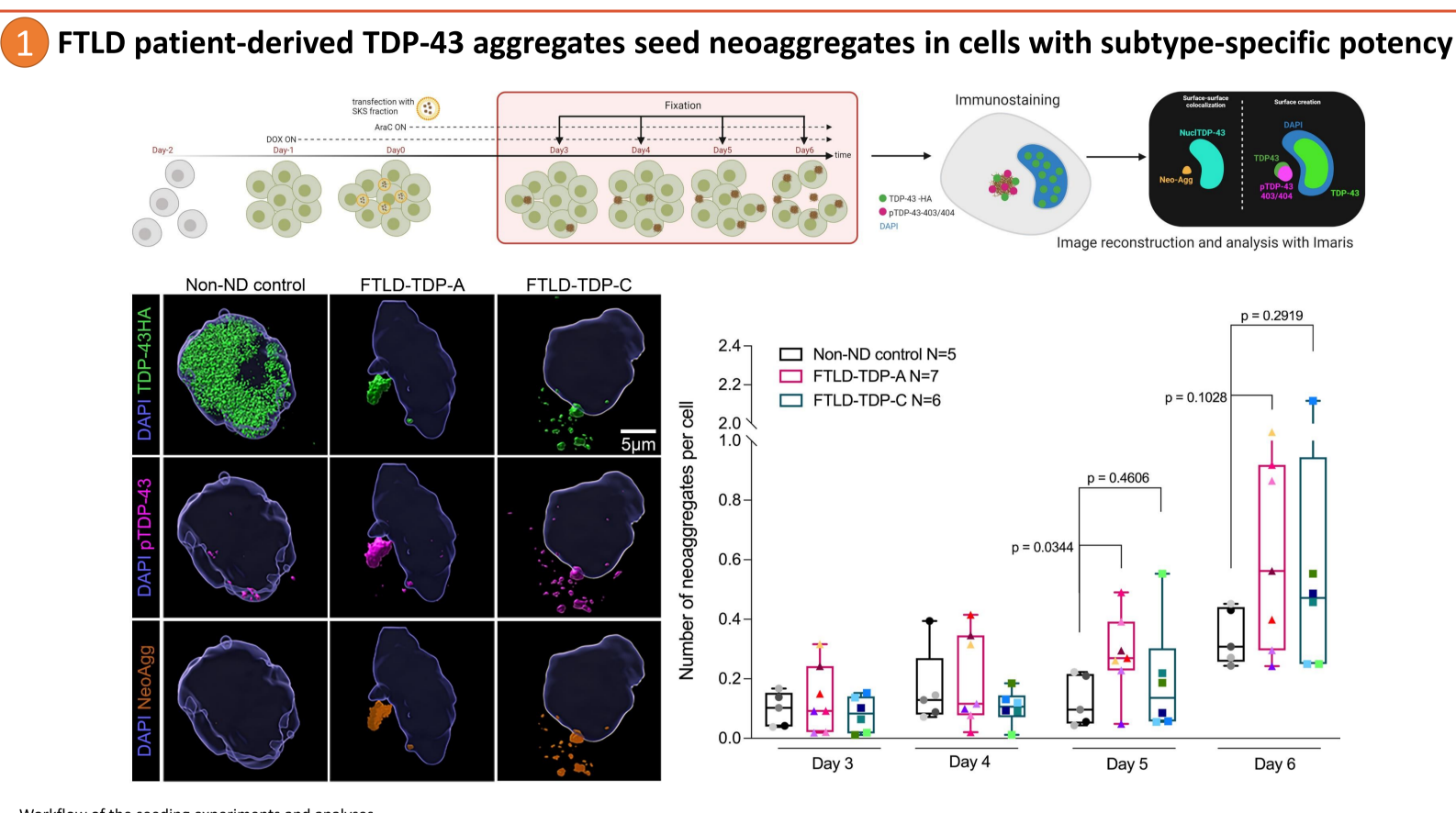


## Template dependent amplification of pathological TDP-43 and roles of phosphorylation

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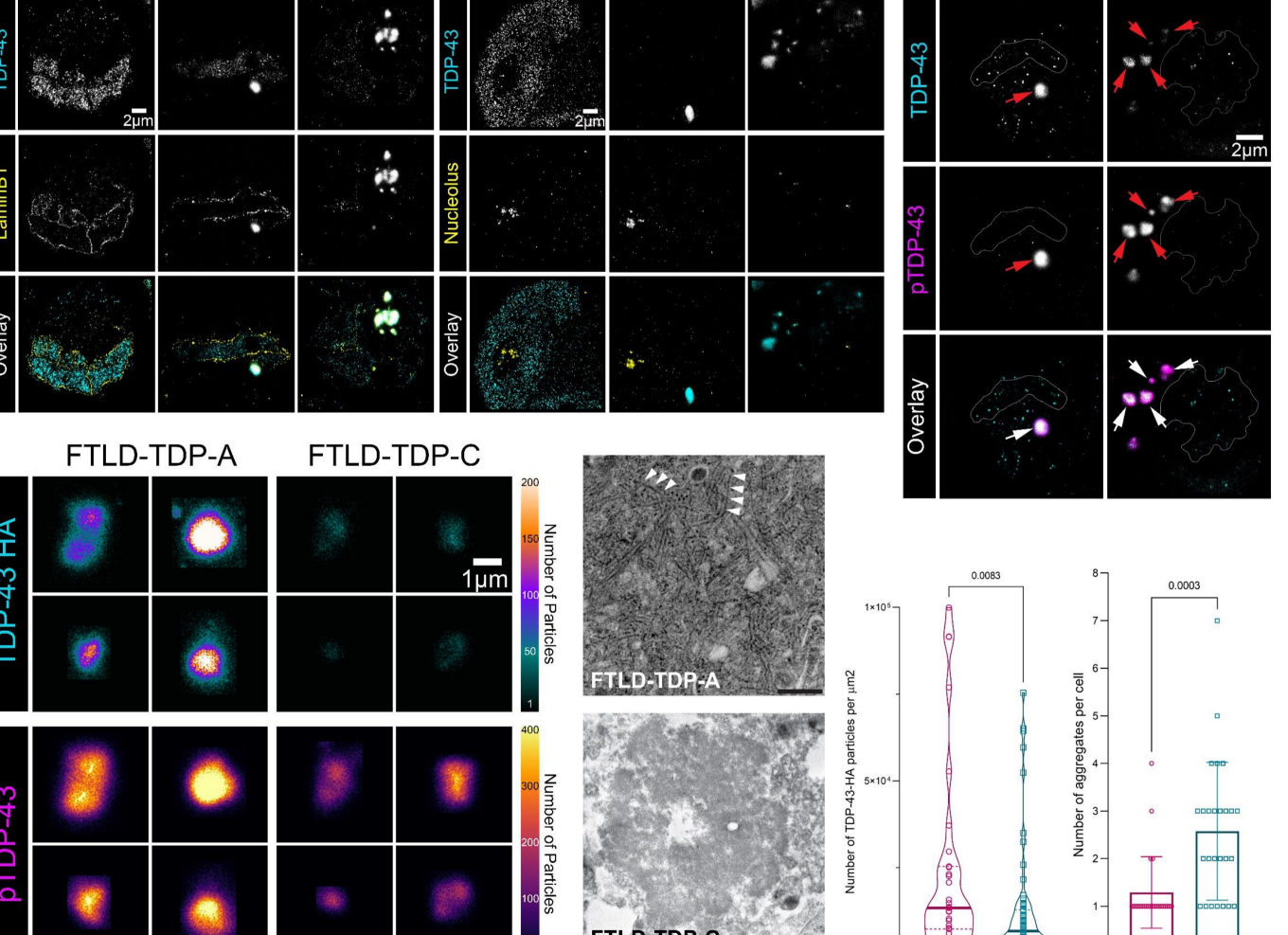
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TDP-43 is an RNA binding protein found aggregated in several neurodegenerative diseases such as frontotemporal lobar degeneration (FTLD), amyotrophic lateral sclerosis (ALS) and Limbic-predominant age-related TDP-43 encephalopathy (LATE). The pathological hallmarks of these diseases are characterized by the presence of hyperphosphorylated TDP-43 within these pathological aggregates. It is assumed that TDP-43 proteinopathies follow the prion-like cascade, but the molecular mechanisms remained unknown. In our study, we demonstrated that isolated pathological seeds from post mortem tissue of patient with FTLD-TDP could trigger de novo aggregation in cells in a template-dependent manner. Our results also suggested that phosphorylation of these neoaggregates was sequential, N to C terminal, with subtype-specific timelines and aggregation profiles. We are currently investigating the role of TDP-43 phosphorylation in neuronal function, as well as disease pathogenesis and progression.



- Workflow of the seeding experiments and analyses. High magnification imaging of cells seeded for 6 days revealed presence of large cytoplasmic TDP-43-HA aggregates in cells seeded with FTLD-TDP-A and multiple cytoplasmic TDP-43-HA aggregates in cells seeded
- with FTLD-TDP-C. Surface representation generated with Imaris showing DAPI, TDP-43-HA (green), phosphorylated TDP-43 (pTDP-43, magenta) and neoaggregates (NeoAgg, orange) Quantification of the number of neoaggregates over time in (n = 3) independent experiments, with multiple biological replicates: non-ND control (n = 5), FTLD-TDP-A (n = 7) and FTLD-TDP-C (n = 6). Whiskers in the box plot represent median and minimum/maximum values. 2-way ANOVA repeated measure, with post-hoc Fisher's LSD, F(3,15) = 1.3, P = 0.3110, NS difference for Day 3 and 4. Day 5: Non-ND control vs FTLD-TDP-A P = 0.0344, non-ND control vs FTLD-TDP-C P = 0.4606. Day 6: non-ND control vs FTLD-TDP-A P = 0.1028, non-ND control vs FTLD-TDP-C P = 0.2919.

## Distinct features of cellular neoaggregates resemble subtype-specific characteristics in FTLD brains



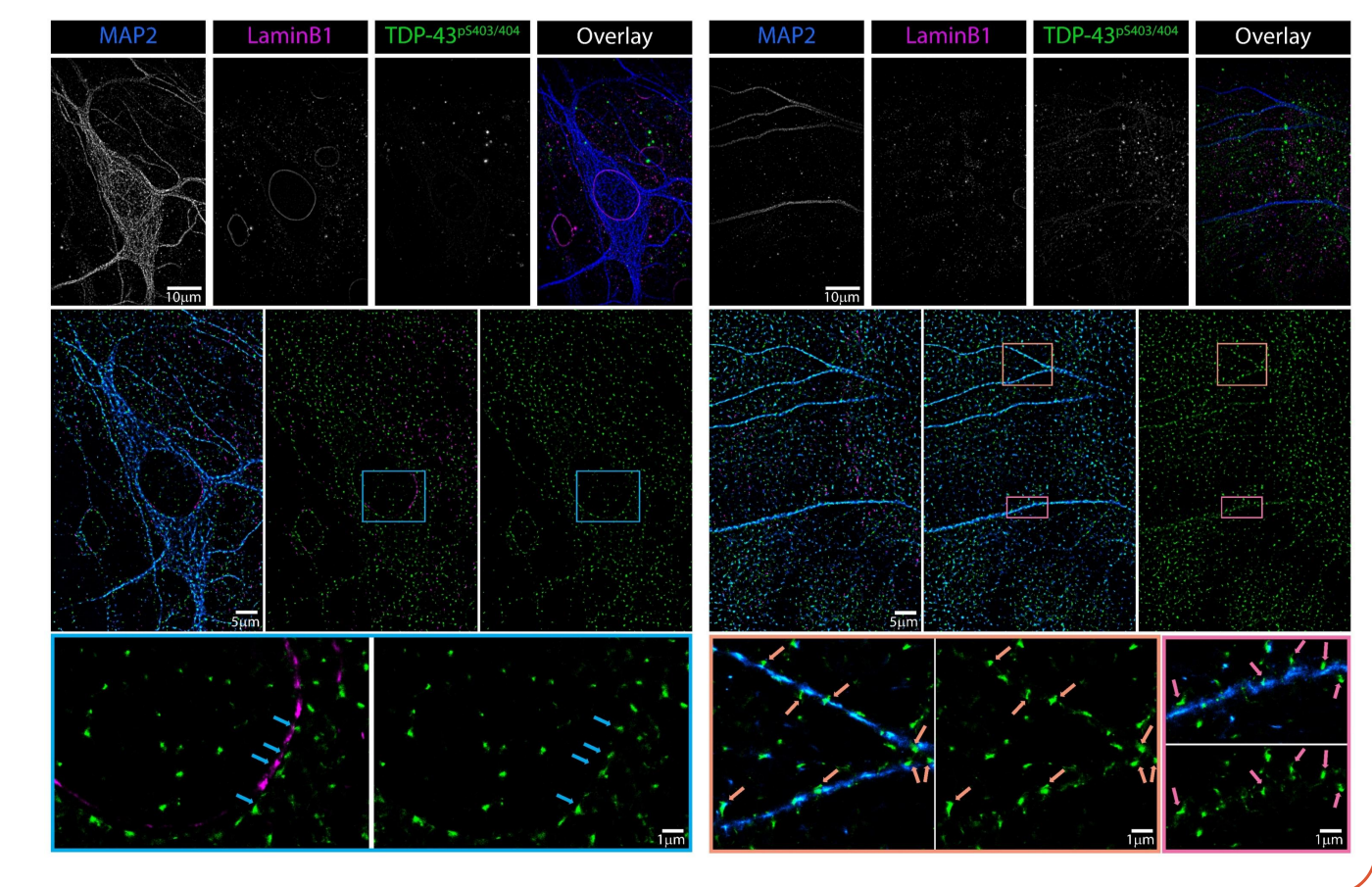
- dSTORM images of cells seeded for 6 days with non-neurodegenerative control (non-ND control), FTLD-TDP-A and FTLD-TDP-C seeds extracted from post-mortem brain tissue. Staining shows localization of TDP-43 (HA tag, cyan) along with LaminB1 and nucleolus (yellow) dSTORM images of cells containing
- aggregates after 6 days of seeding SarkoSpin fractions o postmortem brain from patients with FTLD-TDP-A or FTLD-TDP-C Red arrows point to cytoplasmic aggregates. Cells were labeled with TDP-43 (HA-tag, cyan) phosphorylated TDP-43 pS403/404 (pTDP-43, magenta).
- Heatmap representation of the density of TDP-43-HA and pTDP-43 particles for four representative neoaggregates of FTLD-TDP-A and FTLD-TDP-C seeded cells.
- Violin plots showing the density of particles TDP-43 per µm2. Non parametric unpaired t test FTLD-TDP-A vs FTLD-TDP-C, P = 0.0083(Continuous line represent median dashed lines represent quartiles, n = 3 independent experiments, six biological replicates for each condition).
- Bar graph showing the number of neoaggregates per cell. Nonparametric unpaired t test FTLD-TDP-A vs FTLD-TDP-C, P = 0.0003Error bars represent mean ± SEM, n = 3 independent experiments, six biological replicates for each condition.

## Physiological phosphorylation of TDP-43 in neurons uncovered by superresolution

Primary hippocampal neurons imaged by both confocal (top row) and dSTORM microscopy (middle and bottom rows) and immunostained for MAP2 (dendritic marker, blue), LaminB1 (nuclear membrane marker, magenta) and TDP-43<sup>pS403/404</sup>. Bottom row shows enlargement areas indicated by the boxes on the middle row. Blue arrows point at the TDP-43<sup>pS403/404</sup> structures positive for

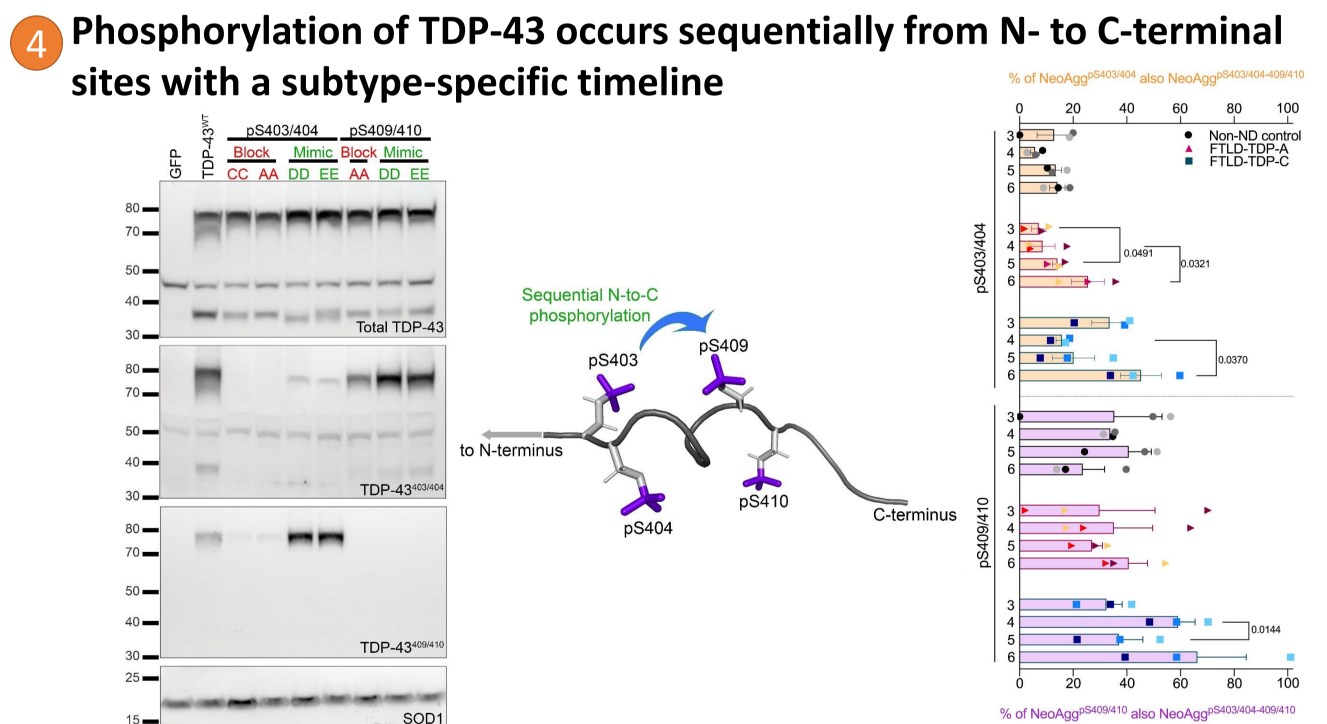
43<sup>pS403/404</sup> along dendrites.

microscopy

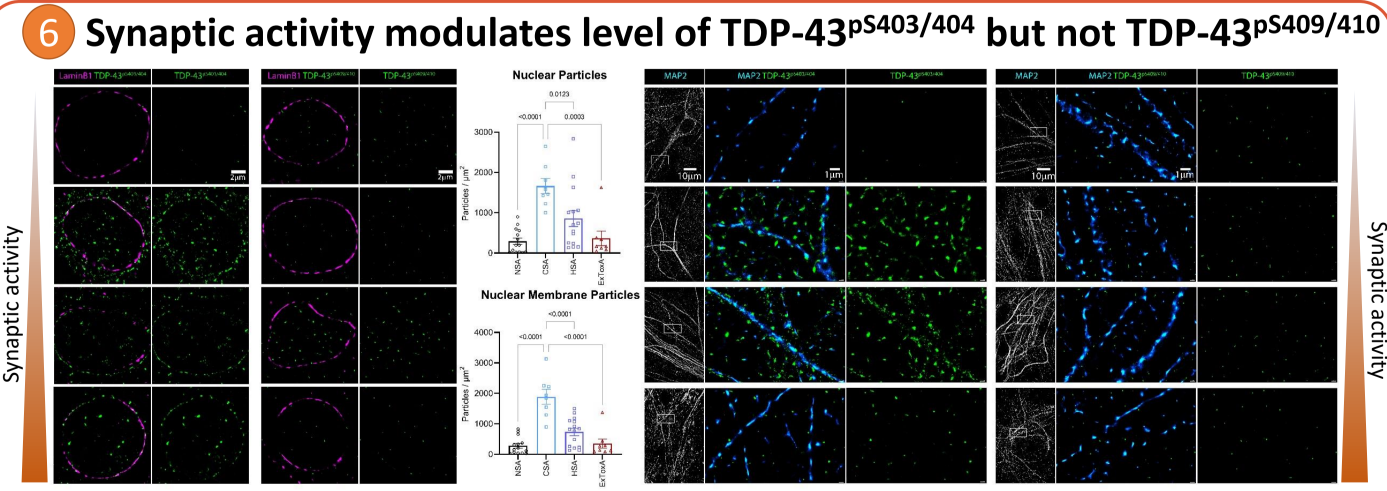


## TDP-43 seeding depends on specific aggregate conformation(s) Non-degenerative control N=3 ▲ FTLD-TDP-A N=3 Normal seeding protocol Denaturation by autoclave Untreated Temperature Autoclave **1.0 —** 20 μM FL-TDP-43-MBP + TEV 20 μM FL-TDP-43-MBP pTDP-43 Neoaggregates p = 0.0385

- Strategy to explore the prion-like properties of TDP-43 seeds. Bar graph showing the seeding effect at day 5 and day 6 post-seeding in untreated and denaturating conditions. One-way ANOVA, with post-hoc Fisher's LSD test, Day 5: F(5,12) = 6.713, P = 0.0033, non-neurodegenerative (non-ND) control vs FTLD-TDP-A P = 0.0016, Day 6: F(5,12) = 4.271, P = 0.0183, non-ND control vs FTLD-TDP-A P = 0.0397
- $(N = 3 \text{ biological replicates for non-ND control, FTLD-TDP-A and FTLD-TDP-C, bar graph showing mean <math>\pm SEM$ ). Aggregation kinetics of full length (fl) recombinant human TDP-43 (rec-hTDP-43) measured as an increase in absorbance. Rec-hTDP-43 aggregates used for seeding were collected 18 h after the addition of TEV protease
- Rec-hTDP-43 aggregates stained by Thioflavin T (ThT). Transmission Electron Microscopy (TEM) image of the rec-hTDP-43 aggregates. Rec-hTDP-43 aggregates were transfected on mitotically arrested cells and fixed after 5 or 6 days post-transfection. Surface masks generated with Imaris depict DAPI (blue)
- Quantification of neoaggregates (left y-axis) and pTDP-43 (right y-axis) at day 5 and day 6 post-transfection for increasing concentrations of rec-hTDP-43. 2-way ANOVA repeated measure, with post-hoc Fisher's LSD, F(7,80) = 5.886, P < 0.0001, to reduce complexity, only comparisons with AraC are marked in the graph. N = 6 biological



- Overexpression of GFP-TDP-43 variants with different mutations that either block (CC and AA mutation, red labels) or mimic phosphorylation (DD and EE, green label) at S403/404 or S409/410 in motor neuron-like NSC-34 cells. Western blot showed a specific signal for total TDP-43 (top panel), TDP-43<sup>pS403/404</sup> (middle panel) and TDP-43<sup>p409/410</sup> (lower panel) and load control SOD1 (bottom panel).
- Schematic of the extreme C-terminal serine phosphorylation found in pathological TDP-43. Part of a whole graph summarizing analysis of double C-terminal phosphorylation of the neoaggregates. N = 3 biological replicates per condition.



Primary hippocampal neurons imaged by dSTORM microscopy and immuno-stained for LaminB1 (nuclear membrane marker, magenta) or MAP2 (dendritic marker, blue) and TDP-43<sup>pS403/404</sup> or TDP-43<sup>pS409/410</sup> (in green). Localization and levels of TDP-43<sup>pS403/404</sup> or TDP-43<sup>pS409/410</sup> were observed under different level of synaptic activity, modulated using pharmacological treatments to obtain no synaptic activity to excitotoxicity levels. Quantification of TDP-43pS403/404 nuclear particles showed synaptic activity dependent changes (Filter precision 30nm, Nuclear particles, one-way ANOVA F=11.25, p<0.0001; Nuclear membrane particles, one-way ANOVA F=23.40, p<0.0001). NSA= no synaptic activity, CSA= control synaptic activity, HSA= high synaptic activity, ExToxA= Excitotoxicity activity



NuclTDP-43 (cyan) and pTDP-43 (magenta).

- 1. Pathological TDP-43 extracted from post mortem tissue of patients with FTLD is able to trigger neoaggregation after transfection
- 2. Neoaggregation is dependent on the seed structure
- 3. Seeds extracted from different subtypes trigger structurally different neoaggregation
- 4. Phosphorylation timeline of the neoaggregates is different between subtypes 5. TDP-43<sup>pS403/404</sup> is present under physiological conditions in different compartments of the neurons
- 6. TDP-43<sup>pS403/404</sup> levels are modulated by synaptic activity

We are now working at understanding what drives these changes and what are the functional roles of TDP-43 phosphorylation







