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## **Multi-influential interactions alters behaviour and cognition through six main biological cascades in Down syndrome mouse models**

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25

26 **Keywords:** Down syndrome, mouse model, partial trisomy 21, Down syndrome critical  
27 region, genotype–phenotype maps, learning, memory, transcriptomics.

28

29 **Short title:** Neuronal altered pathways in Down syndrome mouse models

30

31

## 32 **Abstract**

33 Down syndrome (DS) is the most common genetic form of intellectual disability caused by the  
34 presence of an additional copy of human chromosome 21 (Hsa21). To provide novel insights  
35 into genotype–phenotype correlations, we screened the *in vivo* DS mouse library with  
36 standardized behavioural tests, magnetic resonance imaging (MRI) and digged into  
37 hippocampal gene expression. Altogether this approach brings novel insights into the field.  
38 First, we unravelled genetic interactions between different regions of the chromosome 21 and  
39 how they importantly contribute in altering the outcome of the phenotypes. Then, in depth  
40 analysis of misregulated expressed genes involved in synaptic dysfunction highlighted 6  
41 biological cascades centered around DYRK1A, GSK3 $\beta$ , NPY, SNARE, RHOA and NPAS4.  
42 Finally, we provide a novel vision of the existing altered gene-gene crosstalk and molecular  
43 mechanisms targeting specific hubs in DS models that should become central to advance in our  
44 understanding of DS and therapies development.

45

## 46 **Author Summary**

47 Down syndrome (DS) is the most frequent cause of intellectual disability and is caused by increase in  
48 gene dosage and multiple genetic interaction but not so many have been described so far. Taking  
49 advantage of DS mouse models, we investigated behavior and cognition, brain morphology and  
50 hippocampal gene expression in a controlled environment and we unraveled how multiple genetic  
51 interactions between different regions of the chromosome 21 contribute in altering the outcome of the  
52 behavioural, morphological and molecular/pathways phenotypes. Nonetheless we found  
53 multiple deregulated genes in the hippocampus, where overlapping DS models  
54 show convergence in the biological cascades altered, observed via building protein-protein  
55 interaction and regulatory networks, and centered in 6 main hubs: DYRK1A, GSK3 $\beta$ , NPY,  
56 SNARE, RHOA and NPAS4. Although four of them were already described to be altered in some  
57 DS models, we validated two additional ones, RHOA and NPAS4, and we have built a novel vision of

58 the existing altered gene-gene crosstalk and molecular mechanisms, targeting 6 specific highly  
59 interconnected hubs in DS models, that should become central to advance in our understanding of DS  
60 physiopathology and therapy development.

61

## 62 **Introduction**

63 Down syndrome (DS) is the most common genetic form of intellectual disability and  
64 was first described as a disease by John Langdon Down in 1866. One century later, genetic  
65 studies demonstrated that DS is caused by trisomy of human chromosome 21 (Hsa21) (1).  
66 People with DS have a wide range of phenotypic and physiological features with some  
67 phenotypic variability but always present several disabling features like mental retardation or  
68 Alzheimer disease (2). The leading cause of DS is non-disjunction of chromosome 21 (3).  
69 However, in rare cases, people with partial Hsa21 duplications are observed with a smaller  
70 spectrum of DS features. Studying these rare conditions increased our understanding of the  
71 genotype–phenotype correlations in DS (4-10). In particular, this method led to the hypothesis  
72 of the existence of a DS critical region (DCR) in which a small set of Hsa21 genes (10) could  
73 play a major role in DS phenotypes, including cognitive disabilities. The understanding of DS  
74 has been substantially improved over the years, with the progress of genome analysis  
75 technologies. Recently, two studies showed that not a single trisomic region can be responsible  
76 for all DS features but rather several susceptibility regions when presented in 3 copies can  
77 induce a large variety of features (5, 7). In addition, recent publications showed that several  
78 patients displayed complex rearrangement like contiguous or non-contiguous deletions or  
79 duplications, copy number variations of other regions or duplication of genes located in the  
80 short arm of Hsa21, that can potentially impact the phenotypic outcome of the Hsa21  
81 duplication and add noise to the genetic dissection of human trisomy 21 clinical manifestations.

82 To circumvent the difficulties in studying DS in human, several efforts have been made  
83 to generate DS mouse models (11). Indeed, there are three independent mouse genomic regions

84 carrying altogether 158 protein-coding homologous genes of the 218 protein-coding genes  
85 located on the Hsa21, including two keratin clusters of respectively 37 and 13 genes (12). The  
86 largest region homologous to Hsa21 is found on the mouse chromosome 16 (*Mus musculus* 16,  
87 noted as Mmu16) in a fragment of 22.42 Mb with 119 orthologous genes between *Lipi* and  
88 *Zbtb21* (13). The most telomeric part is splitted into two parts. The first part, on the mouse  
89 chromosome 17 (noted as Mmu17) with 19 homologous genes in 1.1 Mb interval between  
90 *Umodl1* and *Hsf2bp*. Then, the second part, on the the mouse chromosome 10 (Mmu10) with  
91 37 genes included in the 2.3 Mb *Cstb-Prmt2* genetic interval (11, 12, 14). Several DS mouse  
92 models have been generated over the years, most of them were carrying trisomy of the largest  
93 genetic region located on Mmu16, and have built the *in vivo* DS mouse library (11, 14). The  
94 Ts(17<sup>16</sup>)65Dn (noted Ts65Dn) is the most widely used DS model and it is quite unique with a  
95 supplementary mini-chromosome obtained by x-ray irradiation of the male germline and  
96 containing the centromeric region of Mmu17, with genes from *Psid-ps2* to *Pde10a*, and the  
97 13.5 Mb telomeric fragment of Mmu16 containing genes between *Mrpl39* and *Zbtb21* (15-18).  
98 Several models were made by chromosomal engineering (11) and carry segmental duplication  
99 of Mmu16. The Dp(16*Lipi-Zbtb21*)1Yey (noted Dp1Yey) corresponds to the duplication of the  
100 entire Mmu16 region syntenic to Hsa21 (19). The Dp(16*Cbr1- Fam3b*)1Rhr (noted Dp1Rhr)  
101 model carries a duplication from *Cbr1* to *Fam3b* and demonstrate the contribution of the DS  
102 critical region (DCR) (20-23). All the DS mouse models displayed defects in behaviour and  
103 cognition which had been investigated in different laboratories with different protocols and  
104 environmental condition making difficult inter-model comparison (11).

105 In order to improve our knowledge on DS, we analysed a series of old and new  
106 combinations of mouse models with a unique and in depth behaviour, morphological and  
107 transcriptomics pipeline in adults to dissect the contribution of the genes located on Mmu16 to  
108 DS mouse features. The behaviour pipeline relied on assessing specific brain functions found  
109 altered in people with DS and in particular those depending on the hippocampus (24, 25),. Thus,

110 we performed hippocampal-based learning tests together with other standardized protocols  
111 including the Y-maze, Open field (OF), Novel Object Recognition (NOR), Morris Water Maze  
112 (MWM) and contextual and cue Fear Conditioning (FC). All procedures were carried in similar  
113 environmental conditions to reduce any impact (26, 27). Besides, variations in specific brain  
114 regions have been observed in people with DS and mouse models (28-31). Neuroanatomical  
115 changes affect the whole brain volume or specific regions like the frontal region of the limbic  
116 lobe or the hippocampus in people with DS. Thus, we performed an in depth morphological  
117 investigation of the brain by magnetic resonance imaging (MRI). Finally, whole gene  
118 expression was performed on hippocampi isolated from all the models to decipher the genes,  
119 functional pathways and biological cascades affected in the different DS mouse models.  
120 Moreover, we identified the gene expression and functional profiles conserved in post-mortem  
121 patients brain derived datasets publicly available in the GEO database.

122

## 123 **Results**

124

### 125 **Dissecting the contribution of Mmu16 subregions to the DS-related behavioral** 126 **phenotypes in mouse models**

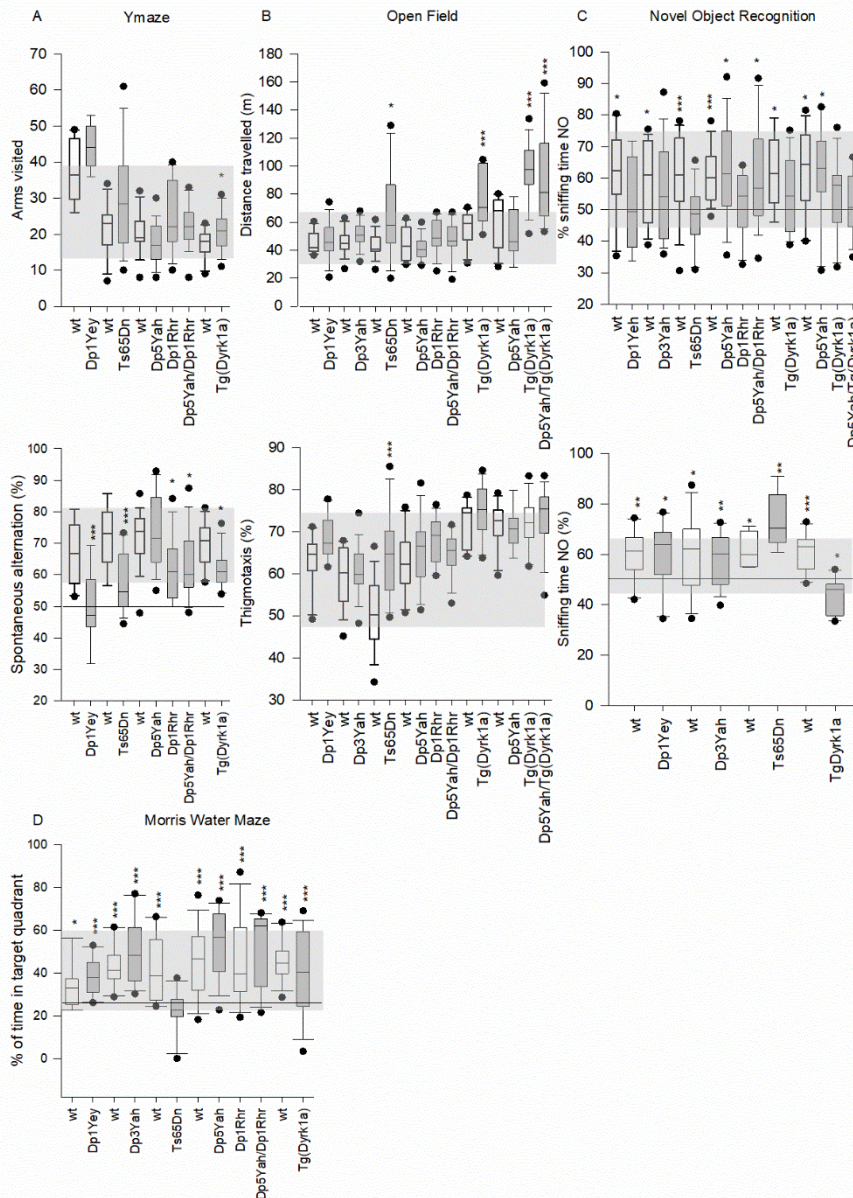
127 We wanted to dissect the contribution of sub-regions located in the telomeric part of  
128 *Mus musculus* chromosome 16 (Mmu16), homologous to human chromosome 21 (Hsa21); (32),  
129 to the DS-related cognitive phenotypes in the mouse. First, we selected 4 DS mouse models:  
130 the Ts65Dn, the most commonly used DS model (15), and three additional models that are  
131 carrying segmental duplications of well-defined sub-regions located on the Mmu16, the  
132 Dp1Yey (19), Dp3Yah (33) and Dp1Rhr (20). In addition, we engineered a new one, the  
133 Dp5Yah, corresponding to three functional copies of the genes included in the genetic interval  
134 between *Cyyr1* to *Clic6*. This model was crossed with the Dp1Rhr one in order to generate the  
135 Dp5Yah/Dp1Rhr (noted Dp5/Dp1) compound transheterozygote carrying a trisomic gene

136 content similar to the Ts65Dn for the genes located on the Mmu16. We also included a model  
137 carrying an additional copy of *Dyrk1a*, one of the driver genes for DS-related phenotypes (30,  
138 34), and the Tg(*Dyrk1a*) combined with Dp5Yah model (noted Dp5-Tg) (Supplementary fig  
139 1). We used standardized behavioral tests to study several aspects of learning and memory in  
140 mice including the Y-maze (working memory), the open field (exploration memory), the novel  
141 object recognition (recognition memory), the Morris water maze (spatial memory) and the fear  
142 conditioning (associative memory). For all the lines, independent cohorts of control and  
143 trisomic mouse littermates went through the pipeline at similar ages, then the resulting data  
144 were processed using standard statistical analyses (See supplementary material and methods for  
145 details). First, we assessed the potential existence of a background effect over the distribution of  
146 the measurements taken in the different tests. No difference between B6J or hybrid B6JC3B  
147 wild-type controls was observed over the Q-Q plots with the cumulative frequency. Then, we  
148 defined a unique wild-type reference range corresponding to 90% of all controls (see  
149 Supplementary Table 1, Supplementary fig 2).

150 Mice activity and working memory were evaluated in the Y-maze (Fig 1A). The number  
151 of arm entries in the Y-maze showed that only Tg(*Dyrk1a*) mutant line was hyperactive in this  
152 test while deficit in spontaneous alternation was found in Dp1Yey (35), Ts65Dn (36), Dp1Rhr  
153 and Tg(*Dyrk1a*) (37). The Dp5/Dp1 also showed a clear deficit in the percentage of spontaneous  
154 alternation as compared to littermate controls while Dp5Yah trisomic animals showed normal  
155 performance.

156 The patterns of exploratory activity and anxiety were assessed in the open field (Fig  
157 1B). Ts65Dn, Tg(*Dyrk1a*) and Dp5-Tg presented hyperactivity with an increased travelled  
158 distance compared to wild-type littermates, and support, for Tg(*Dyrk1A*), results obtained in  
159 the Ymaze. In addition, Ts65Dn mice displayed thigmotaxic behavior with increased distance  
160 travelled at the periphery.





### Figure 1. Standardized behavioural profiling of DS mouse models.

Y-maze spontaneous alternation. Arm visited (A upper panel) and alternation rate (A lower panel) are presented as box plots with the median and quartiles (upper and lower 90% confidence interval are indicated by a grey box). Only the *Tg(Dyrk1a)* mice were showed hyperactivity in this test with increased arms entries compared to the wild type ( $p=0,017$ ). Alternation rate in *Dp1Yey* ( $p=0,002$ ), *Ts65Dn* ( $p<0,001$ ), *Dp1Rhr* ( $P=0,012$ ), *Dp5/Dp1Rhr* ( $p=0,018$ ) and *Tg(Dyrk1a)* ( $P=0,010$ ) mice was significantly lower than respective wild-type mice. Exploratory activity in a novel environment. Distance travelled (B upper panel) and % of distance

194 travelled in peripheral zone recorded in the Open field arena (B lower panel). The total distance  
 195 travelled was significantly higher in *Ts65Dn* ( $p=0,022$ ), *Tg(Dyrk1a)* ( $p=0,008$ ) and  
 196 *Dp5Yah/Tg(Dyrk1a)* ( $p>0,001$ ). Moreover, the % of distance in the peripheral zone was  
 197 increased in *Ts65Dn* ( $p>0,001$ ) mice compared to wild type mice. Novel Object Recognition  
 198 for 24 hour time lapse (C upper panel) and 1 hour time lapse (C lower panel). The results are  
 199 presented as % of sniffing time (as box plots with the median and quartiles) for the novel object  
 200 (NO). For 24 hours time laps, one sample t test vs 50% (hazard) showed that *Dp1Yey* ( $p=0,837$ ),  
 201 *Dp3Yah* ( $P=0,173$ ), *Ts65Dn* ( $p=0,432$ ), *Dp1Rhr* ( $p=0,492$ ), *Tg(Dyrk1a)* ( $p=0,144$ ) and  
 202 *Dp5Yah/Tg(Dyrk1a)* ( $P=0,488$ ) failed to recognize the new object. The *Dp5Yah* genomic  
 203 fragment restored the capacity of the *Dp1Rhr* in the *Dp5Yah/Dp1Rhr* mice ( $p=0,0157$ ). For 1  
 204 hour time laps, all the mice were able to discriminate the NO except for the *Tg(Dyrk1a)*  
 205 ( $p=0,011$  preference for FO). Probe test session in Morris Water Maze (D). The results are  
 206 presented as % of time in the target quadrant. All the mice have spent more time in the target  
 207 quadrant versus non target excepted for the *Ts65Dn* mice ( $p=0,398$ ) (\*  $p<0,05$ , \*\* $p<0,01$ ,  
 208 \*\*\* $p<0,001$ ).



210

211           The spatial reference memory was tested in the standard Morris water maze (MWM)  
212 task, in which mice have to escape from a circular pool of opaque water by localising a hidden  
213 platform at a single fixed location using distal spatial cues. We analysed the velocity, the  
214 distance travelled by the mice to reach the platform and the thigmotaxis over training  
215 (Supplementary fig 3). The velocity of Dp1Yey and Dp5Yah, was slightly lower than the wild-  
216 type mice (Supplementary fig 3B). As previously described (22, 38-44), Ts65Dn mice  
217 displayed a longer distance travelled to find the platform during all the sessions, compared to  
218 the wild type (Supplementary fig 3A). Although Tg(*Dyrk1a*) were able to locate the platform,  
219 they also showed delayed acquisition compared to the control mice. Surprisingly, the Dp1Yey,  
220 Dp1Rhr, Dp5Yah and Dp3Yah completed this test without any difference with the wild type  
221 group. Retention of place location was evaluated during a single probe trial (PT) with no  
222 platform available, 24h after the last training session (Fig 1D). The results confirmed as  
223 previously shown, that all the mouse strains except the Ts65Dn, remembered where the  
224 platform was located after the learning sessions. Interestingly, the Ts65Dn model, and to a  
225 lower extend the Tg(*Dyrk1a*) one, presented marked thigmotaxic behaviour, spending higher  
226 percentage of time in the peripheral zone as compared to controls (Supplementary fig 3C).  
227 Finally, to check the visual ability of mice, we performed a visual training version of the MWM  
228 during which the platform position is indicated by a flag. All mice were able to find the visible  
229 platform without any significant difference with controls except for the Tg(*Dyrk1a*) that  
230 presented a small delay in session 2 (Supplementary fig 3A).

231           We then evaluated non-spatial recognition memory using the novel object recognition  
232 (NOR) paradigm with the retention time of 24h. The percentage of sniffing time for the novel  
233 object was analysed and compared to 50% (hazard). This analysis showed that Dp1Yey,  
234 Dp3Yah, Ts65Dn, Dp1Rhr and Tg(*Dyrk1a*) were not able to discriminate the familiar and the  
235 novel objects unlike the Dp5Yah and more surprisingly the Dp5/Dp1 (Fig 1C). to further

236 characterize the effect/lack of effect of Dp5Yah mutation on novel object recognition, the  
237 Dp5Yah mouse line was crossed with the Tg(*Dyrk1a*) and compared to new sets of wild-type,  
238 Dp5Yah and Tg(*Dyrk1a*) mice. Interestingly, we found that the Dp5Yah/Tg(*Dyrk1a*) and as  
239 expected the Tg(*Dyrk1a*) displayed altered novel object discrimination while the Dp5Yah spent  
240 more time exploring the novel object than the familiar one. We also assessed Dp1Yey, Dp3Yah,  
241 Ts65Dn and Tg(*Dyrk1a*) short term memory performing the NOR with 1 hour delay between  
242 acquisition and retention and only the Tg(*Dyrk1a*) mice showed a deficit.

243 All the trisomic lines were also tested for associative memory in the Pavlovian fear  
244 conditioning test. All the groups showed higher percentage of freezing during the 2 min post-  
245 shock compared to the habituation session, indicating that all groups developed fear responses  
246 during the training session (Supplementary fig 4). When animals were re-exposed 24 h later to  
247 the same context, the level of freezing in all groups was increased compared to the habituation  
248 (PRE2 and PRE4). However, freezing time for Ts65Dn mice was lower compared to the  
249 respective control littermates. When we assessed cued fear conditioning in a new context, all  
250 the mice presented an increased time of immobility with a strong difference between pre-cue  
251 and cue periods (Supplementary fig 4). In addition, Dp1Yey and slightly Ts65Dn showed lower  
252 freezing during presentation of the cues as compared to wildtype counterparts. These data  
253 suggest altered emotional associative memory Ts65Dn.

254

## 255 **Dissecting the contribution of Mmu16 sub-regions to the DS-related brain morphological** 256 **phenotypes in mouse models**

257 DS models have been reported to show brain morphological alterations of specific  
258 regions (30). Thus, we wondered if we could detect changes in the brain morphology using  
259 MRI on these different partial trisomic mice models. Data were first analyzed using a volume  
260 approach and a brain region atlas. We confirmed that the brain of Tg(*Dyrk1a*) mice was larger  
261 ( $p < 0,001$ ) (30) and the brain of Dp1Yey mice was smaller than the respective wildtypes. Then,

262 we analyzed different brain regions/structures taking into consideration/correction the whole  
263 brain volume . Even with this correction, the Tg(*Dyrk1a*) mice were the most affected in terms  
264 of the brain structures volume, and on the contrary, the Dp1Rhr mice did not show any  
265 significant variation compared to the wt mice. Several regions, such as the basal forebrain  
266 septum, central gray matter, the rest of midbrain, and superior colliculi were significantly larger  
267 in Tg(*Dyrk1a*), Ts65Dn and Dp1Yey DS models. Moreover, the cerebellum, hypothalamus,  
268 inferior colliculi and caudate putamen were significantly different in Dp1Yey and Tg(*Dyrk1a*)  
269 compare to the control group (Supplementary Table 2, Supplementary fig 5) and a few  
270 additional areas were altered specifically in certain models (Amygdala, Globus pallidus,  
271 Hippocampus, Neo Cortex, and Thalamus for Tg(*Dyrk1a*); External capsule, Fimbria, and  
272 Ventricles for Dp1Yey). Altogether this brain morphometric analysis showed a greater  
273 similarity between Dp1Yey and *Dyrk1a* overexpression transgenic models with intermediate  
274 overlapp with the Ts65Dn mouse model.

275

## 276 **Dissecting the contribution of Mmu16 subregions to the DS-related transcriptome** 277 **phenotypes in mouse models**

278

279 Various studies have shown the consequences of trisomy on gene expression (45-54).  
280 Here we took the opportunity to dissect the alteration of gene expression and functional  
281 pathways in various DS trisomic models carrying different duplications of the Mmu16. We  
282 analyzed the Ts65Dn, Dp1Yey, Dp3Yah, Dp5/Dp1, Dp5Yah, Dp1Rhr, and we included the  
283 trisomic model for *Dyrk1a* alone, Tg(*Dyrk1a*). Considering the hippocampal formation as a hub  
284 structure involved in learning and memory, we performed gene expression analysis in the adult  
285 hippocampus comparing the DS models with their own littermate controls using a unique  
286 pipeline for all the models. For each DS model, we defined the expressed genes (noted as EGs)  
287 as the genes whose expression level was detected, the differentially expressed genes (noted as

288 DEGs) as the genes whose expression level was found to be significantly altered in the trisomic  
 289 model compared to the controls littermates, and then the trisomic expressed genes (TEGs) as  
 290 the DEGs that are included inside the duplicated chromosomal regions for each model  
 291 (Supplementary table 3, Table 1).  
 292

Mouse lines	Dp1yey	Dp3Yah	Ts65Dn (Mmu16)	Dp5/Dp1	Dp5Yah	Dp1Rhr	TgDyrk1A	Ts65Dn (Mmu17)
Nb of probes detected by the array					35556			
Nb of annotated probes detected (without control probes)					27359			
Number of trisomic genes detected by the array	155	21	130	127	87	40	1	40
Differential expressed genes (DEGs)	711	826	1074	922	736	1306	850	
Differential expressed trisomic genes (DETGs)	66	13	64	54	39	18	1	28
% of **DETGs	43%	62%	49%	43%	45%	45%	100%	70%
% of compensated trisomic genes detected	57%	38%	51%	57%	55%	55%	0%	
Number of GAGE KEGG and GOs (CC,BP,MF) terms disregulated in the trisomic model FDR<0.1	244	67	12	111	318	225	231	
Number of GAGE KEGG and GOs (CC,BP,MF) terms <b>upregulated</b> in the trisomic model FDR<0.1	207	60	3	33	4	132	222	
Number of GAGE KEGG and GOs (CC,BP,MF) terms <b>downregulated</b> in the trisomic model FDR<0.1	37	7	9	78	314	93	9	
Number of GAGE KEGG and GOs (CC,BP,MF) terms in the trisomic model unique to each mice line FDR<0.1	80	1	2	30	195	107	64	

\*DEGs/DETs: Diferential expressed genes/ Diferential expressed

\*\*DETGs/DETTs: Diferential expressed trisomic genes/

293

294 **Table 1.** Differential expression analysis results of the seven models analysed. DEGs are  
 295 Differential Expressed Genes and TEGs for Differential Expressed trisomic genes. \* Analysis  
 296 done with Mmu17 and 16 trisomic genes. GO are Go functional terms involved in cellular  
 297 compartment (CC), molecular function (MF) and biological processes (BP).  
 298

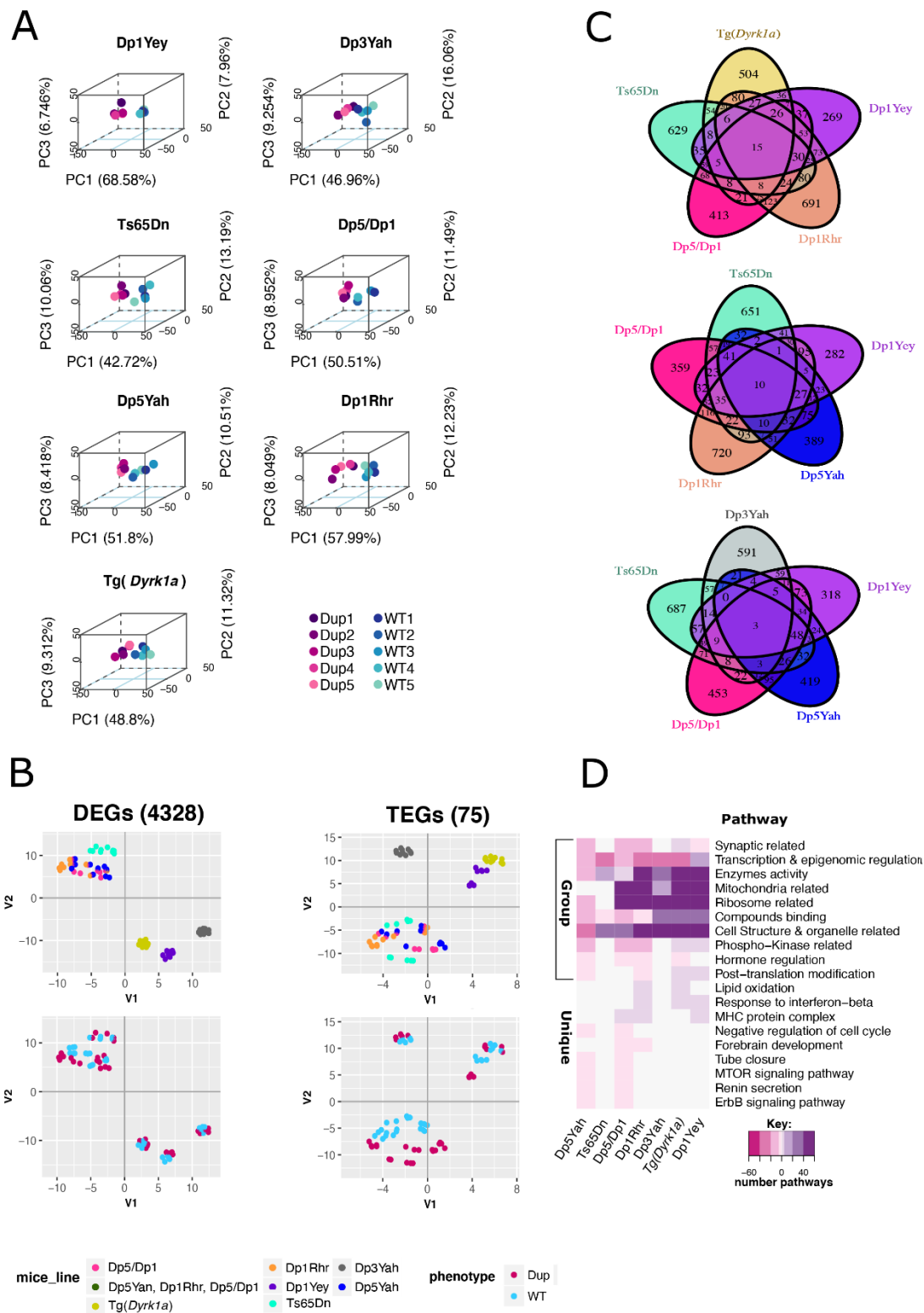
299

300 Although most of the genes in 3 copies were overexpressed in the relevant mouse model-  
 301 derived hippocampi with a ratio around 1.5 (Supplementary fig 6), from 38% to 57% of the  
 302 trisomic genes showed a dosage compensation (Table 1, Supplementary table 3). While this  
 303 compensation is expected, we noticed that most of the compensated genes behave similarly in  
 304 the different trisomic context, even if the experiments were done independently. As such the

305 genes from *Cldn17* to *Krtap11-1*, including the keratin cluster were not overexpressed when  
306 trisomic in any model (Dp1Yey, Ts65Dn, Dp5/Dp1 or Dp5Yah). This could be due to the fact  
307 that this region seems to be under strong regulatory constraints as on the borders two REST sites  
308 plus a LaminB1 peak encompassing this region were found (UCSC browser), while  
309 *Btg3* and *D16Ertd472e*, or *Mrpl39*, *Jam2*, *Atp5*, *Gabpa* and *App* are overexpressed in various  
310 DS models. We also found that the genes located on the trisomic segment not homologous to  
311 Hsa21 on the Mmu17 in Ts65Dn hippocampi, were overexpressed. (Supplementary fig 7) This  
312 fact is due to the trisomy of this region in the mouse line as previously reported in the mouse  
313 heart (16). Looking in detail into the homologous Hsa21 region in Mmu17, we saw two main  
314 genetic effects due to the overdosage of the Mmu16 region homologous to Hsa21. Noticeably,  
315 a constant deregulation of *Cbs* was found in all the models except Dp3Yah and Tg(*Dyrk1a*);  
316 suggesting a direct control of the *Cbs* transcription by at least two loci, one located in the  
317 Dp5Yah, and another one, not due to *Dyrk1a* overdosage, in the Dp1Rhr trisomic region.  
318 Similarly, under-expression of *Glp1r* was observed in Dp1Yey, Ts65Dn, Dp5/Dp1 and Dp5Yah  
319 while this gene was overexpressed in Dp1Rhr and not affected in Tg(*Dyrk1a*). Here too, *Dyrk1a*  
320 dosage was not involved, but at least two loci controlling *Glp1r* expression with opposite and  
321 epistatic effects should be found respectively in the Dp5Yah and Dp1Rhr genetic intervals.

322 The analysis of DEGs in each model separately highlighted the capabilities to separate  
323 the trisomic individuals from the wild-type littermates (Fig 2A) by principal component  
324 analysis (PCA), t-SNE (Fig 2B) or OPLS techniques (See supplementary material and  
325 methods).

326



327

328 **Figure 2: The differential expression analysis discriminates trisomic from disomic**  
 329 **hippocampi and identifies common dysregulated genes and pathways.** (A) 3D-PCA on the  
 330 DEGs for each sample allows to separate trisomic (Dp) from disomic (wt) adult hippocampi.  
 331 (B) Left column: two dimensional Principal Component Analysis (2D-PCA) on the 5599  
 332 transcripts of the 4328 DEGs over all the samples identified using  $f_{cros} 0.025 < \alpha < 0.975$ . Right  
 333 column: 2D-PCA on the 75 trisomic genes with a measured expression in all the models. (C)  
 334 Venn Diagrams showing the overlap in gene expression between the different mouse lines  
 335 represented in different colours. (D) Heatmap representation of the number and regulation sense



336 of the pathways shared at least by two mice lines identified using the genome expression for  
337 each mice line by GAGE R package and filtered by q-value cut off  $< 0.1$ . Grouped in the  
338 categories showed on the ordinate: synaptic related, synaptic related: representation of the  
339 pathways involved in Myelin sheath and SNARE complex formation, synaptic related: all the  
340 synaptic related pathways excluding myelin sheath and SNARE complex formation,  
341 Transcription & epigenomics regulation, Enzymes activity, Ribosome related, Mitochondria  
342 related, Cell Structure & organelle related, Phospho-kinase related... The color key breaks  
343 represents the number of pathways within the categories 60,40,20,5,0.5. The minus or pink  
344 color represents down regulated pathways, the white color represents no pathway found in the  
345 category and the purple or positive numbers stands for up regulated pathways respectively.  
346

347 A genome-wide misregulation was found independently of the model, as DEGs were  
348 spread in all the chromosomes (Supplementary table 3-4, Supplementary fig 8), as shown  
349 previously (51), although with a stronger impact of the Dp1Rhr duplication on the number of  
350 total DEGs detected. The most overexpressed genes in terms of log2FC of expression and  
351 significance in various genetic conditions were visualised using Volcano plots (Supplementary  
352 fig 9, Supplementary table 4). For example, the *listerin E3 ubiquitin protein ligase 1 (Ltn1)*  
353 gene, coding for a major component of the ribosome quality control and causing  
354 neurodegeneration in mice (55), was found overexpressed in Dp1Yey, Ts65dn, Dp5/Dp1, and  
355 Dp5Yah or *Ifnar2*, coding for the Interferon Alpha and Beta Receptor Subunit 2, is  
356 overexpressed as expected in models that carry three copies of this gene (Dp1Yey, Ts65Dn,  
357 Dp5/Dp1 and Dp5Yah). Instead, a more controlled gene like the *neuronal acetylcholine*  
358 *receptor subunit alpha-3 (Chrna3)*, is found upregulated only in Dp1Rhr and Dp1/Dp5,  
359 certainly due to the overexpression of one gene from the *Cbr1- Fam3b* region but not *Dyrk1a*.  
360 Nevertheless, when we performed the intersection between the list of DEGs from the different  
361 models, we only found a few genes in common (Fig 2C, Supplementary table 4). We decided  
362 to combine the analysis of all the lines together using PCA and t-SNE and revealed a strong  
363 clustering of models that share the same number of trisomic genes (Fig 2B). t-SNE analysis,  
364 based on 4328 DEGs, adding all DEGs detected in each mouse model together, showed  
365 different contribution of the various DS models to the transcriptome variation (Fig 2B, left  
366 panel) with 2 distinct groups: one encompassing four overlapping trisomies: Ts65Dn, Dp5/Dp1,

367 Dp5, Dp1Rh and three isolated models: Dp1Yey, Dp3Yah and Tg(*Dyrk1a*) that are closer  
368 together although Dp3Yah is clearly farthest from the other two. Similar distinct groups were  
369 seen when analysing the TEGs (Fig 2B, right panel) and overall, the trisomic and the wild-type  
370 individuals in each mouse line were nicely separated. As expected, the expression level of the  
371 TEGs and the DEGs in the different trisomic conditions were strongly correlated  
372 (Supplementary fig 10). Interestingly, the 4328 DEGs showed a level of mis-regulation strongly  
373 correlated between Dp1Yey and Dp3Yah (33%), Dp5/Dp1 (50%), Dp1Rhr (40%) and  
374 Tg(*Dyrk1a*) 42%. Of the 75 TEGS, the correlation was quite strong between Dp1Yey and  
375 Tg(*Dyrk1a*) with 28%. Thus, the correlation in gene deregulation showed that *Dyrk1a*  
376 overdosage is a key driver of transcriptome deregulation in the Dp1Yey and Dp1Rhr models.  
377 Unexpectedly, the correlation of DEGs mis-expression level was lower between Ts65Dn and  
378 Dp1Yey (29%) or Dp1/Dp5 (28%). On the opposite, a high number of TEGs are mis-regulated  
379 in the same way between Ts65Dn and Dp1Yey (49%) or Dp1/Dp5 (52%; Supplementary fig  
380 10) suggesting that the other region found in 3 copies in the Ts65Dn over Mmu17 must be  
381 affecting the general DEGs landscape. We confirmed by qPCR the mRNA overexpression of  
382 first, *Dyrk1a* and *Sod1* genes in the DS models where they are trisomic; second, of *Synj2* and  
383 *Tiam2* that are located on the Mmu17 centromeric region in the Ts65Dn and third, of *Chrna1* a  
384 gene misregulated in Dp1Yey, Dp5/Dp1, Dp1Rhr and Ts65Dn models (Supplementary fig 15).  
385 As expected *Cbs*, expression was down regulated in all the models except Tg(*Dyrk1a*) and  
386 Dp3Yah. We also detected alterations of the expression of immediate early-response genes as  
387 *Arc*, *FosB*, *Fos* and *Npas4* that are important for cognition.

388

### 389 **Differential functional analysis unravel a few common altered pathways in DS models**

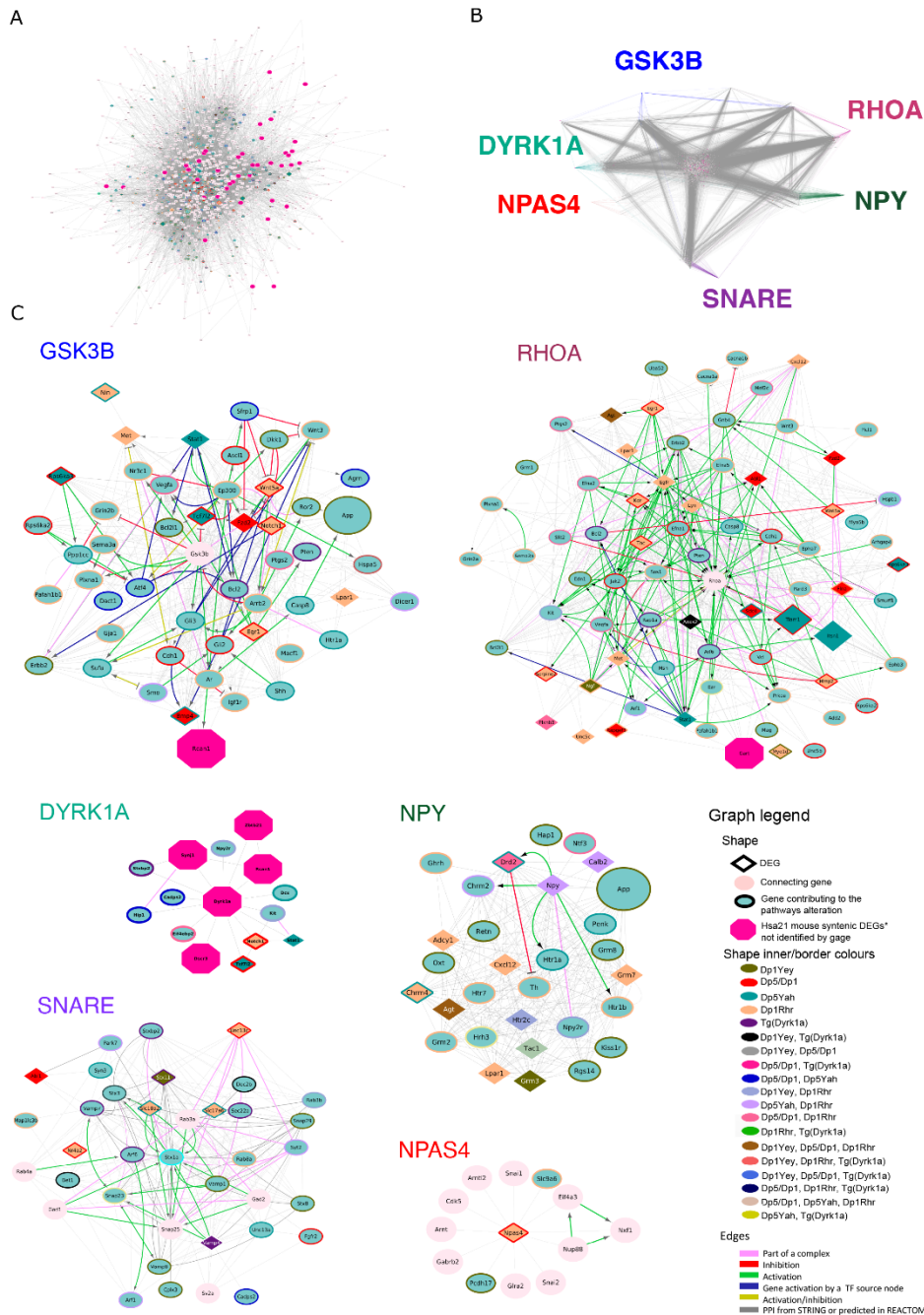
390 To go further, we performed a differential functional analysis and found 12 to 318 mis-  
391 regulated pathways in the DS models (table 1, Supplementary table 5). Interestingly, the sense  
392 of regulation of the overall deregulated pathways is trisomic region-dependent, as the Dp5Yah

393 (99%) region produced an overall downregulation whereas the Dp3Yah (89.5 %) and Dp1Rhr  
394 (56 %) regions together with the full Hsa21 syntenic model Dp1Yey (84.8 %), an upregulation.  
395 To facilitate the understanding, we clustered the broad functional dysregulation into 8 major  
396 functionality groups or meta-pathways. We found ribosomal components and mitochondrial  
397 processes pathways altered in all models with a high number of intermodel shared genes  
398 (Supplementary fig 12). Others meta-pathways like cell structure and organelles, transcription  
399 and epigenetic regulation, interferon and synaptic pathways were more affected in some models  
400 than in the others (Supplementary fig 13 and 14). As such, we observed strong and connected  
401 effects in the control of transcription and epigenetic regulation, enzyme activity and cell  
402 structure and cellular organelles involved in membrane and protein processing (endoplasmic  
403 reticulum, Golgi body, lysosome, peroxisome,...; Fig 2D) in the Dp1Yey, Dp5/Dp1, Dp1Rhr,  
404 and Tg(*Dyrk1a*) models, whereas the myelinization and 10 SNARE components, such as  
405 *Snap25* and *Snap23*, were specifically dis-regulated in the Dp1Yey Dp5Yah and Tg(*Dyrk1a*)  
406 models.

407

408 Interestingly, we saw a high number of shared genes between these pathways and the  
409 models giving rise to a high intermodel pathway connectivity. To assess the gene connectivity  
410 & cross talk, we build a protein-protein interaction (PPI) network of genes involved in synaptic  
411 function (noted MinPPINet) and furthermore we added regulatory PPIs to build the final  
412 RegPPINet. Then, we analysed the betweenness centrality to identify hubs, keys for  
413 maintaining the network communication flow that are most susceptible to targeted drugs attacks  
414 and the main signalling cascades affected. The relevance of the connecting nodes unknown to  
415 play a role in brain dysfunction, was further predicted by the machine learning algorithm  
416 Quack<sup>118</sup> and we found several evidences linking 449 (22 with a high confidence score) DEGs  
417 to GO/KEGG pathways involved in brain dysfunction, not previously known (Supplementary  
418 table 5). In fact, several evidences make us consider that those genes should be included in the

419 pathways:gene associations. Inside the evidences we can point that these DEGs were found dis-  
 420 regulated in hippocampus, were included in the network built by STRING<sup>73</sup> as connecting  
 421 nodes (not seeds) to produce a fully connected DS network, and their relevance to brain  
 422 dysfunction was also predicted by Quack, in some cases, with a high score.



423  
 424  
 425 **Figure 3: Protein-protein interaction networks involving DEGs linked to the synaptic**  
 426 **function.** (A) STRING04 MinPPINet of genes involved in synaptic function visualized using  
 427 the edge weighed spring embedded layout by betweenness index in Cytoscape. The network was  
 428 built by querying STRING and selecting the PPIs with a medium confidence score (CS=0.4)  
 429 coming from all sources of evidence. The shapes of the nodes represent the following  
 430 information: Shapes: i) Pallid pink ellipses: represent connecting proteins added to assure the

431 full connectivity of the network; ii) pink octagons, represent HSA21 syntenic genes in mouse  
432 not identified as contributing to the meta-pathway dysregulation by GAGE; iii) green inner  
433 coloured ellipses, genes identified by GAGE after q-val <0.1 cut off to be contributing even  
434 slightly, to any pathway of those found dysregulated inside the meta-pathway. If the size is  
435 similar to the octagons, they are also HSA21 syntenic genes in mouse. Additionally, the border  
436 colour represents the mouse model multi group where those genes are found altered in; iv)  
437 diamonds, genes identified by GAGE after q-val <0.1 cut off and also by FCROS as DEGs.  
438 (B) Network Structure Decomposition of the STRING04 MinPPINet. Highlighting in different  
439 colors the interactions of GSK3B, NPY, SNARE proteins, DYRK1A and RHOA respectively.  
440 In the case of NPAS4, the interactions coloured correspond up to the first level interactions.  
441 (C) The six RegPPINets were extracted from the selection of each fo the following proteins and  
442 their 2<sup>nd</sup> interactors from STRING04 MinPPINet: RHOA, DYRK1A, GSK3B, NPY, SNARE  
443 proteins and NPAS4. Then, those were further annotated with regulatory information using  
444 REACTOME (See Supplementary material and Methods). The shapes of the nodes represent  
445 the following information: Shapes: i) Pallid pink ellipses: represent connecting proteins added  
446 to assure the full connectivity of the network; ii) pink octagons, represent HSA21 syntenic  
447 genes in mouse not identified as contributing to the meta-pathway dysregulation by GAGE; iii)  
448 green inner coloured ellipses, genes identified by GAGE after q-val <0.1 cut off to be  
449 contributing even slightly, to any pathway of those found dysregulated inside the meta-  
450 pathway. If the size is similar to the octagons, they are also HSA21 syntenic genes in mouse.  
451 Additionally, the border colour represents the mouse model multi group where those genes are  
452 found altered in; iv) diamonds, genes identified by GAGE after q-val <0.1 cut off and also by  
453 FCROS as DEGs. The edges colored represent the type of interaction annotated by following  
454 the PathPPI classification (Tang *et al.* 2015), and ReactomeFIViz annotations as follows i) The  
455 GEl edges indicating expression were colored in blue and repression in yellow. ii) PPre edges  
456 indicating activation were coloured in green, inhibition in red. Iii) Interactions between proteins  
457 known to be part of complexes in violet. Iv) Predicted interactions were represented in grey  
458 including the PPI interactions identified by STRING DB (Szklarczyk *et al.* 2017) after merging  
459 both networks.  
460

461 Looking into the DS synaptic MinPPINet (Fig 3A), first we analysed the DS network  
462 topography and betweenness connectivity and found hubs and genes more central for the  
463 network information flow. As expected from a PPI biological network, after computing 100000  
464 random networks with a similar degree, the likelihood of observing such connectivity in the DS  
465 network was more than one can expect by chance (P-value < 2e-16) and it showed a small world  
466 effect and scale-free topology. Using a network decomposition approach (see supplementary  
467 material and methods), we highlighted 5 major subnetworks or biological cascades that strongly  
468 centralized 5 different proteins: DYRK1A, GSK3B, NPY, RHOA and SNARE proteins (Fig  
469 3B-C). While DYRK1A is known to be central and was shown to have interaction with GSK3B  
470 and NPY in DS models (34, 56-60), the important role of SNARE complex proteins and the



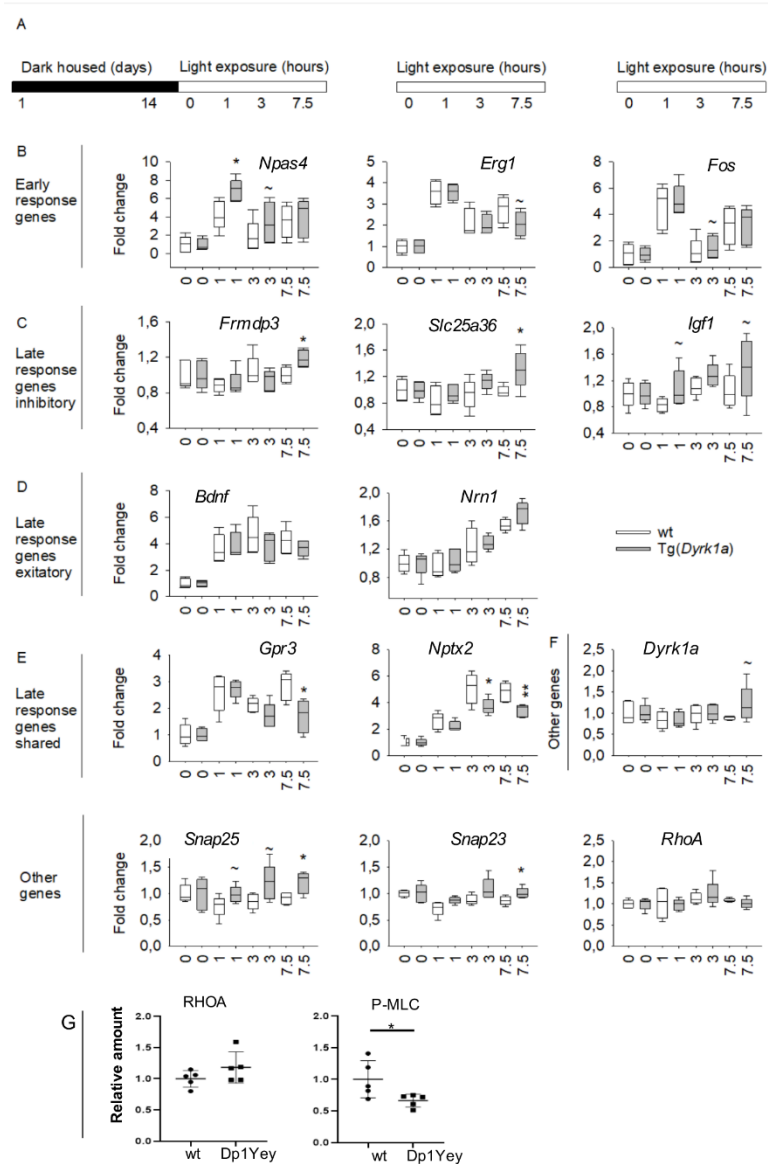
471 biological cascade centred around RHOA for the pathophysiology of DS cognitive dysfunction  
472 is exposed by our analysis. Moreover, the network analysis showed that DYRK1A controls 42,3  
473 % of the network nodes and 69,4 % of the network seeds, (DEGs known to be involved in brain  
474 synaptic pathways) via 2<sup>nd</sup> level interactors. Hence, DYRK1A could control the DS synaptic  
475 network via PPI and regulatory interactions. Furthermore, the biological cascades centred on  
476 GSK3B, DYRK1A and RHOA are highly interconnected (Supplementary fig 15) and in fact  
477 several interactors of RHOA are connected and could somehow modulate a higher percentage  
478 (75 and 68.5%) of the nodes of the network and synaptic seeds. Interestingly, *RhoA* was not  
479 found altered in the DEA analysis, instead was a node introduced in the network to have a fully  
480 connected synaptic PPI network. We checked whether the RHOA pathway was altered in the  
481 Dp1Yey and we found no changes in the expression of RHOA, confirming the transcriptomics  
482 analyses, but a significant decrease of the phosphorylation of the RHOA interactor Myosin light  
483 chain (P-MLC) in the Dp1Yey hippocampi compared to control. Thus, the RHOA pathways  
484 appeared to be down-regulated in the Dp1Yey DS mouse models (Fig 4A).

485

#### 486 ***Npas4* pathway is up-regulated as a consequence of *Dyrk1a* overexpression**

487 In our transcriptomics and network analysis, we found that *Npas4* was down-regulated in  
488 Tg(*Dyrk1a*), Dp1Rhr and Ts65Dn models and this gene level of expression is highly variable  
489 in the models (Supplementary fig 11). *Npas4* belongs to the class of immediate early genes with  
490 *Arc*, *Fosb* and *Fos*. Light deprivation and then exposure is known to drive robust gene  
491 expression, with shared early-response genes and distinct sets of late-response genes in  
492 excitatory and inhibitory (E/I) neurons (61). *Npas4* was recently identified as a key in E/I  
493 balance in the visual cortex after light deprivation (62).





#### Figure 4: Evaluating *Npas4* and *RhoA* pathways in DS models

(A) RHOA pathway was altered in the Dp1Yey. Western blot analysis was revealed no changes in the expression of RHOA but a significant decrease of the phosphorylation of the Myosin light chain (P-MLC) in the Dp1Yey hippocampi compared to control. (B) Mice were housed in total darkness for 14 days and then were subsequently exposed to light for 0, 1, 3 or 7.5 h. Relative expression levels were determined, and fold change were calculated for each condition. Genotypes differences in fold change were assessed by T Test. (C) Only the fold change for early response genes *Npas4* was up-regulated in Tg(*Dyrk1a*) mice compared to wt at 1 hours of light induction. (D) The late responses genes specific to inhibitory neurons *Frmdp3*, *Slc25a36* and *Igf1* were up-regulated after 7.5 hours of light induction. (E) The fold change

of late responses genes specific to excitatory neurons *Bdnf* and *Nrn1* were unchanged. (F) The fold change of late response genes shared by excitatory and inhibitory neurons *Gpr3* and *Nptx2* were downregulated after 3 and/or 7.5 hours of light induction. (G) *Dyrk1a* and *RhoA* showed a similar fold change along the different condition whereas *Snap25* and *Snap23* presented an increased enrichment for the 7.5 hours condition. Data are presented as box plots with the median and quartiles.

526 of late responses genes specific to excitatory neurons *Bdnf* and *Nrn1* were unchanged. (F) The  
 527 fold change of late response genes shared by excitatory and inhibitory neurons *Gpr3* and *Nptx2*  
 528 were downregulated after 3 and/or 7.5 hours of light induction. (G) *Dyrk1a* and *RhoA* showed  
 529 a similar fold change along the different condition whereas *Snap25* and *Snap23* presented an  
 530 increased enrichment for the 7.5 hours condition. Data are presented as box plots with the  
 531 median and quartiles.  
 532

533 To confirm the impact of *Npas4* down regulation in Tg(*Dyrk1a*) mice, we performed  
 534 qRT-PCR experiments to determine the specific early and late response genes altered in the  
 535 visual cortex after light deprivation at 3 time points (1, 3 and 7.5 hours) after de novo light  
 536 exposure (Fig 4B). The results showed that *Npas4* was clearly induced after light deprivation  
 537 following 1 hour of light stimulation. Expression data revealed also a strong overexpression of  
 538 *Npas4* following 1 hour of light stimulation in Tg(*Dyrk1a*) mice (Fig 4C). On the contrary, the

539 expression of late response genes specific for inhibitory neurons (*Frmdp3*, *Slc25a36* and *Igf1*,  
540 Fig 4D) and late response genes (*Grp3* and *Nptx2*, Fig 4F) were altered after 7.5 hours of light  
541 stimulation in Tg(*Dyrk1a*). Interestingly, late response genes specific to excitatory neurons (*Bdnf*  
542 and *Nrn1*, Fig 4E) were not affected. The *Snap25*, *Snap23* candidate genes found in our analysis  
543 showed an altered expression after 7.5hours of light stimulation while *Dyrk1a* and *RhoA* levels  
544 were not affected (Fig 4G). These results indicate that inhibitory pathways controlled by *Npas4*  
545 were affected by *Dyrk1a* overexpression. Furthermore, the network analyses highlighted  
546 NPAS4 as a potential modulator of the synaptic dysfunction as via well connected interactors,  
547 NPAS4 could affect the main altered biological cascades plus the GABA and NMDA receptors  
548 involved in the modulation of the excitatory / inhibitory balance of the brain (61).

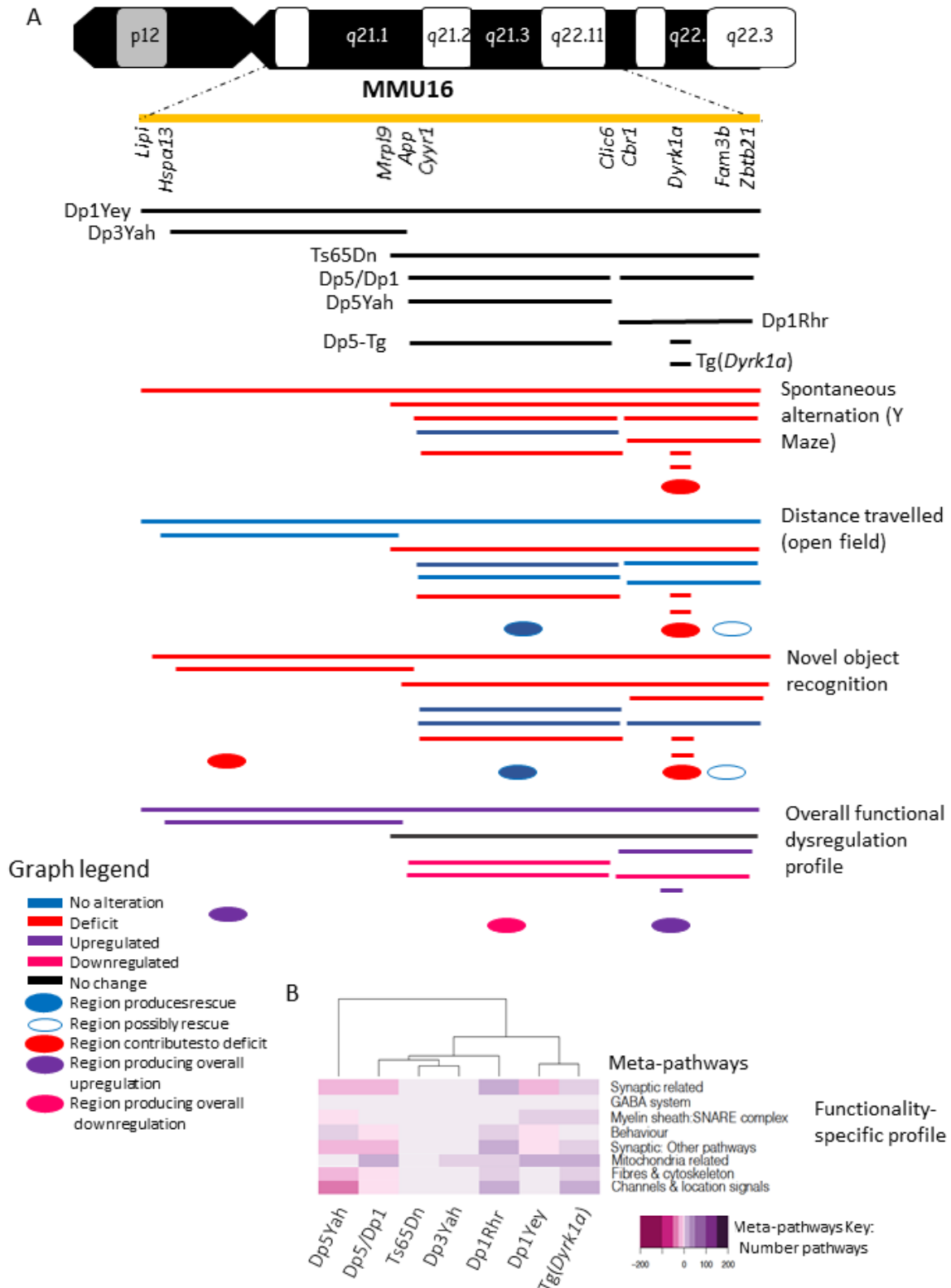
549

## 550 Discussion

551 In this study, we explored the in vivo DS mouse model library focusing on the region  
552 homologous to Hsa21 found on Mmu16, to decode the DS genotype-phenotype relationships  
553 and further underlie genetic interactions between different regions. For this, we assessed 6  
554 partial DS trisomic models and a transgenic model overexpressing one copy of *Dyrk1a* using  
555 a standardized behavioural pipeline focused on hippocampus-dependent memory processes. As  
556 expected, we found deficits in the Ts65Dn mice similar to the ones previously published in the  
557 Y-maze, the open field, NOR, MWM and contextual fear conditioning (36, 63) tests.  
558 Strikingly, thigmotaxis and time in the target quadrant in the probe test of the MWM were two  
559 variables only modified in the Ts65Dn trisomic model while the Dp1Yey, that carry a  
560 duplication of the complete syntenic Mmu16 region, was less affected as described before (64,  
561 65). Interestingly, the Dp5Yah mice, duplicated for a chromosomal region from *Cyrr1* to *Clic6*,  
562 present no deficits in all the tests done and have the lower number of DEGs. This might indicate  
563 that the *Cyrr1-Clic6* region was not sufficient to induce by itself cognitive defects. However,  
564 this syntenic region includes the genes *Sod1*, *Olig1*, *Olig2*, *Rcan1* and *Synj1*, proposed before

565 as inducing early DS cognitive phenotype (66-69). Curiously, no defects were found in the  
566 Dp5/Dp1 mice contrary to the Ts65Dn in the open field or less severe in the NOR test. This  
567 indicates the existence of a key modulator in the *Cyrr1-Clic6* region. The major behavioural  
568 alterations found in the Ts65Dn could result from the influence of different factors. First, the  
569 presence of the freely segregating mini-chromosome (70). Similarly, transcriptomic analysis  
570 showed a different global disruption of the genome expression, compared to the other trisomic  
571 models with segmental duplications. Then, the trisomy of about 60 Mmu17 centromeric genes  
572 non homologous to Hsa21(16), overexpressed in the hippocampus of Ts65Dn mice,  
573 highlighting specially *Tiam2* and *Synj2* that may exacerbate the effect of the overexpression of  
574 genes located on the Mmu17 segment homologous to Hsa21 as *Tiam 1* and *Synj1*. In fact,  
575 previous studies underline the strong role of *Synj1* in cognitive deficits of Ts65Dn (69, 71).  
576 Moreover, the major differences in behavioral alterations could be the result of the syntenic  
577 genes overexpressed in this model that have complex interactions with others. This is  
578 emphasized by the low correlation of DEGs and deregulated pathways between Ts65Dn,  
579 Dp1Yey and Dp5/Dp1. Finally, the presence of modulator genes, in the Dp5 region for example,  
580 could alter the behavioral performance and gene expression landscape. In example, as suggested  
581 by our genotype-phenotype analysis it seems to be a suppression of the effect of the *Mrpl39-*  
582 *Zfp295* trisomy in the Dp1 Yey model. In addition, several genes are deregulated only in Dp1Rhr  
583 and Ts65Dn, and not in Dp5/Dp1, or vice versa, which can explain the rescue of episodic  
584 memory in Dp5/Dp1 and highlights even more the complexity in gene interactions established  
585 between the different trisomic regions.

586 In this unique comparison, we found that the spontaneous alternation in Y-maze, was  
587 altered in all the models (except the Dp5Yah alone; Fig 5).



588

589 **Figure 5: Genotype correlation associated to behaviour phenotype in partial trisomic DS**  
 590 **model.** Here we highlight the duplicated region carried on each model with the corresponding  
 591 syntenic region in the human chromosome 21 together with the main behavioral and  
 592 transcriptomics results pointing to the existence of region specific phenotypes and functional  
 593 alterations. The black lines represents the duplicated syntenic regions to human chromosome  
 594 21 on each model (represented in the yellow line). The blue lines represents the behavioral

595 results where no alteration was found, instead the red lines identified the tests with deficits. Over  
596 the transcriptomics meta-pathways functional profile summary picture, in purple is highlighted  
597 upregulation whereas in pink downregulation. The intensity of the color stands for the number  
598 of pathways included on each meta-pathway from the total number of pathways found altered  
599 on each model.  
600

601         The minimal common genetic part of these lines was the overexpression of *Dyrk1a* and  
602 the result observed for the transgenic Tg(*Dyrk1a*), altogether with results previously obtained  
603 (72-74), support DYRK1A as a main driver of this DS cognitive phenotype (34). Similarly,  
604 DYRK1A overdosage caused the increased exploration measured by the distance travelled in  
605 the open field that was only affected in Ts65Dn, Dp5/Tg(*Dyrk1a*) and Tg(*Dyrk1a*).  
606 Interestingly, the Dp1Rhr Ds model was not affected suggesting that another loci interfere with  
607 *Dyrk1a* overdosage in this model and the situation should be even more complex as no  
608 phenotype is observed for the distance travelled in the Dp5/Dp1 while the Ts65Dn showed an  
609 increase in this variable. The novel object recognition with 24h of retention unravelled  
610 deficiency in most of the models, except in the Dp5Yah and Dp5/Dp1, suggesting that there are  
611 at least two causative loci: one located in the Dp3Yah region and *Dyrk1a*; and presumably two  
612 modifier interacting loci: one located in the Dp5Yah and another in the Dp1Rhr regions (Fig  
613 5). Altogether, the altered behavior and cognition observed in DS models results from the  
614 interaction of several loci, spread along the Mmu16, including *Dyrk1a*. Moreover, as  
615 summarized in Fig 5, our study highlights the existence of different degrees of genetic  
616 interaction complexity for behavioural and brain morphology phenotypes.

617         Surprisingly, brain morphology was less affected in Dp1Rhr compared to the other DS  
618 models, while Dp1Yey, Ts65Dn and Tg(*Dyrk1a*) shared alterations in identical structures.  
619 Changes were found in several regions including the basal forebrain septum, a predominant  
620 source of cortical cholinergic input with an early substantial loss of basal forebrain cholinergic  
621 neurons (BFCN) as a constant feature of Alzheimer's disease and other deficits in spatial  
622 learning and memory (75). Individuals with DS exhibit progressive degeneration of BFCN (76).

623 Similarly, significant loss of BFCN is present in the Ts65Dn (for review see (77)) and  
624 moreover, our results seem to indicate a defect in BFCN in Dp1Yey and Tg(*Dyrk1a*) too.  
625 Several findings suggest that overexpression of amyloid protein precursor (APP) play a major  
626 role and could be strictly necessary for BFCN cell loss (78), thus, future investigation in  
627 Tg(*Dyrk1a*) should reveal if BFCN loss is present even without APP overexpression. Besides,  
628 only the Ts65Dn present an enlargement on the ventricles, which was previously associated  
629 with a decrease of cortical neurogenesis in the brains of Ts1Cje and Ts2Cje mouse models (28).

630 Our comparative genome wide expression profiling in the mouse hippocampus revealed  
631 that not a single gene or a single region can recapitulate the whole dysregulation. The overall  
632 effect results from a complex interplay of a few trisomic overexpressed genes and other genes  
633 spread along the genome, shown by the majority of DEGs not being Hsa21 genes  
634 (Supplementary table 3). Additionally, we identified 34 trisomic DEGs (TEGs) with regulatory  
635 activity (Transcription factors, chromatin modellers...) as *Mir99a*, *Usp16*, *Erg* or *Rcan1* that  
636 may be involved in the models' changed regulatory landscape. Indeed, *RCAN1* and *USP16* were  
637 found upregulated in all human brain datasets (cerebrum and cerebellum) and *USP16* was also  
638 found as DEG upregulated in heart and adult fibroblasts while *MIR99A* was found upregulated  
639 in adult fibroblast (46). Nevertheless, the DEGs expression was strongly correlated and  
640 conserved in Mmu16 based DS models. This is similar to the behavioural results obtained where  
641 related phenotypes were found in models carrying correlated partial duplications.  
642 Unexpectedly, Dp1Yey DEGs correlation is closer to Tg(*Dyrk1a*) than to Ts65Dn (42% against  
643 25%) and there is a negative correlation between Dp3Yah & Dp5Yah (22%) and Tg(*Dyrk1a*)  
644 & Dp5Yah (13%) pointing out to a different gene dysregulation and phenotype development  
645 on these models and to the existence of epistasis with some regulatory trisomic genes countering  
646 the effect of genes in other trisomic regions. Carrying an in depth functional annotation analysis,  
647 we could not find a unique common pathway altered in all models. Nevertheless, due to the  
648 high redundancy in both GO and KEGG terms and the genes associated with each term,



649 grouping the pathways shared at least by two models (materials & methods) in 8 functional  
650 groups unravelled 8 major meta-pathways with ribosome and mitochondrial function,  
651 transcription & epigenomics regulation, and the synapse function categories highly affected.  
652 We also found a strong upregulation of genes involved in the InterferonB pathway  
653 (Supplementary fig 13B) as some interferon receptors are found upregulated in Mmu16 DS  
654 models. As such *Ifnar2* and *Il10rb* were found upregulated in all the mice lines (except  
655 Tg(*Dyrk1a*)) pointing to a potentially critical role in the interferon pathway dysregulation. The  
656 same was observed with other genes as *Irgn1*, *Ifit1*, *Ifit2* or *Ndufa13*. This upregulation of the  
657 interferonB pathways was previously reported in the Ts1Cje mouse model (79, 80) and linked  
658 to a possible increase of activity of the Jak-Stat signaling pathway as recorded here by the up  
659 regulation of *Stat1*.

660 Expression of genes involved in long-term synaptic potentiation (LTP) and synaptic  
661 plasticity were decreased in Dp1Yey, Dp5Yah, Dp1Rhr, Dp5/Dp1 models respectively,  
662 corroborating previous reports in different DS mouse models and in vitro studies. The only  
663 upregulated pathways were the myelin sheath and SNARE; both found in Dp1Yey and  
664 Tg(*Dyrk1A*) models. Ten genes from the SNARE complex were dysregulated, from those, we  
665 validated the dysregulated expression of *Snap25* and *Snap23* by qRT-PCR. Interestingly,  
666 models carrying the Dp1Rhr region duplication showed a dysregulation mainly in synapse  
667 transmission, plasticity and LTP, while models carrying the Dp5Yah region duplication showed  
668 a dysregulation associated with genes involved in stemness and differentiation. Together, the  
669 models with both Dp5Yah & Dp1Rhr duplicated regions, were involved in post-synapse  
670 modulation and transmission. Moreover, there is a high number of shared DEGs between these  
671 pathways and the models giving rise to a high intermodel inter-pathway connectivity with genes  
672 strongly connected into major subnetwork biological cascades centralized over 6 different  
673 proteins DYRK1A, GSK3B, RHOA, NPY, NPAS4 and SNARE complex. These new cascades  
674 may also play a crucial role in the brain dysfunction associated with the DS phenotypes.

675 Organizing the network to follow the betweenness centrality index values, RHOA, DYRK1A,  
676 GSK3B, and their interactors were more closely knitted together and populated the central part  
677 of the network while SNARE, NPAS4 and NPY with their first- and second-layer interactors  
678 were more in the periphery of the network. This strong interconnectivity has a double  
679 interesting effect, first it makes the full network really sensible to targeted attack against these  
680 proteins and at the same time the network would be robust against the attacks if the attack does  
681 not target several proteins simultaneously; for example during a drug trial. Thus, studying  
682 further closely connected altered genes and understanding the interactions could provide novel  
683 insights into the possible molecular mechanism explaining why so many compounds, including  
684 DYRK1A specific kinase inhibitors, can restore learning and memory in DS models (35, 81-  
685 83). Additionally, these nodes show a high number of connections. Indeed, organising the  
686 network using the betweenness index, these nodes and their interactors occupy the centre of the  
687 network showing the extreme importance of these nodes for the stability of the network in terms  
688 of network theory. Moreover, similar to the observation in DS patients, where the  
689 affectation/severity of the gene dysfunction vary for one patient with partial duplications to the  
690 other, we proposed here that different DS mouse models show different signalling cascades that  
691 were affected with several members strongly interconnected and affecting brain dysfunction  
692 leading to similar behavioural phenotypes; may be one possible explanation of the  
693 developmental instability hypothesis, which postulates that the non specific triplication of a  
694 relative small number of genes causes genetic imbalance with wide impact on global gene  
695 expression. This hypothesis should be taken into account when therapeutic assay are planned.  
696 The result in one partial trisomic mouse model should be replicated in more genetically complex  
697 models to test potential genetic influential factor (35, 84, 85). This is probably the limit of the  
698 model, even if the mechanisms of behaviour and memory are common between mice and  
699 human, the complexity of the system is lower. Conducting the same studies in more complex

700 animal models, carrying all the trisomic genes homologous to Has21, probably would  
701 definitely permit a better deciphering of genes impacting on cognitive behaviour.

702

## 703 **Material and Methods**

### 704 **Mouse lines**

705 The duplications of different Mmu 16 regions (Dp(16*Lipi-Zbtb21*)1Yey (or Dp1Yey),  
706 the Dp(16*Hspa13-App*)3Yah (noted Dp3Yah), the Dp(16*Cbr1-Fam3b*)1Rhr (or Dp1Rhr)) and  
707 the BAC transgenic for *Dyrk1a* (Tg(*Dyrk1a*)) models were previously described (19, 20, 33,  
708 86). The genetic background of the DS lines carrying each duplication was pushed toward the  
709 C57BL/6J (B6J) genetic background for more than 7 generation of backcross. The only  
710 exception is the trisomic Ts65Dn (Ts(17<sup>16</sup>)65Dn) mice, initially obtained from the Jax, that was  
711 kept on a F1 B6C3B genetic background (with the C3B line as a C<sub>3</sub>H/HeH congenic line for  
712 the BALB/c allele at the *Pde6b* locus (87)). The Dp(16*Cyrr1-Clic6*)5Yah (noted Dp5Yah) was  
713 generated by the *in vivo* TAMERE technology inserting loxP sites in *App* and *Runx1* (see  
714 Supplementary information). In the Dp3Yah and Dp5Yah models, only 2 complete copies of  
715 *App* and *Runx1* genes are expressed. The Dp5Yah line was crossed with the Dp1Rhr in order  
716 to generate the Dp5Yah/Dp1Rhr (also noted Dp5/Dp1) compound transheterozygote carrying  
717 a similar trisomic Mmu16 gene content to that of the Ts65Dn. Indeed, only 15 Hsa21  
718 homologous genes (*Mrpl39*, *Jam2*, *Atp5j*, *Gabpa*, *App*, *Cyrr1*, *Runx1*, *Setd4*, *Mx2*, *Tmprss2*,  
719 *Ripk4*, *Prdm15*, *C2cd2* and *Zbpb21*) over 174 are not in 3 copies in Dp5/Dp1 compared to  
720 Ts65Dn while the genes located on the Mmu17 telomeric regions trisomic in the Ts65Dn  
721 minichromosome (16) are not trisomic in the Dp5/Dp1. The Dp5Yah model was also combined  
722 with the Tg(*Dyrk1a*) by crossing Dp5Yah/+ and Tg(*Dyrk1a*)/0 animals and generating the four  
723 genotypes (Dp1Yah, Dp5Yah, Tg(*Dyrk1a*) and [Dp5Yah; Tg(*Dyrk1a*)] noted here Dp5-Tg), to  
724 test specific interaction between Dp5Yah and *Dyrk1a* overdosage.

725 All the lines were maintained under specific pathogen-free (SPF) conditions and were  
726 treated in compliance with the animal welfare policies of the French Ministry of Agriculture  
727 (law 87 848) and the phenotyping procedures were approved by our local ethical committee  
728 (Com'Eth, n°17, APAFIS n°2012-069).

### 729 **Behaviour pipeline**

730 A serie of behavioural experiments were conducted in mice with a range of age between  
731 2,5 to 7 months, as described in the Supplementary information. The tests were administered in  
732 the following order: Y-maze, open field, novel object recognition (24h), Morris water maze and  
733 fear conditioning (contextual and cue). Behavioural experimenters were blinded to the genetic  
734 status of the animals. Separate groups of animals were composed for each line (as indicated in  
735 supplementary table 1). Several mouse models found defective for the NOR performed with  
736 24h of retention memory were also tested after 1h of retention. The Dp5Yah crossed with  
737 Tg(*Dyrk1a*) was tested for Y-maze and NOR at 24h. All the standard operating procedures for  
738 behavioural phenotyping have been already described (85, 88-91) and are detailed in the  
739 supplementary information.

740

### 741 **Magnetic Resonance Imaging**

742 A dedicated cohort of animals at the age 102 +/- 7 days was anesthetized and perfused  
743 with 30 ml of room temperature 1X Phosphate Buffer Saline (PBS) complemented with 10%  
744 (% w/v) heparine and 2mM of ProHance Gadoteridol (Bracco Imaging, Courcouronnes,  
745 France) followed by 30ml of 4% PFA complemented with 2mM of ProHance Gadoteridol.  
746 Then the brain structure was dissected and kept in PFA 4% 2mM ProHance over night at 4°C.  
747 The next day, each specimen was transferred into 1X PBS 2mM ProHance until imaging.

748 Just prior to imaging, the brains were removed from the fixative and placed into plastic  
749 tubes (internal diameter 1 cm, volume 13 mL) filled with a proton-free susceptibility-matching

750 fluid (Fluorinert® FC-770, Sigma-Aldrich, St. Louis, MO). Images of excised brains were  
751 acquired on a 7T BioSpec animal MRI system (Bruker Biospin MRI GmbH, Ettlingen,  
752 Germany). Images were reconstructed using ParaVision 6.0. An actively decoupled quadrature-  
753 mode mouse brain surface coil was used for signal reception and a 72-mm birdcage coil was  
754 used for transmission, both supplied by Bruker. The first protocol consisted of a 3D T2-  
755 weighted rapid-acquisition with relaxation enhancement (RARE). The second imaging protocol  
756 consisted of a 3D T2\*-weighted Fast Low Angle (FLASH) sequence. The image matrix for  
757 both sequences was 195 x 140 x 90 over a field of view 19.5 x 14.0 x 9.0 mm<sup>3</sup> yielding an  
758 isotropic resolution of 100 µm and was treated and analysed for anatomical parameters as  
759 detailed in the supplementary information.

#### 760 **Gene expression assay**

761 Hippocampus were isolated from DS trisomic models and their littermate controls (N =  
762 5 per group) and flash frozen in liquid nitrogen. Total RNA was prepared using the RNA  
763 extraction kit (Qiagen, Venlo, Netherlands) according to the manufacturer's instructions. The  
764 samples quality was checked using an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa  
765 Clara, California, USA). Gene expression analysis was carried out using GeneChip® Mouse  
766 Gene 1.0 ST arrays (Affymetrix, Santa Clara, CA). All the procedure and the analysis are  
767 detailed in the supplementary information. Raw microarray data and re-analysed data have been  
768 deposited in GEO (Accession No. GSE149470).

769

#### 770 **Bioinformatic analysis**

771 The gene expression profile of the mouse hippocampi isolated from Dp1Yey, Dp3Yah,  
772 Ts65Dn, Dp5/Dp1, Dp5Yah, Dp1Rhr and Tg(*Dyrk1a*) trisomic mouse models was analysed  
773 with a specific bioinformatic pipeline and controlled for quality prior and after the Data pre-  
774 processing and normalization (see supplementary information in the detailed material and

775 methods section). The identification of the differentially expressed genes (DEGs), was  
776 performed using a method based on fold change rank ordering statistics (FCROS)(92). In the  
777 FCROS method, k pairs of test/control samples are used to compute fold changes (FC). For  
778 each pair of test/control samples, obtained FCs for all genes are ranked in increasing order.  
779 Resulting ranks are associated to genes. Then, the k ranks of each gene are used to calculate a  
780 statistic and resulting probability (f-value) used to identify the DEGs after fixing the error level  
781 at 5% False Discovery Rate (FDR).

782 We performed the functional differential analysis using GAGE (93) and grouped all the  
783 pathways into 8 functional categories (noted meta-pathways). The functional intermodel meta-  
784 pathway connectivity was studied identifying the genes shared between pathways and models  
785 inside the same meta-pathway. Then, to assess the gene connectivity we build a minimum fully  
786 connected protein-protein interaction (PPI) network (noted MinPPINet) of genes known to be  
787 involved in synaptic function as they were associated with synaptic pathways via GO (94) and  
788 KEGG databases (95) and furthermore added regulatory information to build the final  
789 RegPPINet. We used the betweenness centrality analysis to identify hubs, keys for maintaining  
790 the network communication flow. The relevance of the connecting nodes was further predicted  
791 by the machine learning algorithm Quack (96). Finally, we computed 100000 random networks  
792 with a similar degree, to assess if the likelihood of observing such connectivity in the DS  
793 network was more than one can expect by chance using statnet and sna R packages  
794 (<https://cran.r-project.org/web/packages/statnet/index.html>; [https://cran.r-](https://cran.r-project.org/web/packages/sna/index.html)  
795 [project.org/web/packages/sna/index.html](https://cran.r-project.org/web/packages/sna/index.html)).

796

## 797 **Visual stimulation**

798 Mice raised in a standard light cycle were housed in constant darkness for two weeks.  
799 Then, animals in the light-exposed condition group, were consecutively exposed to light for 0,  
800 1, 3, or 7.5 h before being sacrificed. Instead, animals part of the dark-housed condition group



801 were sacrificed in the dark. After euthanasia, their eyes were enucleated before the visual cortex  
802 dissection in the light and flash frozen in liquid nitrogen. cDNA and quantitative PCR were  
803 performed as indicated in the supplementary information. The Ct values were transformed to  
804 quantities by using the comparative Ct method. Hence, all data were expressed relative to the  
805 expression of the most expressed gene. This relative expression levels, were normalized with  
806 Genorm by keeping the more stable reference genes (97). To calculate fold-induction, relative  
807 quantity of gene expression at each time point was divided by the mean of relative level of gene  
808 expression of dark housed mice for corresponding genotype. The mean and standard error were  
809 calculated at each time point from these fold-induction values.

## 810 **Statistical analysis**

811 All data are expressed as mean group value  $\pm$  standard error of the mean (SEM) or as  
812 box plots with the median and quartiles. For each data set, we analysed if the data was normally  
813 distributed by Shapiro–Wilk test and Quantile-Quantile plots (Supplementary fig 2) and the  
814 homogeneity of variances by the Brown-Forsythe test. Differences between groups were inferred  
815 by one-way ANOVA (Open field) and ANOVA for repeated measures, or in case of datasets  
816 where the assumptions of normality or homogeneity of variances were not fulfilled we  
817 performed the Kruskal-Wallis non-parametric test. The post hoc tests (Fisher LSD Method)  
818 were conducted only if the F parameter in ANOVA achieved 0.05 level. All the behavioral  
819 analysis results are found in Supplementary table 1. For MRI data, intergroup comparisons on  
820 region-based data were conducted on the normalized volumes (i.e., ratio between the volume  
821 of the structure and the whole brain volume) of each segmented structure using the Student t-  
822 test while correcting by multiple testing setting up a FDR correction.

823

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833

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## Supporting information

**S1 Fig. Down syndrome mouse models analysed in the study.** In the upper part of the plot the human chromosome 21 is represented, in yellow we highlighted the Hsa21 syntenic region found in mouse from *Lipi* to *Zbtb21* (known as *Zfp295* previously). The eight models analysed on this study Dp1Yey, Dp3Yah, Ts65Dn, Dp5/Dp1, Dp5Yah, Dp1Rhr, Tg(*Dyrk1a*), Dp5yah crossed with Tg(*Dyrk1a*) (noted as Dp5-Tg)) trisomic chromosomal regions were drawn in comparison with the Hsa21 region.

**S2 Fig. Quantile-Quantile plots to assess Normal distribution.** The quantile-quantile plots (Q-Q plots) and Shapiro-Wilk test for the following behavioural datasets: Y-maze, NOR and OF were analysed to study how well our data could be modelled following a normal distribution. Only the distance travelled in OF did not reach the statistical significance level in the Shapiro-Wilk test.

**S3 Figure. DS mouse models and Morris water maze.** The MWM results are presented as distance travelled to reach the platform (A), speed (B) and % of distance travelled in the peripheral zone (C, Mean  $\pm$  SEM). Ts65Dn mice were drastically impaired with an increase in distance travelled to reach the platform and in the distance travelled along the peripheral zone. Less drastically Tg(*Dyrk1a*) mice showed an increased distance travelled to escape the pool and in the peripheral zone (\*  $p < 0.05$ , \*\* $p < 0.01$ , \*\*\* $p < 0.001$ ).

**S4 Fig. DS mouse models and context and cue fear conditioning.** The fear conditioning results are presented as inactivity periods in seconds (Mean  $\pm$  SEM). Only Ts56Dn mice present an impairment in context fear conditioning (\*  $p < 0.05$ , \*\* $p < 0.01$ , \*\*\* $p < 0.001$ ).

**S5 Fig. Comparison of the Z-score calculated from the MRI volume of different brain structures from the DS models.** The structures affected in Tg(*Dyrk1a*) were also affected in Dp1Yey and Ts65Dn, whereas the Dp1Rhr mice. (Dp1Yey n=6wt and 6 Tg ; Ts65Dn n=5 wt and 6 Tg ; Tg(*Dyrk1a*) n=8 wt and 5 Tg ; Dp1Rhr n=7 wt and 7 Tg).

**S6 Fig. Fold change expression levels of the genes analysed by the microarrays homologous to the Hsa21 mouse chromosome 16 (mm16).** The genes are displayed following the order of their genomic start site coordinates. The duplicated areas for each mouse model appear shaded in the following colours, purple, grey, aquamarine, red, blue and golden for Dp1Yey, Dp3Yah, Ts65Dn, Dp1Rhr, Dp5Yah and Tg(*Dyrk1a*) respectively.

**S7 Fig. Fold change expression levels of genes analyzed by the microarrays.** (A) Fold change expression levels of the genes analysed by the microarrays part of the centromeric region of the mouse chromosome 17. The centromeric region of the mouse chromosome 17, is defined by the UCSC table browser as the region between the coordinates chr17:110000-3000000. The genes are displayed following the order of their start site genomic coordinates. The region duplicated in the Ts65Dn model non homologous to the hsa21, appears shaded in aquamarine. (B) Fold change expression levels of the genes homologous to the Hsa21 in mouse

chromosome 10 (mm10) and chromosome 17 (mm17) analysed by the microarrays. The genes are displayed following the order of their start site genomic coordinates.

**S8 Fig. Number of deregulated probes per chromosomes per models.** Trisomic mice have similar patterns with a pic of deregulated genes located on mouse chromosome 16. Tg Dyrk1a at contrary, present a global pattern of deregulated probes.

**S9 Fig. Overview of the top 7 dysregulated genes (higher significance and fold change difference) for each mouse model over the global transformed expression profiles illustrated over volcano plots.** The x axis and y axis represent the log<sub>2</sub> fold change of the genes expression and the log transformed FDR values respectively. The genes whose fcros FDR value <0.05 are considered significant. These genes are represented in purple or pink and labelled as “Significant”. Instead the genes whose abs (FC)> 1.4 are represented in green or pink and labelled “FC”. Finally, the genes that don’t comply these conditions are considered not significant.

**S10 Fig. Gene expression correlation analyses.** (A) Correlation of the 4328 differentially expressed genes (DEGs) expression levels on the different models. B) Correlation of the 75 differentially expressed trisomic genes (TEGs) expression levels on the different models. Each row represents a model pairwise correlation with the rest of the models. The density and histograms distribution of the gene expression values for each model is found in the medial diagonal (Black for DEGs, and pink for TEGs). The LOESS smoothed fit is represented by a red line. The plots were created in R using the pairs.panel function from Psych R CRAN package (28).

**S11 Fig. Weighted networks representing the connectivity of the pathways included in the Cell structure & organelles, and the Transcription & epigenomic regulation meta-pathways.** (A) Representation of the strength connecting the pathways included on the Cell Structure & organelle meta-pathway. (B) Representation of the strength connecting the pathways included on the Transcription & epigenomics regulation meta-pathway. The pathways incorporated in each meta-pathway were identified in the differential functional analysis (DFA) by GAGE after imposing a q-val <0.1 cut off. The connectivity between the pathways, (the edges weight or strength connecting the pathways) is represented by the number of altered genes identified by GAGE shared by each two pathways and is calculated based on the number of common genes within the pathways shared inter and intra mouse model (by each mouse model multi group defined) also grouped by regulation sense using the following formula\* and represented by the thickness of the edges connecting the nodes (pathways) of the network. Each edges colour depends on the mouse models where those genes were identified (mouse model multi groups). Formula\* = number of genes shared by each pathway on each model on each sense of regulation \* factorRes with factorRes= 0.6.

**S12 Fig. Weighted networks representing the connectivity of the pathways included in the Ribosome and Mitochondrial meta-pathways.** (A) Representation of the strength connecting the pathways included on the Ribosome meta-pathway. (B) Representation of the strength connecting the pathways included on the Mitochondria meta-pathway. The pathways incorporated in each meta-pathway were identified in the differential functional analysis (DFA) by GAGE after imposing a q-val <0.1 cut off. The connectivity between the pathways, (the

edges weight or strength connecting the pathways) is represented by the number of altered genes identified by GAGE shared by each two pathways and is calculated based on the number of common genes within the pathways shared inter and intra mouse model (by each mouse model multi group defined) also grouped by regulation sense using the following formula\* and represented by the thickness of the edges connecting the nodes (pathways) of the network. Each edges colour depends on the mouse models where those genes were identified (mouse model multi groups). Formula\* = number of genes shared by each pathway on each model on each sense of regulation \* factorRes with factorRes= 0.6

**S13 Fig. Weighted network representing the connectivity of the pathways included in the Synaptic meta-group and the network representation of the genes linked to Interferon Beta (IFN-B) ungrouped pathway.** (A) Representation of the strength connecting the pathways included on the Synaptic meta-pathway. The pathways incorporated in each meta-pathway were identified in the differential functional analysis (DFA) by GAGE after imposing a q-val <0.1 cut off. The connectivity between the pathways, (the edges weight or strength connecting the pathways) is represented by the number of altered genes identified by GAGE shared by each two pathways and is calculated based on the number of common genes within the pathways shared inter and intra mouse model (by each mouse model multi group defined) also grouped by regulation sense using the following formula\* and represented by the thickness of the edges connecting the nodes (pathways) of the network. Each edges colour depends on the mouse models where those genes were identified (mouse model multi groups). Formula\* = number of genes shared by each pathway on each model on each sense of regulation \* factorRes, factorRes= 0.6. (B) Genes linked to Interferon Beta (IFN-B) ungrouped pathway, found altered in the different mouse model multi groups by GAGE. The genes are represented by the ellipse shapes instead the mouse model multi groups by lilac cuadrangles. If the genes were found up regulated then a red border color was used around the ellipses. The genes also identified by FCROS as differentially expressed (DEG) were identified by a red colored inner ellipse.

**S14 Fig. Protein-protein interaction and regulatory gene connectivity networks (RegPPINets) of genes involved in the synaptic meta-pathway highlighting biological cascades connectivity.**The two sub-networks (A and B) were extracted from the STRING04 MinPPINet. This network was built querying STRING and selecting the PPIs with a medium confidence score (CS), CS=0.4 from all sources of evidence, and was further annotated with regulatory information using REACTOME (See Supplementary material and Methods). The shapes of the nodes represent the following information: Shapes: i) Pallid pink ellipses: represent connecting proteins added to assure the full connectivity of the network; ii) pink octagons, represent HSA21 syntenic genes in mouse not identified as contributing to the meta-pathway dysregulation by GAGE; iii) green inner coloured ellipses, genes identified by GAGE after q-val <0.1 cut off to be contributing even slightly, to any pathway of those found dysregulated inside the meta-pathway. If the size is similar to the octagons, they are also HSA21 syntenic genes in mouse. Additionally, the border colour represents the mouse model multi group where those genes are found altered in; iv) diamonds, genes identified by GAGE after q-val <0.1 cut off and also by FCROS as DEGs. (A) RegPPINet Sub-network extracted from the selection of RHOA 2<sup>nd</sup> interactors from STRING04 MinPPINet, highlighting RHOA 1<sup>st</sup> Interactors and DYRK1A and GSK3B interactors annotated with the regulatory information from REACTOME. (B) RegPPINet Sub-network extracted from the selection of NPAS4 2<sup>nd</sup> interactors from STRING04 MinPPINet. Inside of the ellipsoidal line shapes we highlighted NPAS4 interactors involved in the regulation of GABA receptors (cyan), of NMDA receptors (orange), SNARE complex (purple), RHOA (pink), GSK3B (Blue), both RHOA and GSK3B (black) and the genes linked to mitochondrial, ribosomal and cytoskeleton activity (yellow).



**S15 Fig. Expression level of genes of interest in the different DS models.** Different panels to show the expression level of genes of interest located or not in the region homologous to Hsa21. Data are presented as boxplot plots of the ratio of trisomic on disomic with the median and quartiles (Statistical analysis was done comparing wt and mutant value with a student t test, \*  $p < 0.05$ , \*\* $p < 0.01$ , \*\*\* $p < 0.001$ ).

**S16 Fig. Detection of RHOA and the phosphorylated form of Myosin Light Chain (P-MLC) by Western blot in Dp1Yey hippocampal lysates and their control (wt) littermates.** (A) Western blot of RHOA showed no changes in RHOA protein levels in the Dp1Yey line compared to their wt littermates (wt (n=5) and Dp1Yey (n=5)). (B) Western blot of P-MLC revealed a statistically significant decrease (\*  $p < 0.05$ ) in the amount of this protein in Dp1Yey mice (wt (n=5) and Dp1Yey (n=5)) (Student's t-test: wt vs. Dp1Yey  $t(8) = 2,392$ ;  $p = 0,0437$ ).

**S1 Table: Summary of behavioural results including statistical assessment.**

**S2 Table. Volumes of each brain structure (in mm<sup>3</sup>) normalized by total brain volume derived from MRI region-based analysis.**

**S3 Table. Summary of the DEA and DFA results for each trisomic DS model.** We included the number of trisomic genes with compensated or partially compensated expression levels, approximately half of the trisomic genes are compensated and the overall identity of DEGs are different between the models.

**S4 Table. Fold-change of the gene expression in trisomic (Tg) compared to the controls for each mouse model dataset.** The genes included on those lists were used to produce the supplementary figs 6 and 7 as follows: i) genes included inside the duplicated genomic regions of chr 16, ii) genes included inside the Ts65Dn chr17 duplicated region iii) genes of the centromeric region of chr17.

**S5 Table. Pathways identified by GAGE as dysregulated including the genes contributing to each altered pathway per each mouse model multi group combination and including the name of the meta-pathway assigned.**