1	TRANSCRIPTOMIC SIGNATURES OF TELOMERASE-DEPENDENT AND -INDEPENDENT							
2	AGEING, IN THE ZEBRAFISH GUT AND BRAIN							
3	Running title: Kinetics of ageing in the zebrafish gut and brain							
4								
5	AUTHORS							
6	Raquel R. Martins ¹ , Michael Rera ² and Catarina M. Henriques ¹							
7	Affiliations :							
8	1. The Bateson Centre, Healthy Lifespan Institute and Department of Oncology							
9	and Metabolism, University of Sheffield Medical School, Sheffield, UK.							
10	2. Université de Paris / Inserm- Centre de Recherche Interdisciplinaire (CRI Paris)							
11								
12	Corresponding author: <u>c.m.henriques@sheffield.ac.uk</u>							
13								
14	SUMMARY							
1 ~	The shall find that for the data							

Telomerase is best known for its role in the maintenance of telomere length and its 15 16 implications for ageing and cancer. The mechanisms, kinetics and tissue-specificity 17 underlying the protective or deleterious mechanisms of telomerase, however, remain 18 largely unknown. Here, we sought to determine the telomerase-dependent and -19 independent transcriptomic changes with ageing, in the gut and brain, as examples of high and low proliferative tissues, respectively. We hypothesised this could shed light on 20 21 common telomerase-dependent and -independent therapeutic targets aimed at preventing 22 or ameliorating age-associated dysfunction in both tissues. For this, we used the zebrafish 23 model which, similarly to humans, depends on telomerase for health- and lifespan. We performed whole tissue RNA sequencing of gut and brain, in naturally aged zebrafish 24 alongside prematurely aged telomerase null mutants (*tert^{-/-}*), throughout their lifespan. Our 25 study highlights stem cell exhaustion as the first main hallmark of ageing to be de-regulated 26 27 in WT zebrafish gut and brain. Towards the end of life, altered intercellular communication becomes the main hallmark of ageing de-regulated in both gut and brain, and this is 28 29 accelerated in both tissues, in the absence of telomerase. Finally, we identify 7 key gene 30 changes common between the gut and brain at the early stages of ageing, highlighting

31 potential early intervention therapeutic targets for preventing age-associated dysfunction in

32 both tissues.

33 **KEYWORDS:** Ageing, telomerase, telomeres, gut, brain, zebrafish, transcriptomics,

34 RNA sequencing

35 **1 INTRODUCTION**

36 Ageing is the strongest risk factor for chronic diseases. How and why this is the case 37 remain important questions in the field, especially as key research has shown that targeting 38 common hallmarks of ageing, such as cellular senescence (Baker et al., 2016; Baker et al., 39 2011), can have a positive impact across multiple tissues and ameliorate several chronic 40 diseases of ageing at the same time. There are well-known key hallmarks of ageing, such as genomic instability, telomere attrition, epigenetic alterations, loss of proteostasis, de-41 42 regulated nutrient sensing, mitochondrial dysfunction, cellular senescence, stem cell 43 exhaustion and altered intercellular communication(Lemoine, 2021; López-Otín, Blasco, 44 Partridge, Serrano, & Kroemer, 2013). However, a major challenge in ageing research is to 45 identify where and when these potentially pathological changes start and when the tipping 46 point between homeostasis and loss of function takes place (Rando & Wyss-Coray, 2021). 47 Additionally, several lines of evidence suggest there may be specific tissues where age-48 related changes start earlier, potentially influencing others (de Jong, Gonzalez-Navajas, & 49 Jansen, 2016; Rera, Azizi, & Walker, 2013). One example is the gut, which has been 50 suggested to be a trigger for multiple organ failure (Cardoso et al., 2008). Evidence suggests 51 that the kinetics of ageing can vary dramatically between cells, tissues (Shokhirev & 52 Johnson, 2021; M. J. Zhang, Pisco, Darmanis, & Zou, 2021) and individuals, and that this is 53 influenced not only by intrinsic but also extrinsic factors, recently discussed elsewhere 54 (Rando & Wyss-Coray, 2021). These considerations are of particular importance when the 55 aim is to understand how changes in ageing lead to disease and how, when and where to 56 intervene. This is crucial in order to shift towards a more preventive form of medicine, 57 which is a current global ambition (Rudnicka et al., 2020), aiming to match the dramatic 58 increase of lifespan we have experienced in the past century, with an equivalent increase in 59 years of healthy living, i.e, healthspan (England, 2017).

Tissue-specific transcriptomics analysis over the lifecourse can offer important insights
 into the downstream molecular mechanisms potentially driving the pathology of ageing.

62 Significant research is being dedicated to these approaches in different animal models, 63 including in mice (Schaum et al., 2020; Tabula Muris, 2020; M. J. Zhang et al., 2021). 64 Different animal models may offer different insights into the mechanisms of ageing, and 65 some models may be better suited to explore the role of specific human hallmarks of 66 ageing. The role of telomere attrition in natural ageing can be considered a hallmark of ageing that may benefit from additional and complementary models, beyond the mouse 67 68 (Forsyth, Wright, & Shay, 2002; Gomes et al., 2011; Sullivan et al., 2021). Once such model is 69 the zebrafish that, like humans, age and die in a telomerase-dependent manner (Anchelin et 70 al., 2013; Carneiro, de Castro, & Ferreira, 2016; Madalena C Carneiro et al., 2016; Henriques, 71 Carneiro, Tenente, Jacinto, & Ferreira, 2013; Henriques & Ferreira, 2012). Restricted 72 telomerase expression and function are key determinants of natural ageing in humans, 73 underpinning multiple age-related diseases (Blackburn, Epel, & Lin, 2015). However, the role 74 and the dynamics of telomerase-dependent changes that may contribute to tissue-specific 75 ageing are still poorly understood. This is partially due to the fact that telomerase appears 76 to have multiple functions in the cell, that go beyond the maintenance of telomere length, 77 recently reviewed elsewhere (Segal-Bendirdjian & Geli, 2019).

78 Telomerase is best known for its telomere-dependent function (i.e. canonical 79 functions), acting as a reverse transcriptase, maintaining telomere length through its 80 catalytic domain (TERT protein) and RNA template (TERC) (Greider & Blackburn, 1985). 81 Telomeres are $(TTAGGG)_n$ DNA repeats that together with a complex of proteins (known as 82 Shelterin) create a "cap-like" structure at the end of linear chromosomes (de Lange, 2004). 83 preventing the ends of linear chromosomes from being recognised as deleterious DNA 84 double strand breaks (Ferreira, Miller, & Cooper, 2004). However, in humans, due to time-85 and cell-specific-limited telomerase expression, telomeres shorten with ageing, leading to 86 proliferative exhaustion and replicative senescence (Bodnar, 1998; d'Adda di Fagagna et al., 87 2003). Importantly, there is accumulation of cellular senescence with ageing in humans 88 (Dimri et al., 1995) and senescence has been linked to several age-associated diseases 89 (Ovadya & Krizhanovsky, 2014). Additionally, short telomeres themselves can lead to de-90 regulated gene expression, particularly in genes near the chromosome ends, due to loss of 91 the "telomere positioning effect" (TPE), which is known to regulate gene expression of 92 genes at least up to 10MB away from the chromosome ends (Robin et al., 2014).

93 Growing evidence now suggests that telomerase also has activity independent of its 94 action at telomeres, known as non-canonical (Goodman & Jain, 2011; Romaniuk et al., 2018; 95 Segal-Bendirdjian & Geli, 2019; Sung, Ali, & Lee, 2014). In the nucleus, these non-canonical 96 functions include transcriptional regulation of genes involved in inflammation, including 97 nuclear factor kappa B (NFkB) and tumour necrosis factor alpha (TNF α) (Deacon & Knox, 98 2018; Ghosh et al., 2012; Mattiussi, Tilman, Lenglez, & Decottignies, 2012), as well as genes 99 involved in cell proliferation (Choi et al., 2008; Sarin et al., 2005) and cell survival (Cao, Li, 100 Deb, & Liu, 2002; Rahman, Latonen, & Wiman, 2005). Telomerase can also translocate to 101 the mitochondria, where it has been shown to play a protective role against DNA damage 102 and oxidative stress (Ahmed et al., 2008; Haendeler et al., 2009).

103 As tissues with high cellular turnover present accelerated telomere erosion (Bodnar, 104 1998; H. W. Lee et al., 1998), it is reasonable to think that telomerase functions are likely to 105 primarily affect highly proliferative tissues. Accordingly, premature accumulation of critically 106 short telomeres has been identified in high proliferative tissues such as the gut, in tert-107 deficient animal models. Nonetheless, the role of telomerase and telomeres is not restricted 108 to highly proliferative tissues. In the brain, considered a predominately post-mitotic tissue, 109 telomerase has been shown to have a protective role against excitotoxicity (Eitan et al., 110 2012), oxidative stress (Spilsbury, Miwa, Attems, & Saretzki, 2015), and neuronal death (J. 111 Lee et al., 2010), all involved in neurodegenerative diseases. Studies in late-generation 112 telomerase-deficient mice have suggested that limited telomerase expression is associated with premature accumulation of senescence-associated markers in different cell 113 114 populations including Purkinje neurons, cortical neurons and microglia (De Felice et al., 115 2014; Jaskelioff et al., 2011; Jurk et al., 2012; Raj et al., 2015). Telomerase is therefore a 116 promising target to promote healthy ageing in multiple tissues and so the identification of 117 mechanisms driving telomerase-dependent ageing could enable the identification of 118 targeted therapies to improve healthspan.

In this study, we aimed to determine the telomerase-dependent and -independent transcriptomic changes and their kinetics occurring during ageing, in both brain and gut within the same individuals. We hypothesised that this would allow us to identify key ageassociated genes and pathways that become prematurely de-regulated in both or either tissue, providing key insights into the early stages of ageing in these tissues and likely

124 interactions. We further hypothesised that this may highlight potential common 125 telomerase-dependent and -independent therapeutic targets for early intervention aimed at 126 preventing age-associated dysfunction in both tissues. To address these questions, we 127 performed RNA sequencing in whole tissues (gut and brain) of WT fish (2, 9, 22 and 35 months of age) alongside telomerase mutant fish (tert^{-/-}) (2, 9 and 22 months of age). tert^{-/-} 128 129 zebrafish, extensively characterised elsewhere (Anchelin et al., 2013; Madalena C Carneiro 130 et al., 2016; Henriques et al., 2013), display no telomerase activity and have significantly 131 shorter telomeres from birth, consequently ageing and dying prematurely. Ageing is usually 132 described as a time-dependent change in tissue homeostasis, that increases the probability 133 of disease and death (Hayflick, 2007). However, whether the genes and pathways driving or 134 accompanying these time-dependent changes are also consistently changing in a time-135 specific manner, remains unresolved (Rando & Wyss-Coray, 2021). We therefore decided to 136 combine a time-series analysis (STEM), which allowed the identification of genes and 137 pathways that are consistently up or down-regulated over-time, with the more traditional 138 differential gene expression (DEGs) analysis between young and old animals. This combined 139 strategy allowed the identification of genes that change in a monotonic, time-dependent 140 manner (STEM), versus genes that change at specific stages of life (DEGs).

141 We show that although the gut and brain have distinct transcriptomic signatures of 142 ageing, both tissues display hallmarks of ageing as early as 9 months in the WT zebrafish. 143 Importantly, telomerase depletion accelerates the appearance of such hallmarks in both gut 144 and brain. In particular, we identify stem cell exhaustion as the common principal hallmark 145 of ageing at the early stages of ageing, in both tissues. Further, we identify altered 146 intercellular communication, in which immunity and inflammation play a central role, as the 147 main telomerase-dependent hallmark of ageing common between the gut and brain. Finally, 148 we conclude that the gut displays telomerase-dependent hallmarks of ageing at an earlier 149 age than the brain and that these include changes in several key genes that have also been 150 included in the GenAge database, a benchmark curated database that included genes 151 involved in ageing across different organisms, including in humans (Tacutu et al., 2018). 152 Finally, we identify 7 key gene changes common between the gut and brain at the early 153 stages of ageing, namely ccnb1, kif2c, serpinh1a, temem37, si:ch211-5k11.8, cfap45 and 154 eif4ebp31.

155

156 **2 RESULTS**

157 2.1 Identification of monotonic, time-dependent gene signatures and process 158 changes with ageing in the zebrafish gut and brain

159 In order to identify telomerase-dependent and -independent transcriptional 160 signatures of ageing in the zebrafish gut and brain, we performed RNA-Sequencing of whole tissues, throughout the lifespan of WT and telomerase-deficient (tert^{-,-}) fish (Fig. 1 and 161 source data). While $tert^{-/-}$ fish have a lifespan of c. 12-20 months, WT fish typically die 162 between c. 36-42 months of age (Madalena C Carneiro et al., 2016). The data shown here 163 164 include 4 age-groups of WT (2, 9, 22 and >30 months), corresponding to young, adult, median lifespan and old. As the *tert*^{-/-} fish have a shorter lifespan compared with their WT 165 166 siblings, the data include 3 age-groups of telomerase-deficient fish (2, 9, and 22 months), 167 which correspond to young, medium lifespan and old. Each group has a sample size of 3 168 animals and the brains and guts from within the same groups of animals were used (Fig 1A). 169 The reads were aligned to the latest zebrafish genome build GRCz11 (Lawson et al., 2020) 170 and resulted in uniquely mapped read percentages ranging from 92.1% to 94.4%, which is a 171 readout of good quality (Lawson et al., 2020). All samples had at least 10 million uniquely 172 mapped reads, except G11 which had around 8 million. Further quality control, using a 173 principal component analysis (PCA) including all the samples, revealed that one of the gut 174 samples clustered with the brain samples, and not with the gut samples (Fig. 1B). This was 175 considered to be a technical error and this sample (G7) was therefore excluded from further 176 analysis. To analyse the overall impact of the genotype and age on transcriptomic 177 regulation, we then performed a PCA in the samples from the gut and brain, separately. We 178 further observed that some samples cluster *per* age, but there are some genotypes that 179 separate quite distinctly, despite being of the same age (Fig 1C and D). As an example, the WT and tert^{-/-} gut samples are guite distinct, and the tert^{-/-} 2-month samples cluster closely 180 181 to the WT at 9 months than the WT at 2 months (Fig 1C), providing the first hint of an 182 acceleration of the ageing transcriptomic profile in the tert /-. We found that there was 183 higher variability in the gut than in the brain, between samples within each age group (Fig 184 **1C and D**). A summary of the number of significant differentially expressed genes (DEGs) in 185 all samples is represented in Fig. 1E. How these DEGs relate to which other, how many

186 overlap and how many are in common or accelerated in the absence of telomerase (tert^{-/-}),

187 will be explored later in the manuscript.

188 To identify genes and pathways that are consistently up- or down-regulated in a time-189 dependent manner in natural ageing in the zebrafish gut and brain, we grouped the genes 190 into temporal expression profiles, by time-series analysis using Short Time-series Expression 191 Miner (STEM) software (Ernst & Bar-Joseph, 2006). To determine whether the temporal 192 profiles were associated with specific biological processes and pathways, pathway over-193 representation analysis (ORA) were performed for the genes assigned to the significant 194 STEM profiles. Enrichments of GO Biological Process (GOBP), GO Molecular Function 195 (GOMF), GO Cellular Compartment (GOCC), Kyoto Encyclopedia of Genes and Genomes 196 (KEGG) terms and REACTOME pathway terms were therefore analysed for each profile. 197 Time-series analysis of WT gut identified 9 different profiles, with 2 of them containing up-198 regulated genes (profiles 7 and 21. Total of 523 genes), and 7 containing down-regulated 199 genes (profiles 32, 31, 23, 9, 12, 22 and 34. Total of 11,594 genes) (Fig. 2A1 and source data). Interestingly, in the *tert*^{-/-} gut, all the profiles identified by time-series analysis contain 200 201 up-regulated genes (profiles 8, 6, 15 13, 12 and 11. Total of 10,317 genes) (Fig. 2A2 and 202 source data). Enrichment analysis showed that the profiles containing up-regulated genes 203 are associated with immune response, while profiles containing down-regulated genes are 204 largely associated with proliferation, cellular response to DNA damage, and DNA damage repair, in both WT and tert^{-/-} gut (Fig. 2 A1.1, 1.2 and A2.1). To help contextualise our 205 206 analysis, we performed a further classification of enriched processes according to the 207 hallmarks of ageing, which have been previously identified (Lemoine, 2021; López-Otín et 208 al., 2013). This classification further strengthened the observation that the up-regulated 209 profiles include genes mostly involved in altered intercellular communication, in which 210 immunity and inflammation play a key role, whereas the down-regulated profiles identify 211 stem cell exhaustion and genomic instability as the main hallmarks of ageing, to which the 212 genes affecting proliferation, DNA damage and repair are likely to contribute (Fig. 2 A1.1, 213 1.2 and A2.1 and source data).

In the WT brain, we identified 9 temporal profiles, 6 of them including up-regulated genes (profiles 42, 29, 40, 30, 21 and 48. Total of 7,230 genes), and 3 including downregulated genes (profiles 1, 12 and 26. Total of 561 genes) (**Fig. 2B1 and source data**). In the

217 $tert^{-1}$ brain, time-series analysis revealed 6 different profiles. Profiles 4, 0 and 3 containing 218 down-regulated genes (total of 5,374 genes) and profiles 15, 12 and 11 containing up-219 regulated genes (total of 1,155 genes) (Fig. 2B2 and source data). As in the gut, up-220 regulated profiles reveal genes mostly involved in immune regulation and inflammation and 221 down-regulated profiles are mostly involved in cell cycle, genome stability and DNA damage 222 responses, in both genotypes This is further highlighted when placed into context by the 223 analysis based on the hallmarks of ageing, where up-regulated profiles identify altered 224 intercellular communication whereas the down-regulated ones identify stem cell exhaustion 225 as the main hallmarks affected (Fig 2 B1.1, 1.2; B2.1, 2.22 and source data).

In summary, STEM analysis and enrichment pathways in both gut and brain ageing show a general trend towards up-regulation of genes involved in immune response and down-regulation of genes involved in cell cycle, DNA damage and repair. This general trend is recapitulated in the absence of telomerase. These mechanisms are all known contributors to the well-described altered intercellular communication, genome stability and stem cell exhaustion hallmarks of ageing, respectively (Lemoine, 2021; López-Otín et al., 2013).

232

233 2.2 Comparing the hallmarks of ageing over time, in WT and *tert^{-/-}* zebrafish gut and 234 brain

235 More than just identifying the signatures of natural WT ageing in the gut and brain 236 and identifying telomerase-dependent and -independent changes, we wanted to understand if there were particular changes occurring before others, and how their kinetics 237 238 compared between the gut and the brain. In order to contextualise our analysis in the light 239 of the well-described hallmarks of ageing, (Lemoine, 2021; López-Otín et al., 2013), we used 240 the main hallmarks of ageing as headers in which we could group the different changes in 241 processes that were enriched from both the STEM profiles and all DEGs. This allowed us to 242 compare the effects of age, genotype and tissue on the evolution of the key hallmarks of 243 ageing. When we combine all the gene changes (up- and down-regulated) and associated 244 biological processes affected (Fig. 3 and source data), we observe distinct tissue-specific 245 signatures of ageing, namely in the gut (Fig. 3A) and brain (Fig. 3B), particularly when 246 considering the time-series analysis on its own (STEM) (Fig 3. A1 and B1). Whereas ageing in 247 the WT gut seems to be predominantly affected by the de-regulated nutrient sensing

248 initially, the brain displays stem cell exhaustion as the main hallmark affected at the early 249 age of 9 months (Fig. 3A1, B1, respectively). When we combine the different analysis, 250 though (Fig. 3A3 and B3), both gut and brain have stem cell loss as the main hallmark of 251 ageing associated with the enriched processes found at 9 months of age, highlighting this as 252 a potential contributor to the early stages of ageing in both tissues. Towards the end of life, 253 both gut and brain have altered intercellular communication as the main hallmark of ageing identified. Importantly, independently of the analysis, the *tert^{-/-}* zebrafish show accelerated 254 hallmarks of ageing. In specific, a tert^{-/-} 2-month-old gut profile is very similar to a WT 35 gut 255 month profile (Fig. 3A). In the brain, at 2 months of age the *tert*^{-/-} mutant also displays some 256 257 of the hallmarks of ageing that also become altered in WT ageing at later ages, particularly altered intercellular communication, but the $tert^{-/-}$ brain becomes much more similar to the 258 259 aged WT from the age of 9 months onwards (Fig. 3B). This suggests that hallmarks of ageing 260 accelerated in the absence of telomerase are developing earlier in the gut than in the brain.

261 Finally, when we look at the overall number of gene expression changes, i.e, not just 262 the ones associated with the hallmarks of ageing, we observe that there is a general 263 increase in the number of changes in gene expression with ageing (Fig 3 A4 and B4). 264 However, this increase does not appear to be linear. In specific, in the gut, the number of 265 DEGs is fairly low until 9 months of age, after which there seems to be an inflexion point and 266 the number of DEGs increases up to 5-fold in the oldest WT (>35 months) and 3 fold in the oldest $tert^{-/-}$ (22 months). In the brain, there seems to be a more gradual, consistent 267 increase over time, both in WT and $tert^{-1}$. This increase in number of DEGs with ageing may 268 269 be a consequence of the known de-regulation of gene expression with ageing due to de-270 repression of heterochromatin and/or changes in epigenetic markers.

271

272 **2.3** When does a *tert*^{-/-} resemble an aged WT the most?

11 is of note that, even though the *tert*^{-/-} accelerates hallmarks of ageing in the gut and 224 brain, the set of genes identified are not necessarily the same as in WT ageing. In specific, 275 looking at the genes at the early stages of ageing, the most significant hallmark of ageing 276 shared between the 2 months old *tert*^{-/-} and WT 9 months gut is nutrient sensing, and only 2 277 up-regulated genes are in common, namely *lipca (Hepatic triacylglycerol lipase precursor)* 278 and *pla1a (phospholipase 1a)*. Interestingly, hepatic lipase has been involved in

279 atherogenesis (Santamarina-Fojo, Gonzalez-Navarro, Freeman, Wagner, & Nong, 2004), and 280 age-related macular degeneration (Neale et al., 2010), diseases that have been 281 hypothesised to have a parallel response to tissue injury induced by multiple factors, 282 including impaired immune responses, and oxidative stress (Neale et al., 2010). In line with 283 the general de-regulated inflammatory response we see in the gut, with ageing (Figs. 2 and 284 **3** and source data), phospholipase 1a has been reported to be up-regulated in inflamed gut 285 tissue of Crohn's disease patients (Hong et al., 2017) and has a complex role in the 286 regulation of immunity and inflammation (recently reviewed in (Zhao, Hasse, & Bourgoin, 2021)). In the brain, there are no genes in common between the 2 months old $tert^{-7}$ and WT 287 288 9 months associated with altered intercellular communication or genomic instability, the 2 289 main hallmarks of ageing shared between the genotypes at the early stages of ageing. Following this type of analysis, we could then ask at what age does the *tert*^{-/-} best mimic 290 291 naturally aged WT, at the level of gene expression from the shared hallmarks of ageing and 292 these "other" genes we found to be associated with ageing but not obviously associated 293 with the described hallmarks of ageing. We therefore analysed the overlap between all DEGs identified in the old WT (35 months) and the $tert^{-1/2}$ at the different aged (2, 9 and 22) 294 295 months), using Venn diagrams created using the Venny 2.1 online platform (Oliveros, 2007-2015) to ask this question. We observe that the 2 months old $tert^{-/-}$ has the most genes 296 297 shared with the 35 months old WT gut (59 genes in common) (Fig. 4 A1 and Supp. Fig. 1) whereas it is the *tert^{-/-}* at 22 months that has the most genes shared with the 35 months old 298 299 brain (112 genes in common) (Fig. 4 B1 and Supp. Fig.1). This similarity between the 2 month and 22 months *tert^{-/-}*, with the gut and brain, respectively, is also apparent when we 300 just look at the pattern of the main hallmarks of ageing accelerated in the tert $^{-1}$ (Fig. 3, 4 301 302 and Graphical Abstract). Together, these data suggest that the gut is displaying telomerase-303 dependent hallmarks of ageing at an earlier age than the brain, consistent with what would 304 be expected for a high versus low proliferative tissue. Of relevance, In the brain, of the genes de-regulated in the tert^{-/-} 22 month that are in common with the WT aged at 35 305 306 months ccna2 (cyclin a2), cdk6 (cyclin-dependent kinase 6), chek1 (checkpoint kinase 1), 307 mad2l1(Mitotic spindle assembly checkpoint protein mad2a), tacc3 (Transforming, acidic coiled-coil containing protein 3), top2a (DNA topoisomerase ii alpha) and mcm2 (DNA 308 309 helicase, MCM2 minichromosome maintenance deficient 2) have also been included in the 310 Human Ageing Resources databases (Tacutu et al., 2018) and are mostly located within the

311 same cluster (green), identified using *k*-means clustering in STRING analysis (Szklarczyk et 312 al., 2021) (Fig 4 and source data). Of the proteins encoded by these genes, chek1 313 (Poehlmann et al., 2011), cdk6 (Morris, Hepburn, & Wynford-Thomas, 2002), mad2l1 314 (Lentini, Barra, Schillaci, & Di Leonardo, 2012), tacc3 (Schmidt et al., 2010)have been 315 reported to be involved in cellular senescence, . Additionally, Top2a has also been involved in neuron proliferation (Watt & Hickson, 1994) and mcm2 depletion in mice leads to 316 317 decreased proliferation in various tissue stem cell progenitors (Pruitt, Bailey, & Freeland, 318 2007).

- 319
- 320

2.4 Analysis of telomerase (*tert*)-dependent gene changes of old age

321 Once we identified what WT ageing looked like at the level of time-dependent gene 322 expression changes over the life-course and the main biological processes affected, we 323 sought to determine how much of these were likely to be telomerase-dependent. If a gene 324 expression change or a biological process alteration is accelerated in the absence of telomerase (i.e becomes significant at an earlier age in the tert^{-/}), we consider it to be 325 326 telomerase dependent, as has been described before (Madalena C Carneiro et al., 2016; 327 Henriques et al., 2013). Conversely, if none of these pre-requisites are met, we consider the 328 gene/process alteration to be telomerase-independent. With this in mind we performed 329 Venn diagrams (Oliveros, 2007-2015) to identify the telomerase-dependent significant gene 330 alterations of old age, i.e., DEGs present in the WT at 35months old, when compared with 331 the WT young control. For this, we used the DEGs identified in the STEM profiles (i.e., genes 332 that change monotonically, in a time-dependent manner), combined with the more 333 traditional DEG analysis, which include all gene changes, whether they change consistently 334 in a time-dependent manner across the lifecourse or not (called "all DEGs" from hereafter) 335 (Fig. 5 and source data). From these analyses, we identified 50 significant DEGs (out of 491; c. 10%) (present in old age (WT 35 months) that are prematurely de-regulated in the tert $^{-/-}$ 336 337 gut (Fig. 5 A) and 100 genes (out of 428; c.23%) to be prematurely de-regulated in the tert^{-/-} 338 brain (Fig. 5 B). Importantly, most of these genes are directly or indirectly involved in known 339 hallmarks of ageing (Lemoine, 2021; López-Otín et al., 2013), namely altered intercellular 340 communication (including immunity, inflammation, extra-cellular matrix), genome stability 341 (including DNA replication and repair), stem cell exhaustion and mitochondrial dysfunction,

342 as was also evident in Fig 3 and now further detailed below. In the gut, telomerase-343 dependent gene expression changes with ageing include up-regulation of genes involved in 344 immune response, such as tlr18 (Toll-like receptor 18 precursor), sytl1 (Synaptotagmin-like 345 protein 1) and down-regulation of genes involved in metabolism, such as cyp8b1 (Bertaggia 346 et al., 2017) (Cytochrome P450, family 8, subfamily B, polypeptide 1) and iqfbp2a (Insulinlike growth factor-binding protein 2). Within the down-regulated genes, there are other 347 348 well-known genes such as sox6 (involved in stem cell function), vav3b (involved in wound 349 healing), (Fig 5 and source data). Additionally, we identify a number of non-annotated 350 genes, which may represent novel tert-dependent genes, that would require further 351 investigation. Importantly, amongst these *tert*-dependent genes of "old age", there are a 352 number of homolog or closely related genes which have been previously identified in ageing 353 datasets (Tacutu et al., 2018) such as several igfbps (insulin-like growth factor binding 354 proteins), iqf (insulin growth factor), mapks (mitogen-activated protein kinases), tlr3 (toll-355 like receptor 3) and nrg1 (neuroregulin 1).

356 In the brain, tert-dependent ageing gene expression changes include mostly down-357 regulation of genes involved in cell cycle or neurogenesis, such as aurkb (aurora kinase B), 358 chek1 (checkpoint kinase 1), ccnb1 (cyclin b1), cdk2 (cyclin-dependent kinase 2) and neurod4 359 (neuronal differentiation 4), dld (delta d), nog1 (noggin 1), respectively, as well as genome 360 stability and DNA repair, such as rad54l (rad54 like), mcm5 (minichromosome maintenance 361 complex component 5) and smc4 (structural maintenance of chromosomes 4). Up-regulated 362 genes are mostly involved in immune response or inflammation, such as cxcl18b (chemokine 363 (C-X-C motif) ligand 18b), mhc2a (major histocompatibility complex class II integral 364 membrane alpha chain gene), socs1a (suppressor of cytokine signaling 1a), irf8 (interferon 365 regulatory factor 8) and csf1b (colony stimulating factor 1b (macrophage)). Of note, 366 amongst these tert-dependent DEGs of old age, we identified dre-mir-29b-1, which encodes 367 for a regulatory micro RNA 29 (mir29), widely described to be up-regulated in ageing, in 368 response to DNA damage, alongside P53 (Ugalde et al., 2011) and can act as a protective 369 response, limiting excessive iron-exposure and damage in neurons (Ripa et al., 2017). As for 370 the gut, there are a number of homolog or closely related genes which have been previously 371 identified in ageing (Tacutu et al., 2018), such as chek1, mad2l1 (MAD2 mitotic arrest 372 deficient-like 1 (yeast)), dl3 (delta like 3, nog (noggin), ifnb1 (interferon beta), socs2

373 (suppressor of cytokine signaling 2), amongst many others, which can be found in the 374 GeneAge database.

375 Since it is known that short telomeres themselves can lead to de-regulated gene 376 expression, particularly in genes near the chromosome ends due to loss of the "telomere 377 positioning effect" (TPE) (Robin et al., 2014), we proceeded to map the genes identified to 378 be de-regulated prematurely in the absence of telomerase to their chromosome location, 379 with the aim of determining whether they located to within up to 10MB of either of the 380 telomeric ends (Fig. 5C1). We found that whereas most tert-dependent gene changes of old 381 age do not locate to the end of chromosomes, in both gut and brain (Fig. 5C2 and C3). 382 However, there are significantly more *tert*-dependent genes located at the end of 383 chromosome in the gut than in the brain, (Fig. 5 C4). This is consistent with gut being a more 384 proliferative tissue, where telomeres are likely to shorten more, which would be particularly 385 exacerbated in the absence of *tert*, leading to a higher probability of TPE contributing to 386 gene transcription alterations.

387

2.5 Transcriptional changes in common between the gut and brain at the early and 388 late stages of ageing

389 Even though the aged phenotype is something usually observed late in life, the 390 underlying molecular and cellular mechanism driving these changes can, arguably, start 391 from the moment you are born (Gladyshev, 2021). To understand what significant 392 transcriptional changes are taking place in the gut and brain and, in particular, in common 393 between the gut and brain, we compared all DEGs and STEM DEGs in these tissues at the 394 earliest time point we detect significant changes (9 months) and at the late stages of ageing, 395 i.e, at the latest time point analysed of 35 months (Fig 6, Supp Fig 3 and source data). We 396 identified 7 gene changes in common between WT gut and brain at the early stages of 397 ageing (9 months of age). In specific, we identified 5 down-regulated genes, namely ccnb1, 398 kif2c, serpinh1a, temem37 and si:ch211-5k11.8, and 2 up-regulated genes, namely cfap45 399 and *eif4ebp31*. Of these, ccnb1 (*G2/mitotic-specific cyclin-B1*) and *kif2c* (*Kinesin-like protein*) 400 are both proteins involved in cell-cycle progression. Whereas *Ccnb1* has been reported to be 401 involved in normal stem-cell/progenitor maintenance in the brain (Hagey et al., 2020) and 402 gut (Basak et al., 2014); kif2c can act as a DNA damage repair protein, and, accordingly, its 403 depletion leads to significant accumulation of DNA damage (Zhu et al., 2020). STRING

analysis suggests these proteins are likely to be co-expressed in a variety of organisms
including zebrafish and humans, highlighting potential functional links (Fig 6 B and source
data). The downregulated *elf4ebp3l* gene (Eukaryotic translation initiation factor 4E-binding
protein 3-like) encodes a repressor of translation and is inhibited in response to TORC1
(mammalian target of rapamycin complex 1) (Boutouja, Stiehm, & Platta, 2019).

409 At "late stages" of ageing, we identified 23 gene changes in common between the gut 410 and brain. Of these, STRING analysis identified 2 main protein network interactions, namely 411 a link between cd59 and cd99l2 and link between il2rb and b2m. In specific, cd59 and cd99l, 412 which our data show to be downregulated in old age, in both gut and brain, have both been 413 cited in the literature as markers of new-born neurons and oligodendrocyte progenitor cells, 414 respectively, in a single-cell transcriptomic analysis of the adult zebrafish brain (Lange et al., 415 2020). In the gut, cd59, also known as protectin, has been shown to be downregulated in 416 the gut epithelium of ulcerative colitis and Chron's disease patients and thought to render 417 the tissue susceptible to inflammatory damage (Scheinin et al., 1999). cd99 has been shown 418 to be a key molecule in modulating monocyte migration through endothelial junctions 419 (Schenkel, Mamdouh, Chen, Liebman, & Muller, 2002) and monocyte differentiation to 420 macrophages is known to be triggered by endothelial transmigration (Gerhardt & Ley, 2015; 421 Li et al., 2020), including in the brain via migration through the blood-brain barrier (Ifergan 422 et al., 2007). Il2rb and b2m, which our data show to be up-regulated in both gut and brain at 423 the "late stages" of ageing, are key molecules involved in adaptive and innate immune 424 function. Whilst the IL2R beta is important for T-cell mitogenic response to IL-2, the b2m 425 (Beta-2-microglobulin) protein is a component of the major histocompatibility complex I 426 (MHCI), involved in antigen presentation. Intriguingly, b2m has been shown to be present in 427 a soluble free-form, and has been found to be systemically increased with ageing in humans 428 and in individuals with neurodegenerative diseases (Smith et al., 2015). Importantly, 429 heterochronic parabiosis experiments between young and old mice suggest that increased 430 b2m with ageing leads to cognitive impairment and neurodegeneration, and has hence been 431 identified as a systemic pro-ageing factor (Smith et al., 2015). Additionally, increased IL2 432 receptor expression has been shown to contribute to CD4 differentiation and exhaustion of 433 their naïve pool and therefore ability to respond to infection with ageing, in humans 434 (Pekalski et al., 2013; H. Zhang, Weyand, & Goronzy, 2021).

435

436 **3 DISCUSSION**

In this study, we used RNA sequencing analysis to determine the kinetics of telomerase-dependent and -independent transcriptomic changes occurring during WT ageing in brain and gut tissues, using the zebrafish as a model. We hypothesised that this would allow us to identify key age-associated genes and pathways that become prematurely de-regulated in both or either tissue, providing key insights into the early stages of ageing in each tissue and highlight potential interactions.

- 443
- 444

3.1 STEM versus all DEGs analysis

445 Ageing can be described as a time-dependent change in tissue homeostasis, that 446 increases the probability of disease and death (Hayflick, 2007). Whether the genes and 447 pathways driving these time-dependent changes are also consistently changing in a time-448 specific manner, remains unresolved. To account for both possibilities, we performed a 449 time-series analysis (STEM) and then combined this with the more traditional differential 450 gene expression (DEGs) analysis between young and old animals. Even though the gene 451 changes identified with the STEM analysis were also picked up by the traditional DEGs 452 analysis, the STEM analysis provided a much tighter, restrictive view of the transcriptomic 453 signatures of ageing. In the gut, particularly, the main hallmarks of ageing identified using 454 the STEM analysis are quite different from the ones using the traditional all DEGs. In 455 particular, STEM analysis identifies mitochondrial dysfunction and de-regulated nutrient 456 sensing as the main hallmarks of WT ageing at the earlier stages of ageing in the gut (WT 9 457 months), whereas all DEGs analysis identified stem cell dysfunction as the principal hallmark 458 de-regulated at that age. The significance of these differences is difficult to judge, but it can 459 suggest that changes in genes affecting mitochondrial function and nutrient sensing have a 460 mostly monotonic trajectory in gut ageing, whereas the ones involved in stem cell 461 maintenance can have different behaviours at different times throughout life. Nevertheless, this difference was not observed in the WT brain or in the *tert^{-/-}* ageing, where STEM and all 462 463 DEGs analysis led to very similar conclusions regarding the identity or kinetics of the main 464 hallmarks of ageing affected. Importantly, the kinetics of gene changes and processes 465 identified in our data match very well to the phenotypes of ageing previously reported in the tert^{-/-} and WT ageing. In particular, in the gut, impaired cell proliferation in the gut is one 466

467 of the first *tert*-dependent phenotypes of ageing identified, followed by cellular senescence 468 and inflammation later in life (Madalena C Carneiro et al., 2016; Henriques et al., 2013). 469 Moreover, the recently reported *tert*-dependent changes in macrophage immune activation 470 and increased gut permeability (Pam S. Ellis, 2022) are consistent with key tert-dependent 471 gene changes of old age identified here. An example of such genes are cd99, potentially 472 involved in macrophage differentiation via trans-endothelial migration (Gerhardt & Ley, 473 2015; Li et al., 2020; Schenkel et al., 2002); and *cldn5a* (claudin 5), involved in tight-junctions 474 (Lu, Ding, Lu, & Chen, 2013).

- 475
- 476

3.2 Main hallmarks of ageing

477 A simplistic prediction of how the kinetics of the hallmarks of ageing would behave 478 over the lifecourse could be that, at early ages, we would detect more changes affecting the 479 primary hallmarks of ageing, i.e, the "causes of damage", namely genomic instability, 480 telomere attrition, epigenetic alterations and loss of proteostasis. At the last stages, one 481 could predict that we would detect more significant changes in the integrative hallmarks, 482 i.e, the "culprits of the phenotype", namely stem cell exhaustion and altered intercellular 483 communication, of which inflammation is a key component, as described in (López-Otín et 484 al., 2013). However, either separately or combined, neither STEM nor all DEGs analysis show 485 this. Our combined analyses show that most of the gene changes occurring at the early 486 stages of WT ageing are observed from 9 months and are mostly involved in stem cell 487 maintenance, in both gut and brain. One potential explanation for this observation, is that 488 our qualitative analysis was not able to distinguish between such hallmarks or is under-489 estimating the primary hallmarks. Another explanation comes from the relatively low 490 sample size used for each time-point and the heterogeneity of individuals physiology 491 amongst each population. As recently showed in (Dambroise et al., 2016), zebrafish age 492 following the two-phase model first proposed flies (Tricoire & Rera, 2015), based on the 493 age-related intestinal permeability assessed using the Smurf assay they previously described 494 (Rera, Clark, & Walker, 2012). Moreover, we have recently shown that gut permeability with ageing is accelerated in the absence of telomerase (*tert^{-/-})* (Pam S. Ellis, 2022). Following this 495 496 model and considering the longevity curve from the population we sampled, the proportion 497 of Smurfs might have been approximately, <10% at 2 month, 25% at 9 month, 50% at 22

498 months and >80% at 35 months. The chances to have selected at least 1 Smurf by accident 499 are approximately 27% at 2 months, 57% at 9, 86% at 22 and 99% at 35%. This could 500 contribute to the heterogeneity or potential bias towards having more of a specific ageing 501 signature over another (smurf versus non-smiurf).

502 At first glance, it is surprising that telomere dysfunction is not picked up as a main 503 hallmark of ageing in the telomerase mutant model. However, this may simply be the 504 reflection of the number of genes that have been directly implicated in each of these 505 hallmarks. In specific, there are a lot less genes that one would identify as directly involved 506 in telomere dysfunction, when compared to stem cell dysfunction, for example. The main 507 culprits for telomere dysfunction would be telomerase and the shelterin components, 508 whereas for stem cell dysfunction, all the cell cycle and DNA damage repair proteins can 509 play a role. For example, *chek1*, *fxr1 and daxx*, which are all de-regulated in the old WT gut 510 and brain (chek1) (see source data), have all been identified as potential regulators of 511 telomeres (Nersisyan et al., 2019). Yet, because that is not the main function one would 512 attribute to such genes, these would have been missed as part of the telomere dysfunction 513 hallmark of ageing. Additionally, it is not possible to distinguish from just looking at lists of 514 DEGs if such gene was de-regulated due to telomere dysfunction in the first place, or if its 515 dysfunction will affect telomere function indirectly. These are some of the considerations 516 that highlight the complexity of ageing and the non-linearity of how the hallmarks of ageing 517 may drive ageing as well as each other and it is important to have them in mind when interpreting our analyses. Nevertheless, at the later stages of WT ageing (>35 months), 518 519 altered intercellular communication, a previously described "culprit of the phenotype" is 520 indeed the most significant hallmark of ageing detected in common between the gut and 521 brain. Finally, in terms of the progression hallmarks of ageing, different hallmarks may play 522 more prominent roles at specific times of life, and may be replaced by others at other times, 523 explaining why we don't necessarily always see a linear accumulation of the different 524 hallmarks of ageing over-time. This is particularly evident in the gut, where stem cell 525 exhaustion is the main hallmark identified by the combined STEM and ALL DEGs analysis, 526 and it is barely represented at the later stages of life. In the brain, the progression seems to 527 be more linear, though, and most hallmarks present at the early time point of 9 months 528 remain until old age, when other hallmarks are further added.

529

530

3.3 Telomerase- and/or telomere-dependent changes with ageing

531 If a gene expression changes or a biological process alteration is accelerated in the absence of telomerase (i.e becomes significant at an earlier age in the tert^{-/-}), we considered 532 it to be telomerase dependent. If we take a step back and look at the processes and, in turn, 533 534 hallmarks of ageing affected in the absence of telomerase, it becomes clear that the tert - is 535 indeed accelerating many of these changes. It is particularly evident when we look at the pie 536 charts depicting the different hallmarks of ageing affected at each time point in WT versus tert^{-/-}. Here, it is striking how a 2-month-old tert^{-/-} gut resembles a WT gut at the old age of 537 35 months, and how a 9- and 22-month-old $tert^{-/-}$ brain resembles an old WT brain at 35 538 539 months. This is further highlighted by the further comparison we performed, where we asked at which age does the tert^{-/-} share more gene expression changes with the WT old 540 (>35 months). In this analysis, we show that the 2 months old $tert^{-/-}$ has the most gene 541 expression changes shared with the 35 months old WT gut, whereas it is the tert $^{-/}$ at 22 542 543 months that has the most gene expression changes shared with the 35 months old WT 544 brain. This suggests that the gut is displaying telomerase-dependent hallmarks of ageing at 545 an earlier age than the brain, which is consistent with what would be expected for a high 546 versus low proliferative tissue. Accordingly, when looking at the specific gene changes 547 accelerated in the absence of tert, i.e, we identify significantly more tert-dependent genes 548 located near the ends of chromosomes in the gut than in the brain, and therefore more 549 liable to have been altered due to telomere shortening. In the future, it would be insightful 550 to test how many of the *tert*-dependent gene changes occur due to non-canonical functions 551 of telomerase involved in transcriptional regulation.

552 Nevertheless, *tert*-dependent gene changes in both tissues are still a minority, serving 553 as a reminder that "all roads lead to Rome", and one should exercise caution when trying to 554 identify genes influencing the natural process of ageing using premature ageing models. It is 555 not necessarily the same genes influencing the processes of ageing in these models, even if 556 the processes and phenotypes are accelerated. We should also have this in mind when we 557 compare the sets of genes identified in this study with those previously identified as 558 implicated in human ageing, and not necessarily be surprised if only a small percentage of 559 those are shared. One could argue that it is more important that the processes are shared.

560 Nevertheless, we do identify many gene changes shared between zebrafish and human 561 ageing databases as highlighted throughout the results' section.

3.4 Which genes to focus on if we were to target age-associated dysfunction in thegut and brain.

564 As a final step in our analysis, we wanted to identify common gene expression 565 changes between the gut and the brain at the earliest stages of ageing, in our case, from the 566 age of 9 months. We hypothesised that this may highlight potential common therapeutic 567 targets for early intervention to prevent age-associated dysfunction in both tissues. We 568 identified 8 significant DEGs in common between the gut and brain at 9 months of age, 5 569 down-regulated (ccnb1, kif2c, serpinh1a, si:ch211-5k11.8 and tmem37) and 2 up-regulated 570 (cfap45 and eif4ebp31). We could then hypothesise that restoring expression of these genes 571 to youthful levels would have a positive impact on delaying or ameliorating the 572 development of ageing phenotypes in both these tissues at the same time. Of note, we 573 identified that, amongst these, ccnb1 and eif4ebp31 were telomerase-dependent changes in 574 the brain. If so, one could hypothesise that telomerase re-activation in the brain should 575 restore these genes to young WT levels and potentially contribute to amelioration of ccnb1-576 and *eif4ebp31*-dependent ageing phenotypes in the brain.

577

578 4 CONCLUSIONS

579 We provide the first systematic analysis of transcriptomic changes throughout the 580 lifespan of zebrafish in the gut and brain of the same group of individuals, leading to the 581 identification of key genes and processes likely involved in driving the ageing process in 582 these tissues. Many of these genes have previously identified in human ageing databases 583 and many of them are potentially new genes of ageing, which will have to be experimentally 584 tested in relevant model organisms. This analysis and the open access availability of its 585 source and raw data should provide a key steppingstone and framework supporting future 586 work for understanding the ageing process and using zebrafish for studying ageing.

587

588 **5 MATERIALS AND METHODS**

589 5.1 Zebrafish husbandry, genotypes and ages

Zebrafish were maintained at the standard conditions of 27-28°C, in a 14:10 hour light-dark cycle, and fed twice a day with *Artemia* (live rotifers) and Sparus (dry food). All the experiments were performed in the University of Sheffield. All animal work was approved by local animal review boards, including the Local Ethical Review Committee at the University of Sheffield (performed according to the protocols of Project Licence 595 70/8681).

596 Two strains of adult zebrafish (Danio rerio) were used for these studies: wild-type (WT; AB strain) and *tert^{-/-}* (*tert^{AB/hu3430}*). Wild-type fish were obtained from the Zebrafish 597 International Resource Center (ZIRC). The *telomerase* mutant line *tert*^{AB/hu3430} was generated 598 599 by N-ethyl-nitrosourea mutagenesis (Utrecht University, Netherlands(Wienholds et al., 600 2003)). It has a $T \rightarrow A$ point mutation in the *tert* gene and is available at the ZFIN repository, 601 ZFIN ID: ZDB-GENO-100412–50, from ZIRC. The fish used in this study are direct descendants of the ones used previously^{29,30}, by which point it had been subsequently outcrossed five 602 603 times with WT AB for clearing of potential background mutations derived from the random ENU mutagenesis from which this line was originated. The *tert*^{hu3430/hu3430} homozygous 604 mutant is referred to in the paper as $tert^{-/-}$ and was obtained by incrossing 605 our *tert*^{AB/hu3430} strain. Genotyping was performed by PCR of the *tert* gene^{13,14}. In order to 606 607 study age-related phenotypes in the zebrafish gut and brain, we used >30 months old fish 608 for what we consider old in WT (in the last 25-30% of their lifespan), and we considered the $tert^{-/-}$ old fish at the equivalent age (>22 months), which approximately corresponds to the 609 610 last 25-30% of their lifespan. In specific, 'Old' was defined as the age at which the majority 611 of the fish present age-associated phenotypes, such as cachexia, loss of body mass and 612 curvature of the spine. These phenotypes develop close to the time of death and are observed at >30 months of age in WT and at >22 months in *tert*^{-/}(Madalena C Carneiro et 613 614 al., 2016; Henriques et al., 2013).

615

616

5.2 Tissue dissection and RNA extraction

Animals from various age-groups were used for RNA-Sequencing (WT at 2, 9, 22 and >30 months of age; and *tert^{-/.}* at 2, 9 and 22 months of age). Three biological replicates were used per group. Fish were culled by overdose of MS-222, followed by confirmation of death. Then, the whole tissues were rapidly dissected in cold PBS (Sigma-Aldrich), transferred to a

621 microcentrifuge tube containing 100 μ l of Trizol (Thermo Fisher Scientific), snap-frozen in 622 dry ice and stored at -80°. To isolate the RNA, extra 50 μ l of Trizol was added to each 623 sample, and the tissue was homogenized with a mechanical homogenizer (VWR 624 International) and a 1.5 pestle (Kimble Chase, Vineland, NJ, USA). After 5 min incubation at 625 room temperature (RT), 30 μ l of chloroform (1:5, VWR International) was added and the 626 samples were incubated for further 3 min at RT before centrifuged at 13,000g, for 30 min, at 627 4°C. Isopropanol (Thermo Fisher Scientific) was then added to the aqueous phase of the 628 solution, and the resultant mix was incubated for 10 min at RT, before centrifuged (13,000g 629 for 15 min at 4°C). Finally, the pellet was twice washed in 250 μ l of ice cold 75% ethanol and 630 left to air-dry, before resuspended in 14 μ l of nuclease-free water. RNA integrity was 631 assessed with the bioanalyzer Agilent 2100 Bioanalyzer (Agilent, Santa Clara, CA, USA). All 632 the samples had a RNA integrity number (RIN) ≥ 9 .

633

634 **5.3 RNA-Sequencing**

635 The RNA-Sequencing was performed at the Genomics and Sequencing facility of 636 Sheffield Institute for Translational Neuroscience (SITraN). Library preparation was 637 performed following the Illumina methodology. mRNA was extracted from 500 ng of RNA 638 with oligo-dT beads, capturing poly(A) tails, using the NEBNext Poly(A) mRNA Magnetic 639 Isolation Module (New England Biolabs Inc). cDNA libraries were made with the NEBNext® 640 Ultra[™] RNA Library Prep Kit for Illumina, following the manufacturer's instructions (New 641 England Biolabs Inc). The samples were then sequenced on an Illumina HiSeg SQ, using a 642 high output run and sequencing by synthesis V3 chemistry on a single 100 bp run. The data 643 was imported into a FASTQ file format in order to perform the analysis.

644

645 **5.4 RNA-Sequencing analysis**

646 5.4.1 Data processing. Quality control was performed using MultiQC version 1.9. 647 Cutadapt version 3.0 was used for trimming the first 13 bases from the reads to remove 648 poor quality base pairs in the reads. Read alignment was performed as follows. Single-end 649 reads were aligned against the reference genome Danio rerio.GRCz11.dna. 650 primary assembly.fa using STAR. A bespoke alignment index was built using annotation file

Danio_rerio.GRCz11.103.gtf and an expected read length of 88 bp. Ht-seq count was run innon-stranded mode to obtain *per* gene read counts.

5.4.2 Differential expression. To identify signatures of ageing, WT gut and brain samples were subjected separately to DESEq2 analysis, comparing the time points 9, 22 and >30 months with the time-point of 2 months. Then, *tert*^{-/-} samples at 2, 9 and 22 months were contrasted with WT at 2 months, in order to identify telomerase-dependent ageing processes.

658 5.4.3 Time-series analysis. Short Time-series Expression Miner (STEM) software was 659 used to assign genes to predefined expression profiles genes based on their expression 660 across the time points. For this, the 2 months' time-point was used as the zero time point 661 for the analysis. The maximum number of model profiles was set to 50; the maximum unit 662 change in model profiles between time points was set to 2; and the minimum absolute log 663 ratio of expression was set to 1.0, with change based on maximum - minimum. Significance 664 level of the model profiles was set to 0.05 with Bonferroni correction. Minimum correlation 665 for profile clustering was set to 0.7. The statistically significant temporal profiles were 666 visualised as line plots using ggplot2. Median expression fold change values of the genes in 667 each profile were shown on the plots with a thicker line.

668 5.4.4 Enrichment analysis of temporal profiles from STEM. Enrichment analysis was 669 performed using all the differentially expressed genes (DEGs) and using the genes identified 670 in the STEM analysis, separately. Gee-set over-representation analysis (ORA) of GO 671 Biological Process (GOBP), GO Molecular Function (GOMF), and GO Cellular Compartment 672 (GOCC) terms were performed using the enrichGO function of clusterProfiler package 673 version 3.18.0. Minimum and maximum gene set sizes were set to 10 and 500, respectively. 674 Results were simplified using the simplify function of clusterProfiler with the similarity cut-675 off set to 0.7 and minimum adjusted p-value used as the feature for selecting the 676 representative terms. Enrichment results with adjusted p-values below 0.05 and at least 3 677 core enrichment genes were considered significantly enriched. ORA of KEGG and 678 REACTOME pathway terms were performed using the enrichKEGG and enrichPathway 679 functions of clusterProfiler. Minimum and maximum gene set sizes were set to 10 and 500, 680 respectively. The same significance criteria for the enrichments were used as for the GO 681 term enrichments. Results of the enrichment results were visualised as barplots or as pie

charts. Barplots were made using the ggbarplot function of R package (R studio version
2021.09.2), ggpubr version 0.4.0, showing a maximum of five pathways with adjusted pvalue below 0.05 *per* pathway category. Pie charts were made using Prism GraphPad version
9.0.

686 **5.4.5 Protein-protein interaction network analysis**

The search tool for retrieval of interacting genes (STRING) (https://string-db.org) database, which integrates both known and predicted PPIs, was used to predict functional interactions of proteins(Szklarczyk et al., 2021). Active interaction sources, including text mining, experiments, databases, and co-expression as well as species limited to "Danio rerio" and an interaction score > 0.7 (high confidence) were applied to construct the PPI networks. The network was further clustered using *k-means* clustering to a specific number of up to 3 clusters.

694

695 **5.4.6 Venn Diagram analysis**

696 Venn Diagram analysis was performed using the online tool Venny 2.1.0 – BioinfoGP
697 (https://bioinfogp.cnb.csic.es/tools/venny/)(Oliveros, 2007-2015).

698

699 **5.4.7** Statistical analysis

A chi-square test was used in **Fig 5** to compare between the relative chromosome location of telomerase-dependent genes in the gut and brain, using raw data values, even though it is the % that is represented in the graphs, to ensure accurate analysis. P value www.enalty.com 20.05 was considered significant and denoted with a *, whereas ns was used to denote nonsignificant differences.

705

706 **ACKNOWLEDGEMENTS**

The authors would like to acknowledge Genevia Technologies Oy for the main bulk of RNA sequencing analysis service. We are very grateful to Joao Ribeiro for valuable advice regarding the many functionalities of Microsoft Excel, which greatly facilitated data analysis. This work was generously funded by a University of Sheffield PhD studentship to RRM, a

711 Sheffield University Vice Chancellor's Research Fellowship and a Wellcome Trust/Royal

712 Society Sir Henry Dale Fellowship (UNS35121) to CMH.

713

714 **AUTHOR CONTRIBUTIONS**

RRM and CMH conceived and designed this work; RRM performed additional RNA sequencing beyond that provided by Genevia services; RRM, MR and CMH analysed and interpreted RNA sequencing results and co-wrote the manuscript; CMH designed the figures with input from co-authors.

719

720 **COMPETING INTERESTS**

- 721 The authors declare no competing interests
- 722
- 723

724 DATA AVAILABILITY STATEMENT

725 The RNA sequencing data from this experiment were deposited in gene expression

726 omnibus (GEO) and will be made available when in peer review.

727

728 **ORCID**

- 729 Raquel Rua Martins: <u>https://orcid.org/0000-0003-3952-8649</u>
- 730 Michael Rera: <u>https://orcid.org/0000-0002-6574-6511</u>
- 731 Catarina Martins Henriques: <u>https://orcid.org/0000-0003-1882-756X</u>
- 732

733 SUPPORTING INFORMATION

734 Supporting information in the form of figures, tables and source data can be found

- online in the Supportive information section at the end of the article
- 736

737 **REFERENCES**

- Ahmed, S., Passos, J. F., Birket, M. J., Beckmann, T., Brings, S., Peters, H., . . . Saretzki, G.
 (2008). Telomerase does not counteract telomere shortening but protects
 mitochondrial function under oxidative stress. *J Cell Sci*, *121*(Pt 7), 1046-1053.
 doi:10.1242/ics.019372
- Anchelin, M., Alcaraz-Perez, F., Martinez, C. M., Bernabe-Garcia, M., Mulero, V., & Cayuela,
 M. L. (2013). Premature aging in telomerase-deficient zebrafish. *Dis Model Mech*,
 6(5), 1101-1112. doi:10.1242/dmm.011635
- Baker, D. J., Childs, B. G., Durik, M., Wijers, M. E., Sieben, C. J., Zhong, J., . . . van Deursen, J.
 M. (2016). Naturally occurring p16(Ink4a)-positive cells shorten healthy lifespan. *Nature*, 530(7589), 184-189. doi:10.1038/nature16932
- Baker, D. J., Wijshake, T., Tchkonia, T., LeBrasseur, N. K., Childs, B. G., van de Sluis, B., ...
 van Deursen, J. M. (2011). Clearance of p16Ink4a-positive senescent cells delays
 ageing-associated disorders. *Nature*, 479(7372), 232-236. doi:10.1038/nature10600
- Basak, O., van de Born, M., Korving, J., Beumer, J., van der Elst, S., van Es, J. H., & Clevers, H.
 (2014). Mapping early fate determination in Lgr5+ crypt stem cells using a novel
 Ki67-RFP allele. *EMBO J*, 33(18), 2057-2068. doi:10.15252/embj.201488017
- Bertaggia, E., Jensen, K. K., Castro-Perez, J., Xu, Y., Di Paolo, G., Chan, R. B., . . . Haeusler, R.
 A. (2017). Cyp8b1 ablation prevents Western diet-induced weight gain and hepatic
 steatosis because of impaired fat absorption. *Am J Physiol Endocrinol Metab*, *313*(2),
 E121-E133. doi:10.1152/ajpendo.00409.2016
- Blackburn, E. H., Epel, E. S., & Lin, J. (2015). Human telomere biology: A contributory and
 interactive factor in aging, disease risks, and protection. *Science*, *350*, 1193-1198.
 doi:10.1126/science.aab3389
- 762Bodnar, A. G. (1998). Extension of Life-Span by Introduction of Telomerase into Normal763Human Cells. Science, 279(5349), 349-352. doi:10.1126/science.279.5349.349
- Boutouja, F., Stiehm, C. M., & Platta, H. W. (2019). mTOR: A Cellular Regulator Interface in
 Health and Disease. *Cells*, 8(1). doi:10.3390/cells8010018
- Cao, Y., Li, H., Deb, S., & Liu, J. P. (2002). TERT regulates cell survival independent of
 telomerase enzymatic activity. *Oncogene*, *21*(20), 3130-3138.
 doi:10.1038/sj.onc.1205419
- Cardoso, B. a., Gírio, a., Henriques, C., Martins, L. R., Santos, C., Silva, a., & Barata, J. T.
 (2008). Aberrant signaling in T-cell acute lymphoblastic leukemia: Biological and therapeutic implications. *Brazilian Journal of Medical and Biological Research*, *41*, 344-350. doi:10.1590/S0100-879X2008005000016
- Carneiro, M. C., de Castro, I. P., & Ferreira, M. G. (2016). Telomeres in aging and disease:
 lessons from zebrafish. *Dis Model Mech*, *9*(7), 737-748. doi:10.1242/dmm.025130
- Carneiro, M. C., Henriques, C. M., Nabais, J., Ferreira, T., Carvalho, T., & Ferreira, M. G.
 (2016). Short Telomeres in Key Tissues Initiate Local and Systemic Aging in Zebrafish.
 PLoS Genet, 12, e1005798. doi:10.1371/journal.pgen.1005798
- Choi, J., Southworth, L. K., Sarin, K. Y., Venteicher, A. S., Ma, W., Chang, W., . . . Artandi, S. E.
 (2008). TERT promotes epithelial proliferation through transcriptional control of a
 Myc- and Wnt-related developmental program. *PLoS Genet, 4*(1), e10.
 doi:10.1371/journal.pgen.0040010
- d'Adda di Fagagna, F., Reaper, P. M., Clay-Farrace, L., Fiegler, H., Carr, P., Von Zglinicki, T., . .
 Jackson, S. P. (2003). A DNA damage checkpoint response in telomere-initiated senescence. *Nature*, *426*(6963), 194-198. doi:10.1038/nature02118

- Dambroise, E., Monnier, L., Ruisheng, L., Aguilaniu, H., Joly, J. S., Tricoire, H., & Rera, M.
 (2016). Two phases of aging separated by the Smurf transition as a public path to
 death. *Sci Rep, 6*, 23523. doi:10.1038/srep23523
- De Felice, B., Annunziata, A., Fiorentino, G., Manfellotto, F., D'Alessandro, R., Marino, R., ...
 Biffali, E. (2014). Telomerase expression in amyotrophic lateral sclerosis (ALS)
 patients. J Hum Genet, 59(10), 555-561. doi:10.1038/jhg.2014.72
- de Jong, P. R., Gonzalez-Navajas, J. M., & Jansen, N. J. (2016). The digestive tract as the
 origin of systemic inflammation. *Crit Care, 20*(1), 279. doi:10.1186/s13054-016-14583
- de Lange, T. (2004). T-loops and the origin of telomeres. Nature reviews. Molecular cell
 biology, 5, 323-329. doi:10.1038/nrm1422
- Deacon, K., & Knox, A. J. (2018). PINX1 and TERT Are Required for TNF-alpha-Induced Airway
 Smooth Muscle Chemokine Gene Expression. J Immunol, 200(4), 1283-1294.
 doi:10.4049/jimmunol.1700414
- Dimri, G. P., Lee, X., Basile, G., Acosta, M., Scott, G., Roskelley, C., . . . et al. (1995). A
 biomarker that identifies senescent human cells in culture and in aging skin in vivo. *Proc Natl Acad Sci U S A, 92*(20), 9363-9367.
- Eitan, E., Tichon, A., Gazit, A., Gitler, D., Slavin, S., & Priel, E. (2012). Novel telomeraseincreasing compound in mouse brain delays the onset of amyotrophic lateral
 sclerosis. *EMBO Mol Med*, 4(4), 313-329. doi:10.1002/emmm.201200212
- 805England, P. H. (2017). Chapter 1: life expectancy and healthy life expectancy. Retrieved from806https://www.gov.uk/government/publications/health-profile-for-england/chapter-1-807life-expectancy-and-healthy-life-expectancy#references.
- 808 Ernst, J., & Bar-Joseph, Z. (2006). STEM: a tool for the analysis of short time series gene 809 expression data. *BMC Bioinformatics, 7*, 191. doi:10.1186/1471-2105-7-191
- Ferreira, M. G., Miller, K. M., & Cooper, J. P. (2004). Indecent exposure: when telomeres
 become uncapped. *Mol Cell*, 13(1), 7-18.
- Forsyth, N. R., Wright, W. E., & Shay, J. W. (2002). Telomerase and differentiation in
 multicellular organisms: turn it off, turn it on, and turn it off again. *Differentiation*,
 69(4-5), 188-197. doi:10.1046/j.1432-0436.2002.690412.x
- 815 Gerhardt, T., & Ley, K. (2015). Monocyte trafficking across the vessel wall. *Cardiovascular* 816 *Research*, 107(3), 321-330. doi:10.1093/cvr/cvv147
- Ghosh, A., Saginc, G., Leow, S. C., Khattar, E., Shin, E. M., Yan, T. D., . . . Tergaonkar, V.
 (2012). Telomerase directly regulates NF-kappaB-dependent transcription. *Nat Cell Biol, 14*(12), 1270-1281. doi:10.1038/ncb2621
- Gladyshev, V. N. (2021). The Ground Zero of Organismal Life and Aging. *Trends Mol Med*,
 27(1), 11-19. doi:10.1016/j.molmed.2020.08.012
- Gomes, N. M. V., Ryder, O. a., Houck, M. L., Charter, S. J., Walker, W., Forsyth, N. R., . . .
 Wright, W. E. (2011). Comparative biology of mammalian telomeres: hypotheses on ancestral states and the roles of telomeres in longevity determination. *Aging Cell*, 10, 761-768. doi:10.1111/j.1474-9726.2011.00718.x
- Goodman, W. A., & Jain, M. K. (2011). Length does not matter: A new take on telomerase
 reverse transcriptase. *Arteriosclerosis, thrombosis, and vascular biology, 31*, 235 236. doi:10.1161/ATVBAHA.110.220343
- Greider, C. W., & Blackburn, E. H. (1985). Identification of a specific telomere terminal
 transferase activity in Tetrahymena extracts. *Cell*, 43(2 Pt 1), 405-413.
 doi:10.1016/0092-8674(85)90170-9

- Haendeler, J., Drose, S., Buchner, N., Jakob, S., Altschmied, J., Goy, C., . . . Dimmeler, S.
 (2009). Mitochondrial telomerase reverse transcriptase binds to and protects
 mitochondrial DNA and function from damage. *Arterioscler Thromb Vasc Biol, 29*(6),
 929-935. doi:10.1161/ATVBAHA.109.185546
- Hagey, D. W., Topcic, D., Kee, N., Reynaud, F., Bergsland, M., Perlmann, T., & Muhr, J.
 (2020). CYCLIN-B1/2 and -D1 act in opposition to coordinate cortical progenitor selfrenewal and lineage commitment. *Nat Commun*, *11*(1), 2898. doi:10.1038/s41467020-16597-8
- Hayflick, L. (2007). Biological aging is no longer an unsolved problem. Ann N Y Acad Sci,
 1100, 1-13. doi:10.1196/annals.1395.001
- Henriques, C. M., Carneiro, M. C., Tenente, I. M., Jacinto, A., & Ferreira, M. G. (2013).
 Telomerase is required for zebrafish lifespan. *PLoS Genet, 9*(1), e1003214.
 doi:10.1371/journal.pgen.1003214
- Henriques, C. M., & Ferreira, M. G. (2012). Consequences of telomere shortening during
 lifespan. *Curr Opin Cell Biol, 24*(6), 804-808. doi:10.1016/j.ceb.2012.09.007
- Hong, S. N., Joung, J. G., Bae, J. S., Lee, C. S., Koo, J. S., Park, S. J., . . . Kim, Y. H. (2017). RNAseq Reveals Transcriptomic Differences in Inflamed and Noninflamed Intestinal
 Mucosa of Crohn's Disease Patients Compared with Normal Mucosa of Healthy
 Controls. Inflamm Bowel Dis, 23(7), 1098-1108.
 doi:10.1097/MIB.00000000001066
- Ifergan, I., Kébir, H., Bernard, M., Wosik, K., Dodelet-Devillers, A., Cayrol, R., . . . Prat, A.
 (2007). The blood-brain barrier induces differentiation of migrating monocytes into
 Th17-polarizing dendritic cells. *Brain*, 131(3), 785-799. doi:10.1093/brain/awm295
- Jaskelioff, M., Muller, F. L., Paik, J.-H., Thomas, E., Jiang, S., Adams, A. C., . . . Depinho, R. a.
 (2011). Telomerase reactivation reverses tissue degeneration in aged telomerasedeficient mice. *Nature*, 469, 102-106. doi:10.1038/nature09603
- Jurk, D., Wang, C., Miwa, S., Maddick, M., Korolchuk, V., Tsolou, A., . . . von Zglinicki, T.
 (2012). Postmitotic neurons develop a p21-dependent senescence-like phenotype
 driven by a DNA damage response. *Aging Cell*, *11*(6), 996-1004. doi:10.1111/j.14749726.2012.00870.x
- Lange, C., Rost, F., Machate, A., Reinhardt, S., Lesche, M., Weber, A., . . . Brand, M. (2020).
 Single cell sequencing of radial glia progeny reveals the diversity of newborn neurons
 in the adult zebrafish brain. *Development*, *147*(1). doi:10.1242/dev.185595
- Lawson, N. D., Li, R., Shin, M., Grosse, A., Yukselen, O., Stone, O. A., . . . Zhu, L. (2020). An
 improved zebrafish transcriptome annotation for sensitive and comprehensive
 detection of cell type-specific genes. *Elife*, *9*. doi:10.7554/eLife.55792
- Lee, H. W., Blasco, M. A., Gottlieb, G. J., Horner, J. W., 2nd, Greider, C. W., & DePinho, R. A.
 (1998). Essential role of mouse telomerase in highly proliferative organs. *Nature*,
 392(6676), 569-574. doi:10.1038/33345
- Lee, J., Jo, Y. S., Sung, Y. H., Hwang, I. K., Kim, H., Kim, S. Y., . . . Lee, H. W. (2010).
 Telomerase deficiency affects normal brain functions in mice. *Neurochem Res*, *35*(2),
 211-218. doi:10.1007/s11064-009-0044-3
- Lemoine, M. (2021). The Evolution of the Hallmarks of Aging. *Front Genet, 12*, 693071.
 doi:10.3389/fgene.2021.693071
- Entini, L., Barra, V., Schillaci, T., & Di Leonardo, A. (2012). MAD2 depletion triggers premature cellular senescence in human primary fibroblasts by activating a p53

878 pathway preventing an euploid cells propagation. J Cell Physiol, 227(9), 3324-3332. 879 doi:10.1002/jcp.24030 880 Li, L., Song, J., Chuquisana, O., Hannocks, M. J., Loismann, S., Vogl, T., . . . Sorokin, L. (2020). 881 Endothelial Basement Membrane Laminins as an Environmental Cue in Monocyte 882 Differentiation to Macrophages. Front Immunol, 11, 584229. 883 doi:10.3389/fimmu.2020.584229 884 The hallmarks of aging, 153 (2013). 885 Lu, Z., Ding, L., Lu, Q., & Chen, Y. H. (2013). Claudins in intestines: Distribution and functional 886 health and d ise ases. e24978. significance in Tissue Barriers, 1(3), 887 doi:10.4161/tisb.24978 888 Mattiussi, M., Tilman, G., Lenglez, S., & Decottignies, A. (2012). Human telomerase 889 represses ROS-dependent cellular responses to Tumor Necrosis Factor- α without 890 affecting NF-ĸB activation. Cellular signalling, 24. 708-717. 891 doi:10.1016/j.cellsig.2011.11.004 Morris, M., Hepburn, P., & Wynford-Thomas, D. (2002). Sequential extension of proliferative 892 893 lifespan in human fibroblasts induced by over-expression of CDK4 or 6 and loss of 894 p53 function. Oncogene, 21(27), 4277-4288. doi:10.1038/sj.onc.1205492 895 Neale, B. M., Fagerness, J., Reynolds, R., Sobrin, L., Parker, M., Raychaudhuri, S., . . . Seddon, 896 J. M. (2010). Genome-wide association study of advanced age-related macular 897 degeneration identifies a role of the hepatic lipase gene (LIPC). Proc Natl Acad Sci U S 898 A, 107(16), 7395-7400. doi:10.1073/pnas.0912019107 899 Nersisyan, L., Hopp, L., Loeffler-Wirth, H., Galle, J., Loeffler, M., Arakelyan, A., & Binder, H. 900 (2019). Telomere Length Maintenance and Its Transcriptional Regulation in Lynch 901 and Sporadic Colorectal Carcinoma. Front Syndrome Oncol, 1172. 9, 902 doi:10.3389/fonc.2019.01172 903 Oliveros, J. C. (2007-2015). Venny. An interactive tool for comparing lists with Venn's 904 diagrams. Retrieved from https://bioinfogp.cnb.csic.es/tools/venny/index.html 905 Ovadya, Y., & Krizhanovsky, V. (2014). Senescent cells: SASPected drivers of age-related 906 pathologies. Biogerontology, 15, 627-642. doi:10.1007/s10522-014-9529-9 907 Pam S. Ellis, R. R. M., Emily J. Thompson, Asma Farhat, Stephen A. Renshaw, Catarina M. 908 Henriques. (2022). A subset of gut leukocytes have telomerase-dependent "hyper-909 long" telomeres and require telomerase for function in zebrafish. Biorxiv. 910 Pekalski, M. L., Ferreira, R. C., Coulson, R. M., Cutler, A. J., Guo, H., Smyth, D. J., . . . Wicker, 911 L. S. (2013). Postthymic expansion in human CD4 naive T cells defined by expression 912 of functional high-affinity IL-2 receptors. J Immunol, 190(6), 2554-2566. 913 doi:10.4049/jimmunol.1202914 914 Poehlmann, A., Habold, C., Walluscheck, D., Reissig, K., Bajbouj, K., Ullrich, O., . . . Schneider-915 Stock, R. (2011). Cutting edge: Chk1 directs senescence and mitotic catastrophe in 916 recovery from G(2) checkpoint arrest. J Cell Mol Med, 15(7), 1528-1541. 917 doi:10.1111/j.1582-4934.2010.01143.x 918 Pruitt, S. C., Bailey, K. J., & Freeland, A. (2007). Reduced Mcm2 expression results in severe 919 stem/progenitor cell deficiency and cancer. Stem Cells, 25(12), 3121-3132. 920 doi:10.1634/stemcells.2007-0483 921 Rahman, R., Latonen, L., & Wiman, K. G. (2005). hTERT antagonizes p53-induced apoptosis 922 independently of telomerase activity. Oncogene, 24(8), 1320-1327. 923 doi:10.1038/sj.onc.1208232

- Raj, D. D. A., Moser, J., van der Pol, S. M. A., van Os, R. P., Holtman, I. R., Brouwer, N., . . .
 Boddeke, H. W. G. M. (2015). Enhanced microglial pro-inflammatory response to
 lipopolysaccharide correlates with brain infiltration and blood-brain barrier
 dysregulation in a mouse model of telomere shortening. *Aging Cell*, 1003-1013.
 doi:10.1111/acel.12370
- Rando, T. A., & Wyss-Coray, T. (2021). Asynchronous, contagious and digital aging. Nat
 Aging, 1(1), 29-35. doi:10.1038/s43587-020-00015-1
- 931Rera, M., Azizi, M. J., & Walker, D. W. (2013). Organ-specific mediation of lifespan extension:932more than a gut feeling? Ageing Res Rev, 12(1), 436-444.933doi:10.1016/j.arr.2012.05.003
- Rera, M., Clark, R. I., & Walker, D. W. (2012). Intestinal barrier dysfunction links metabolic
 and inflammatory markers of aging to death in Drosophila. *Proc Natl Acad Sci U S A*,
 109(52), 21528-21533. doi:10.1073/pnas.1215849110
- Ripa, R., Dolfi, L., Terrigno, M., Pandolfini, L., Savino, A., Arcucci, V., . . . Cellerino, A. (2017).
 MicroRNA miR-29 controls a compensatory response to limit neuronal iron accumulation during adult life and aging. *BMC Biol*, *15*(1), 9. doi:10.1186/s12915-017-0354-x
- Robin, J. D., Ludlow, A. T., Batten, K., Magdinier, F., Stadler, G., Wagner, K. R., ... Wright, W.
 E. (2014). Telomere position effect: regulation of gene expression with progressive
 telomere shortening over long distances. *Genes Dev, 28*(22), 2464-2476.
 doi:10.1101/gad.251041.114
- Romaniuk, A., Paszel-Jaworska, A., Toton, E., Lisiak, N., Holysz, H., Krolak, A., . . . Rubis, B.
 (2018). The non-canonical functions of telomerase: to turn off or not to turn off. *Mol Biol Rep.* doi:10.1007/s11033-018-4496-x
- Rudnicka, E., Napierala, P., Podfigurna, A., Meczekalski, B., Smolarczyk, R., & Grymowicz, M.
 (2020). The World Health Organization (WHO) approach to healthy ageing. *Maturitas, 139*, 6-11. doi:10.1016/j.maturitas.2020.05.018
- Santamarina-Fojo, S., Gonzalez-Navarro, H., Freeman, L., Wagner, E., & Nong, Z. (2004).
 Hepatic lipase, lipoprotein metabolism, and atherogenesis. *Arterioscler Thromb Vasc Biol, 24*(10), 1750-1754. doi:10.1161/01.ATV.0000140818.00570.2d
- Sarin, K. Y., Cheung, P., Gilison, D., Lee, E., Tennen, R. I., Wang, E., . . . Artandi, S. E. (2005).
 Conditional telomerase induction causes proliferation of hair follicle stem cells. *Nature*, 436(7053), 1048-1052. doi:10.1038/nature03836
- Schaum, N., Lehallier, B., Hahn, O., Palovics, R., Hosseinzadeh, S., Lee, S. E., . . . Wyss-Coray,
 T. (2020). Ageing hallmarks exhibit organ-specific temporal signatures. *Nature*,
 583(7817), 596-602. doi:10.1038/s41586-020-2499-y
- Scheinin, T., Bohling, T., Halme, L., Kontiainen, S., Bjorge, L., & Meri, S. (1999). Decreased
 expression of protectin (CD59) in gut epithelium in ulcerative colitis and Crohn's
 disease. *Hum Pathol*, 30(12), 1427-1430. doi:10.1016/s0046-8177(99)90163-6
- Schenkel, A. R., Mamdouh, Z., Chen, X., Liebman, R. M., & Muller, W. A. (2002). CD99 plays a
 major role in the migration of monocytes through endothelial junctions. *Nature immunology*, 3(2), 143-150. doi:10.1038/ni749
- Schmidt, S., Schneider, L., Essmann, F., Cirstea, I. C., Kuck, F., Kletke, A., . . . Piekorz, R. P.
 (2010). The centrosomal protein TACC3 controls paclitaxel sensitivity by modulating
 a premature senescence program. *Oncogene, 29*(46), 6184-6192.
 doi:10.1038/onc.2010.354

970 Segal-Bendirdjian, E., & Geli, V. (2019). Non-canonical Roles of Telomerase: Unraveling the 971 Imbroglio. Front Cell Dev Biol, 7, 332. doi:10.3389/fcell.2019.00332 972 Shokhirev, M. N., & Johnson, A. A. (2021). Modeling the human aging transcriptome across 973 tissues, health status, and sex. Aging Cell, 20(1), e13280. doi:10.1111/acel.13280 974 Smith, L. K., He, Y., Park, J. S., Bieri, G., Snethlage, C. E., Lin, K., . . . Villeda, S. A. (2015). 975 beta2-microglobulin is a systemic pro-aging factor that impairs cognitive function 976 and neurogenesis. Nat Med, 21(8), 932-937. doi:10.1038/nm.3898 977 Spilsbury, A., Miwa, S., Attems, J., & Saretzki, G. (2015). The role of telomerase protein TERT 978 in Alzheimer's disease and in tau-related pathology in vitro. J Neurosci, 35(4), 1659-979 1674. doi:10.1523/JNEUROSCI.2925-14.2015 980 Sullivan, D. I., Jiang, M., Hinchie, A. M., Roth, M. G., Bahudhanapati, H., Nouraie, M., . . . 981 Alder, J. K. (2021). Transcriptional and Proteomic Characterization of Telomere-982 Induced Senescence in a Human Alveolar Epithelial Cell Line. Front Med (Lausanne), 983 8, 600626. doi:10.3389/fmed.2021.600626 984 Sung, Y. H., Ali, M., & Lee, H. W. (2014). Extracting extra-telomeric phenotypes from 985 models. telomerase mouse Yonsei Medical Journal, 55, 1-8. 986 doi:10.3349/ymj.2014.55.1.1 987 Szklarczyk, D., Gable, A. L., Nastou, K. C., Lyon, D., Kirsch, R., Pyysalo, S., . . . von Mering, C. 988 (2021). The STRING database in 2021: customizable protein-protein networks, and 989 functional characterization of user-uploaded gene/measurement sets. Nucleic Acids 990 Res, 49(D1), D605-D612. doi:10.1093/nar/gkaa1074 991 Tabula Muris, C. (2020). A single-cell transcriptomic atlas characterizes ageing tissues in the 992 mouse. Nature, 583(7817), 590-595. doi:10.1038/s41586-020-2496-1 993 Tacutu, R., Thornton, D., Johnson, E., Budovsky, A., Barardo, D., Craig, T., . . . de Magalhaes, 994 J. P. (2018). Human Ageing Genomic Resources: new and updated databases. Nucleic 995 Acids Res, 46(D1), D1083-D1090. doi:10.1093/nar/gkx1042 996 Tricoire, H., & Rera, M. (2015). A New, Discontinuous 2 Phases of Aging Model: Lessons from 997 Drosophila melanogaster. PLoS One, 10(11), e0141920. 998 doi:10.1371/journal.pone.0141920 999 Ugalde, A. P., Ramsay, A. J., de la Rosa, J., Varela, I., Marino, G., Cadinanos, J., . . . Lopez-1000 Otin, C. (2011). Aging and chronic DNA damage response activate a regulatory 1001 pathway involving miR-29 and p53. EMBO J, 30(11), 2219-2232. 1002 doi:10.1038/emboj.2011.124 1003 Watt, P. M., & Hickson, I. D. (1994). Structure and function of type II DNA topoisomerases. 1004 Biochem J, 303 (Pt 3), 681-695. doi:10.1042/bj3030681 1005 Wienholds, E., van Eeden, F., Kosters, M., Mudde, J., Plasterk, R. H., & Cuppen, E. (2003). 1006 Efficient target-selected mutagenesis in zebrafish. Genome Res, 13(12), 2700-2707. 1007 doi:10.1101/gr.1725103 Zhang, H., Weyand, C. M., & Goronzy, J. J. (2021). Hallmarks of the aging T-cell system. FEBS 1008 1009 J, 288(24), 7123-7142. doi:10.1111/febs.15770 1010 Zhang, M. J., Pisco, A. O., Darmanis, S., & Zou, J. (2021). Mouse aging cell atlas analysis 1011 reveals cell type-specific signatures. Elife, global and aging 10. 1012 doi:10.7554/eLife.62293 1013 Zhao, Y., Hasse, S., & Bourgoin, S. G. (2021). Phosphatidylserine-specific phospholipase A1: A 1014 friend the devil in disguise. Prog Lipid Res, 83, 101112. or 1015 doi:10.1016/j.plipres.2021.101112

1016 Zhu, S., Paydar, M., Wang, F., Li, Y., Wang, L., Barrette, B., . . . Peng, A. (2020). Kinesin Kif2C
1017 in regulation of DNA double strand break dynamics and repair. *Elife, 9*.
1018 doi:10.7554/eLife.53402
1019

1020

1021 FIGURE LEGENDS

1022

1023 Graphical Abstract: Although the gut and brain have distinct transcriptomic signatures of 1024 ageing, both tissues display hallmarks of ageing as early as 9 months in the WT zebrafish. 1025 Importantly, telomerase depletion accelerates the appearance of such hallmarks in both gut and brain. In specific, the tert^{-/-} gut at 2 months and the tert^{-/-} brain at 9 months display 1026 1027 similar distribution of the mainly affected hallmarks of ageing, in the WT old gut and brain at 1028 35 months of age. We identified stem cell exhaustion (light red) as the common principal 1029 hallmark of ageing at the early stages of ageing, in both tissues. Finally, we further identified 1030 altered intercellular communication (light green), in which immunity and inflammation play 1031 a central role, as the main telomerase-dependent hallmark of ageing common between the 1032 gut and brain.

1033 Fig 1. Summary of the experimental design and principal component analysis (PCA). (A) 1034 RNA-Sequencing was performed in whole gut and brain tissues from WT and tert '-1035 zebrafish, at different timepoints throughout their lifespan. Age-associated transcriptomic 1036 changes were analysed using two different methods: time-series analysis (genes that change 1037 consistently overtime) and all differentially expressed genes (ALL DEGs; genes whose 1038 expression is altered over-time, in both genotypes, as compared with the WT young 1039 baseline. (B) PCA representing the variation in the data from gut (gold) and brain (pink) tissues, in both WT and tert^{-/-} fish. (C-D) PCA showing the variation in the data from fish at 1040 different ages (2 months, pink; 9 months, green; 22 months, blue; 35 months, purple), in WT 1041 (circle) versus tert^{-,-} fish (triangle), in (C) gut and (D) brain samples. PCA was performed 1042 1043 using the plotPCA function of DESeq2 and considering the top 500 genes with highest 1044 variance across the samples. (E) Summary of the number of significantly de-regulated genes 1045 at each time-point, in both genotypes, in gut (E1) and brain (E2) tissues, over-time, as 1046 compared with the WT young baseline.

Fig 2. Identification of time-dependent signatures of ageing in the WT and *tert^{-/-}* zebrafish 1048 gut and brain. (A, B) Transcriptomic temporal profiles in the (A) gut and brain (B) of WT 1049 (black) and tert^{-/-} fish (red) were identified using the Short Time-series Expression Miner 1050 1051 (STEM) and are represented in line plots. The thicker lines on the plots represent the 1052 median fold change of each profile. For each temporal profile, enrichment analysis was 1053 performed using the enrichGO, enrichKEGG, and enrichPathway functions of clusterProfiler 1054 package version 3.18.0. Processes and pathways from each temporal profile were further 1055 classified and grouped according to the main hallmarks of ageing (Lemoine, 2021; López-1056 Otin et al., 2013), which is represented in pie charts. (A1, B1) A summary of the enrichment 1057 analysis of one of the profiles is represented in the bar plots, where processes and pathways 1058 are represented in different colours according to the classification into hallmarks of ageing. 1059 This summary contains the top enriched processes and pathways from each database (p-1060 value >0.05 and at least 3 core enrichment genes; up to 5 terms per database: GOBP, GOCC, 1061 GOMF, KEEG, and REACTOME).

1062

1063 Fig 3. Qualitative changes in the hallmarks of ageing over-time, comparing WT and tert^{-/-} 1064 zebrafish gut and brain. The age-associated enriched processes and pathways identified in 1065 the previous figures were further re-classified and grouped into the main well-known 1066 hallmarks of ageing. (A-B) Pie charts represent the hallmarks of ageing identified in the (A) gut and (B) brain, at different ages throughout WT (black) and *tert^{-/-}* (red) zebrafish lifespan. 1067 This analysis was performed considering the (A1, B1) genes identified in the temporal 1068 1069 profiles (within STEM), (A2, B2) all the genes differentially expressed at any timepoint 1070 (within ALL DEGs), and (A3, B3) STEM and ALL DEGs combined. (A4, B4) The number of 1071 transcriptomic changes increases with ageing in both (A4) gut and (B4) brain, independently 1072 of the phenotype, when considering either the genes within STEM or the genes within ALL 1073 DEGs.

1074

Fig 4. Determining at which age tert^{-/-} share more genes associated with the hallmarks of ageing, with the naturally aged WT. (A) Venn diagrams highlight the number of genes associated with hallmarks of ageing in common between old WT (35 months) and tert^{-/-} at the different ages tested, in (A.1) gut and (A.2) brain tissues. Data show that the tert^{-/-} gut at 2 months has the most number genes associated with hallmarks of ageing in common with

old WT gut. The *tert*^{-/-} brain at 22 months has the most number genes associated with hallmarks of ageing in common with old WT brain. The respective lists of genes shared between old WT gut and *tert*^{-/-} gut at 2 months and old WT brain and *tert*^{-/-} 22-month brain are shown in B (B.1 gut, B.2, brain). (B1.1, B1.2) Gene networks with the genes identified in B.1 and B.2 were performed (B1.1 and B1.2, respectively) by K-means clustering using String software. These include the genes identified in the temporal profiles (from STEM) and in ALL DEGs.

1087

1088 Fig 5. Identifying tert-dependent gene changes in zebrafish gut and brain ageing and their 1089 chromosome location in relation to the telomeric end. (A-B) Gene alterations that are anticipated in the *tert^{-/-}* when comparing with WT at the same age (i.e., telomerase-1090 1091 dependent). (A1, B1) Protein-protein interaction network of these genes highlights clusters 1092 of genes associated with (A1) metabolic processes in the gut and clusters of genes 1093 associated with (B1) cell cycle, genome instability and immune system in the brain. (C)Genes 1094 located at the end of the chromosome (i.e., <10MB from chromosome end) are likely to be 1095 directly affected by telomere shortening due to the telomere positioning effect (TPE). The 1096 data show that there is no significant difference between telomerase-dependent and -1097 independent genes, in what concerns their proximity to the chromosome end, in either gut 1098 (C2) or brain (C3). However, there is a significantly higher number of telomerase-dependent 1099 genes located at the end of chromosomes in the gut than in the brain (C4).

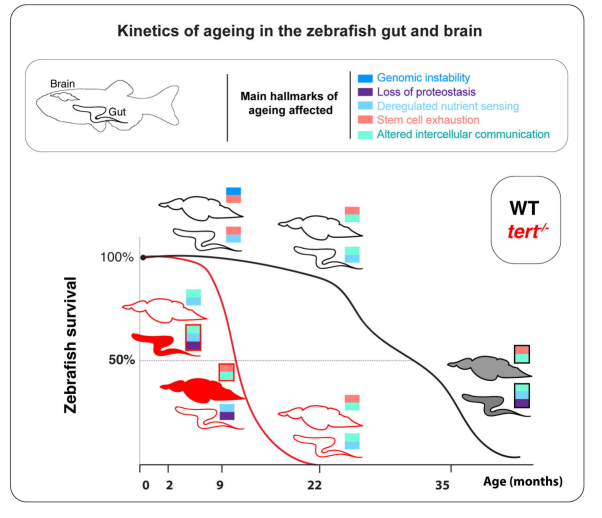
1100

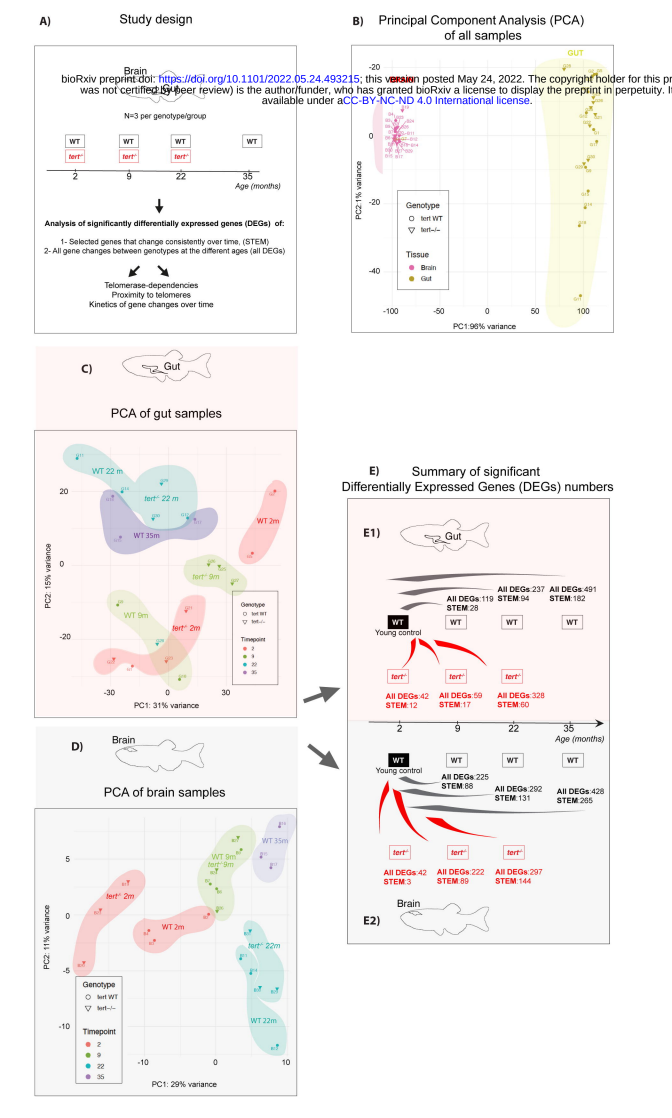
1101 Fig 6. Determining genes and pathways altered with ageing that are in common between 1102 gut and brain. (A) Graph represents WT zebrafish lifespan and highlights 'early' and 'late' 1103 stages of ageing. (A.1) All genes identified in common between gut and brain at early (9 1104 months) and late (35 months) stages of ageing. *G and *B represent telomerase-dependent 1105 genes in the Gut or in the Brain, respectively. (B-C) Protein-protein interaction networks 1106 with the genes found in common between the gut and brain at (B) early and (C) late stages 1107 of ageing were performed using STRING software. These include the genes identified in the 1108 temporal profiles (from STEM) and in ALL DEGs.

- 1109
- 1110
- 1111

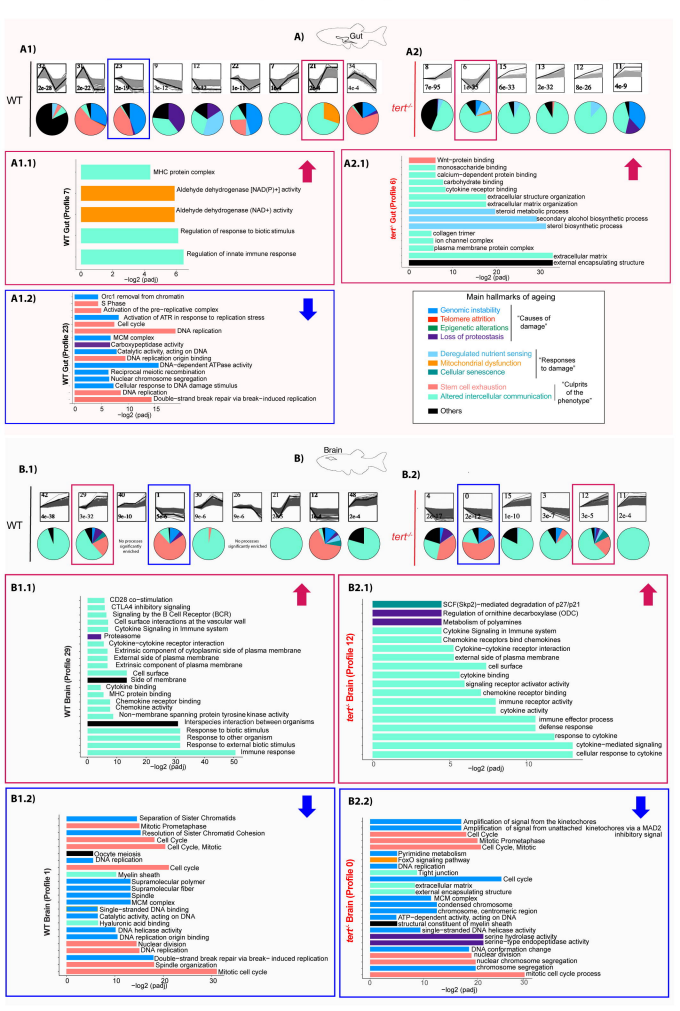
1112 SUPPLEMENTARY FIGURE LEGENDS

Supplementary Fig 1. Genes associated with the hallmarks of ageing that are in common between 35-months-old WT and tert^{-/-} at different ages, in the (A1) gut and (A2) brain. The different colours represent different hallmarks of ageing: purple, loss of proteostasis; yellow, mitochondrial dysfunction; green, altered intercellular communication. These include the genes identified in the temporal profiles (from STEM) and in ALL DEGs. Supplementary Fig 2. Protein-protein interaction network and cluster analysis of gene changes in WT at the 'origins' versus 'later' stages of ageing. (A) Genes identified in common between gut and brain at 9 and 35 months of age. B) Genes significantly altered at "early" and "late" stages of ageing in the gut (B1, 2) and brain (C1, 2), respectively. Network analysis was performed in STRING software and included the genes identified in the temporal profiles (from STEM) and in ALL DEGs. Red squares highlight the protein-protein interaction network of the gene changes that are accelerated in the *tert*^{-/-} (tert-dependent).

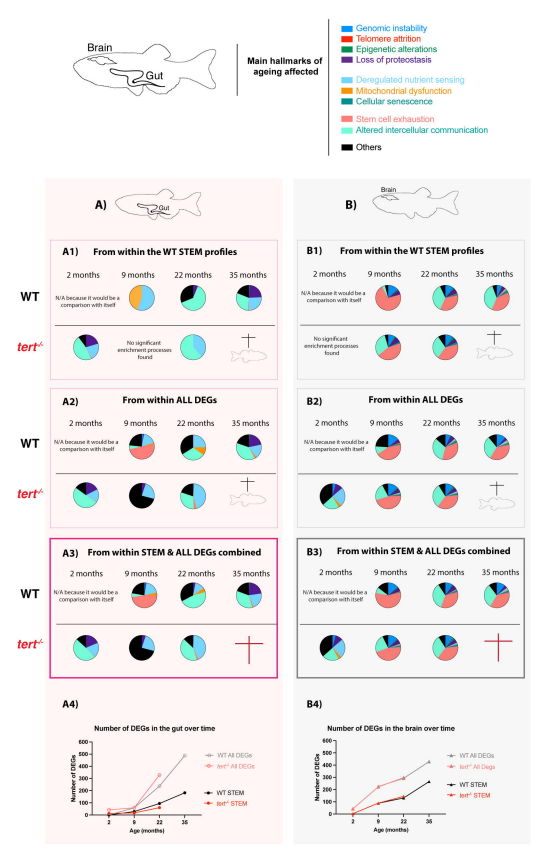




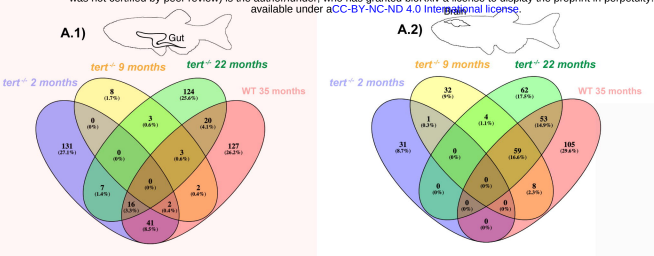
was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It Significant STEM profiles and main processes associated, categorised according to known halfmarks of ageing



Kinetics of ageing in the zebrafish gut and brain

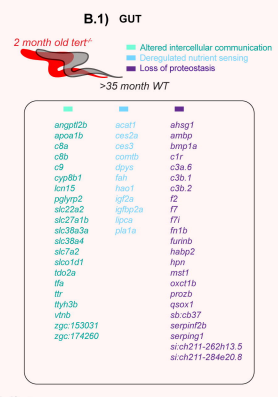


A) Genes associated with the copyright holder for this provide the provided and the provided as a social with the copyright holder for this provided associated with the provided as the provided as a social with the provided as the provide

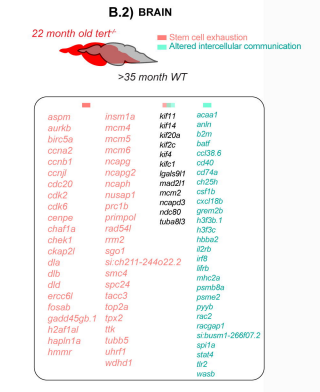


B)

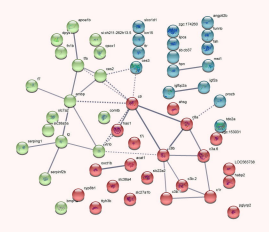
Most gene changes shared between tert $^{\prime\prime}$ and 35 month old WT, related to the main hallmarks of ageing affected

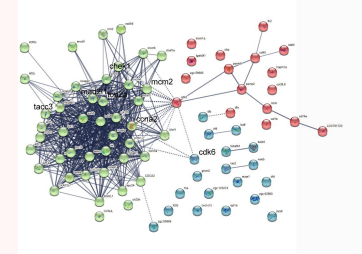


B1.1) String Network analysis (K means clustering) GUT

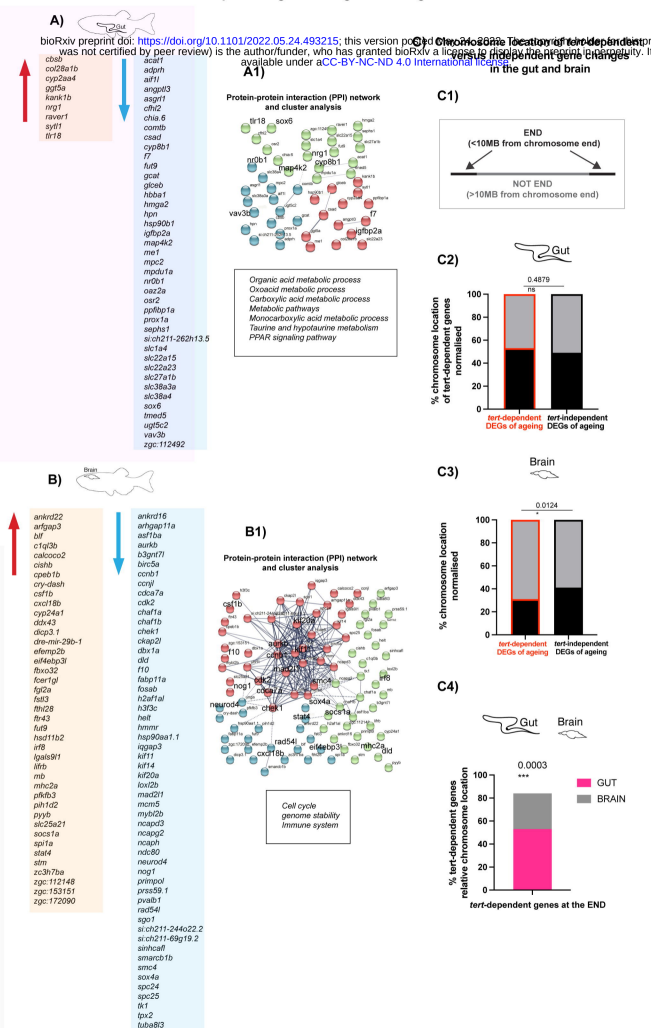


B.2.1) String Network analysis (K means clustering) BRAIN



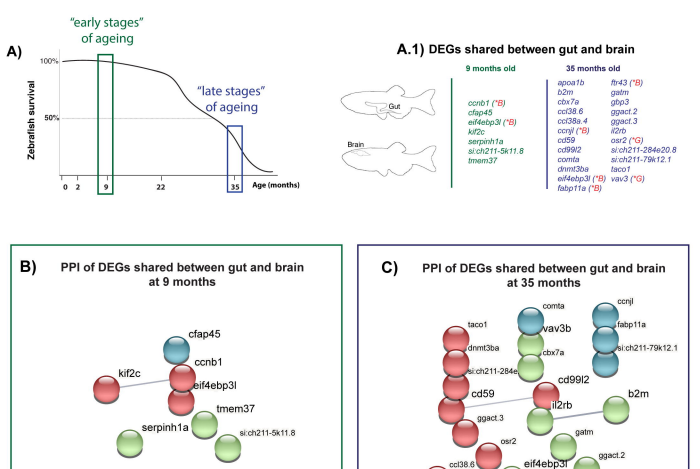


tert-dependent gene changes of old age



tyms uhrf1 unga zwi

Protein-protein interaction (PPI) network and cluster analysis of changes in WT ageing (STEM and All DEGs combined)



apoa1b

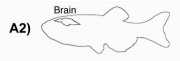
gbp3

ftr43

A) Genes associated with the hallmarks of ageing in common between tert[/] at the different ages and the aged WT at 35 months



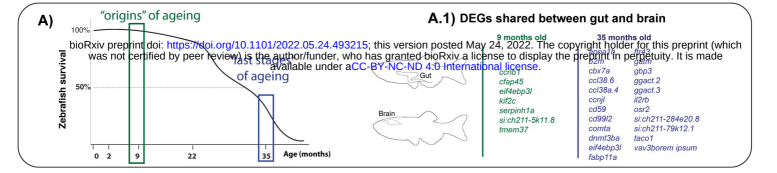
WT 35 months x

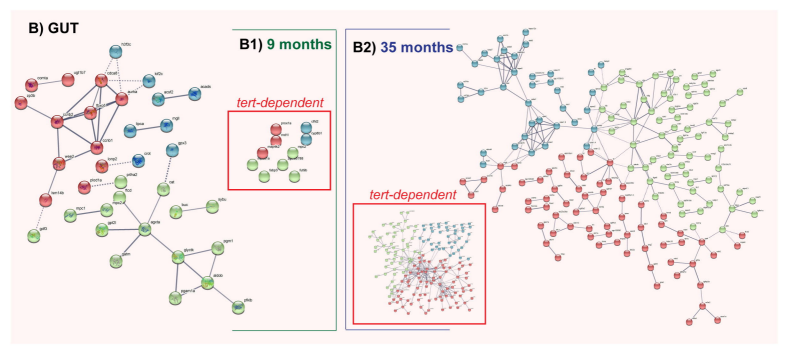


WT 35 months x

tert 2	2 months	tert -/- 9 months	tert 🗠 22 months	tert * 9 months tert * 22 months			hs	
acat1	sb:cb37	acsf2	acat1	anIn	тст6	acaa1	h3f3b.1	plp1b
ahsq1	serpinf2b	cyp8b1	angpt/3	arhgap11a	mybl2b	anIn	h3f3c	ppp1r14ba
mbp	serping1	hbaa2	comtb	aspm	ncapd3	asf1ba	hapin1a	prc1b
	si:ch211-262h13.	5 mgll	f10	aurkb	ncapg	aspm	hbaa2	prdm8
poa1b	si:ch211-284e20.8		f7	ccna2	ncapg2	aurkb	hbba2	primpol
mp1a	slc22a2	pla1a	fam83d	ccnb1	ncaph	b2m	hmmr	psmb8a
1r	slc27a1b	sephs1	gatm	cd40	ndc80	batf	igf2bp1	psme2
3a.6	slc38a3a		gbp3	cdc20	neurod4	birc5a	il2rb	pyyb
3b.1	slc38a4		gcat	cdk2	nusap1	ccl38.6	inab	rac2
3b.2	slc7a2		ggt5a	cdk6	orc3	ccna2	insm1a	racgap1
8a	slco1d1		glyctk	cenpe	plp1b	ccnb1	irf8	rad54l
8b	tdo2a		gpt	chaf1b	primpol	ccnjl	kif11	rrm2
9	tfa		hpn	cyp24a1	rrm2	cd40	kif14	sgo1
es2a	ttr		itIn2	dbx1a	sgo1	cd74a	kif20a	si:busm1-266f07.
es3	ttyh3b		me1	dld	si:ch211-244o22.2	cdc20	kif2c	si:ch211-244o22.
omtb	vtnb		mlsl	dnmt3aa	smc4	cdk2	kif4	smarcb1b
yp8b1	zgc:153031		mpc1	dnmt3ba	socs1a	cdk6	kifc1	smc4
pys	zgc:174260		nr0b1	ercc6l	sox4a	cenpe	Igals9I1	sox11a
2	0		nrg1	f10	sox4b	ch25h	lifrb	sox4a
7			oaz2a	gadd45gb.1	spc24	chaf1a	mad2l1	sox4b
7i			pcbd1	hapIn1a	spc25	chaf1b	marcksa	spc24
ah			, pcxa	hsp90aa1.1	spi1a	chek1	mbpb	spi1a
n1b			pfklb	igf2bp1	tacc3	ckap2l	mcm2	spinb
urinb			prox1a	insm1a	top2a	csf1b	mcm4	stat4
abp2			si:ch211-262h13.5	kif11	tpx2	cxcl18b	mcm5	tacc3
ao1			si:ch211-284e20.8	kif14	ttk	dbx1a	<i>тст</i> 6	tap2a
bba1			slc16a6a	kif20a	tubb5	dla	mhc2a	tlr2
pn			slc1a4	kif2c	tyms	dlb	mybl2b	top2a
f2a			slc27a1b	kif4	uhrf1	dld	ncapd3	tpx2
gfbp2a			slc38a3a	kifc1	unga	dnmt3aa	ncapg	ttk
cn15			slc38a4	lgals9l1		dnmt3ba	ncapg2	tuba8l3
pca			slc43a1b	lifrb		ercc6l	ncaph	tubb5
nst1			slc7a2	mad2l1		fosab	ndc80	tyms
r0b1			slco1d1	marcksa		foxn4	nefmb	ube2c
xct1b			ttr	mbpb		gadd45gb.1	neurod4	uhrf1
glyrp2			ttyh3b	mcm2		grem2b	nusap1	wasb
la1a			urahb	mcm5		h2af1al	plp1a	wdhd1
prox1a			vkorc1					zwi
orozb			zgc:92040					
sox1			0					

Protein-protein interaction network and cluster analysis of changes in WT ageing (STEM and All DEGs combined)





C) BRAIN

