Title: Dietary intervention improves health metrics and life expectancy of the genetically obese DU6 (Titan) mouse

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Abstract

Suitable animal models are essential for translational research, especially in the case of complex, multifactorial conditions, such as obesity. The outbred mouse line Titan (DU6) results from the world's longest selection experiment for high body mass and was previously described as a model for metabolic healthy (benign) obesity. The present study deeper characterized the genoand phenotypes of this outbred mouse line and tested its suitability as an interventional obesity model. In contrast to previous findings, our data suggests that Titan mice are metabolically unhealthy obese and short-lived. Line-specific patterns of genetic invariability are in accordance with observed phenotypic traits. Titan mice show modifications in the liver transcriptome, proteome and epigenome that are linked to metabolic (dys)regulations. However, dietary intervention partially reversed the metabolic phenotype in Titan mice and significantly extended their life expectancy. Therefore, the Titan mouse line is a valuable resource for translational and interventional obesity research.

Keywords

Obesity, metabolic diseases, translational animal model, dietary intervention, lifespan, transcriptomics, proteomics, histone acetylation

Introduction

Animal models are essential tools for translational biomedical research. The identification of suitable animal models is critical for the successful translatability of pre-clinical data^{1–3}. Each animal model presents limitations and that the use of complementing models might highlight particular aspects of a disease. This is critical to model complex, multifactorial diseases, such as metabolic disorders and obesity. The genetic homogeneity of inbred lines, for instance, represents both a strength and a limitation for animal models. While they are useful experimentally by limiting the inter-animal variability, inbred lines might not reflect the genetic heterogeneity characterizing some human diseases^{1–3}. A model based on the selection of phenotypic traits (e.g. obesity) may provide novel insight into complex disorders^{1.4}.

An example of such selection line is the Dummerstorf "Titan" mouse. Titan (also called DU6) is a unique mouse model resulting from an ongoing longterm breeding scheme (currently over 180 generations) that selected mice based on high body mass at six weeks of age⁵. The initial population of mice was created in 1969/1970 by crossing four outbred and four inbred populations^{6,7} creating a non-inbred mouse line (FZTDU) with a polygenetic background. Based on this polygenetic mouse line, the selection for high body mass started in 1975 with every generation consisted of 60-80 mating pairs^{8,9} (see Methods). Importantly, an unselected population was maintained as a control group throughout the selection process⁹.

Several phenotypic and molecular analyses of the Titan mice were conducted over the course of the selection experiment^{5,10–13}. It was reported that 6-week-

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old Titan mice from earlier selected generations displayed high body mass, as well as elevated levels of fat, insulin, leptin and growth hormone^{5,10,13}. An epididymal fat gene expression study identified 77 differentially expressed genes (between Titan and control mice) involved in regulatory and metabolic pathways¹¹. A more recent study showed that Titan mice (generation 146) displayed impaired glucose tolerance six weeks of age, while surprisingly, alucose tolerance progressively improved with age compared to unselected control mice⁵. Within the scope of this observation, the Titan (DU6) mice have been proposed as a mouse model to specifically study benign obesity⁵. While metabolically healthy obesity (MHO) is characterized by preserved insulin sensitivity, further characteristics are important to distinguish between MHO to metabolically unhealthy obesity (MUO)¹⁴. This includes, but is not limited to, lipid profile, liver function, cardiovascular disease, physical activity and healthy life span¹⁴. Importantly, various obesity-associated comorbidities develop over time and become more prevalent in the later stages of life. Comprehensive analyses of Titan mice throughout their life are lacking¹⁵.

In this work, we set out to test the hypothesis that the obese Titan mice developed typical hallmarks of pathological detrimental obesity (MUO)¹⁴. To address this hypothesis, we conducted a deep phenotypic characterization of the mice and investigated factors involved in detrimental obesity in humans or other model species such as plasma levels of cholesterol and triglycerids¹⁶, Leptin^{16–18}, FGF21^{19,20}, heart morphology²¹ and whitening of adipose tissue²². Since body mass and lifespan are two traits known to be inversely correlated to each other²³, we also assessed whether the lifespan of Titan mice was shortened compared to control mice. In addition, we used recently published

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genomic sequence data to link several phenotypic features of the mouse line to their specific genetic differentiation caused by the selection procedure⁹.

Furthermore, since the liver is an organ central to lipid and carbohydrate metabolism, we evaluated possible molecular alterations in liver tissue. Specifically, we examined transcriptomic²⁴, epigenetic and proteomic alterations that might be associated with obesity and diet. Importantly, dietary intervention is a well-studied and robust intervention to promote healthy lifespan^{25–27} and to improve obesity phenotypes^{4,28} in various animal models. Since not all mouse strains are responsive to caloric restriction^{29,30}, we also tested whether the Titan mice might be amenable to therapeutic intervention, using a late and moderate diet intervention as a proof-of-principle.

Our data support the notion that Titan mice exhibit detrimental obesity (MUO), rather than benign obesity (MHO). Titan mice display genetic variability in genomic regions containing genes associated with several regulatory pathways, notably metabolic regulation. The Titan mice exhibit a shorter lifespan, along with molecular changes in liver. Importantly, dietary intervention in Titan mice partly reversed the obesity, transcriptomics phenotypes and lifespan supporting the idea that Titan mice might be suitable as a translational and interventional model of obesity.

Results

This study was conducted in male Titan (DU6) mice to avoid sex-specific confounding factors. Titan mice aged between 10 and 21 weeks were used for genomic and phenotypic analyses.

Titan mice display increased size, hallmarks of metabolically unhealthy obesity and reduced lifespan

Weight development, gross morphology and abdominal fat distribution

We first conducted a thorough characterization of the phenotype in order to support or refute the notion that the obesity in Titan is benign (MHO vs MUO)⁵.

We started by revisiting the phenotypic characterization of the Titan mice, now at generation 180 of the breeding scheme. Titan mice had an average weight of 90 g at the time of selection (6 weeks of age) vs. 35 g for unselected control mice (Figure 1A). Adult Titan mice further grew to an average of 110-115 g (Figure 1B) and their body and skeleton were substantially larger than those of controls (Figure 1C). At 10–11 weeks of age, Titan mice reached 14.75 cm in body length (Figure 1D). Notably, Titan mice exhibited an increase in both total fat and lean mass (Figure 1E), as well as in fat percentage (Figure 1F). Fat distribution was also altered, with Titan mice accumulating as much as 4% intra-abdominal fat of total body weight compared to 1% in control mice (Figure 1G). Overall, our data confirmed the large and obese phenotype of the mice but also demonstrate an increase in body mass compared to previous data^{5,10}.

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Hallmarks of detrimental obesity

Since obesity is often linked with hyperlipidemia¹⁶, we measured the cholesterol and triglyceride plasma levels in Titan mice. We detected significantly higher fasting plasma levels of triglycerides, non-HDL and HDL cholesterol in Titan compared to control mice at 18-20 weeks of age (Figure S1A-C). Consistent with the notion that hyperlipidemia usually occurs in the presence of insulin resistance, insulin levels were more than 3-fold higher in ad libitum 16-17-week-old Titan mice (Figure S1D). We also measured average levels of leptin, a hormone secreted by adipocytes and known to modulate food intake and fat storage¹⁶⁻¹⁸. Leptin levels were significantly higher (more than 5-fold; P < 0.001) in 16-17-week-old Titan mice compared to control mice (Figure S1E). Finally, we found that the plasma concentration of fibroblast growth factor 21 (FGF21), a factor associated with metabolic syndrome (MetS) in humans^{19,20}, was elevated in the serum (Figure S1F).

As metabolically unhealthy obesity (MUO) can cause heart disease, we also assessed histological alterations in the heart of the Titan mice^{14,21}. Using Sirius Red staining, we detected fibrotic tissue in the hearts of three out of six 16-17-week-old and five out of six 24-26-week-old Titan mice (Figure S2). Furthermore, whitening of adipose tissue has been associated with obesity and obesity-related diseases²². Indeed, at 16-17 weeks of age, Titan mice showed a substantial whitening of brown adipose tissue (BAT), not observed in control animals (Figure 1H). In addition, Titan mice showed multifocal fatty cells in the pancreas, termed pancreatic lipomatosis or steatosis, without atrophic or inflammatory changes of the adjacent parenchyma, a phenotype not presented in control mice (Figure 1I).

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The lifespan of Titan mice was dramatically shorter than that of control mice (Log Rank test; p<0.0001, χ^2 =124.9) (Figure 1J). Under Standard Breeding Food (SBF), Titan mice reached 10% and 50% mortality at 125 and 325 days, respectively. Their average and maximal lifespans were 317.5 and 614 days (Table S3). In comparison, control mice showed double mean lifespan, reaching 10% and 50% death at 307 and 645 days, respectively. The shortened lifespan of Titan mice combined with obese metrics suggest that the Titan mice suffer from MUO rather than exhibiting MHO.

Association between phenotypic features of Titan mice and regions of distinct genetic differentiation

Based on the phenotypic analysis, we hypothesized that several genetic alterations in Titan mice would translate to the observed phenotype (metabolic regulation, weight and size). We first assessed the genetic differentiation associated with the Titan mouse, in comparison to the control unselected line but also to other Dummerstorf long-term selected mouse lines⁹. For this, we used the whole genome sequencing data recently generated from the 'Dummerstorf mouse lines'⁹. We first re-analyzed regions of distinct genetic differentiation (RDD) unique to Titan mice and identified 84 genes (Table S1). Since both Titan and DU6P are giant mice (see ref ⁹), an overlap of RDD was expected between both lines. A second analysis excluding the DU6P mice was conducted and identified 173 protein coding genes (335 including non-coding genes) in RDD for Titan mice (Table S2). Gene Ontology (GO) enrichment analysis (Biological Process) of these Titan-specific RDD genes

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revealed an association with several biological processes, including metabolic regulation (*Kat2a*, *Sfrp5*, *Agpat4* and *Hcrt*), growth factor signaling (*Stat3*, *Stat5a*, *Stat5b* and *Igf2r*), skin differentiation and immune regulation (Table 1).

Next, we sought to further explore the link between genotype and phenotype. Increased skin thickness has been shown to positively correlate with increased BMI³¹. As expected, Titan mice displayed a thicker dermis of the skin compared to the control group (Figure S3A). We also found that the inflammatory cytokines interleukin 6 (IL-6) and tumor necrosis factor-alpha (TNF α) were significantly increased in the plasma of Titan mice compared to unselected control mice (Figure S3B and C). As a further indication of inflammation, histological analysis of the thymus revealed the presence of nodular thymic medullary hyperplasia in Titan mice (Figure S3D). Immunohistochemistry of these medullar nodes revealed that they were composed mainly of B cells (Figure S3D). Thus, our data supports a link between Titan-specific RDD such as metabolic regulation, growth, skin differentiation and immune regulation genes with Titan-specific phenotypes of obesity, size, thicker dermis and increased hallmarks of Inflammation.

Altered histone 4 acetylation levels in Titan mice

Genetics and epigenetics functions are intimately linked, and their alterations are associated with disease³². We identified the histone acetyl transferase *kat2a* (*Gcn5*), as one of the genes associated with specific genetic differentiation in Titan mice (Figure 2A, Table S1; Allele frequency = 0.97). Kat2a is a histone acetyl-transferase, which has also been implicated in metabolic and energy regulation³³, thus raising the possibility that histone

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acetylation might be altered in Titan mice. To address this possibility, we compared histone 4 (H4) acetylation levels of 10-week-old and 19-20-week-old Titan vs. control mice by mass spectrometry^{34,35}. For this analysis, we focused on the liver as it is an organ central to lipid and carbohydrate metabolism and KAT2a is involved in regulation of metabolic activity in the liver³⁶.

At 19-20 weeks of age, we observed significantly lower levels of H4 acetylation in Titan vs. control mice (Fig 2B). Highly abundant H4 acetylation motifs in mammalian such as H4K16 and H4K12K16³⁵, as well as poly-acetylated motifs were significantly reduced in Titan vs. control mice (Fig 2C,D). By contrast, H4K5 acetylation motif, which was shown to be moderately regulated at least in part by Kat2a³⁴, was significantly higher in Titan mice compared to control mice (Fig 2C). Of note, comparable results were obtained in younger Titan mice (10 weeks of age, data not shown). Taken together, Titan mice display considerable alterations in various histone acetylation motifs compared to the unselected mouse line which might manifest in altered transcriptomes³⁷.

Hybrids of Titan and unselected mice show intermediate phenotypes

We next assessed the potential contribution of the maternal impact and phenotypic features of Titan mice. In order to address this we generated hybrids by crossing Titan and control unselected mice (mother control/Titan x father Titan/control respectively). Male F1 hybrid of 10-11 weeks of age

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showed similar weight and fat % independently of the genotype of the mother (Fig 3A).

Notably, the combined population of male F1 hybrids showed intermediate body weight and fat percentage compared to control and Titan mice at the same age (Figure 3B). Similarly, the hybrid mice showed intermediate H4 acetylation levels in the liver, including at H4K5 and H4K16 (Figure 3C). Furthermore, the lifespan of hybrid mice, although improved compared to Titan mice, was reduced compared to that of control mice (Hybrid vs Titan Log Rank test; p <0.0001, χ^2 =96.45 Control vs hybrid, Log Rank test; p =0.0362, χ^2 =4.385). The median life span of 546 days vs 325 (Titan) and vs 645.5 days (control)) (Figure 3D). These observations illustrate the quantitative genetic background determining mice obesity and life span.

Titan mice display altered liver metabolism along with changes in gene and protein expression

We next evaluated possible alterations in liver function and metabolism in Titan mice. First, alanine aminotransferase (ALAT) and alkaline phosphatase (ALP) activity, markers of potential liver damage³⁸, were significantly elevated in Titan mice at 16-17 weeks of age (Figure 4A and B). Combined Oil red O and Hematoxylin staining of the liver also revealed numerous small fat droplets in Titan mice (Figure 4C). However, no clear signs of hepatic steatosis were found by H&E staining of 16-17-week-old Titan liver tissue (Figure S4). Bilirubin, a waste product of heme metabolism in the liver, was significantly lower in Titan mice compared to controls (Figure 4D). In addition,

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plasma urea levels were lower in Titan than in control mice (Figure 4E), pointing to the dysregulation of the urea cycle and protein metabolism in the liver. Altogether, these findings suggest an altered liver metabolism in Titan compared to control unselected mice.

We hypothesized that Titan mice, unlike control mice, would display transcriptome profiling that is linked to increased body fat phenotype and altered metabolism. To identify pathways altered at the transcriptional level in the liver, we performed RNA sequencing on the liver from 11-week-old and 19-21-week-old mice. Principal component analysis (PCA) revealed distinct Titan and control transcriptomes (Figure 5A). Gene expression in younger and older controls clustered together, but in Titan mice, the expression profile shifted between the age groups (Figure 5A), indicating age-dependent effects in Titan but not in control mice.

Compared to age-matched controls, 11-week-old Titan mice exhibited 1150 upregulated and 1227 downregulated genes (Figure 5B, S5A and Table S4) and 19-21-week-old Titan mice 693 upregulated and 828 downregulated genes (Figure 5B and S5D). We observed a significant overlap of transcriptomic alterations between both age group comparisons, with 402 upregulated and 461 downregulated genes.

We next compared 11-week-old Titan to control mice. The top 10 upregulated GO terms highlighted genes involved in several central metabolic processes (Figure 5C). Central metabolic regulators such as *mTOR*, *ElovI5*, *ElovI6*, and acetyl-CoA carboxylase were upregulated (Table S4), suggesting altered fat metabolism in the liver. The top 10 downregulated GO terms included

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xenobiotic, amino acid and sulfur metabolism (Figure 5C). For example, many cytochrome P450 family genes, including *Cyp2c37*, were downregulated in 11-week-old Titan mice compared to controls at the same age (Table S4). Interestingly, cytochrome P450 family genes are downregulated in a high-fat diet²⁴. Furthermore, several genes related to the methionine, folate cycle and H_2S production [e.g., *Bhmt, Gnmt, Cth (Cgl), Dmgdh*], as well as *Hcrtr2* were downregulated in 11-week-old Titan mice.

The top 10 upregulated GO terms in 19- 21 week-old Titan mice compared with control revealed the upregulation of genes involved in neuronal differentiation and peptidyl-tyrosine phosphorylation activity. (Figure 5D). The top 10 downregulated GO terms in 19-21-week-old Titan mice revealed genes involved in the regulation of xenobiotic, amino acid and sulfur metabolism (Figure 5D), similar to the comparison at 11 weeks of age (Figure 5C). Of note, genes involved in mitochondrial organization and mitochondrial respiratory complex assembly were downregulated as well (Figure 5D). We found that *Acss2*, an acetyl-CoA synthesis enzyme linked with histone acetylation³⁹, was downregulated in Titan mice.

Compared to 11-week-old Titan mice, 19-21-week-old Titan mice had increased expression of several enzymes involved in glucose and insulin response (See supplementary data). Also, *Cyp7b1*, which encodes an enzyme involved in cholesterol catabolism that converts cholesterol to bile acids and is upregulated by high-fat diet²⁴, was upregulated in 19-21-week-old Titan mice compared to 11-week-old Titan mice. On the other hand, the top 10 downregulated GO term highlighted genes involved in fatty acid, coenzyme and carbohydrate metabolism (Figure S5C). Taken together, liver

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transcriptomic analysis demonstrate several metabolic alterations in Titan mice.

Next, we compared the liver proteomes of Titan and control mice. As with the RNAseq data, larger proteomic changes were seen in the 11-week-old Titan to control comparison than in the 19-21-week-old comparison (Figure S6). Relative to age-matched controls, 11-weeks-old Titan mice exhibited 207 upregulated and 289 downregulated proteins, while 19-21-week-old Titan mice showed 125 upregulated and 142 downregulated proteins (Figure S6). Consistent with the transcriptome results, pathway analysis revealed increased fatty acid metabolism, lipid catabolism, and coenzyme metabolism, notably pathways involved in fat utilization (ACACA, IDH1 and ACOX1) and in lipid biosynthesis (FASN and ACSL5), while downregulation in xenobiotic, amino acid and sulfur metabolism in 11-week-old Titan mice (Figure S6).

Proteins involved in lipid metabolism were also enriched in 19-21-week-old Titan mice compared to controls (Figure S6), although this was not observed in the RNAseq top 10 GO terms (Figure 5D). We also found that proteins upregulated in 19-21-week-old Titan mice were associated with carbohydrate catabolism and coenzyme metabolism (Figure S6), while downregulated proteins were enriched in amino acid and sulfur metabolism (Figure S6), in line with the RNAseq analysis (Figure 5D). As with the transcriptomic data, we observed a reduction in the abundance of BHMT, GNMT, and CTH (CGL).

Collectively, both transcriptomics and proteomics data hint towards increased fat metabolism in Titan mice compared to control mice.

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Late dietary intervention at 12 weeks of age improves the obesity and lifespan phenotypes in Titan mice

Finally, we investigated the impact of dietary restriction on measures of obesity, epigenetic markers, liver gene expression patterns and lifespan of Titan mice. We compared the impact of a diet with moderate energy reduction using Energy Reduced Food, hereafter referred to as ERF (see Methods), to that of Standard Breeding Food (SBF). We introduced ERF at 12 weeks of age onward and termed this diet intervention as late intervention due to the fact that 10% and 25% of the Titan population are already dead by 18 and 33 weeks, respectively.

ERF intervention resulted in a weight reduction of 5–10% in Titan mice, compared to SBF-fed siblings (Figure 6A; MWU-test, P=0.0017). By 21 weeks, ERF-fed Titan mice had significantly reduced abdominal fat percentage compared to SBF-fed mice (Figure 6B; MWU-test, P=0.0313). We also showed that the ERF regime lowered the levels of plasma cholesterol, HDL and glucose in Titan mice (Figure 6C). Furthermore, plasma levels of leptin, glycerol and non-esterified fatty acids (NEFA) were decreased under ERF regime (Figure 6C).

Late ERF intervention in Titan also resulted in a general increase in polyacetylated motifs of histone 4 levels, which resembled the levels observed in control mice (Fig. 6D). In general, motifs containing K12K16 acetylation motifs (Significant H4K12K16 and H4K5K12K16) were increased in ERF-fed compared to SBF-fed Titan mice (Figure 6D). We next tested whether the ERF regime affected the expression level of liver genes that were already identified in

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our omics experiments as significantly altered in Titan compared to control mice (Figures 4 and S6, See supplement list of genes). In addition, late ERF intervention reversed the expression profiles of some of the investigated genes. In fact, genes that were downregulated in Titan versus control mice (e.g. *Dmdgh, Gnmt, Cth, Cyp2c37*) were upregulated in ERF-fed vs SBF-fed Titan mice (Figure 6E)²⁴. Similarly, genes which were upregulated in Titan versus control mice (e.g. *Cyp7b1*) were downregulated in ERF-fed vs SBF-fed Titan mice (Figure 6E).

Finally, we investigated the impact of the ERF diet on the lifespan of Titan and control mice. The ERF diet significantly increased the lifespan of both Titan and control mice (Log Rank test; for P=0.0087, χ^2 =6.892 Titan mice and P=0.01, χ^2 =6.66 for control mice) (Figure 6F). ERF-fed Titan mice reached the 10% and 50% death at 219 and 374 days, respectively, which is significantly longer than for the SBF-fed Titan mice (10% death at 125 days and 50% death at 325 days). In comparison, ERF-fed control mice reached 10% death at 377 days vs 307 days for SBF-fed control mice. The average lifespan of ERF-fed Titan mice (Table S3).

Taken together, our results indicate that a late ERF intervention could partly revert the phenotypes of Titan mice associated with MUO and lifespan.

Discussion

This study provides a comprehensive phenotypic analysis of the Titan mouse line DU6 and investigated the link between Titan's unique phenotype to their

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genotype. We demonstrate that Titan mice display various physiological and molecular alterations that may underlie their metabolically unhealthy obesity phenotype. We also show that Titan mice are short-lived, and that the obesity and lifespan phenotypes are responsive to and can be partly reversed by diet intervention.

We provide several links between Titan mice's genotype and their unique phenotypes. Gene Ontology (GO) enrichment of regions of distinct genetic differentiation (RDD) highlighted genetic variability associated with genes involved in multiple pathways including metabolic regulation and growth control. Similarly, RDD genes involved in skin differentiation correlated with our observation that Titan mice display a thicker dermis of the skin compared to control mice, which is in line with previous observations linking increased BMI with thicker skin³¹. Several RDD genes linked to immune regulation and inflammation (notably Stat3 and Stat5 family members) were also identified⁴⁰, in correlation with the immune and inflammatory phenotype of Titan mice. For example, Titan mice expressed markers of systemic inflammation, such as high plasma levels of IL-6 and TNF α or thymic medullary hyperplasia. Besides being associated with the initiation and progression of multiple diseases^{41,42}. elevated circulating IL-6 and TNFa were documented in obese mice and humans^{43,44}. Supporting this finding, several mouse models of obesity revealed increased accumulation of macrophages in the liver^{45,46} or in adipose tissue⁴⁷. Increased tissue inflammation in Titan mice might contribute to tissue damage³⁸.

Notably, one identified RDD contained the *Hcrt* gene, which is a hypothalamic neuropeptide that can regulate feeding behavior and metabolic

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homeostasis^{48,49}. While we did not evaluate gene expression levels in the brain area that may account for metabolic alterations in the Titan mice, we observed a significant reduction of *Hcrtr2* in the liver. Intriguingly, *Kat2a*, which encodes the histone acetyltransferase GCN5, was also part of a RDD specific to Titan mice and is involved in metabolic regulation³³. This correlated with an increase in H4K5 acetylation in the liver of Titan mice, suggesting a possible causal link between the two phenotypes³⁴. However, we did not detect significant changes in the expression level of *Kat2a* in the liver of Titan mice. Although it is unlikely that changes in H4K5 acetylation are the consequence of changes in *Kat2a* enzymatic activity or to alterations in other GCN5-related genes. Additional investigations will be necessary to address this question and further assess alterations in expression and activity of histone acetyltransferases in Titan mice, in correlation with changes in histone acetylation levels.

Notably, F1 hybrids show intermediate body weight and fat percentage, H4 acetylation levels and lifespan. The intermediate phenotype is independent of the mother's genotype (Control or Titan), thus demonstrating a weaker link between maternal inheritance/ care to weight and fat levels. Further analyses of F2 hybrids are required to link specific genes to the specific phenotypes of the Titan mice described in this study^{4,50}.

Previous research at earlier generations suggested that Titan mice display benign obesity (MHO) without additional comorbidities⁵. In contrast, we now show that Titan mice indeed exhibit detrimental obesity, including several hallmarks that are attributed to metabolic syndrome (MetS), such as high

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levels of fat in tissues, high plasma levels of fasting triglycerides, fasting cholesterol, FGF21^{19,20,51} and insulin^{16,52} as well as early signs of heart fibrosis²¹. Titan mice also showed whitening of BAT, and ectopic fat in the pancreas, both being linked to obesity and MetS^{22,53}. Increased pancreatic fat might have implications for the onset of type 2 diabetes⁵⁴, which is frequently associated with insulin resistance²¹ .Importantly, obesity is known to be associated with reduced life expectancy and various conditions related to aging, such as inflammation^{55–60}. Indeed, we found that Titan mice exhibited a shorter lifespan compared to control mice, which is in line with a recent study showing that female Titan mice show a reduced lifetime fecundity compared to the unselected control mice⁶¹.

Alterations in liver function were evidenced by epigenomics, transcriptomic and proteomic changes detected in the liver of Titan mice. Genes and proteins involved in lipid metabolism and biosynthesis were consistently upregulated in Titan mice compared to control mice, which might account for the high plasma levels of cholesterol and fasting triglycerides observed in this model. On the other hand, compared to control mice, Titan mice exhibited several downregulated genes and proteins associated with xenobiotic metabolism, such as cytochrome P450. These downregulations might indicate altered endogenous metabolite metabolism⁶², increased liver inflammation^{63,64} and/or a generally reduced capacity to deal with toxic compounds in the liver. Xenobiotic metabolism strongly impacts the ability of the organism to maintain homeostasis and cope with disease⁶⁵, which may contribute to the increased morbidity of these mice. Also, Titan mice showed a reduced expression of genes related to the folate cycle and its downstream effectors, such as *Ahcy*,

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Gsta family *Gnmt* and *Cth*. Such genes were also shown to be downregulated in the mouse liver upon high-fat diet²⁴. Similarly, genes involved in mitochondrial organization and function were downregulated in 19-21-weekold Titan mice, which may hint towards altered mitochondrial activity in older Titan mice⁵⁸. Altogether, these findings suggest an altered liver metabolism in Titan compared to control mice, along with their obese phenotype and shorter lifespan.

Interestingly, the liver of Titan mice showed overall decreased H4 acetylation at the 4-17 motif, specifically histone acetylation motifs such as H4K16 and H4K12K16 acetylation, which is in correlation with a reduction in the expression of acetyl-CoA synthases (*Accs2* and *pdhx*) which may imply reduced acetyl-CoA availability. This may underlie the general alteration in histone acetylation levels^{39,66} and might also link this epigenetics phenotype to lifespan regulation⁶⁷.

Our data support the notion that Titan mice are a potential translational model, as ERF feeding introduced at 12 weeks of age could partially reverse the phenotypes associated with obesity, lifespan and histone H4 acetylation patterns. Notably, intra-abdominal fat was reduced at 21 weeks and plasma levels of leptin, glycerol and non-esterified fatty acids (NEFA), which are linked to obesity^{16,17,68,69}, were decreased in ERF-fed Titan mice. This phenotype improvement is reminiscent of other time-restricted fasting animal models or patients with metabolic syndrome⁷⁰. ERF feeding appears to specifically increase levels of poly acetylated histone 4 motif which include K12K16 acetylation. These modification might impact transcription regulation that are important for mediating the benefits of healthy diet in obese animals

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and maybe novel therapeutic targets to treat obesity. In line with these data, we also found that ERF feeding partially reversed the expression pattern in the liver, including metabolic key enzymes (e.g. *Dmdgh, Gnmt, Cth, Cyp2c37*)²⁴. Several of these genes were implicated in mediating the benefits of caloric restriction^{71–73}. For example, CTH (CGL) mediates the dietary restriction response in mice via H₂S production⁷⁴. Similarly, food intake restriction increases GNMT levels, promotes energy homeostasis, and increases healthy lifespan^{73,75}. Altogether, our data demonstrate that Titan mice are responsive to diet intervention, and thus may represent a suitable translational mouse model to assess the impact of diet or other therapeutic interventions on manifestations of obesity^{16,29,76}.

The poly-genetic Titan mouse model may present several advantages over other currently used mouse models of unhealthy obesity. Notably, no further genetic alterations or elimination of regulatory genes are needed to achieve the obesity phenotype in Titan mice. Most genetic rodent models of obesity are based on the disruption of the leptin signaling pathway, including the mouse Leptin-deficient (ob/ob), the mouse leptin receptor-deficient (db/db), the rat Zucker Diabetic Fatty and DahlS.Z-Lepr^{fa}/Lepr^{fa} (DS/obese), the Koletsky rats carrying a nonsense mutation in the leptin receptor, or the POUND mice (C57BL/6NCrl-Lepr db-lb/Crl)^{16,76,77}. A major disadvantage of these rodent models is that they do not closely reflect human obesity, which is multifactorial and not always affecting leptin signaling, thus rendering it difficult to translate findings in these rodents to human patients. Another advantage of the Titan mice, notably when studying the impact of obesity on lifespan phenotype, is that ERF diet intervention could be already assessed in

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4-5 months Titan mice, whereas similar intervention would take much longer in other mouse models²⁹

In conclusion, the Titan mouse line represents an important scientific achievement in breeding of mice, making Titan mice one of the biggest laboratory mouse lines ever described. The detailed genomic, epigenetic and phenotypic analyses described here strongly support the Titan line as a relevant model to study various aspects of interventions in unhealthy obesity. The present phenotypic analysis provides a strong basis for further characterization of the Titan model as a novel tool for studying and potentially developing pharmaceutical interventions targeting obesity and associated disorders.

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Materials and methods

Animals and housing conditions

All procedures were performed in accordance with national and international guidelines and approved by our own institutional board (Animal Protection Board from the Leibniz Institute for Farm Animal Biology). At the German Mouse Clinic, mice were maintained in IVC cages with water and standard mouse chow according to the directive 2010/63/EU, German laws and GMC housing conditions (www.mouseclinic.de). All tests were approved by the responsible authority of the district government of Upper Bavaria.

At the FBN, the animals were maintained in a specific pathogen-free (SPF) environment with defined hygienic conditions at a temperature of 22.5±0.2°C, at least 40% humidity and a controlled light regime with a 12:12-h light-dark cycle. The mice were kept in Polysulfon-cages of 365 x 207 x 140 mm (H-Temp PSU, Type II L, Eurostandard, Tecniplast, Germany) and had free access to pellet concentrate and water. Mice were fed ad libitum using a standard breeding food (SBF, GE=16.7 MJ/kg, ME= 14.0 MJ/kg) including 12% fat, 27% protein and 61% carbohydrates (ssniff® M-Z autoclavable, Soest, V1124-300, Germany).

For the energy-reduced survival experiment, the mice were fed with a mouse maintenance energy-reduced diet (ERF, GE=15.7 MJ/kg, ME= 11.7 MJ/kg) characterized by a low energy density and high fiber contents including 10% fat, 21% protein and 69% carbohydrates (ssniff® M-H autoclavable, Soest, V1574-300, Germany). Mice were placed under the ERF diet starting at 12 weeks of age.

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In both control and Titan lines, the litter size was standardized to 10 (5 male / 5 female pubs) immediately after birth and until weaning occurs at the age of 21 days. At the age of three weeks, males of different litters were grouped in a cage at three animals per cage. The first survival experiment was done with n=104 Titan males from generation 181 and n=96 control males originating from generation 192. The second survival experiment under the ERF diet started with n=103 Titan and n=102 control males, one generation later. Additionally, 27 Titan male full siblings pairs of generation 183 were divided into two contemporaneous groups and were fed with SBF or ERF after 12 weeks of age to generate the samples to analyze metabolic parameters. Individual weights of all animals were taken regularly every three weeks, starting with the age of 21 days. Hybrid mice were generated by crossing unselected control mice to Titan mice (Titan mother x control father or control mother x Titan father).

During the survival experiments, all included males were observed daily for their health condition. If physical impairments were detected, which would cause considerable suffering or substantial pain to the animals, they were sacrificed and such incidents were documented accordingly.

Origins of the growth-selected strain and the control strain.

We used mice of an unselected strain (FZTDU) as control and a strain selected for high body mass at day 42 of age (DU6/Titan), both bred at the Leibniz Institute of Farm Animal Biology (FBN), Dummerstorf, Germany.

The initial population of mice was created in 1969/1970 by crossing four outbred (NMRI orig., Han/NMRI, CFW, CF1) and four inbred (CBA/BIn,

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AB/BIn, C57BL/BIn, XVII/BIn) populations⁷. Mice of the control line FZTDU used in this experiment were mated randomly over about 192 generations with a population size of 100 to 200 mating pairs per generation, respectively. Four generations of the control line are generated yearly using a rotation procedure of Poiley (1960) to avoid inbreeding⁷⁸.

The growth selection started in 1975, thus creating the Dummerstorf growth line DU6 (Titan) by selection for high body mass by sib selection in a conventional environment. In every generation, four generations per year, 60-80 mating pairs were made at the age of 63 \pm 3 days^{8,9}. The selection procedure as described above was maintained for 153 generations. Only seven males and seven females of the DU6 line belonging to generation 154 and 155 (year 2011-2012), respectively, were used as new founder animals after embryo transfer in a newly built animal facility. Of note, in 2011/2012, the Titan mouse lines (generations 154 and 155) were transferred into a new state-of-the-art pathogen-free animal facility and their diet composition changed at that point¹². Over the entire term of the following five generations the new breeding population of at least 60 pairs of Titan DU6 mice was established⁹, taking care of an equal distribution of the alleles of the 14 founder animals. In opposition to the former sib selection in generation number 161, breeding value estimation started introducing a two-trait BLUP (Best Linear Unbiased Prediction) animal model for male and female body mass. The raising inbreeding coefficient in the selection line was then controlled by the 'Optimal Genetic Contributions to the Next Generations' method of Meuwissen⁷⁹.

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Histology, plasma, DEXA – GMC

Importantly, because Titan mice are not inbred, we increased the number of animals from 15 (standard GMC animal number per group and sex for inbred mice) to 20, focusing on one sex only (males) to avoid sex-specific confounding factors.

Two cohorts of 20 DU6/Titan and 20 control (FZTDU) male mice were subjected to an extensive phenotypic screening at the German Mouse Clinic (GMC), including standardized phenotyping in e.g. the areas of energy metabolism, clinical chemistry, pathology⁸⁰ (see also www.mouseclinic.de). The phenotypic tests were part of the GMC screen and were performed according to standardized protocols as described before^{81–83}. Variations in protocols are specified. Depending on the test performed, animal numbers varied, as indicated in the respective Figure/Table.

Plasma clinical chemistry analyses included determination of blood lipid and glucose levels on freshly collected Li-heparin-plasma samples collected after overnight food withdrawal and measurement of a broad spectrum of parameters including ALAT and ALP activity, urea and bilirubin levels in samples collected in ad libitum fed state during final blood withdrawal. Analyses were performed using an AU480 clinical chemistry analyzer (Beckman-Coulter, Germany) as described previously. Samples collected during final blood withdrawal were frozen and stored at -80°C until the day of analysis.

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The determination of the IL-6, $TNF\alpha$, Insulin, leptin and FGF21 plasma levels was performed with a combined electrochemiluminescence multiplexed assay system (Meso Scale Discovery, MSD, Rockville, MD USA).

X-ray imaging was performed in an UltraFocus DXA system (Faxitron Bioptics, LLC) with automatic exposure control.

Macroscopic examination performed was in combination with histopathological analyses using hematoxylin and eosin (H&E) staining on formalin-fixed paraffin-embedded sections (3 µm) of tissues from 29 described in www.mouseclinic.de/screens/pathology. organs as Immunohistochemistry was carried out in a Leica Bond III (Leica Biosystems) automatic stainer. Heat-induced antigen retrieval was performed with citrate buffer (pH 6) for 30 minutes (AR9961; Bond TM Epitope Retrieval Solution; Leica Biosystems) in 2-µm-thick sections. For the identification of specific cells in the thymus, antibodies against CD3 (Clone SP7; ZYT-RBG024; Zytomed systems) and CD45R/B220 (Clone RA3-6B2; 550286; BD Pharmingen) were employed and the staining was detected with DAB chromogen. The slides were scanned using a Hamamatsu NanoZoomer 2.0HT digital scanner and analyzed by two independent pathologists using NDP.view2 software (Hamamatsu Photonics).

Analysis of genetic differentiation

The genomes of the Dummerstorf mouse lines were sequenced in order to conduct genomic analyses which included the detection of line-specific patterns of genetic differentiation with respect to the control line FZTDU⁹. This was done by averaging the per-SNP genetic differentiation (Fst⁸⁴) in 50Kb

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windows (sliding window mode with size=50Kb and step=25Kb) containing at least 10 SNPs using vcftools v0.1.13⁸⁵. Per-window Fst scores were standardized into z-scores, in order to evaluate the data as deviations from the genomic mean (z-Fst). The X-chromosome was standardized separately. Regions of distinct genetic differentiation (RDD) were defined as highly differentiated windows in DU6 (top 15% of the z-Fst distribution) that also displayed low genetic differentiation (bottom 25% of the z-Fst distribution) in a) all remaining lines or b) the rest of the lines, except DU6P. Genes overlapping this windows were detected and used for enrichment analysis using WebGestaltR⁸⁶ Genomic regions around genes of interest were visualized with Gviz ⁸⁷.

Histone acetylation analysis

Histone acetylation analysis was done as previously described^{34,35,88}. Briefly, 25 mg of liver was used for acid extraction for histone. The resuspended samples were loaded on gels and were stained overnight followed by distaining with water. Gel pieces were processed for mass spectrometry analysis using heavy labeling of d6-deuterated acetic anhydride. Histone peptides were desalted using C18-StageTips, eluted in 80% acetonitrile/0.25% TFA, vacuum concentrated, reconstituted in 25 µL 0.1% TFA and stored at 4°C (short) or 20°C (long).

Desalted peptides (5µl) were injected and separated on an Ultimate 3000 RSLCnano (ThermoFisher) with a gradient from 3 to 32% acetonitrile in 0.1% formic acid over 40 min at 400 nL/min in a 25-cm analytical Aurora C18

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nanocolumn (75µm ID 120 Å, 1.6 µm, Ion Opticks). The effluent from the HPLC was directly electrosprayed into a Q Exactive HF instrument operated in data-dependent mode to automatically switch between full-scan MS and MS/MS acquisition. Survey full-scan MS spectra (from m/z 250–1600) were acquired with resolution R=60,000 at m/z 400 (AGC target of 3x10⁶). The ten most intense peptide ions with charge states between 2 and 5 were sequentially isolated to a target value of 1x10⁵ and fragmented at 27% normalized collision energy. Typical mass spectrometric conditions were: spray voltage, 1.5 kV; no sheath and auxiliary gas flow; heated capillary temperature, 250°C; ion selection threshold, 33,000 counts.

Peak integration was performed with Skyline (Skyline-daily (64bit) 4.2.1.19004) using doubly o triply charged MS1 precursor ions. A list of all precursor ions can be found in the supplementary Table S1. Peaks were selected manually and the integrated peak values (Area) were exported as .csv file for further calculations. After peak integration, the data summarization and statistical analysis were performed in Excel and R³⁴.

Staining of triglycerides in liver tissues

Liver tissue samples embedded in Tissue Tek (Weckert, Kitzingen, Germany) were cryosectioned (8 µm thick) using a Leica CM3050 S (Leica, Bensheim, Germany) cryostat microtome. After fixation in 4% paraformaldehyde / PBS for 1 min at RT, the slides were stained in RedOil solution (1 mg/ml RedOil (#A12989, Alfa Aesar, Karlsruhe, Germany) in 60% Isopropanol) for 10 min and then washed three times in distilled water. The slides were further stained in hematoxylin for 5 minutes and washed for 3 min in fresh tap water before

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embedding with Aquatex (Roth, Germany) and dried overnight. The staining of the triglycerides was visualized with a Nikon Microphot-Fxa microscope (Nikon Instruments Europe B.V., The Netherlands) and an image analysis system (Nikon Digital Sight, DS-L2).

RNAseq

For RNA extraction and library preparation, 50 mg of tissues (liver) were homogenized in Trizol (Thermo Fisher; cat. no. 15596026) and processed according to the manufacturer's instructions. RNA concentration and A_{260/280} ratio were measured with NanoDrop, followed by Bioanalyzer using RNA pico assay kit using manufacturer's protocol. rRNA depletion was performed using NEBNext rRNA Depletion Kit (Human/Mouse/Rat) [NEB #E6310] and library preparation for RNA-sequencing was performed using NEBNext Ultra II Directional RNA Library Prep Kit for Illumina [NEB #E7760] following manufacturer's protocol. Libraries were sequenced on an Illumina HiSeq 1500 instrument at the Laboratory of Functional Genomic Analysis (LAFUGA, Gene Center Munich, LMU).

For RNA-seq data analysis, read mapping of mouse tissue samples to the mouse genome (GRCm38) and counting of reads mapped to genes were performed using STAR v2.5.3a⁸⁹ using parameters --quantMode GeneCounts and providing annotation –sjdbGTFfile Mus_musculus.GRCm38.97.gtf. Aligned reads were filtered for unmapped, multi mapped and ambiguous reads. Reads from histones and Y chromosome were removed. Reads were also filtered if they had low read counts in at least two samples. Differential expression analysis was carried out using DESeq2 v1.24.0⁹⁰ at an adjusted p-

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value cut-off of 0.05. GO term analysis was performed using ClusterProfiler v3.12.0⁹¹ at FDR of 0.05 using Benjamini-Hochberg procedure and with a log fold change cut-off of 0.5. GO terms containing at least a minimum of 10 genes were considered.

All the plots generated for RNA sequencing data were obtained using ggplot2 v3.2.1⁹² unless otherwise stated. For heatmaps and Venn diagram for RNA sequencing and proteomics data, pheatmap v1.0.12 (Kolde, R. (2013). pheatmap: Pretty Heatmaps. R package version 0.7.7. http://CRAN.R-project.org/package=pheatmap) were used respectively with genes (or proteins) passing the adjusted p-value significance of 0.05.

Proteomics

The proteome protocol was adopted from^{88,93} with the following modifications. A total of 200 mg of frozen mice liver was homogenized in 500 µl lysis buffer [50 mM Tris-HCl pH 7.5, 500 mM NaCl, 1 ml EDTA, 0.1% NP-40 and 20% glycerol, 15 mM sodium butyrate, 60 mM of sirtinol and one protease inhibitor tablet (Roche)] and then supplemented with 200 µl 6 M urea/2 M thiourea and 900 µl lysis buffer. To reduce disulfide bonds, samples were treated with 1 mM DTT for 45 min at 4°C, followed by a treatment with 550 mM IAA for 30 min at 4°C in the dark. 1 M ammonium bicarbonate (Ambic) was added to the samples to get a final concentration of 1 M urea. The proteins were digested for 5 hours with Lys-C (Wako) at room temperature and overnight with trypsin (Worthington). Samples were acidified and diluted with TFA to a final concentration of 1% TFA before being loaded on the Sep-Pak Light C18 cartridges (Waters). Columns were washed with 0.1% TFA and eluted with

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60% acetonitrile (ACN)/0.25% TFA. The eluates were dried out by speed vacuum. The pellets were re-dissolved in buffer [50 mM Hepes pH 8.0 and 50 mM NaCl] and the protein concentration was measured by Nanodrop.

LC-MS/MS measurements were performed on an Ultimate 3000 RSLCnano system coupled to a Q-Exactive HF-X mass spectrometer (ThermoFisher Scientific). For full proteome analyses, ~0.25 µg of peptides were delivered to a trap column (ReproSil-pur C18-AQ, 5 µm, Dr. Maisch, 20 mm × 75 µm, self-packed) at a flow rate of 5 µL/min in 100% solvent A (0.1% formic acid in HPLC grade water). After 10 minutes of loading, peptides were transferred to an analytical column (ReproSil Gold C18-AQ, 3 µm, Dr. Maisch, 450 mm × 75 µm, self-packed) and separated using a 110 min gradient from 4% to 32% of solvent B (0.1% formic acid in acetonitrile and 5% (v/v) DMSO) at 300 nL/min flow rate. Both nanoLC solvents (solvent A = 0.1% formic acid in HPLC grade water and 5% (v/v) DMSO) contained 5% DMSO to boost MS intensity.

The Q-Exactive HF-X mass spectrometer was operated in data-dependent acquisition (DDA) and positive ionization mode. MS1 spectra (360–1300 m/z) were recorded at a resolution of 60,000 using an automatic gain control (AGC) target value of 3e6 and maximum injection time (maxIT) of 45 msec. Up to 18 peptide precursors were selected for fragmentation in the case of the full proteome analyses. Only precursors with charge state 2 to 6 were selected and dynamic exclusion of 30 sec was enabled. Peptide fragmentation was performed using higher energy collision-induced dissociation (HCD) and a normalized collision energy (NCE) of 26%. The precursor isolation window width was set to 1.3 m/z. MS2 resolution was

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15,000 with an automatic gain control (AGC) target value of $1e^5$ and maximum injection time (maxIT) of 25 msec (full proteome).

Peptide identification and quantification were performed using MaxQuant (version 1.6.3.4) with its built-in search engine Andromeda^{94,95}. MS2 spectra were searched against the Uniprot mus musculus proteome database (UP000000589, 54,208 protein entries, downloaded 22.3.2019) supplemented with common contaminants (built-in option in MaxQuant). Trypsin/P was specified as the proteolytic enzyme. Precursor tolerance was set to 4.5 ppm, and fragment ion tolerance to 20 ppm. Results were adjusted to 1% false discovery rate (FDR) on peptide spectrum match (PSM) level and protein level employing a target-decoy approach using reversed protein sequences. The minimal peptide length was defined as 7 amino acids, the "matchfunction was disabled. full between-run" For proteome analyses. carbamidomethylated cysteine was set as fixed modification and oxidation of methionine and N-terminal protein acetylation as variable modifications.

For the statistical analysis of the proteomics data, six biological replicates were measured in young and old Titan as well as young and old control mice. Protein abundances were calculated using the LFQ algorithm from MaxQuant⁹⁶. Before further downstream analyses, protein LFQ values were logarithm (base 10) transformed. Next, Limma⁹⁷ was used to identify the differentially expressed proteins between young control vs young Titan; young control vs old control; young Titan vs old Titan, and old control vs old Titan. The resulted p-values were adjusted by the Benjamini-Hochberg algorithm⁹⁸ to control the false discovery rate (FDR). The differential analyses were

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performed on proteins that were identified in at least four out of six biological replicate samples in both groups under comparison.

Gene set annotations were downloaded from MSigDB⁹⁹, including the Gene Ontology annotation (C5 category) and pathway annotation (C2 category). The gene IDs of differentially expressed proteins were mapped to the gene set annotations. The significance of over-representation was evaluated using fisher's exact test.

Total RNA extraction for quantitative RT-PCR

RNA extraction was performed from 50 mg deep-frozen liver tissue. The tissue was homogenized in 1 ml TRItidy G[™] via pestle and incubated at room temperature for 5 minutes. After the addition of 200 µl chloroform, the sample was mixed 15 seconds with the Vortex and again incubated for 2 min at room temperature with a following centrifugation step at 12,000 x g, 4°C for 15 min. The aqueous phase was transferred into a new vial and mixed with 500 µl - 20°C isopropanol and incubated 10 minutes at -20°C. The samples were then centrifuged at 8,000 x g, 4°C for 10 minutes and the supernatant was discarded. Two washing steps with 1 ml -20°C cold 70% ethanol were applied and the pellet was air-dried and dissolved in 100 µl nuclease-free water. RNA concentration was measured with NanoDrop 2000.

Quantitative Real-time PCR

A total of 800 ng RNA was used for cDNA synthesis, using the SensiFAST™cDNA Synthesis Kit (Bioline). Quantitative PCR was performed with the SensiFAST™SYBR® No-Rox Kit (Bioline) using 1.25 ng per reaction,

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except for the testing of mTOR (2.5 ng), and 4 pmol of forward and reverse primers. Quantitative PCR (qPCR) reactions were run on the Roche Lightcycler 96. The annealing temperature was 60°C. The qPCR data were normalized to the levels of beta-actin. Primers are listed in Table S5. PCR products were cleaned up using the High Pure PCR Product Purification Kit (Roche) and sequenced by LGC Genomics GmbH. Each sample was measured using two technical replicates.

Statistics and visualization

Unless stated otherwise in the RNA-seq and proteomics experimental method, statistics and graphing were conducted on GraphPad Prism 9. Unpaired two-tailed *t*-tests with Welch's correction (parametric) or MWU testing (non-parametric) were used for calculating the *P*-values unless stated otherwise. A multiple correction (FDR) was performed for SBF vs ERF qPCR and plasma analysis. A p-value ≤ 0.05 was used as level of significance.

Importantly, according to the GMC analysis outline, correction for multiple testing was not performed when analyzing the GMC data set (See GMC website).

Data availability

The proteomic raw data and MaxQuant search files have been deposited to the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the PRIDE partner repository and can be accessed using the dataset identifier PXD019030 (reviewer account details: username reviewer73331@ebi.ac.uk; password rvqDlf58). All RNAseq data have been deposited to GEO. bioRxiv preprint doi: https://doi.org/10.1101/2020.05.11.088625; this version posted June 9, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

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Author contributions

SP conceptualized the project. ML prepared all the mice work, conducted the survival curves with the weight measurements and assisted with

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conceptualizing this work. ML also maintains the Dummerstorf selection line. AH contributed to this work by the management of pilot studies on life expectancy in the mouse model used in this study. ID-D extracted the RNA for RNAseq. AV-V prepared the libraries for transcriptomic analysis. AM-E prepared the proteomics and histone acetylation. AS-M, RG, BR, AA-P, TK-R, JC-W performed mouse phenotyping tests and analyzed the data, LB supported data analyzes and manuscript preparation. CS coordinated and VG-D, HF and MHdA supervised the project at the GMC and conceived phenotyping test pipeline. AV-V analyzed the RNA-seq and histone acetylation and generated the data visualization. Al supervised AV-V's work. SPV and JS conducted and analyzed the RDD data. CM and CL ran, analyzed and visualized the proteome data. FK isolated the blood for the ERF experiments. VC ad JB prepared and analyzed the Oil red O/ hematoxylin straining. ID-D, FC, AK and SP wrote the initial manuscript. All authors contributed to the final version of the manuscript. AM-E generated all the final figures for the paper. BG assisted with graph analysis and manuscript preparation.

Declaration of interests

Authors declare no competing interests.

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Figure legends

(A) Representative images of unselected control mice (left) and 180 generation-selected Titan mice (right) at 6 weeks, at the time of the selection. (B) After the first 16-19 weeks of life, Titan mice reached 110–115 g, whereas average control animals weighed 45 g. (C) Representative X-ray images of control and Titan mice at 16-17 weeks of age. Control (n = 38) and Titan (n = 31) mice. (D) Body length (cm) of 10–11-week-old Titan mice compared to age-matched control mice. Control (n = 20) and Titan (n = 17) mice. (E) Total fat and lean mass of control (n = 20) and Titan (n = 17) mice at the age of 10– 11-weeks. (F) Total percentage of fat in control (n = 20) and Titan (n = 17) mice. (G) Percentage of intra-abdominal fat (n = 10 per group) at the age of 10–11-weeks. (H) Representative images of hematoxylin and eosin (H&E)stained brown adipose tissue (BAT) from control (n = 6) and Titan (n = 5) mice (20x magnification) at 16–17 weeks of age. (I) Representative images of H&Estained pancreas from control and Titan (n = 6) mice (2.5x magnification) at 16–17 weeks of age. (J) Comparison of lifespans and mortality rates of Titan and control male mice showing significantly reduced lifespan of Titan mice (Log Rank test; p <0.0001, χ^2 =124.9). ****P* < 0.001. Error bars indicate SEMs. Unpaired two-tailed t-tests with Welch's correction were used to calculate P-values.

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Figure 2. Histone 4 acetylation levels are altered in Titan compared with control unselected mice.

(A) Scheme of the RDD of the *kat2a* gene in Titan and control mice. (B) Total levels of histone 4 4-17 levels via mass spectrometry (p =0.0111, MWU-test) (n = 7 per group). (C) Quantification of monoacetylation of histone 4 4-17. P-MWU-test (H4K5K8)=0.0175, (H4K8)=0.901, (H4K12)=0.455, values. (H4K16)=0.0023 (n = 7 per group). (D) Quantification of polyacetylation of histone 4 4-17. P-values, MWU-test (H4K5K8)=1, (H4K12K16)=0.0379, (H4di-ac)=0.0379, (H4K5K8K12)=0.62, (H4K5K8K16)=0.208, rest (H4K5K12K16)=0.053, (H4K8K12K16)=0.208, (H4 tetra-ac)=0.62 (n = 7 per group). *P < 0.05, **P < 0.01. Error bars indicate SEMs.

Figure 3. Hybrid offsprings of control and titan mice display intermediate obesity and life span

(A) Body weight and visceral fat percentage in F1 hybrid (Offsprings of either mother control x father mother vs mother titan x father control) n = 15 each, MWU-test was used to calculate p-value.(B) Quantification of total histone 4 4-17 acetylation, H4K5ac and H4K16ac in control (n = 6), Titan (n = 7) and hybrid (n = 10) mice. One way ANOVA followed by Tukey test P-value: control vs hybrid (H4ac)=0.0765 (H4K5)=0.116, (H4K16)=0.0184 and Titan vs hybrid (H4ac)=0.0383, (H4K5)=0.330, (H4K16)=0.0065. (C) Body weight and visceral fat percentage in control (n = 20), Titan (n = 19) and hybrid (n = 30). One way ANOVA followed by Tukey test P-value: control vs hybrid (Weight)<0.0001, (%visceral fat)=0.0021 and Titan vs hybrid (Weight)<

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0.0001 (%visceral fat)<0.0001. (D) Comparison of lifespans and mortality rates of Titan and control male mice to Hybrid (N=66). *P < 0.05, **P < 0.01. ***P < 0.001. Error bars indicate SEMs.

Figure 4. Titan mice display altered liver metabolism

(A) Alanine aminotransferase (ALAT) (p = 0.0262) and (B) alkaline phosphatase (p = 0.0019) levels in control (n = 20) and Titan mice (n = 18). (C) Oil Red O with Hematoxylin staining of fat in liver of control and Titan mice (n = 4 per group). (D) Bilirubin levels in control (n = 20) and Titan mice (n = 18) (p = 0.0019). (E) Urea levels of control (n = 20) and Titan (n = 18) mice (p = 0.0132). **P* < 0.05, ***P* < 0.01. Error bars indicate SEMs. Unpaired two-tailed *t*-tests with Welch's correction were used to calculate *P*-values.

Figure 5. Liver transcriptome analyses reveal differential expression of genes associated with metabolic pathways in Titan mice.

(A) Principal component analysis (PCA) of liver transcriptomes revealed clustering differences between Titan mice and controls. Control mice at 11 weeks and 19-21 weeks had similar transcriptomes, while younger and older Titan mice had distinct transcriptomes. (B) Heat map showing significantly altered genes in control and Titan mice at 11 weeks and 19-21 weeks of age. Color gradient in each cell in the heatmap represents the scaled normalized log counts from each replicate and condition. (C) GO term analysis in 11-week-old Titan mice compared with corresponding control. (D) GO term in 19-

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21-week-old Titan mice compared with corresponding control. (n = 5 per group).

Figure 6. Late dietary intervention by switching to energy-reduced food (ERF) improves health metrics and extends the life span of Titan mice.

(A) Switching standard feeding (SBF) to ERF at 12 weeks resulted in a persistent average weight loss in Titan siblings (n = 23 per group). MWU-test, P=0.0017. (B) Compared to age-matched control Titan mice, ERF-fed Titan mice had a lower percentage of intra-abdominal fat at 21 weeks (n= 7 per group). Paired MWU-test, p = 0.0313. (C) Late ERF feeding at 12 weeks of age decreased the levels of plasma cholesterol (pval.adj=0.00013), HDL (pval.adj=0.0013), Glucose (pval.adj=0.043), Leptin (pval.adj=0.043), Glycerol (pval.adj=0.0165) and NEFA (pval.adj=0.00132) in 21 weeks old mice. (n= 10 per group). MWU-test, by multiple correction to calculate P-values. (D) Quantification of polyacetylation of histone 4 4-17 following ERF feeding. Pvalues, MWU-test (H4K5K8)=0.8125, (H4K12K16)=0.0469 (H4-rest diac)=0.1563, (H4K5K8K12)=0.9375, (H4K5K8K16)=0.0781, (H4K5K12K16)=0.0313, (H4K8K12K16)=0.0781, (H4 tetra-ac)=0.375 (n = 7 per group) Paired MWU-test, to calculate P-values. (E) RT-gPCR comparing gene expression of candidate genes (from table S4) of ERF- and control-fed Titan mice siblings at 21 weeks of age (n = 7 per group). Paired t-test was performed followed by multiple correction to calculate P-values pval.adj (Dmdg)=0.032,(*Gnmt*)=0.012, (Bhmt)=0.736,(Cyp7b1)=0.008,(*Elov*/5)=0.736, (Cyp2c37)=0.012,(Acaca)=0.0587, (*Cth*)=0.012. (F) Switching to ERF feeding at 12 weeks of age (vertical dashed line) increased the lifespan of both the Titan and control mice. The ages at 25%, 75% and

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90% death, and the median lifespan are presented in Table S1. $\ ^{*}P$ < 0.05,

P < 0.01, *P < 0.01. Error bars indicate SEMs.

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Supplementary figures

(A-C) Plasma analyses comparing control and Titan mice after fasting at 18–20 weeks. Titan mice showed higher levels of triglycerides (p = 0.0055) and both non-HDL (p = 0.0022) and HDL cholesterol (n= 20 control vs 19 Titan). (D) Insulin levels in control (n = 19) and Titan (n = 18) mice at 16–17 weeks of age. (E) Leptin levels in control (n = 20) and Titan (n = 18) mice and (F) higher FGF21 levels in the Titan mice line compared to the control mice line (N=20 control vs 18 Titan). **P < 0.01 ***P < 0.001. Error bars indicate SEMs. Unpaired two-tailed *t*-tests with Welch's correction were used to calculate *P*-values.

Figure S2. Titan mice show signs of heart fibrosis.

H&E and Sirius Red stainings of the heart from 16-17-weeks-old control and Titan mice are shown (0.45x magnification). Sirius Red staining allows a better visualization of fibrotic tissue. Higher magnification pictures from rectangles were taken at 10x.

Figure S3. Titan mice show thickened dermis and early signs of increased inflammation.

Representative images of hematoxylin and eosin (H&E) staining of the skin. Comparison of Titan and control mice at 16–17 weeks of age. (n = 4–6 per group). (A) The dermis of the Titan mice is substantially thicker compared to control mice. (B and C) IL-6 plasma levels (p = 0.0086) in control (n = 13) and Titan (n = 17) mice, and TNF α plasma levels in control (n = 20) and Titan (n =

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18) mice. (D) Representative images of hematoxylin and eosin (H&E) staining and B-cell immunohistochemistry (IHC) of the thymus of control (left) and Titan (right) mice (2.5x magnification). IHC of thymic medullar nodes revealed that they are composed mainly of B-cells (CD45R/B220-positive) instead of Tcells (CD3-positive). Images in squares were taken at 10x magnification (n = 4–6 per group). **P* < 0.05, ****P* < 0.001. Error bars indicate SEMs. Unpaired two-tailed *t*-tests with Welch's correction were used to calculate *P*-values.

Figure S4. Liver H&E staining comparison of control vs Titan mice.

Shown are representative H&E-stained images of the liver from 16-17-weeks old control and Titan mice (10x and 40x magnifications are shown).

Figure S5. Transcriptome changes in the Liver of younger vs older Titan mice.

(A) Heat map showing significantly altered genes between 11-week-old control and Titan mice. (B) Heat map showing significantly altered genes between 11- and 19-21-week-old Titan mice. (C) GO term analysis of (B) revealed an increase in insulin-related pathway and glucose homeostasis and a decrease in various metabolic processes including fatty acid metabolism.
(D) Heat map showing significantly altered genes between 19-21-week-old control and Titan mice. n=5 per group.

Figure S6. Liver proteome analyses reveal differences in metabolic protein levels between control and Titan mice.

(A) Heat map comparing proteomes of 11-week-old control and Titan mice.(B) Heat map comparing proteomes of 19-21-week-old control and Titan mice.

(C) GO term analysis of 11-week-old mice indicated increased fatty acid metabolism, various lipid catabolism, and coenzyme metabolism while downregulation in xenobiotic, amino acid and sulfur metabolism. (D) GO term analysis of 19-21-week-old mice showed upregulation of proteins involved in fatty acid metabolism, carbohydrate catabolism and coenzyme metabolism and downregulation of proteins involved in amino acid and sulfur metabolism (n = 6 per group).

Table S1. Unique RDD in Titan mice

 Table S2. Unique RDD in Titan mice excluding DU6P mice

Table S3. Lifespans in days for Titan mice receiving SBF and ERF, respectively

Survival	Titan SBF	Titan ERF
Minimum	68	71
10% Percentile	123	214.8
25% Percentile	231.5	264
75% Percentile	405	447
90% Percentile	475	504
Maximum	614	660
Median	323.5	374
Mean	317.4	359.5
Std. Error	11.49	11.45

Tables S4-5. List of significantly altered genes (RNAseq)

Tables S6. Proteome

 Table S7. Key resources and Real-time PCR primer sequences.

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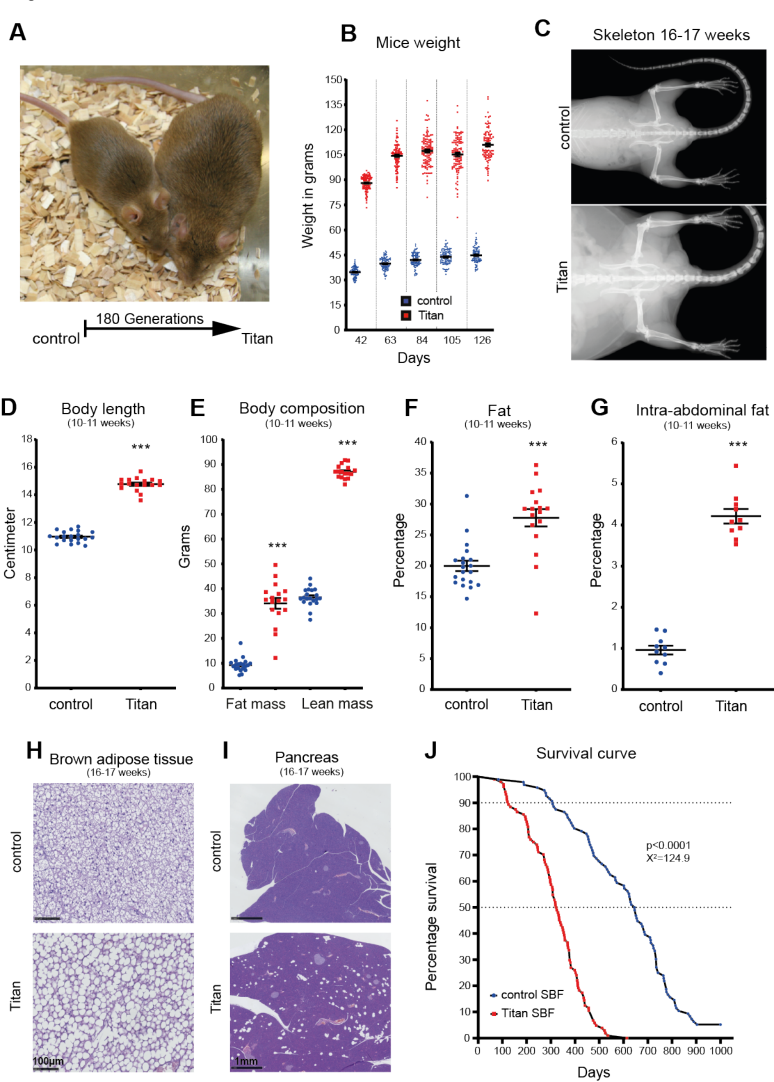
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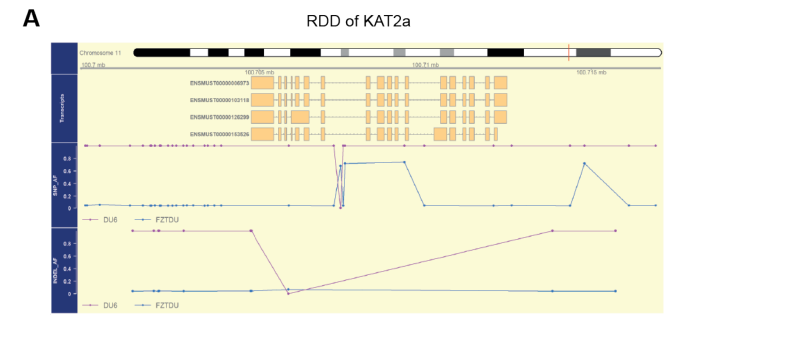
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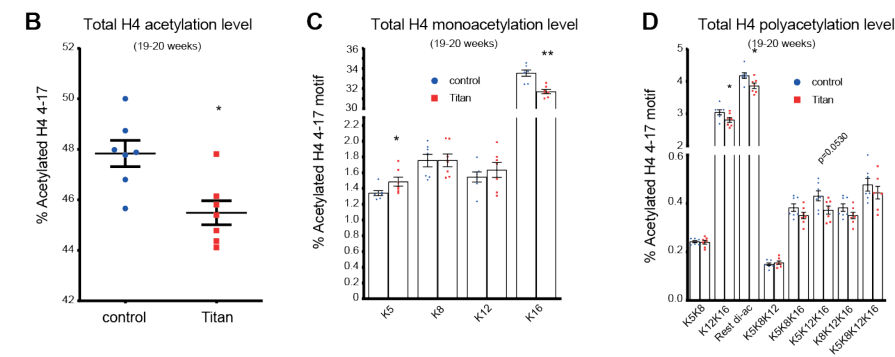
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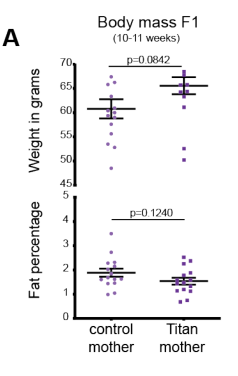
geneSet	description	size	overlap	expect	enrichmentRatio	pValue	FDR	overlapId	userld
<u>GO:0031424</u>	keratinization	35	5	0.2304330	21.698286	0.0000032	0.0196906	20754; 20762; 20763; 20765; 20766	Sprr2h; Sprr2i; Sprr2k; Sprr1b; Sprr3
<u>GO:0044262</u>	cellular carbohydrate metabolic process	285	10	1.8763826	5.329403	0.0000193	0.0504800	11677; 12015; 14042; 14534; 15531; 20848; 50798; 50873; 58246; 66646	Rpe; Gne; Slc35b4; Akr1b3; Kat2a; Stat3; Ext1; Prkn; Ndst1; Bad
<u>GO:0060343</u>	trabecula formation	27	4	0.1777626	22.501926	0.0000279	0.0504800	14225; 15214; 21687; 170826	Fkbp1a; Tek; Hey2; Ppargc1b
<u>GO:0060416</u>	response to growth hormone	28	4	0.1843464	21.698286	0.0000324	0.0504800	20848; 20850; 20851; 227231	Cps1; Stat5b; Stat5a; Stat3
<u>GO:0009913</u>	epidermal cell differentiation	199	8	1.3101759	6.106050	0.0000522	0.0650244	15214; 20754; 20762; 20763; 20765; 20766; 22283; 67694	Ush2a; Sprr2h; Sprr2i; Sprr2k; Sprr1b; Sprr3; lft74; Hey2
<u>GO:0010918</u>	positive regulation of mitochondrial membrane potential	13	3	0.0855894	35.051077	0.0000759	0.0675235	12015; 50873; 269523	Vcp; Prkn; Bad
<u>GO:0072044</u>	collecting duct development	13	3	0.0855894	35.051077	0.0000759	0.0675235	11677; 71228; 240025	Akr1b3; Dlg5; Dact2
<u>GO:0060396</u>	growth hormone receptor signaling pathway	14	3	0.0921732	32.547429	0.0000962	0.0744756	20848; 20850; 20851	Stat5b; Stat5a; Stat3
<u>GO:0060347</u>	heart trabecula formation	15	3	0.0987570	30.377600	0.0001196	0.0744756	14225; 15214; 21687	Fkbp1a; Tek; Hey2
<u>GO:0045579</u>	positive regulation of B cell differentiation	15	3	0.0987570	30.377600	0.0001196	0.0744756	12015; 20850; 20851	Stat5b; Stat5a; Bad
<u>GO:0046629</u>	gamma-delta T cell activation	16	3	0.1053408	28.479000	0.0001465	0.0751777	20850; 20851; 270152	Jaml; Stat5b; Stat5a
<u>GO:0071378</u>	cellular response to growth hormone stimulus	16	3	0.1053408	28.479000	0.0001465	0.0751777	20848; 20850; 20851	Stat5b; Stat5a; Stat3
<u>GO:0050891</u>	multicellular organismal water homeostasis	42	4	0.2765195	14.465524	0.0001651	0.0751777	11677; 20276; 22393; 229574	Flg2; Wfs1; Akr1b3; Scnn1a
<u>GO:0035966</u>	response to topologically incorrect protein	124	6	0.8163910	7.349419	0.0001711	0.0751777	22393; 50873; 56323; 69310; 69601; 269523	Dab2ip; Dnajb5; Vcp; Wfs1; Pacrg; Prkn
<u>GO:0072528</u>	pyrimidine-containing compound biosynthetic process	43	4	0.2831033	14.129116	0.0001812	0.0751777	29807; 68262; 69692; 227231	Cps1; Tpk1; Hddc2; Agpat4
<u>GO:0030216</u>	keratinocyte differentiation	131	6	0.8624776	6.956702	0.0002306	0.0863868	20754; 20762; 20763; 20765; 20766; 67694	Sprr2h; Sprr2i; Sprr2k; Sprr1b; Sprr3; lft74

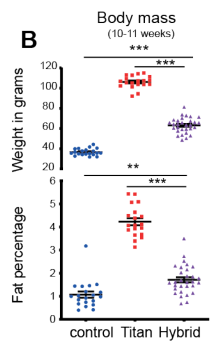
<u>GO:0030104</u>	water homeostasis	46	4	0.3028547	13.207652	0.0002359	0.0863868	11677; 20276; 22393; 229574	Flg2; Wfs1; Akr1b3; Scnn1a
<u>GO:0045838</u>	positive regulation of membrane potential	19	3	0.1250922	23.982316	0.0002499	0.0864356	12015; 50873; 269523	Vcp; Prkn; Bad
<u>GO:0030856</u>	regulation of epithelial cell differentiation	136	6	0.8953966	6.700941	0.0002824	0.0925190	12015; 15214; 20850; 20851; 21937; 74123	Tnfrsf1a; Hey2; Stat5b; Stat5a; Foxp4; Bad
<u>GO:0061383</u>	trabecula morphogenesis	49	4	0.3226061	12.399020	0.0003017	0.0938981	14225; 15214; 21687; 170826	Fkbp1a; Tek; Hey2; Ppargc1b
<u>GO:0043588</u>	skin development	259	8	1.7052038	4.691521	0.0003204	0.0949844	20754; 20762; 20763; 20765; 20766; 67694; 229574; 240025	Sprr2h; Sprr2i; Sprr2k; Sprr1b; Sprr3; Flg2; lft74; Dact2

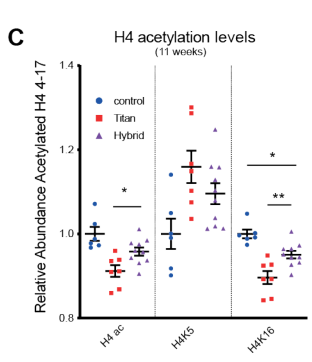


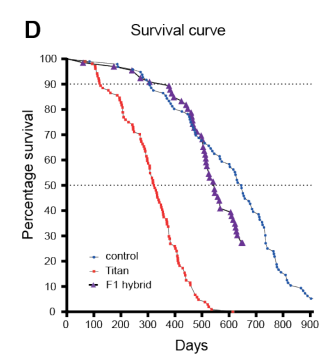


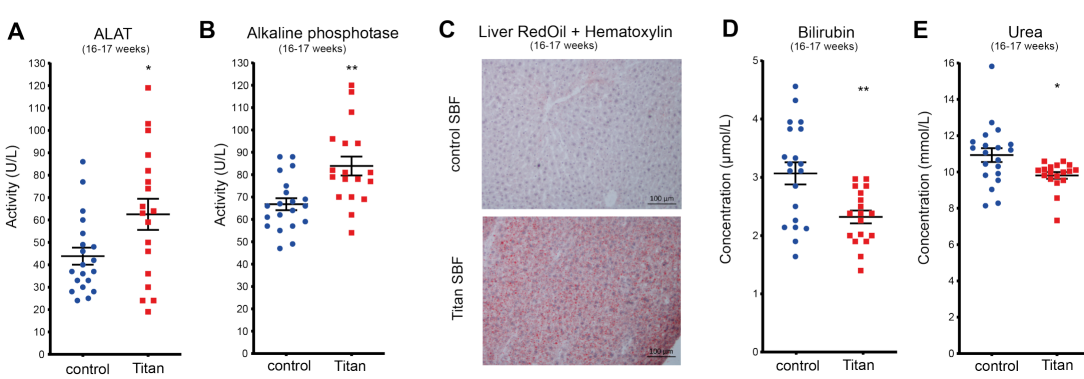




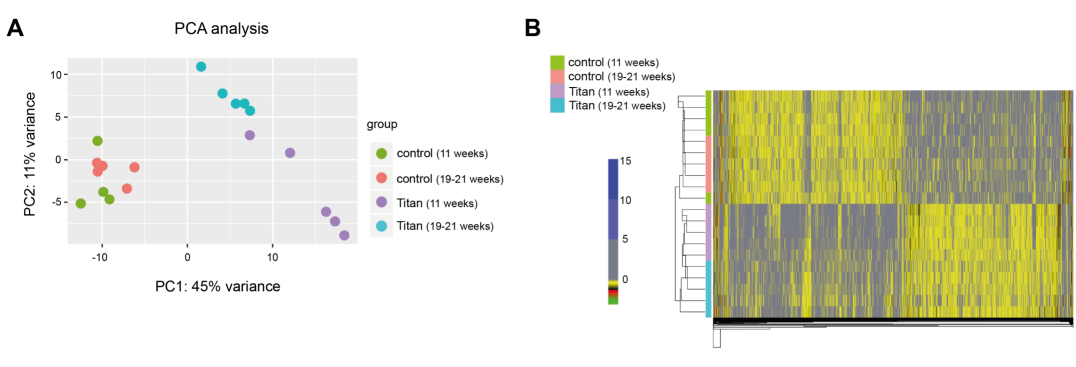




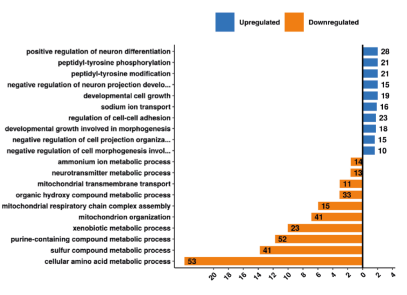




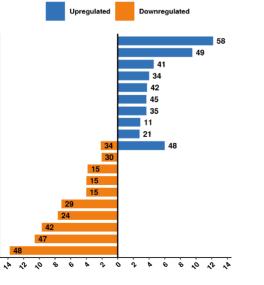
С



19-21 weeks GO Term comparison



D



11 weeks GO Term comparison

fatty acid metabolic process coenzyme metabolic process lipid localization dendrite development Ras protein signal transduction positive regulation of cell projection organiza.. maintenance of location protein localization to microtubule cytoskeleton ammonium ion metabolic process organic anion transport negative regulation of proteolysis regulation of coagulation regulation of hemostasis regulation of blood coagulation cellular modified amino acid metabolic process xenobiotic metabolic process sulfur compound metabolic process small molecule catabolic process cellular amino acid metabolic process

-log10(p.adjust)

-log10(p.adjust)

