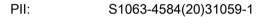
Identification of TGF β signatures in six murine models mimicking different osteoarthritis clinical phenotypes

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Identification of TGF β signatures in six murine models

mimicking different osteoarthritis clinical phenotypes

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Running title: TGF β expression in 6 murine models of OA

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Objective. TGF β is a key player in cartilage homeostasis and OA pathology. However, few data are available on the role of TGF β signalling in the different OA phenotypes. Here, we analysed the TGF β pathway by transcriptomic analysis in six mouse models of OA.

Method. We have brought together seven expert laboratories in OA pathophysiology and, used inter-laboratories standard operating procedures and quality controls to increase experimental reproducibility and decrease bias. As none of the available OA models covers the complexity and heterogeneity of the human disease, we used six different murine models of knee OA: from post-traumatic/mechanical models (meniscectomy (MNX), MNX and hypergravity (HG-MNX), MNX and high fat diet (HF-MNX), MNX and seipin knock-out (SP-MNX)) to aging-related OA and inflammatory OA (collagenase-induced OA (CIOA)). Four controls (MNX-sham, young, SP-sham, CIOA-sham) were added. OARSI-based scoring of femoral condyles and RNA extraction from tibial plateau samples were done by single operators as well as the transcriptomic analysis of the TGFβ family pathway by Custom TaqMan[®] Array Microfluidic Cards.

Results. The transcriptomic analysis revealed specific gene signatures in each of the six models; however, no gene was deregulated in all six OA models. Of interest, we found that the combinatorial *Gdf5-Cd36-Ltbp4* signature might discriminate distinct subgroups of OA: *Cd36* upregulation is a hallmark of MNX-related OA while *Gdf5* and *Ltbp4* upregulation is related to MNX-induced OA and CIOA.

Conclusion. These findings stress the OA animal model heterogeneity and the need of caution when extrapolating results from one model to another.

Keywords: osteoarthritis, signature, murine model, transcriptomic

A collaborative study between 7 expert laboratories in osteoarthritis pathophysiology rose the challenge to identify common gene signatures in six murine models of osteoarthritis representative of clinical phenotypes in osteoarthritic patients using standard operating procedures and centralized analyses.

The main findings of the study are the absence of one common deregulated gene in all six osteoarthritis models but the identification of a combinatorial *Gdf5-Cd36-Ltbp4* signature that might discriminate distinct subgroups of OA.

Journal Prevention

1 INTRODUCTION

2

3 Although osteoarthritis (OA) is the most prevalent joint disease worldwide, there is still 4 not a single disease-modifying OA drug on the market. The current treatment options usually result in poorly predictable outcomes due to the high interpatient variability in OA clinical 5 6 and structural features. Indeed, some studies have reported OA phenotype heterogeneity 7 among patients [1-3]. Recently it has been proposed to use advanced techniques to identify 8 combinatorial biomarkers for distinguishing the different OA phenotypes [4], and also to identify patients at higher risk of disease progression, or with different underlying 9 pathophysiologic mechanisms and risk factors [5]. This will help to improve clinical research 10 and to develop targeted treatments and prevention strategies based on a phenotype-guided 11 approach. The advantage of searching for targets based on differences between risk factors 12 13 is the simplicity then of selecting patients for future personalized medicine.

Currently, OA research relies on the use of various animal models (mainly mice and rats, 14 and more rarely large animals) that mimic mechanical, metabolic or inflammatory OA. 15 However, none of these models covers the complexity and heterogeneity of the human 16 17 disease but different models likely reflect the heterogeneity of human OA. Moreover, it is difficult to compare the results of different experimental studies due to the heterogeneity of 18 19 animal backgrounds and experimental protocols. Several studies have analysed global gene 20 expression in OA samples using RNA-seq [6-8] and have generated huge amounts of datasets. However, only small subsets of data are validated and large amounts of data are 21 commonly not investigated. To try to tackle some of these limitations, seven French 22 23 academic laboratories experts in OA animal models formed a Research on OsteoArthritis Disease (ROAD) consortium to centralize many experimental steps and to put in place 24

1 standard operating procedures (SOP) in order to minimize bias and increase reproducibility. The first objective of the ROAD consortium was to investigate the TGF^β pathway in various 2 OA phenotypes. Indeed, recent findings have shown that TGF β is a central player in cartilage 3 homeostasis and OA pathology [9]. However, few data are available on the 4 pathophysiological role of TGFβ family members in the different OA phenotypes. Therefore, 5 6 the consortium analysed the TGF^β pathway by transcriptomic analysis in six murine models 7 of knee OA that reproduce the main phenotypes of the human disease: surgical 8 meniscectomy (MNX) to mimic mechanical or post-traumatic OA, hypergravity and MNX (HG-MNX) to mimic overweight-induced mechanical OA, high fat diet and MNX (HF-MNX) to 9 mimic obesity-induced OA, seipin knock-out and MNX (SP-MNX) to mimic metabolic 10 syndrome-induced OA, aging to mimic age-related OA, and collagenase-induced OA (CIOA) 11 12 to mimic inflammatory OA.

13

14 MATERIAL AND METHODS

15

16 Animal models

17 Animal models and controls (ten mice/group) were generated using C57BL/6JR6 males that are known to display more severe and reproducible disease [10] and were supplied by the 18 19 same company (Janvier Labs, France). Bscl2-/- mice (SP-MNX and SP-sham controls; C57BL/6J background) were from CEA (Direction des Sciences du Vivant/Genoscope 20 /LABGEM). Six animals per group were calculated to be required to demonstrate significance 21 22 at the 5% level with a power of 80% using the G*power software but 10 animals were included to have 6 animals with an OA score \geq 3 at the end of the experiment. MNX was 23 performed in one joint of 10 weeks old mice by the use of partial meniscectomy as described 24

1 [11, 12] and done by a single trained operator in all laboratories. All animal procedures were approved by the local institutions' animal welfare committees and were performed in 2 accordance with the European guidelines for the care and use of laboratory animals 3 (2010/63/UE). Surgery and euthanasia were performed after anaesthesia with isoflurane 4 gas, and all efforts were made to minimize suffering. Mice were housed in solid bottomed 5 6 plastic cages in quiet rooms at 22° ± 1°C, 60% controlled humidity, and 12h/12h light/dark 7 cycle. Animals were used after an adaptation period of 7 days and had free access to tap water and standard pelleted chow (except the HF model). Mice were sacrificed at week 6 8 after OA induction to have a comparable disease time induction although we were aware 9 that OA severity can vary according to the model. 10

11

• Joint instability model

MNX was selected as the reference model of joint instability-related OA [11]. Knee joint 12 instability was induced surgically in the right knee by medial partial meniscectomy. Surgery 13 was performed under a binocular magnifier (X15) using a Sharpoint microsurgical stab knife. 14 Mice were placed in dorsal position, knee flexed and right foot taped. After skin incision, the 15 medial femoro-tibial ligament was cut, a short incision of the medial side of quadriceps 16 17 muscle was performed, the knee capsule was cleaved and the patella was sub-luxated laterally. After section of the meniscotibial ligament, the medial meniscus was gently pulled 18 19 out and ¾ of its anterior horn removed. Then, the patella was replaced, the quadriceps muscle and the skin plan sutured. Control animals underwent sham surgery (ligament 20 visualization but not dissection). 21

Hypergravity model

Hypergravity mimics the overweight-associated mechanical strain on joints without
 metabolism dysregulation. In mice with MNX, hypergravity induces large OA lesions that are

not observed without surgical induction [13, 14]. MNX was performed on the right knee, and 1 mice were put back in their box for 48 hours. Then, cages were transferred in the gondolas 2 of the centrifuge (COMAT Aérospace, Flourens, France) to maintain a permanent level of 3 hyper gravity [14]. This device has four 1.4m-long arms that hold at their distant end a 4 5 mobile octagonal gondola ($56.2 \times 52.0 \times 59.2$ cm). All gondolas are equipped with an infra-6 red video surveillance system to monitor the animals' condition and food/water stocks. In 7 the centrifuge, temperature and light conditions were identical to that of control cages. At 8 the start of centrifugation, acceleration was smoothly and gradually increased over a period of 40 sec. The final acceleration was 2 g (29.6 rpm), and animals were kept at 2 g for 6 9 weeks. Animals were provided with enough food and water for 4 weeks. Then, the 10 centrifuge was transiently stopped to allow litter change, animal weighing, and chow and 11 water supply refilling. Control mice with MNX were not exposed to hypergravity. 12

13

Metabolic disorder model

Seipin (SP) knock-out mice are representative of metabolism disorder, which is a feature associated with OA [15]. *Bscl2* deficiency in mice recapitulates the main features of the phenotype of patients with Berardinelli-Seip Congenital Lipodystrophy (BSCL), including almost complete absence of adipose tissue, hyperglycaemia, hyperinsulinemia, and insulin resistance. MNX and Sham surgery were performed in 10 week-old *Bscl2-/-* mice.

19 • High Fat Diet model

The high fat diet model reproduces the effect of obesity and dysregulated metabolism on OA onset [16]. At the age of six weeks, mice were fed with High Fat Diet (HFD, 60% of calories from fat, Ssniff, EF D12492 (II) mod. Soest, Germany) that was provided ad libitum for 10 weeks with the chow changed twice per week. A number of mice 20% higher than the final group size was included to ensure statistical power of the experimentation. MNX was

performed at the age of 10 weeks. In absence of surgical induction, mice did not develop
spontaneous lesions of OA. The average weekly weight gain ranged from 1 g to 1.5 g, leading
to a final weight gain of 73% (mean: 14.6 g) associated with insulin resistance (HOMA-IR:
+246%). Considering the large variability generally observed in the final body weight and fat
mass, only animals with a final weight gain higher than 70% were analysed.

6 •

Collagenase-induced OA model

7 The collagenase-induced model (CIOA) is characterized by low grade inflammation of the 8 synovial membrane, leading to OA lesions [10]. A solution of 1 U/5µL type VII collagenase 9 from Clostridium histolyticum (Sigma-Aldrich) was prepared in saline solution. At day 0, a small skin incision was performed on top of the patellar tendon. The knee was bended and 10 the collagenase solution (5 μ L) was injected in the intra-articular space using a 10 μ L syringe 11 (Hamilton) with a 25 gauge needle. On day 2, a second collagenase injection was performed 12 according to the same procedure. Six weeks later, animals were sacrificed. Control animals 13 were injected with saline solution. 14

15 • Age-related model

Ageing is the main risk OA factor [17]. C57BL/6JRj mice exhibit mild OA lesions in the knee at the age of 18 to 24 months [18]. Mice were housed with free access to food and water and euthanized at the age of 24 months. Control young mice were kept in the animal facility and euthanized at the age of 16 weeks.

20

21 Sample preparation for histology and mRNA extraction

After sacrifice, femora and tibiae from 10 knee joints (one joint/mouse) per model were dissected. Skin and muscles were removed and the knee joint was isolated by sectioning the distal extremity of tibiae and proximal part of the femurs. The tibial plateau was isolated

from bone at the growth plate interface, by cutting 2-3 mm beneath the cartilage surface.
The remaining soft tissues (meniscus, ligaments and synovium) were removed. The tibial
plateau was immediately placed in 1mL of TRIzol® Reagent (Life Technologies), snap-frozen
in liquid nitrogen, and stored at -80°C till RNA extraction. After isolation, femoral condyles
were fixed in 4% paraformaldehyde for 36 hours, and then decalcified in 0.5M EDTA at room
temperature for 15 days.

7

8 Histology

9 After dehydration in a graded series of alcohol, femoral condyles were embedded in paraffin at 60°C in a tissue processor. On average, 30 serial sagittal sections of 5 µm were cut, and three were chosen at the upper, medium and lower levels every 50 µm from cartilage surface. OA scoring was performed after Safranin O-Fast Green staining, according to the OsteoArthritis Research Society International (OARSI) recommendations [19]. For each animal, the OA score was the highest score obtained at one of the three levels. For each model, all sections were blindly scored by the same three readers.

16

17 **RNA isolation**

Tibial plateau samples were prepared in each consortium laboratory and then shipped for centralized RNA extraction that was performed by crushing thawed samples with ceramic beads (Precellys[®] Lysing kit CK28R), using a Precellys[®] 24 tissue homogenizer equipped with the Cryolis[®] cooling unit (Bertin Technologies). Samples underwent three successive lysis cycles at 6500 rpm for 15 sec, spaced by a 5 min lag phase at 4°C, before addition of 200 µL chloroform. After incubation at room temperature for 3 min, the aqueous phase was recovered, 600 µL of 70% ethanol was added, and the solution was transferred to an

1 RNeasy[®] spin column (Qiagen) and the next steps were performed according to the 2 supplier's recommendations. Total RNA was quantified with a Nanodrop[®] instrument and 3 aliquots were frozen at - 80°C. RNA integrity was confirmed with the Agilent[®] RNA 6000 kit 4 on an Agilent Bioanalyzer 2100[®].

5

6 **Transcriptomic analysis**

7 Transcriptomic analysis was performed on Custom TagMan[®] Array Microfluidic Cards (TAC) that were designed to perform 384 real-time PCR reactions on a ViiA[™] 7 Fast Real-Time PCR 8 System (Applied Biosystems[®]). Custom TAC were designed for the analysis of TGFβ family 9 members (table 1). Reverse transcription was performed using 250 ng of total RNA and the 10 High capacity cDNA Reverse Transcription Kit (Life Technologies). Quantitative PCR was done 11 using cDNA (150 ng) mixed with TaqMan Fast Advanced Master Mix (Life Technologies). 12 After 40 cycles of amplification (95°C for 20 sec and then 95°C for 1 sec and 60°C for 20 sec), 13 14 data were analysed with the Applied Biosystems[®] Relative Quantification Analysis Module. Amplification curves for each target were individually checked and baselines adjusted, when 15 necessary, to determine the cycle threshold (CT) values. Gene expression was normalized to 16 the mean CT value of four housekeeping genes (Gusb, Hprt, Rps9, Ppia) and expressed as 17 relative gene expression using the $2^{-\Delta CT}$ formula or as a fold change expression using the 2^{-} 18 ΔΔCT formula. 19

20

21 Statistical analysis

22 Unsupervised two-dimensional hierarchical clustering was generated with mean-centred 23 relative expression values $(2^{-\Delta Ct})$ of 91 genes per sample using XLStat software. Distances 24 between samples were calculated based on the ΔCT values using Pearson's Correlation and

average linkage method. The vertical height of the dendogram shows the Euclidean 1 distances between samples. The two-dimensional scatter plot of Principal Component 2 Analysis (PCA) was performed using XLStat and represents the expression pattern of $(2^{-\Delta Ct})$ 3 sample values of the ten subgroups. When plotting the sample data points, F1 (PCA 4 Component 1 (32.85% variance)) was used as the x-axis and F2 (PCA Component 2 (12.69% 5 6 variance)) as the y-axis. Data did not assume a Gaussian distribution and were considered 7 unpaired. The statistical analysis was performed between 2 groups for each OA model versus its respective control (MNX vs MNX-sham, CIOA vs CIOA-sham, Aged vs Young, SP-MNX vs 8 MNX, HG-MNX vs MNX, and HF-MNX vs MNX) using the Mann-Whitney test and GraphPad 9 7 (San Diego, CA, USA). Data were expressed as relative expression $(2^{-\Delta Ct})$ or as fold change 10 (fold change of gene expression in one OA sample as compared to its respective control 11 normalized to 1) and represented as median with interguartile range. Differences were 12 considered significant at p<0.05 and p<0.01. 13

14

15 **RESULTS**

16

Defining the Standard Operating Procedures. One important feature in the study 17 design was to define the SOP after the harmonization of the experimental protocols (from 18 animal models to transcriptomic analysis) in three consensus meetings of the ROAD 19 consortium. A study workflow was designed (Figure 1). At each step, the analysis technique 20 was performed in a single laboratory by the same operator to avoid experimental bias. The 21 centralized OA scoring of the different models and controls (Figure 2A) showed that OA 22 scores were significantly higher in all models (≥ 3 on a scale of 0 to 5) than in their respective 23 control (score ≤ 2), although variability in control samples was observed (Figure 2B). The 24

1 concentration of total RNA isolated from tibial plateau samples was not homogeneous among samples, and was significantly higher in the aged, HG-MNX and HF-MNX models than 2 in their controls (young and MNX mice, respectively) (Figure 2C). The RIN score, which 3 estimates RNA quality and integrity, was heterogeneous among samples, with significantly 4 5 lower scores in samples from the aged and CIOA animals than from their controls (young and 6 CIOA-sham) (Figure 2D). Among all samples, six out of the ten samples per group that met 7 the criteria of selection were analysed by TAC. The mean CT values for the housekeeping 8 genes were significantly higher in the CIOA, HG-MNX and HF-MNX samples than in their controls (Figure 2E). However, the mean CT values for the housekeeping genes were 9 positively correlated with the mean CT values for all genes (Figure 2F). This indicated that 10 the lower expression of housekeeping genes in some samples could be attributed to a lower 11 12 amount of cDNA loaded in the TAC and not to a differential regulation of the housekeeping genes. We also detected the expression of genes specific for cartilage (type II collagen, 13 aggrecan) or bone (Runx2, Sp7) in all OA models (data not shown), indicating that both 14 tissues were represented in our samples. 15

16

TGFβ signatures according to the experimental OA phenotypes. Hierarchical 17 clustering and average linkage clustering of the mRNA expression data in the 10 groups of 18 19 mice (6 OA models and 4 controls) revealed marked differences among groups (Figure 3A). 20 Three main subgroups could be detected: a cluster that included samples from mice with Aging-, HF-MNX-, HG-MNX-related OA; a cluster that included mainly samples from SP-21 sham, SP-MNX mice; and a cluster of samples from MNX, CIOA, and CIOA-sham. Sham 22 23 samples did not cluster together, even though most of them are distributed in the last group with the exception of SP-sham, which is closer to SP-MNX. PCA revealed distinct 24

transcriptional profiles among groups that allowed gathering them in three distinct clusters
(Figure 3B). One (HG-MNX and HF-MNX) was clearly separated from the other two clusters
that included i) CIOA, MNX and young animals, and ii) control groups (sham, SP-sham, CIOAsham). Conversely, old and SP-MNX animals were set apart from the others.

To determine whether a specific gene signature could be associated with the different 5 6 OA phenotypes, the gene expression profile of each OA group was compared with that of its control: MNX vs MNX-sham, CIOA vs CIOA-sham, Aged vs Young, SP-MNX vs MNX, HG-7 MNX vs MNX, and HF-MNX vs MNX. The number of significantly deregulated genes was 8 similar in the MNX, SP-MNX, HG-MNX and HF-MNX groups (around 30 genes) (Figure 3C). 9 Conversely, 15 and 47 genes were deregulated in the samples from CIOA and Aged 10 11 animals, respectively. We identified genes that were common to two or more groups and a gene signature that was specific for each OA model (see Venn diagram in Figure 3D and 12 table 2). Importantly, no gene was deregulated in all six OA models. 13

14

OA model-specific TGF β signatures. To further analyse the specific gene signatures, 15 16 we visualized the genes that were significantly dysregulated (fold change >1.5) in each OA model using Volcano plots. In the MNX model, gene expression profiling revealed that all 30 17 modulated genes were upregulated compared with control (Figure 4A). In the CIOA model, 18 14 of the 15 deregulated genes were significantly upregulated (Figure 4B). Conversely, in the 19 Aging- and SP-MNX-related OA, most genes were downregulated (44/47 and 25/28 genes, 20 21 respectively) (Figure 4C-D). Finally, in the HG-MNX and HF-MNX models, 70% and 71% of 22 genes were upregulated (Figure 4E-F). Only four genes were differentially regulated between these models: Smurf2 and Id2 were upregulated, Tgfbrap1 and Lefty were downregulated 23 only in the HF-MNX model. Altogether, our data revealed that many TGFβ family members 24

were deregulated in the different OA subtypes, supporting the key role of the TGFβ
 pathway, whatever the OA risk factor.

3

A Gdf5, Ltbp4, Cd36 combinatorial gene signature for OA. Then, we split the six 4 OA models in two groups. The first group included the OA models related to obesity or fat 5 6 metabolism (SP-MNX, HG-MNX, and HF-MNX) and/or MNX. The number of shared and 7 specific genes is shown in the Venn diagram (Figure 5A). Most of the modulated genes 8 were common to two or three models, and few genes were specific to each model. However, only Cd36 was deregulated in all four models. The second group included MNX 9 and the two other most common OA models: inflammation (CIOA) and aging (Figure 5B). 10 Approximately 50% of all deregulated genes were specific to each model and only two 11 genes were deregulated in all three models: Gdf5 and Ltbp4. Analysis of these three genes 12 13 in all models and their respective controls showed that Cd36 was significantly upregulated in MNX, SP-MNX, HG-MNX and HF-MNX samples (Figure 5C). Gdf5 was significantly 14 upregulated in the MNX and CIOA models and significantly downregulated in the Aging 15 model. Ltbp4 was significantly upregulated in all models, but for the Aging model where it 16 was significantly downregulated. These data suggest that Cd36 upregulation is a hallmark 17 of trauma-related OA, while the deregulation of Gdf5 and Ltbp4 is related to different OA 18 stimuli. 19

20

21 **DISCUSSION**

22

The first objective of the ROAD consortium was to identify specific gene signatures for the main OA clinical phenotypes using six relevant murine models by focusing on the

1 transcriptomic analysis of the TGFβ pathway. Although this pathway has been extensively studied in some OA murine models [18, 20, 21], it is quite impossible to compare these 2 3 results from independent laboratories due to potential biases that may influence gene expression, such as mouse genetic background, age, sex, housing conditions, 4 histopathological scoring subjectivity and inter-investigator variability. Here, we wanted to 5 6 limit these potential biases by defining SOPs and by centralizing each step of data acquisition and processing, thereby minimizing the risks of failure to identify relevant 7 targets [22-25]. The resulting data allowed the accurate comparative analysis of six models 8 using their respective controls. 9

The main finding of our transcriptomic analysis is the unexpected lack of deregulated 10 genes common to all murine models of OA, although many TGF^β family members were 11 deregulated pointing out the critical role played by the TGF β pathway in OA [26]. This 12 might reflect the heterogeneity of responses to the different stimuli leading to similar 13 symptoms, as observed in patients with OA. Differences in the expression pattern 14 between the different models likely relate to the peculiarities and distinct natures of the 15 models. We are also aware that transcriptional regulation of genes may not be reflected 16 17 at the protein level. Analysis of these differences at the protein level were beyond the scope of the present study but likely warrants further studies. Some genes, such as type II 18 19 collagen, may be differently regulated depending on the OA model suggesting possible different timings or mechanisms of regulation that warrant further investigation. 20 Heterogeneity may also be emphasized by the individual responses within the same 21 22 model, thus highlighting the interest of classifying OA phenotypes using relevant 23 biomarkers in the clinic. Heterogeneity might also reflect different stages of OA in the 24 different models but this is unlikely since the OA scores are similar in all models. The

absence of a common signature could also be due to the late time point (6 weeks after OA
induction and 24 months of age for old mice) chosen for the transcriptomic analysis when
the gene expression profile might reflect an adaptive response. However, this time point
is relevant for patients in whom OA is generally diagnosed long after disease initiation.

Another important finding is the identification of the combinatorial Gdf5-Cd36-Ltbp4 5 6 signature that might discriminate distinct subgroups of OA phenotypes. Indeed, Cd36 was 7 upregulated in all mice that underwent surgical MNX. CD36 is a membrane-bound protein and the receptor of thrombospondin-1, fatty acid translocase (FAT), platelet glycoprotein 8 4 (PG4) and scavenger receptor class B member 3 (SCARB3). It is expressed in adipocytes 9 and mesenchymal stromal cells isolated from fat tissue, and its expression level correlates 10 with poor differentiation into the chondrogenic lineage [27]. CD36 expression is increased 11 at sites of cartilage injury and co-localizes with developing hypertrophic chondrocytes and 12 the aggrecan NITEGE neo-epitope [28]. In patients with OA, CD36 expression has been 13 significantly associated with the presence of osteophytes, of joint space narrowing, and 14 higher Kellgren-Lawrence score [29]. Moreover, in chondrocytes from patients with OA, 15 expression of thrombospondin 1 (a CD36 ligand) is strongly decreased concomitantly with 16 17 the increase in CD36 expression [30]. More recently, the anti-inflammatory and analgesic effects of serum albumin in patients with knee OA was related to inhibition of CD36 in 18 19 synoviocytes, macrophages and chondrocytes [31]. In addition, our study suggests that CD36 might be a specific biomarker of post-traumatic OA. CD36 expression should be thoroughly 20 21 investigated in cartilage and bone samples from patients with different OA phenotypes.

We also found that *Gdf5* expression was deregulated in three of the six OA models under study. It was previously shown that a loss-of-function *GDF5* gene mutation results in joint fusions, and a single-nucleotide polymorphism is associated with higher

1 susceptibility to OA [32]. GDF5 deficiency has also been associated with abnormal ligament laxity and subchondral bone remodelling [33]. Several genome-wide association 2 3 studies (GWAS) have reported the significant association between knee OA and the GDF5 locus [29, 34-36]. Very recently, a GWAS using the United Kingdom OA Biobank cohort 4 reported that GDF5 genetic variants were the strongest predictor of knee pain [37]. In the 5 6 present study, Gdf5 expression was upregulated in the CIOA and MNX models that are 7 characterized by ligament laxity and pain [10, 38]. Our data strongly suggest that GDF5 expression is a biomarker of painful OA phenotypes, as also suggested by genomic studies 8 in humans. 9

Finally, we found that Ltbp4 was deregulated in all six OA models (Figure 5C), although 10 it was not identified as a deregulated gene common to all models in the statistical analysis 11 (Figure 3D). In the bioinformatic analysis, SP-MNX samples were compared with MNX 12 samples (Figure 4) to investigate the impact of the genetic background on OA. Conversely, 13 in the data presented in Figure 5C, all groups were analysed independently of their 14 control. LTBP4 is a key molecule required for the stability of the TGF^β receptor (TGF^βR) 15 complex via interaction with TGFβR2, thereby preventing its endocytosis and lysosomal 16 17 degradation [39]. However, LTBP4 has not been associated with cartilage or OA and unlike its paralogues, LTBP4 is not regulated during chondrogenic differentiation of mesenchymal 18 19 stromal cells [40]. Like Gdf5, Ltbp4 expression was decreased in old mice and not upregulated as observed in the murine models of induced OA. This suggests that 20 spontaneous aging-related OA might involve different mechanisms. 21

In conclusion, the originality of the present study was to rely on relevant murine models of OA to understand the complexity of OA phenotypes in humans through investigation of the TGFβ pathway and based on rigorous SOPs. We did not identify a unique gene signature

common to all six OA phenotypes. This highlights the huge heterogeneity of the animal
models and the need of caution when extrapolating results from one model to another. But
this also highlights that the diversity of the mouse models likely reflects the heterogeneity in
human OA. Further studies are needed to validate these potential signatures.

5

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18 AUTHOR CONTRIBUTIONS

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All authors were involved in revising critically the manuscript and approved the final version.
MM: Data analysis, manuscript writing; DN: Experiment design, data analysis, manuscript
writing; HKE, DM, MR, EH, XH, DC, MCS, CJa, JYJ, MHLP, PR, JS, CV: Experimental work; FR,
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10 COMPETING INTERESTS

- 11
- 12 The authors declare that they have no competing interests.

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Journal Prevention

1 FIGURE LEGENDS

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Fig. 1. Study workflow. Samples were collected from six OA models and their respective 3 4 controls: OA induced by Destabilization of the Medial Meniscus (MNX) versus sham, OA associated with age (Ageing) versus young animals, inflammation (Collagenase-induced OA; 5 CIOA) versus sham, obesity [High Fat (HF) versus Normal Diet (ND)] overweight [Hypergravity 6 7 (HG-MNX) versus MNX] (these models were generated in C57BL/6JRj male mice), and OA 8 associated with metabolic syndrome (Seipin knock-out; SP-MNX versus SP-sham). At the experiment end, knee joints were harvested; femoral condyles and tibial plateaus were 9 prepared for histological analyses and RNA isolation, respectively. OA severity was scored 10 after Safranin-O-Fast Green staining using the OARSI grading system. The expression of 11 genes involved in TGF-β signalling was analysed using custom-made Taqman[™] Array Cards. 12

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Fig. 2. Sample quality controls. (A) Sagittal views of femoral condyles from CIOA-sham (left) 14 and CIOA (right) mice as representative of control and OA cartilage (OA score= 1 and 4, 15 respectively). (B) Distribution of OARSI scores for cartilage destruction in histological 16 sections of femoral condyles. Results are presented as median with interquartile range 17 (n=6). (C) Total RNA concentration (ng/ μ L) after extraction from tibial plateau samples of OA 18 19 models and controls. (D) RNA Integrity Number (RIN) for total RNA extracted from tibial 20 plateau samples of OA models and controls. (E) Mean CT values for the housekeeping genes Ppia, Hprt, Gusb, and Rps9 obtained using Taqman® Array Cards. Data are represented as 21 median with interquartile range; *p <0.05, **p <0.01 (Mann-Whitney test). (F) Correlation 22 23 between the mean CT value of the four housekeeping genes and the mean CT value of all genes. Each dot represents a sample (n=59): Pearson's r=0.8471, p <0.001. 24

Fig. 3. Global gene expression analysis. (A) Unsupervised hierarchical clustering (Pearson correlation, average linkage) of the six samples for each OA model and for each of the four controls. (B) Principal component analysis of the same data as in A. Squares represent the centroid of each group (n=6 mice per group). (C) Number of significantly deregulated genes in the six OA murine models compared with their controls. (D) Venn diagram showing the number of significantly deregulated genes identified in each OA models in a set of 91 targets (Table 1).

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Fig. 4. Analysis of differentially expressed genes in the OA murine models. Results are 10 shown for DMM (A), CIOA (B), Ageing (C), SP-DMM (D), HG-DMM (E) and HF-DMM (F). 11 12 Volcano plots (left panels) show the up- and downregulated genes in each OA model versus its control. For each plot, the x-axis represents the log 2-fold change (FC), and the y-axis 13 represents the log 10 p-values. Genes with an exact p-value <0.05 were considered as 14 differentially expressed. Scatter plots (right panels) show the expression FC of significantly 15 16 deregulated genes (p<0.05) in the six OA models compared with their control group. Results 17 are expressed as the median with interquartile range (n=6).

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Fig. 5. Genes significantly deregulated in the various OA models. (A) Venn diagrams showing the number of differentially expressed genes identified in the four OA models related to overweight or metabolism disorder (MNX, SP-MNX, HG-MNX, HF-MNX), and (B) in the three most common OA models (MNX, CIOA, Aged), among a set of 92 targets (Table 1). (C) Relative gene expression of *Cd36, Gdf5, Ltbp4* in the four controls and six OA models are

- 1 expressed as the median with interquartile range (n=6/group); * p <0.05, ** p <0.01 (Mann-
- 2 Whitney test).
- 3
- 4 Supplementary Fig. Representative photographs of sagittal histological sections of femoral
- 5 condyles from the different OA models.

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