Increased Expression of Matrix Extracellular Phosphoglycoprotein (MEPE) in Cortical Bone of the Rat Tibia after Mechanical Loading: Identification by Oligonucleotide Microarray

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Abstract

Skeletal integrity in humans and animals is maintained by daily mechanical loading. It has been widely accepted that osteocytes function as mechanosensors. Many biochemical signaling molecules are involved in the response of osteocytes to mechanical stimulation. The aim of this study was to identify genes involved in the translation of mechanical stimuli into bone formation. The four-point bending model was used to induce a single period of mechanical loading on the right tibia, while the contra lateral left tibia served as control. Six hours after loading, the effects of mechanical loading on gene-expression were determined with microarray analysis. Protein expression of differentially regulated genes was evaluated with immunohistochemistry. Nine genes were found to exhibit a significant differential gene expression in LOAD compared to control. MEPE, Gam1, V2R2B, and QFG-TN1 olfactory receptor were up-regulated, and creatine kinase (muscle form), fibrinogen-B beta-polypeptide, monoamine oxidase A, troponin-C and kinesin light chain-C were down-regulated. Validation with real-time RT-PCR analysis confirmed the up-regulation of MEPE and the down-regulation of creatine kinase (muscle form) and troponin-C in the loaded tibia. Immunohistochemistry showed that the increase of MEPE protein expression was already detectable six hours after mechanical loading. In conclusion, these genes probably play a role during translation of mechanical stimuli six hours after mechanical loading. The modulation of MEPE expression may indicate a connection between bone mineralization and bone formation after mechanical stimulation.

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Introduction

The ability of the skeleton to form bone after mechanical stimulation is a very complicated process, which consists of mechanosensing, mechanotransduction and the bone formation response. It is well-known that osteocytes function as mechanosensors via canalicular processes and communicating gap junctions in the early stage of bone remodeling [1,2]. Many biochemical signal molecules are involved during mechanical transduction, finally, resulting in bone formation.

In vivo studies in animals report glucose-6-phosphate dehydrogenase (G6PD) [3,4], c-fos [5,6], cAMP [7], COX-2 [8], NO and prostanoid [9], insulin-like growth factors (IGFs) [6,10,11], transforming growth factor-β (TGF-β) [6,10], protein kinase B (PKB or Akt) [12], glutamate transporter (GLAST) [12], wnt/β-catenin [13,14] and sclerostin [14,15] as biochemical signal molecules during the translation of mechanical loading into bone formation. The responses to in vivo mechanical loading are time-dependent. Early strain related changes within 5 minutes after loading are shown in osteocytes in which the glucose-6-phosphate dehydrogenase (G6PD) activity is increased [3,4]. The cAMP level is also rapidly increased after loading [2,7], COX-2 [8], nitric oxide (NO), and prostanoid [9] are also important for bone formation elicited by mechanical...
In vivo mechanical loading experiments within 6 hours after a single loading session, followed by an enhanced c-fos mRNA expression in cortical and trabecular bone surface [11]. In addition, the concentrations of IGF-I mRNA expression is observed on trabecular surfaces and in osteocytes of the diaphyseal cortex (cortical and trabecular osteocytes) of rat caudal vertebrae within 6 hours after a single loading session, followed by an increased expression of type I collagen and osteocalcin on the bone surface [11]. In addition, the concentrations of IGF-I protein within the humerus increased after 6 weeks of weightbearing exercise (5 days/week) in the rat-with-backpack model, whereas the concentrations of TGF-β protein decreased [10].

All of the above mentioned studies observed one single or a few mediators of the osteogenic response, but the microarray technology, used in this study, allows exploring multiple genes due to bending, not squeezing of the tibia and sham-loading only in squeezing (sham-loading). Since loading will result in bending strain. These early events are followed by changes in gene expression level. Compression of caudal vertebrae results in enhanced c-fos mRNA expression in cortical and trabecular osteocytes 30 minutes after loading [5]. Four-point-bending of the tibia shows an up-regulation of c-fos mRNA expression in the periosteum within 2 hours after loading and an increase of IGF-I and TGF-β mRNA in the periosteum within 4 hours after loading [6]. Increased IGF-I mRNA expression is observed on trabecular surfaces and in osteocytes of the diaphyseal cortex (cortical and trabecular osteocytes) of rat caudal vertebrae within 6 hours after a single loading session, followed by an increased expression of type I collagen and osteocalcin on the bone surface [11]. In addition, the concentrations of IGF-I protein within the humerus increased after 6 weeks of weightbearing exercise (5 days/week) in the rat-with-backpack model, whereas the concentrations of TGF-β protein decreased [10].

All of the above mentioned studies observed one single or a few mediators of the osteogenic response, but the microarray technology, used in this study, allows exploring multiple genes at once. The aim of this study was to identify candidate genes that are involved in the translation of mechanical stimuli into bone formation. Load was applied using four-point bending. This model was chosen because adequate bone adaptation was demonstrated previously [16,18,26]. The results from the microarray experiment were confirmed both at RNA level by RT-PCR and at protein level by immunohistochemistry (IHC).

### Materials and Methods

#### In vivo mechanical loading experiments

Three separate mechanical loading experiments were performed to obtain tibia material suitable for detection of RNA expression in microarray and for detection of protein expression by immunohistochemistry, (table 1). Mechanical loading was applied using the 4-point bending system of Forwood and Turner [16,17] and was performed under general anesthesia (2% isoflurane in 1 l/min O2, and 2 l/min N2O). The animal experiments were in accordance with the governmental guidelines for care and use of laboratory animals and approved by the Institutional Animal Care and Use Committee (IACUC) of the VU University Medical Center Amsterdam, the Netherlands. The animals were kept under standard laboratory conditions.

In experiment A nineteen female 12-week-old Wistar rats (Harlan, Zeist, The Netherlands) were randomly assigned to two weight-matched groups: LOAD (223 ± 14 g, n = 9) and SHAM (228 ± 10 g, n = 10). The right tibiae of rats in the LOAD group underwent "medio-lateral" loading (distance between the centers of the loading pads: upper pads: 11 mm and lower pads: 23 mm). The left tibia served as contra lateral control to adjust for systemic effects of the bending. The right tibia of rats in the SHAM group underwent sham loading (opposed pads were placed at the inner position: 11 mm). The left tibia served as contra lateral control to adjust for systemic effects of the squeezing (sham-loading). Since loading will result in bending and squeezing of the tibia and sham-loading only in squeezing of the tibia; the SHAM group was used as control for the LOAD group. A single loading episode constituted 300 cycles (2 Hz) using a peak magnitude setting of 60 N as described before [18]. Measurement with an external force reader before and after bending revealed that the actually used force was 72 ± 3 N (mean ± SD). All left tibia served as contra lateral control. The rats were sacrificed by CO2 asphyxiation exactly six hours after loading. This time-point was based on a previous study of Lean and colleagues [11] and on our own in situ hybridization studies [18,19]. The tibiae were dissected and after removal of the proximal and distal end, periosteum and bone marrow, immediately frozen in liquid nitrogen and stored at -80 °C until RNA isolation.

In experiment B ten female 12-week-old Wistar rats (Harlan, Zeist, The Netherlands) weighing 235 ± 12 g were randomly assigned to two groups: LOAD (n=5) and CONTROL (n=5). The right tibiae of rats in the LOAD group underwent "medio-lateral" loading identical to the rats in the LOAD group of experiment A, while the left tibiae of these rats served as a non-loaded contra lateral control. The rats in the CONTROL group did not receive any mechanical loading. All rats were sacrificed by CO2 asphyxiation six hours after loading. The
tibiae were dissected and the proximal condyles were sawn off. The tibiae were then fixated in 4% (w/v) paraformaldehyde buffered in PBS at 4 °C for 24 hours and decalcified in 10% EDTA with 0.5% paraformaldehyde in PBS at 4 °C for 30 days. The tibiae were dehydrated with ethanol and xylene and embedded in paraffin.

In experiment C nine female 12-week-old Wistar rats (Harlan, Zeist, The Netherlands) weighing 234 ± 12 g were assigned to the REPEATED LOAD group. The right tibia of these rats underwent for two weeks, five days a week, a single episode of "medio-lateral" loading comprising 300 cycles (2 Hz) using a peak magnitude setting of 40 N. Six hours after the final loading episode the rats were sacrificed by CO2 asphyxiation. The tibiae were dissected, the proximal condyles were sawn off and the tibiae were prepared for histology identically to experiment B.

**RNA isolation**

The diaphysis of the tibia from experiment A was pulverized in presence of Trizol (Invitrogen) using the Freezer mill 6750 (Spex Certiprep, Metuchen, NY, USA) subsequently followed by a Trizol extraction, a chloroform/isooamyalcohol extraction, and a second Trizol extraction according to the manufacturer’s instructions. The RNA pellet was dissolved in RNase-free water and stored at -80°C prior to use. Thus the sample included total RNA from the tibia shaft, containing osteocytes, osteoblasts, lining cells and intracortical blood vessel cells. For the real-time RT-PCR analysis the samples were additionally treated with RNase-free DNase to eliminate DNA contamination.

The quality of the RNA samples was determined with the RNA 6000 Nano Assay Kit (Agilent Technologies). The yield of RNA was measured with the spectrophotometer (A260) (NanoDrop® ND-1000 Spectrophotometer). The mean yield of total RNA samples used for the microarray analysis was 1604 ± 501 ng/µl (mean ± SD) ranging from 18.6 µg to 48.9 µg per tibia with an A260/A280 ratio of 1.97 ± 0.05 (mean ± SD).

**Microarray, probe generation, hybridization and washing**

Custom made microarrays were used [20]. The total number of spotted oligonucleotides was 4803 per region, including 52 spots of the housekeeping gene glyceraldehyde-3-phosphate dehydrogenase (GAPDH). Furthermore, there were 138 empty spots. The slides exhibited a crossreactivity of 10%; every spotted oligonucleotide has a chance of 10% that a part of the spotted sequence interacts with a different gene than mentioned in the gene name list.

RNA from five rats of each group was used for microarray analysis cDNA probes were generated from 15 µg total RNA with an oligo-dT ((dT)20-VN) primer (Isogen, Maarssen, The Netherlands) and SuperScript™ II Reverse Transcriptase (Invitrogen), with incorporation of aminoallyl-dUTP (Ambion). Probes were indirectly labeled with Fluorolink Cy3 or Cy5 mono-functional dyes according to a dye-swap design. The dye-swap design was used to avoid a possible bias caused by the molecular structure of Cy3 and Cy5. The treated tibia (LOAD or SHAM) was hybridized together with its own contra lateral control tibia creating unique intra-rat hybridization.

The hybridization protocol was adapted from Snijders and colleagues [21] with minor modifications. Pre-hybridization was performed in a hybridization mixture containing 60 µg yeast tRNA (ribonucleic acid transfer, Sigma), 12 µg polyA (Amersham Biosciences) and 24 µg human Cot-1 DNA (Invitrogen) and 30 µg salmon sperm DNA (Invitrogen) in a total volume of 29.2 µl per array. After precipitation the pellet was dissolved in mastermix (50% formamide, 2xSSC, 9.4% dextran sulphate) and 0.2% SDS in a total volume of 130.2 µl per array. Pre-hybridization was maintained for 45 min at 37°C in the Hybridization Station (Genetac™ hybridization station or Hyb3Array12 (Perkin Elmer)). The pre-hybridization mix was gently removed and slides were dried by centrifugation at 200 g for 3 min. The probe-hybridization mixture contained 16.0 µl Cy3 and 16.0 µl Cy5 labeled samples and 26.2 µl hybridization mixture (60 µg yeast tRNA, 12 µg polyA and 24 µg human Cot-1 DNA). After precipitation the pellet was dissolved in mastermix and 0.2% SDS in a total volume of 130.2 µl per array. Probe mix was denatured at 70°C for 15 min and incubated at 37°C for 60 min. Hybridization was initiated and maintained for 16 h at 30°C in the hybridization station, while the hybridization mixture was agitating. After hybridization, excess hybridization mixture was automatically rinsed off with 50% formamide, 2xSSC, pH 7.0 at 35°C and followed by a wash step with PN buffer (0.1 M sodium phosphate, 0.1% Igepal CA630, pH 8) at 25°C. Excess salt was removed by subsequently rinsing in 0.2xSSC, 0.1xSSC and 0.01xSSC at 25°C. Slides were dried by centrifugation at 200 g for 3 min.

After drying, the slides were scanned at 10 μm resolution for Cy3 and Cy5 intensities using the microarray scanner (Agilent Technologies) operated by Agilent Scan Control software and Feature Extraction software. Array images were processed with BlueFuse version 3.0 for microarrays (BlueGnome, Cambridge, UK). The dataset is submitted to GEO gene expression omnibus (GSE50707) with platform accession number: GPL: 1397.

**Microarray statistics differential gene-expression**

Automatically flagged genes (spots with a confidence value < 0.11) and manually flagged genes (dirty spots and spots with a confidence value between 0.11-0.15 with a low intensity or bad morphology) were excluded from further analysis. The duplicates of the Cy3 and Cy5 intensities were not averaged, but were treated as separate spots in the analysis. The log2 values of the Cy5 to Cy3 ratios were normalized in an intensity dependent fashion (lowess). Genes with more than one missing value across arrays were excluded from the statistical analysis.

Differentially expressed genes were identified by fitting a separate linear model to the expression data for each gene using the language R (http://www.r-project.org). Duplicate spots for each gene, printed on each array, were taken into account [22]. Differential expression was analyzed using moderated t-statistics based on empirical Bayes estimation [22] and P-values were adjusted according to the linear step-up method of Benjamini and Hochberg to reflect the false discovery rate [23,24]. Genes with a false discovery rate <20% were considered to reflect statistical significance.
Estimated relative differential expression for LOAD (i.e., relative to its contra lateral control) was compared to the estimated relative differential expression for SHAM (i.e. relative to its contra lateral control). Thus, aspects of the experimental intervention bending and squeezing were compared with squeezing resulting in the net effect of bending.

**Microarray statistics single channel analysis**

Possible differences between the contra lateral control of the LOAD group and the contra lateral control of SHAM group were investigated. Analyzing the differences between contra lateral control of both groups necessitated single channel analysis to compare the intensities of the control samples between arrays rather than the ratios between the two channels. This analysis was performed on 1944 spots, because missing values could not be handled by the single analysis [25].

**Real-Time RT-PCR**

Quantitative RT-PCR analysis was performed on the nine differentially expressed genes to validate the microarray results. RNA from LOAD (n=8) and SHAM (n=10) was used for quantitative RT-PCR analysis. One hundred ng of total RNA was reverse-transcribed using 10 ng/μl random primers (Roche, Basel, Switzerland) and 5 U/μl M-MLV Reverse Transcriptase (Promega) in a mixture containing 5 mM MgCl₂, 1x RT-buffer, 1 mM dNTPs each, 1M betaine and 0.40 U/μl Transcriptase (Promega) in a mixture containing 5 mM MgCl₂, 1x RT-buffer, 1 mM dNTPs each, 1M betaine and 0.40 U/μl in a total volume of 20 μl. All RNA samples were assayed in triplicate. Three μl of cDNA was amplified by PCR using the primers as described in Table 2. For every gene two primer sets were developed. The first primer set amplified a fragment of the mRNA which overlaps with the oligonucleotide that was spotted on the microarray and the second primer set amplified a fragment of the mRNA that was not present on the array. Both primer sets were used, because the microarray had a crossreactivity of 10%.

In each case, the PCR consisted of an initial denaturation step for 3 min at 95°C, followed by 40-50 cycles (15 sec at 95°C, 1 min at 60°C) in a total volume of 25 μl containing 300nM primers and SYBR Green Supermix (BioRad, Hercules, CA, USA). After the PCR, a melting curve was run from 50°C to 95°C to check the specificity of the reactions.

**Real-Time RT-PCR statistics**

All mean Ct (threshold cycle) values were normalized for the house keeping gene PGBD (porphobilinogen deaminase)(2−ΔΔCt value; ΔCt = Ct GENE OF INTEREST − Ct PGBD), followed by a natural log-transformation to obtain a normal distribution, which was shown with a Kolmogorov-Smirnov test (SPSS, version 9.0). The paired samples t-test (SPSS, version 9.0) was used to examine differences between the treated right and control left tibia in the same rat. The independent samples t-test (SPSS, version 9.0) was used to compare indirect differences between the LOAD and SHAM group (2−ΔΔCt value; ΔΔCt = ΔCt TREATED − ΔCt CONTROL). A P-value < 0.05 was considered to reflect statistical significance.

**Immunohistochemistry**

From both tibiae of the rats from experiment B and C, sections of 5 μm thick were cut. Sections were rehydrated and endogenous peroxidase was quenched with 3% H₂O₂ in 40% methanol/PBS. Antigen retrieval was performed by incubation with 1% trypsin for 15 minutes at 37 °C. After blocking of the non-specific binding sites with blocking reagents for 1 hour the sections were incubated overnight at 4°C with 1/200 rabbit-anti MEPE antibody (Santa Cruz sc-68923, CA, USA). The sections were incubated 1 hour with 1/100 biotinylated goat-anti-rabbit antibody (Dako) and 1 hour with horse radish peroxidase labeled streptavidin (Invitrogen). For color development the sections were incubated with AEC reagent (Zymed) and the sections were counterstained with hematoxilin. Quantitative evaluation of the specific staining for MEPE was performed with NIS Elements digital imaging software (Nikon). Digital images were made of the medial cortex of two sections of each tibia at 200x magnification. In the images the area of MEPE positively stained osteocytes was measured and expressed as a percentage of cortical area to obtain the percentage MEPE positive area. Positive staining in capillary blood vessels was excluded from the measurements.

**Results**

**Identification of differentially expressed mechanosensitive genes with microarray analysis**

After severe filtering of spot quality, 2375 spotted genes of the 4803 genes, including 51 housekeeping genes, were retained in the analysis. Comparison of the LOAD tibia and its contra lateral control tibia resulted in nine significant differentially expressed genes with a false discovery rate < 20%. Four genes showed an up-regulation ranging from 1.4-fold to 1.8-fold and five genes were down-regulated (table 3).

Comparison of the SHAM tibia and its contra lateral control tibia did not result in significant differentially expressed genes (data not shown). Furthermore, the nine genes that were differentially expressed between the LOAD and control comparison were not present in the 189 highest ranked genes of the SHAM versus control comparison. This ranking was based on the posterior odds.

**Validation of differentially expressed genes with real-time RT-PCR**

Real-time RT-PCR analysis confirmed the significant up-regulation of MEPE (P < 0.05) and the tendency to down-regulation of creatine kinase (muscle form) and troponin-C (0.05 < P < 0.10) in the LOAD tibia compared to its contra lateral control tibia (n=8) using the two different primer sets per gene (Figure 1; table 4). No significant differences of MEPE,
| Accession nr | Target Array | Primer set I | Primer set II | Product (bp) |
|--------------|--------------|-------------|--------------|-------------|
| AF269822     |              | Forward(5'-3') | Reverse (5'-3') | 129         |
|              |              | Agaacaagccaccctacacg | Cccactggatgatgactcact | 100         |
|              |              | 1214        | 1342         |             |
| NM_020083    |              | Forward(5'-3') | Reverse (5'-3') | 95          |
|              |              | Nucleotide Position | 126          |             |
|              |              | 3604        | 3698         |             |
| NM_012530    |              | Forward(5'-3') | Reverse (5'-3') | 115         |
|              |              | Nucleotide Position | 126          |             |
|              |              | 3184        | 3290         |             |
| U05675       |              | Forward(5'-3') | Reverse (5'-3') | 105         |
|              |              | Nucleotide Position | 126          |             |
|              |              | 324         | 3290         |             |
| AF053990     |              | Forward(5'-3') | Reverse (5'-3') | 105         |
|              |              | Nucleotide Position | 126          |             |
|              |              | 324         | 3290         |             |
| D00688       |              | Forward(5'-3') | Reverse (5'-3') | 115         |
|              |              | Nucleotide Position | 126          |             |
|              |              | 324         | 3290         |             |
| J00793       |              | Forward(5'-3') | Reverse (5'-3') | 111         |
|              |              | Nucleotide Position | 126          |             |
|              |              | 324         | 3290         |             |
| AF091565     |              | Forward(5'-3') | Reverse (5'-3') | 106         |
|              |              | Nucleotide Position | 126          |             |
|              |              | 324         | 3290         |             |
| M75148       |              | Forward(5'-3') | Reverse (5'-3') | 106         |
|              |              | Nucleotide Position | 126          |             |
|              |              | 324         | 3290         |             |
| PBGD         |              | Forward(5'-3') | Reverse (5'-3') | 135         |
|              |              | Nucleotide Position | 126          |             |
|              |              | 324         | 3290         |             |

Primer set I contained the sequence of the oligonucleotide as spotted on the microarray. Primer set II contained a fragment of the gene which was not present on the array. Target array, position oligonucleotide fragment as spotted on the microarray. PBGD, house keeping gene porphobilinogen deaminase.
creatinine kinase (muscle form) and troponin-C were found in the SHAM tibia compared to its contra lateral control tibia (n=10) (table 4). The other genes did not show significant differences in gene-expression between the LOAD tibia and its contra lateral control tibia (n=8) or SHAM tibia and its contra lateral control tibia (n=10) (table 4). Fibrinogen B beta chain was undetectable in the tibia irrespective of their treatment (LOAD, SHAM, control) as measured with both primer sets.

Comparison between the LOAD (n=8) and the SHAM group (n=10) showed a significant up-regulation of MEPE with primer set II of the 2^-ΔΔCt value in the LOAD group (independent samples t-test: P < 0.05) (table 4). No differences in expression between the LOAD (n=8) and the SHAM (n=10) group were shown in the other genes (table 4).

**Single channel analysis**

When comparing the contra lateral control tibia of the LOAD group with the contra lateral control tibia of the SHAM group, a false discovery rate < 20% yielded 3 significant down-regulated genes: p21 (c-Ki-ras), phosphate regulating neutral endopeptidase on the X chromosome (X-linked hypophosphatemia XLH)(PHEX) and amino acid transporter system A (ATA2).

**Immunohistochemistry for MEPE**

Protein expression of MEPE was seen primarily in the osteocytes of the cortex and the trabeculae and not in the osteoblasts on the endocortical bone surface or in the growth plate (Figure 2d). In the CONTROL group there was no significant difference in the percentage MEPE positive area between the right and left tibia (1.02 ± 0.31 vs 0.99 ± 0.12; Figure 2b). In the LOAD group the percentage MEPE positive area was higher in the tibia that had received mechanical loading compared to its contralateral left tibia as is shown in Figure 2a, the difference between the loaded tibia and the control tibia being borderline significant (1.48 ± 0.59 vs 0.92 ± 0.33; p = 0.06) After repeated mechanical loading of the right tibia for two weeks no difference in percentage MEPE positive area between the loaded tibia and the control tibia was observed (0.62 ± 0.36 vs 0.62 ± 0.33; Figure 2c).

**Discussion**

This study examined the effect of mechanical loading on the expression of multiple genes in vivo in rats using microarray technology. This in vivo study showed differential gene expression in non-pooled and non-amplified samples of rat tibia RNA six hours after a single mechanical loading session using the 4-point-bending system. Hybridization of loaded or sham-loaded tibia versus the contra lateral control tibia of the same rat was performed. Compared to its contralateral control group, the LOAD group showed a significant up-regulation of the following genes: MEPE, Gami1, putative pheromone receptor V2R2B and isolate QFG-TN1 olfactory receptor. Furthermore, five genes were significantly down-regulated i.e., creatine kinase (muscle form), fibrinogen B beta polypeptide, monoamine oxidase A, troponin-C and kinesin light chain C.

Validation with real-time RT-PCR analysis confirmed the up-regulation of MEPE and the down-regulation of creatine kinase.
Figure 1. Real-time RT-PCR results of MEPE (A-B), creatinine kinase (muscle form) (C-D) and troponin-C (E-F) mRNA expression in the loaded tibiae and their contralateral controls represented per rat. Different markers represent different rats. Lines connect the right and left tibia of the same rat. For each gene, two primers were used, primer set I, which contained a sequence of the oligonucleotide as spotted on the microarray (A, C, E) and primer set II, which contained a fragment of the gene that was not present on the microarray (B, D, F).

* P < 0.05; 0.05 < P < 0.10.

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### Table 4. VALIDATION OF DIFFERENTIALLY EXPRESSED GENES WITH REAL-TIME RT-PCR.

| Gene name               | 2−ΔΔCt (MEAN ± SD) control LOAD (n=8) | 2−ΔΔCt (MEAN ± SD) LOAD (n=8) | P-value | 2−ΔΔCt (MEAN ± SD) control SHAM (n=10) | 2−ΔΔCt (MEAN ± SD) SHAM (n=10) | P-value | SHAM (n=10) LOAD (n=8) | P-value |
|------------------------|---------------------------------------|--------------------------------|---------|---------------------------------------|--------------------------------|---------|------------------------|---------|
| MEPE                   | II 8.43 ± 5.24                        | 14.33 ± 4.59                  | 0.009a  | II 11.39 ± 5.90                      | 12.13 ± 4.02                  | 0.455   | II 1.20 ± 0.53         | 2.03 ± 0.53 |
| Garnl/Tulip 1          | I 0.65 ± 0.124                        | 0.63 ± 0.124                   | 0.971   | I 0.73 ± 0.26                        | 0.787 ± 0.314                  | 0.836   | I 0.14 ± 0.45          | 1.09 ± 0.53 |
| Creatine kinase (muscle form) | I 8.43 ± 5.24                        | 14.33 ± 4.59                  | 0.009a  | II 11.39 ± 5.90                      | 12.13 ± 4.02                  | 0.455   | II 1.20 ± 0.53         | 2.03 ± 0.53 |
| V2R2B                  | I 0.010 ± 0.005                       | 0.008 ± 0.002                  | 0.717   | I 0.010 ± 0.004                      | 0.010 ± 0.004                  | 0.763   | I 1.24 ± 0.58          | 1.11 ± 0.77 |
| Monoamine oxidase A    | II 0.019 ± 0.009                      | 0.017 ± 0.005                  | 0.941   | II 0.021 ± 0.006                     | 0.021 ± 0.006                  | 0.892   | II 1.03 ± 0.28         | 0.96 ± 0.36 |
| Troponin-c             | I 1.18 ± 1.700                        | 0.241 ± 0.532                  | 0.074b  | I 1.105 ± 1.516                      | 0.660 ± 1.091                  | 0.376   | I 0.76 ± 12.14         | 1.89 ± 4.30 |
| QFG olfactory receptor | I 0.640 ± 0.827                       | 0.163 ± 0.369                  | 0.080b  | I 1.353 ± 2.260                      | 0.518 ± 0.823                  | 0.387   | I 9.56 ± 22.36         | 2.31 ± 5.42 |
| Kinesin light chain    | I 0.005 ± 0.003                       | 0.002 ± 0.007                  | 0.882   | I 0.013 ± 0.007                      | 0.012 ± 0.005                  | 0.871   | I 1.15 ± 0.72          | 1.12 ± 0.48 |
| I, primer set I: contained sequence of the oligonucleotide as spotted on the microarray; II, primer set II: contained a fragment of the gene which was not present on the array; ΔΔCt = ΔCt GENE OF INTEREST − ΔCt HOUSE KEEPING GENE (PGBD); ΔΔCt = ΔΔCt TREATED − ΔΔCt control; a P < 0.05; b 0.05 < P < 0.1; doi: 10.1371/journal.pone.0079672.t004

### Figure 2. Immunohistochemistry for MEPE on sections from experiment B and C. Figures show the percentage MEPE positive area in the right and left tibiae of the individual rats after mechanical loading of only the right tibia. (A) LOAD group, (B) CONTROL group, (C) REPEATED LOAD group (D) representative MEPE staining of a left control tibia and (E) representative MEPE staining of the loaded right tibia of the same rat; arrows indicate positively stained osteocytes. bar represents 100 µm.

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(muscle form) and troponin-C in the LOAD tibia compared to the contralateral control tibia six hours after a single mechanical stimulation. Because the microarrays exhibited a crossreactivity of 10%, two pairs of primers for the validation analysis were used. Both primer sets showed a similar pattern. We therefore conclude that at least three of the differentially expressed genes, MEPE, creatine kinase (muscle form) and troponin-C of the microarray analysis were specific.

MEPE protein, as measured with immunohistochemistry, increased six hours after loading, the increase being borderline significant. This suggests that the increase in MEPE mRNA is quickly translated into protein, but it is possible that the peak in protein expression occurs at a later time point. To our surprise the level of MEPE protein in the loaded tibia was not changed after repeated loading of the tibia for two weeks, compared to the control tibia. As we did not isolate RNA from the REPEATED LOAD experiment we do not know if MEPE gene expression follows the same pattern. It is possible that the diminishing response of MEPE to mechanical loading is the result of a feedback mechanism.

MEPE is a member of the Small Integrin Binding Ligand N-linked Glycoprotein (SIBLING) family of proteins (i.e. bone sialoprotein, osteopontin, dentin sialophosphoprotein, and dentin matrix protein)[31,32]. Members of the SIBLING family share features like the RGD motif (arginine-glycine-asparagine) and the ASARM motif (Acidic-Serine-Aspartate-Rich-MEPE) [33,34]. In rodents, MEPE mRNA and protein are expressed by osteoblasts and osteocytes [35], and in human bone by osteocytes within mineralized bone [6] and by bone-marrow cells [33]. MEPE plays a role in the regulation of bone mineralization, dentin mineralization, skeletal development, regulation of bone remodeling, renal phosphate handling and vitamin D metabolism [31,33,35]. The importance of MEPE during the bone formation after mechanical stimulation has been described earlier [27-30]. Mechanical loading induces MEPE mRNA expression in the osteocytes in a unique time-course and spatial pattern in mice [28-30]. Our results confirm these studies and show that both MEPE mRNA and protein are both increased six hours after mechanical loading. Nagel and colleagues reported that MEPE requires inducible COX-2 to exert potent anabolic effects on normal human bone marrow osteoblast precursors [37]. It is conceivable that up-regulation of COX-2, which is an early response to mechanical loading [8,38-41], is followed by an increased MEPE mRNA expression six hours after loading in vivo. Although this micro-array study did not find sclerostin to be increased after loading this has been demonstrated in literature [14,15,30]. In addition it is suggested that sclerostin is locally regulating mineralization through a MEPE-Asarm dependent mechanism [42].

The role of MEPE in bone has not yet been elucidated. MEPE-derived ASARM peptide has been shown to directly inhibit mineralization of bone tissue and also appears to inhibit sodium-dependant phosphate uptake in the kidney, thereby reducing serum phosphate concentration. On the other hand Hayashibara reported that a synthetic fragment of human MEPE (AC-100) can stimulate new bone formation in vivo [36]. Overexpression of MEPE in transgenic mice resulted in decreased BMD and bone remodeling while MEPE knockout mice showed increased trabecular bone mass and increased osteoblast activity[43,44]. Whether increased MEPE acts to locally limit mineralization to prepare for increased bone formation or whether it stimulates bone formation itself needs to be further investigated. In bone tissue the actions of MEPE appear to be regulated by the presence of phosphate regulating neutral endopeptidase on the X chromosome (PHEX). This protein can bind MEPE and inhibits thereby the formation of the ASARM peptide. Interestingly, PHEX is one of the three genes that were significantly down regulated when differences between the contra lateral control tibia of the LOAD group and the contra lateral control tibia of the SHAM group were investigated in the single channel analysis. This suggests the possibility that loading of one tibia leads to simultaneous changes in the contra lateral tibia.

Both, the microarray study and the validation with real-time RT-PCR showed that creatine kinase (muscle form) and troponin-C were negatively regulated by mechanical stimulation. Creatine kinase (muscle form) is expressed in adult skeletal and cardiac muscle of mice [45]. It is involved in phosphocreatine biosynthesis and metabolism (Rat Genome Database). Calbindins or vitamin D dependent calcium binding proteins are members of the troponin-C (the calcium-binding subunit of the troponin complex) superfamily [46]. Vitamin D deficiency will result in a decreased troponin-C concentration in rabbits [47]. Our study showed that creatine kinase (muscle form) and troponin-C are associated with the regulation of bone formation six hours after mechanical stimulation. This opens new insight into the process of bone formation after mechanical stimulation, especially for creatine kinase (muscle form) and troponin-C.

This study showed a low number of significant differentially expressed genes at a false discovery rate of 20%. However, in microarray studies many differentially expressed genes are commonly missed when their discovery rates are slightly above the threshold [48]. Our statistical analysis shows that 22 genes are differentially expressed at a false discovery rate of 50%, out of the 2375 genes that could be included in the analysis. This means that about 11 of these genes are differentially expressed and about 11 are false positives. This suggests that the number of genes that change in expression is appreciable larger than the number of genes that were discovered at a false discovery rate of 20% and confirmed by PCR. Other relevant genes besides the 2375 are not included in the analysis because they are not on the microarray or did not meet filter criteria, it is likely that many genes are involved in the response to loading of the bone. This is in agreement with the notion that many biological factors are involved in the process of bone formation (see Introduction).

Unless gene expression changes are very large, differentially expressed genes are often difficult to detect with microarrays. This is among others caused by biological variability commonly encountered in gene expression, the noise in microarray measurements which necessitates extensive filtering of the spots to maintain quality and the need to control statistically for multiple testing of the many thousands of probes. Our experimental set up is unique since we have an intra-animal control instead of a reference RNA control pool. In this model
the treated right tibia is compared to the contra lateral control tibia which resulted in less noise. On the other hand the selection criteria of the spots were very severe. Of the 4803 genes, 2375 genes were left after selection. Some mechanosensitive candidate genes, as mentioned in the introduction, were lost due to filtering criteria. However the IGF-I spot was present after selection. Apparently, it is not possible with the microarray analysis to detect the 2-fold increase of IGF-I as shown with in situ hybridization [18]. This 2-fold increase was restricted to a specific location in the tibia, i.e. in the osteocytes at the endosteal side of the shaft, whereas with the microarray analysis the RNA of a larger part of the shaft is used and therefore a local difference may not be detected. Xing and colleagues [49] also described the effects of 4-point-bending using the microarray technique. They used the mouse as animal model, which has the advantage of exploring 20,280 transcripts at once; however, a disadvantage of their model system is the low amount of RNA per tibia and data of single tibia were therefore not available. The experimental setup of their study was different from our study. They analyzed gene expression 24 hours after 4 days of multiple loading sessions, whereas we studied the osteogenic response six hours after a single loading session. So, the genes in their study may include genes up- or down-regulated between 24 hours and 5 days after loading, i.e. a complex cascade of genes, whereas in our study direct is restricted to the effects 6 hours after one single mechanical loading session were analyzed. Only one gene, (mono) amine oxidase A, was differentially expressed in both studies.

Since the loading consisted of bending and squeezing and the sham-loading consisted of squeezing, comparing the results of loaded tibia with the sham-loaded tibia will reveal the pure effect of bending. The real-time RT-PCR analysis showed a significant up-regulation of MEPE and down-regulation of creatine kinase (muscle form) and troponin-C in the LOAD tibia compared to the contra lateral control tibia and did not show a significant change of expression between the SHAM tibia and the contra lateral control tibia. Thus, the increase of MEPE expression and decrease of creatine kinase (muscle form) and troponin-C expression appears to be due to bending.

A limitation of our study is the fact that MEPE protein detection by IHC was not investigated after SHAM loading. Since comparison of the SHAM tibia and the contralateral control did not reveal any differentially expressed genes, we decided to exclude this group in the IHC experiments. Staining of capillary blood vessels was also excluded from our measurements, because osteocytes are thought to be responsible for relaying the mechanical loading signal. To detect differences in the level of MEPE protein production by osteocytes we measured MEPE positive stained bone area instead of MEPE positive osteocytes, since this is a more accurate method.

We used the 4-point bending system which results in lamellar bone formation at the endosteal side of the shaft. Validation of the 4-point-bending system at our laboratory showed bone formation at the endosteal surface of the rat tibia 15 days after a single loading session [18] An advantage is the non-invasive application of a mechanical load to the tibia. The magnitude of the force, frequency, duration and total cycles of loading can be adjusted. However a disadvantage is the woven bone formation at the periosteal side of the shaft. An alternative for the 4-point bending model is the non-invasive ulna model, an axial compression model, in which woven bone formation at the periosteal side of the ulna is not observed.

In conclusion, the present study shows that expression of MEPE mRNA and protein, are changed during translation of mechanical stimuli six hours after mechanical loading. This could indicate that modulation of bone mineralization might play a role in the response of bone after mechanical stimulation. This study also shows that microarray experiments are helpful to identify new genes such as creatine kinase (muscle form) and troponin-C, which could have a role in bone formation after mechanical stimulation.

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Author Contributions

Conceived and designed the experiments: NB PL. Performed the experiments: CMAR HWvE. Analyzed the data: CMAR BTTMvR JHGMvB. Contributed reagents/materials/analysis tools: JHGMvB BY. Wrote the manuscript: CMAR NB PL HWvE JHGMvB MAB.

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