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3 **Automated MRI assessment confirms cartilage thickness modification in patients**
4 **with knee osteoarthritis: post-hoc analysis from a phase II sprifermin study.**

5
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23
24 **Running headline:** Post-hoc analysis of sprifermin study

28 **ABSTRACT**

29 *Background:* Sprifermin is under investigation as a potential disease-modifying osteoarthritis drug.
30 Previously, 2-year results from the FORWARD study showed significant dose-dependent modification of
31 cartilage thickness in the total femorotibial joint (TFTJ), medial and lateral femorotibial compartments
32 (MFTC, LFTC), and central medial and lateral TFTJ subregions, by quantitative magnetic resonance
33 imaging (qMRI) using manual segmentation.

34 *Objective:* To determine whether qMRI findings from FORWARD could be reproduced by an independent
35 method of automated segmentation using an identical dataset and similar anatomical regions in a post-
36 hoc analysis.

37 *Method:* Cartilage thickness was assessed at baseline and 6, 12, 18 and 24 months, using automated
38 cartilage segmentation with active appearance models, a supervised machine learning method. Images
39 were blinded for treatment and timepoint. Treatment effect was assessed by observed and adjusted
40 changes using a linear mixed model for repeated measures.

41 *Results:* Based on automated segmentation, statistically significant, dose-dependent structural
42 modification of cartilage thickness was observed over 2 years with sprifermin vs placebo for TFTJ (overall
43 treatment effect and dose response, both $P<0.001$), MFTC ($P=0.004$ and $P=0.044$), and LFTC (both
44 $P<0.001$) regions. For highest dose, in the central medial tibial ($P=0.008$), central lateral tibial ($P<0.001$)
45 and central lateral femoral ($P<0.001$) regions.

46 *Conclusions:* Cartilage thickness assessed by automated segmentation provided a consistent dose
47 response in structural modification compared with manual segmentation. This is the first time that two
48 independent quantification methods of image analysis have reached the same conclusions in an
49 interventional trial, strengthening the conclusions that sprifermin modifies structural progression in knee
50 osteoarthritis.

51

52 **Keywords:**

53 Osteoarthritis

54 DMOAD

55 Cartilage

56 Machine learning

57 Active appearance models

58

59

1 Introduction

2 Cartilage is a key tissue of interest in structure-modification trials of osteoarthritis (OA). Although
3 radiographic joint space width (JSW), a surrogate for structural progression, is one of the regulatory
4 endpoints in these trials, there is increasing evidence of the benefits of direct measures of cartilage
5 morphology using quantitative magnetic resonance imaging (qMRI)¹. Techniques employing manual
6 segmentation of cartilage have been explored with respect to a number of morphological characteristics,
7 including volume and thickness. These techniques have been extensively validated, including construct
8 validity against invasive measurement of cartilage volume and thickness, radiographic JSW, predictive
9 and concurrent validity, and clinical outcomes².

10 However, manual segmentation of cartilage morphology is time consuming and challenging, as
11 careful attention must be paid to detecting the eroding outer margin of the cartilage. To address these
12 issues, various methods of semi-automated or fully automated segmentation have been developed. Fully
13 automated methods based on active appearance modeling (AAM) have demonstrated good
14 measurement accuracy for a number of MRI-assessed tissues including knee cartilage³.

15 Many previous disease-modifying osteoarthritis drug (DMOAD) studies have focused on use of
16 anticatabolic agents to delay progression of cartilage breakdown⁴. An alternative approach is to
17 stimulate cartilage development and repair. Sprifermin, a novel recombinant human fibroblast growth
18 factor-18, is currently under investigation as a potential DMOAD. Sprifermin induces hyaline cartilage
19 formation in vitro and in vivo by increasing chondrocyte proliferation, resulting in increased overall
20 extracellular matrix production^{5,6}.

21 FORWARD (NCT01919164) is a 5-year, multicenter, randomized, placebo-controlled Phase II
22 study, evaluating the efficacy and safety of intra-articular sprifermin in patients with symptomatic
23 radiographic knee OA. The primary 2-year results from FORWARD showed significant dose-dependent
24 modification of cartilage thickness change in the total femorotibial joint (TFTJ), medial and lateral
25 femorotibial compartments (MFTC, LFTC), and central medial and central lateral TFTJ subregions⁷.
26 Cartilage thickness was measured by qMRI; images were analyzed at a single center by manual cartilage
27 segmentation.

28 Here, we conducted a retrospective analysis of MR images from FORWARD, to determine
29 whether qMRI findings assessed by manual segmentation could be reproduced using a previously
30 validated independent method of automated segmentation³.

31

32 Methods

33 Patients aged 40–85 years with symptomatic radiographic knee OA, Kellgren-Lawrence Grade 2
34 or 3, and medial JSW ≥ 2.5 mm in the target knee were randomized (1:1:1:1:1) to receive double-blinded
35 3-weekly intra-articular injections of sprifermin 100 μ g every 6 months (q6mo); 100 μ g every 12 months
36 (q12mo); 30 μ g q6mo; 30 μ g q12mo; or placebo. MR images were acquired at baseline, 6, 12, 18 and 24
37 months using 1.5 or 3 Tesla clinical MRI scanners using a coronal spoiled gradient echo sequence with fat
38 saturation or water excitation, and 1.5 mm slice width with 0.31 mm x 0.31 mm in-plane resolution, as
39 previously reported⁷. The study protocol was approved by independent ethics committees or
40 institutional review boards at all study sites. Written informed consent was obtained from all

41 participants, and the study was performed in accordance with the ethical principles of the Declaration of
42 Helsinki.

43 The manual segmentation method has been presented previously^{2, 7, 8}. As with the original
44 analysis⁷, all images were blinded with regard to acquisition order and active treatment/placebo status.

45 Automated cartilage segmentation was performed as a retrospective analysis using a previously
46 validated method³. AAM, a supervised machine learning method (Imorphics Ltd, Manchester, UK), was
47 used to produce maps of cartilage thickness for femoral and tibial cartilage surfaces³. Each timepoint was
48 analyzed independently. As for the previously-employed manual method, total cartilage thickness was
49 computed as total volume divided by total surface area (i.e., average cartilage thickness) for the TFTJ,
50 MFTC and LFTC regions. These regions replicated those used for manual segmentation by
51 Chondrometrics by following published region descriptions^{8, 9} (**Supplementary Figure 1**) and were
52 automatically projected out to each image segmentation during image search by the AAM.

53 In addition, regions representing the central medial tibial (cMT), central medial femoral (cMF),
54 central lateral tibial (cLT) and central lateral femoral (cLF) plates were produced according to previously
55 described manual definition on the mean bone shape¹⁰. These central regions are similar, but not
56 identical to the segmentation regions previously used in the FORWARD trial (**Supplementary Figure 1**)^{7, 8}.
57 Again, these regions were automatically projected out to each image segmentation during image search
58 by the AAM. The cMT, cMF, cLT and cLF regions were defined by Imorphics based on independent data
59 from the Osteoarthritis Initiative and were based on regions that changed most in an Osteoarthritis
60 Initiative data set¹¹. Dense pointwise maps of cartilage changes were produced as standardized response
61 means (SRMs) of change over 2 years in all available knees. The central regions corresponding to SRM
62 >0.7 were defined by smoothing with an enclosing ellipsoid shape (cLT and cMT) or rectangle (cMF and
63 cLF). Of note, these regions correspond closely to the meniscal windows of the joint at 15° of flexion.
64 Average cartilage thickness in these regions was calculated by taking the mean of a set of thickness
65 measures orthogonal to the bone surface and located at each of the AAM correspondence landmarks.

66 As in the previous analyses⁷, the treatment effect on change from baseline was assessed for each
67 method by observed and adjusted changes using a linear mixed model for repeated measures. The
68 analysis was implemented in SAS (PROC MIXED) using treatment group, time, and pooled country as class
69 variables, baseline as covariate, and treatment by time interaction. An unstructured covariance matrix
70 was used to account for the repeated measures within each patient. The linear dose relationship was
71 tested at Year 2 at the two-sided 5% significance level. If the null hypothesis of no linear relation was
72 rejected, the effect of treatment was assessed by pairwise comparisons of absolute change from
73 baseline in cartilage thickness (sprifermin treatment groups versus placebo). Dunnett's approach was
74 used to account for comparison of four dose groups to placebo at a given timepoint. No direct statistical
75 comparisons between the manual and automated methods were prespecified or conducted, as this was
76 not the aim of the study. Statistical significance was set at $P < 0.05$.

77

78 Results

79 550 patients were recruited at 12 sites in the EU, USA and Hong Kong, and 549 were randomized
80 as an intention to treat (ITT) set. The modified ITT (mITT) analysis set included all patients from the ITT
81 analysis set who had a baseline (prior to first injection) and at least one post-treatment qMRI assessment

82 available up to Year 2. mITT subject numbers were 101 (100 µg q6mo); 99 (100 µg q12mo); 99 (30 µg
83 q6mo); 99 (30 µg q12mo); and 96 (placebo).

84 Using the automated method, statistically significant, dose-dependent structural modification of
85 cartilage thickness was observed over 2 years with sprifermin vs placebo for the TFTJ region (**Figure 1,**
86 **top panel**) (overall treatment effect and dose response across all doses, both $P < 0.001$), and also for the
87 MFTC ($P = 0.004$ and $P = 0.044$) and LFTC (both $P < 0.001$) regions. Statistically significant, dose-
88 dependent structural modification of cartilage thickness over 2 years was also observed for sprifermin vs
89 placebo in the cMT (100 µg q6mo [$P = 0.008$]), cLT (100 µg q6mo, q12mo [both $P < 0.001$]) and cLF
90 (100 µg q6mo, q12mo [both $P < 0.001$]) regions. In the cMF region, there was no significant treatment
91 effect ($P = 0.149$), but there was a linear trend for dose responsiveness ($P = 0.013$).

92 Statistically significant differences in the mean change from baseline in cartilage thickness at Year 2
93 for the highest sprifermin dose (100 µg q6mo) vs placebo were obtained for the TFTJ, MFTC and LFTC
94 regions for both methods (**Table 1**). Statistically significant differences were observed from baseline for
95 the highest sprifermin dose vs placebo in the cMT, cLT and cLF regions using the automated method:

- 96 • cMT (mean difference [95% confidence interval (CI)]: 0.09 [0.02, 0.16], $P = 0.008$)
- 97 • cMF (linear trend: $P = 0.013$; mean difference [95% CI]: 0.06 [0.00, 0.12], $P = 0.061$)
- 98 • cLT (mean difference [95% CI]: 0.15 [0.08, 0.22], $P < 0.001$)
- 99 • cLF (mean difference [95% CI]: 0.10 [0.06, 0.13], $P < 0.001$)

100 Scatter plots and regression lines showed consistency of results between the manual and automated
101 methods of MR image analysis assessing change in cartilage thickness from baseline to 2 years in the
102 TFTJ, MFTJ and LFTJ regions (**Supplementary Figure 2**).

103

104 Discussion

105 Measures of cartilage morphometry have been utilized previously in DMOAD trials⁷. However,
106 this is the first time that two independent methods of image analysis have been applied to the same
107 interventional trial population and have reached the same conclusions regarding structural modification,
108 demonstrating beneficial effects of sprifermin on cartilage thickness.¹

109 Careful manual segmentation and review by an expert reader is the gold standard for
110 morphometric analysis of cartilage using MR images¹². MRI cartilage thickness measures are associated
111 with OA progression and joint replacement and provide more responsive measures of progression than
112 radiographic JSW¹³⁻¹⁷. Automated methods of cartilage segmentation based on machine learning and
113 artificial intelligence are showing increasing promise¹². Here we used a previously published method
114 based on AAMs to provide independent comparator measurements. Although this automated method
115 has been shown to be highly correlated with manual segmentation, Bland-Altman analysis of agreement
116 does indicate systematic bias, and in measurement of longitudinal change, the automated method
117 produced almost twice the cartilage thickness change of the manual method³. This, and the fact that
118 correlation may vary depending on the datasets that are compared, means that some differences in
119 measurements would be expected between the automated and manual methods applied to this image
120 dataset.

121 This automated method produced a consistent and similar pattern of structural modification in
122 the FORWARD study compared with previously reported manual segmentation (**Figure 1**)⁷. Although a
123 larger change in the mean cartilage thickness from baseline was observed for all dose groups, as noted
124 previously³, it was associated with greater measurement variance compared to the manual method. It
125 should be noted that the average cartilage thickness loss that was seen in this study (around 30 µm over
126 two years) is much less than one voxel width. Cartilage loss is therefore a change in the margin of
127 cartilage that is defined by the partial volume in an MR image sampling voxel, which is typically 5–10
128 times the magnitude of the change over 2 years. It is likely that the human operator and the automated
129 algorithm make different decisions about where the cartilage edge actually lies within the image, and
130 this may explain the differences in cartilage thickness changes and measurement error seen here.

131 There were some limitations in this study. There was no direct comparison of methods, as this
132 was not the aim of the study. Further, this was a post-hoc analysis; however, all the available image data
133 were utilized even though some may not have been optimal for automated analysis. Although the TFTJ,
134 MFTJ and LFTJ regions used for the previous manual segmentation analysis were replicated for the
135 automated analysis, the cMT, cLT, cMF and cLF regions were not used originally, but rather aggregate
136 measures of medial and lateral femorotibial subregions (cMFTC and cLFTC, respectively) were used, and
137 therefore these subregion results could not be compared directly.

138 In summary, this post-hoc analysis is unique in that two independent quantitative image analysis
139 methods demonstrated the same results, and strengthens the conclusions that the investigational agent,
140 sprifermin, modifies cartilage loss/structural progression in knee OA.

141

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148

149 **Author contributions**

150 FM, HG and CL were responsible for the conception and design of the study. AB was responsible for
151 drafting the article. AB, PC, MB and FE provided critical revision of the article for important intellectual
152 content. FM provided statistical expertise. All authors were responsible for analysis and interpretation of
153 the data and for final approval of the article and take responsibility for the integrity of the work.

154

155 **Role of the funding source**

156 This study is sponsored by Merck KGaA, Darmstadt, Germany. The study sponsor was involved in the
157 study design, collection, analysis, and interpretation of data and in the writing of the manuscript and
158 decision to submit for publication.

159 **Conflict of interest**

160 **AB** and **MB** are employees of Imorphics, Manchester, UK. **PGC** has done consultancies or speakers
161 bureaus for AbbVie, Bristol Myers Squibb, EMD Serono, Flexion Therapeutics, Galapagos,
162 GlaxoSmithKline, Novartis, Pfizer and Stryker. **CL** and **HG** are employees of Merck KGaA, Darmstadt,
163 Germany. **FM** is an employee of EMD Serono (a business of Merck KGaA, Darmstadt, Germany). **FE** is an
164 employee and shareholder of Chondrometrics GmbH, and has received consulting fees from Merck
165 KGaA, Samumed LLC, Abbvie, Bioclinica, TissueGene, Servier, Galapagos, Roche, and Novartis

166

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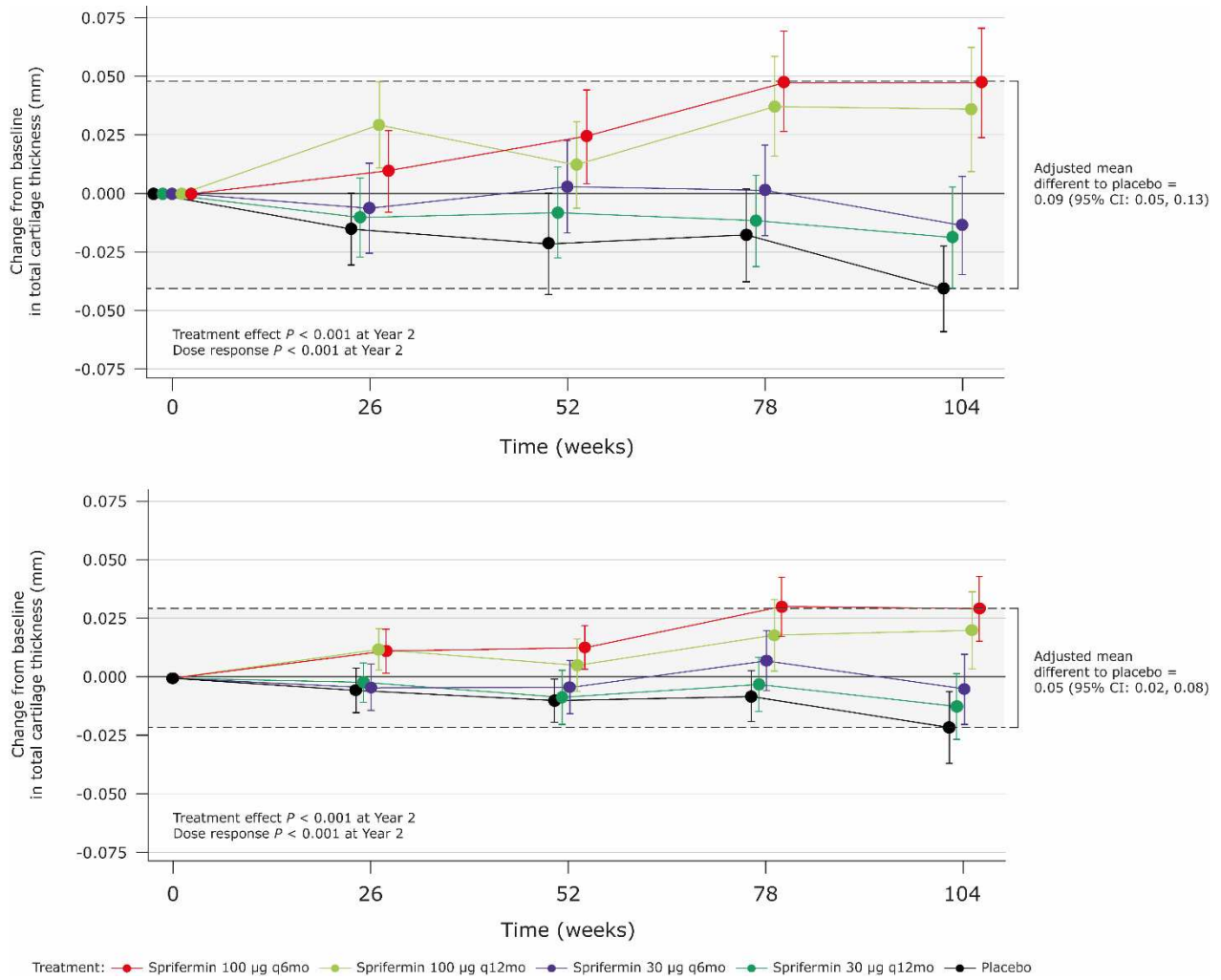
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218

219 **Figures**



220
221 **Fig. 1.** Cartilage thickness in the total femorotibial joint determined by automated segmentation (top)
222 and manual segmentation (bottom)

223
224 CI, confidence interval; q6mo, every 6 months; q12mo, every 12 months.

225
226
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228
229

230 **Table 1**

231 Change from baseline at Year 2 in cartilage thickness (mm) using automated segmentation and manual
 232 segmentation – mITT

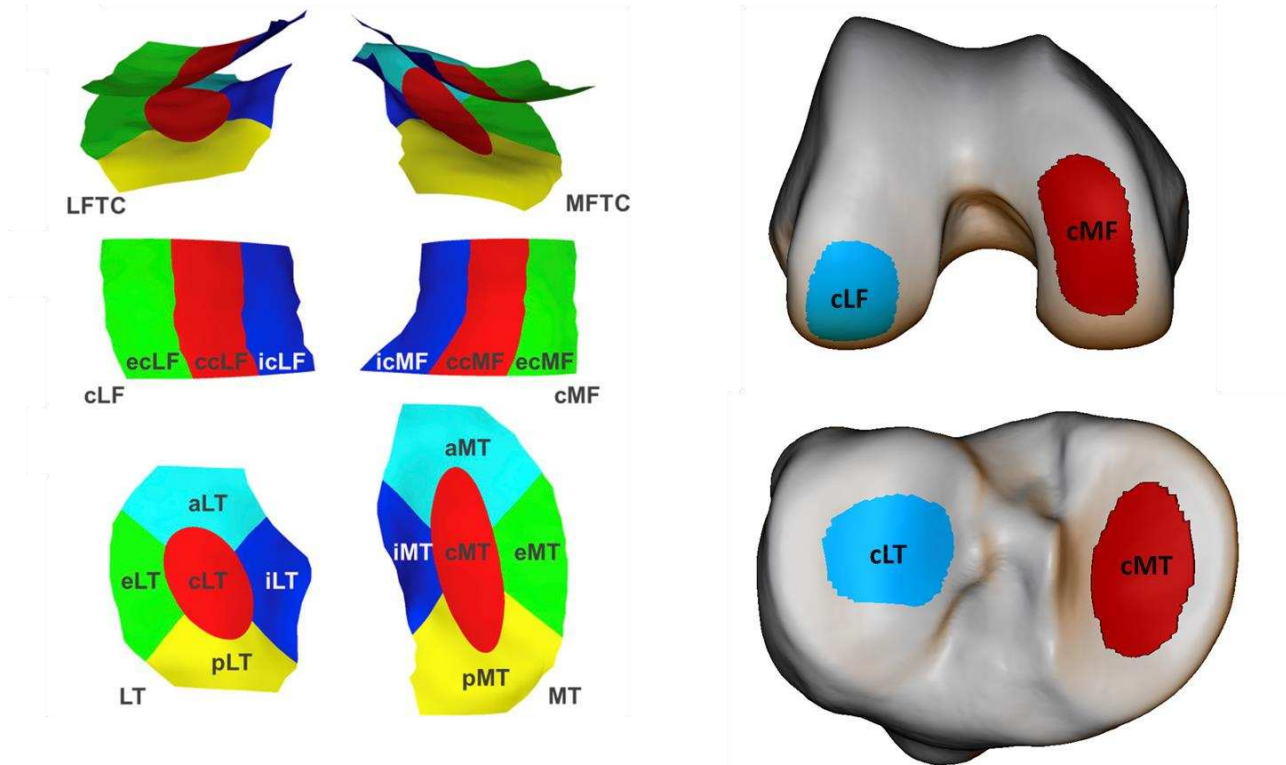
Region	Automated segmentation				Manual segmentation ⁷			
	Sprifermin 100 µg q6mo (n = 101)		Placebo (n = 96)		Sprifermin 100 µg q6mo (n = 101)		Placebo (n = 96)	
	Observed mean (SD) change from baseline, mm	Adjusted mean difference (95% CI) ^a	P-value	Observed mean (SD) change from baseline, mm	Adjusted mean difference (95% CI) ^a	P-value		
TFTJ	0.05 (0.11)	-0.04 (0.08)	0.09 (0.05, 0.13)	< 0.001	0.03 (0.07)	-0.02 (0.07)	0.05 (0.02, 0.08)	< 0.001
MFTC	0.03 (0.15)	-0.04 (0.14)	0.07 (0.01, 0.12)	0.011	0.02 (0.08)	-0.03 (0.12)	0.05 (0.01, 0.08)	0.003
LFTC	0.07 (0.12)	-0.04 (0.10)	0.11 (0.07, 0.15)	< 0.001	0.04 (0.06)	-0.01 (0.05)	0.05 (0.03, 0.08)	< 0.001

^aAdjusted using ANCOVA on change from baseline, including treatment group, timepoint, and (pooled) country as fixed factors, baseline value as covariate, and treatment by timepoint as interaction.
 CI, confidence interval; LFTC, lateral femorotibial compartment; MFTC, medial femorotibial compartment; SD, standard deviation; TFTJ, total femorotibial joint.

233

234

235 **Supplementary Figure 1**



236

237 **Supplementary Fig. 1.** Regions of interest defined for manual segmentation and automated
 238 segmentation (left panel) and additional regions for automated segmentation (right panel)

239

240 aLT, anterior lateral tibial; aMT, anterior medial tibial; cLF, central lateral femoral; ccLF, central subregion
 241 of the central lateral femoral; cLT, central lateral tibial; ccMF, central subregion of the central medial
 242 femoral; cMF, central medial femoral; cMT, central medial tibial; ecLF, external subregion of the central
 243 lateral femoral; ecMF, external subregion of the central medial femoral; eLT, external lateral tibial; eMT,
 244 external medial tibial; icLF, internal subregion of the central lateral femoral; icMF, internal subregion of
 245 the central medial femoral; iLT, internal lateral tibial; iMT, internal medial tibial; LFTC, lateral
 246 femorotibial compartment; LT, lateral tibial; MFTC, medial femorotibial compartment; MT, medial tibial;
 247 pLT, posterior lateral tibial; pMT, posterior medial tibial

248

249 Figure reused with permission from Wirth W et al. Osteoarthritis Cartilage 2017;25(8):1313-1323¹⁸

250

251 **Supplementary Figure 2**

252

253

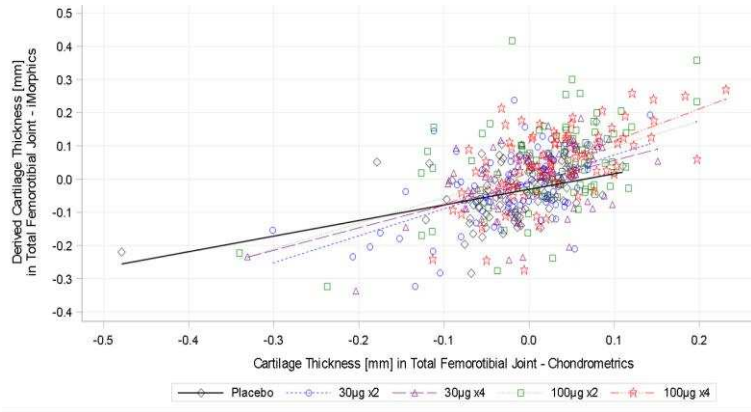
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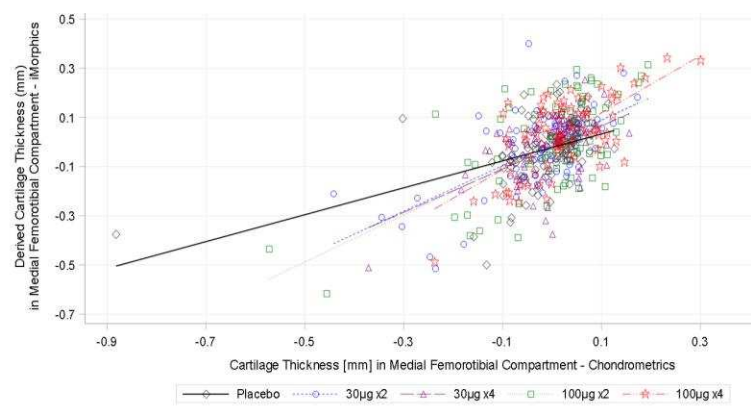
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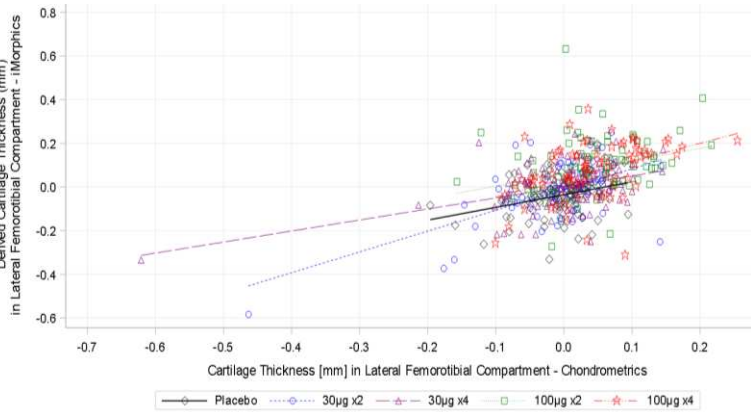
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274 **Supplementary Fig. 2.** Scatter plot and regression lines for change from baseline at Year 2 of cartilage

275 thickness in the total femorotibial joint (top), medial femorotibial joint (middle) and lateral femorotibial

276 joint (bottom): manual vs automated segmentation

277

278

279

280