

Title	Effects of switching weekly alendronate or risedronate to monthly minodronate in patients with rheumatoid arthritis: a 12-month prospective study
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1 **Original Article**

2 Effects of switching weekly alendronate or risedronate to monthly minodronate in
3 patients with rheumatoid arthritis: a twelve-month prospective study

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24 **Abstract**

25 *Purpose*

26 The aim of this prospective, observational study was to evaluate the effects of switching
27 weekly alendronate (ALN 35 mg) or risedronate (RIS 17.5 mg) to monthly minodronate
28 (MIN 50 mg) in patients with rheumatoid arthritis (RA).

29 *Methods*

30 Patient characteristics were as follows: n=172; 155 postmenopausal women; age 65.5
31 (44-87) years; T-score of lumbar spine (LS), -1.4; total hip (TH), -1.8; femoral neck
32 (FN), -2.1; dose and rate of oral prednisolone (2.3 mg/day), 69.1%; prior duration of
33 ALN or RIS, 46.6 months; were allocated, based on their preference, to either the (1)
34 continue group (n=88), (2) switch-from-ALN group (n=44), or (3) switch-from-RIS
35 group (n=40).

36 *Results*

37 After 12 months, increase in BMD was significantly greater in group 3 compared to
38 group 1: LS (4.1 vs 1.2%; $P < 0.001$), TH (1.9 vs -0.7%; $P < 0.01$), and FN (2.7 vs
39 -0.5%; $P < 0.05$); and in group 2 compared to group 1: LS (3.2 vs 1.2%; $P < 0.05$) and
40 TH (1.5 vs -0.7%; $P < 0.01$). The decrease in bone turnover markers was significantly
41 greater in group 3 compared to group 1: TRACP-5b (-37.3 vs 2.5%; $P < 0.001$), PINP

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2 42 (-24.7 vs -6.2%; $P < 0.05$), and ucOC (-39.2 vs 13.0%; $P < 0.05$); and in group 2
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5 43 compared to group 1: TRACP-5b (-12.5 vs 2.5%; $P < 0.05$) at 12 months.
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8 44 *Conclusions*

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11 45 Switching weekly ALN or RIS to monthly MIN in patients with RA may be an effective
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14 46 alternative treatment option of oral bisphosphonate treatment.
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19 20 21 48 **Keywords**

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24 49 Rheumatoid arthritis; osteoporosis; minodronate; alendronate; risedronate.
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29 30 31 51 **Mini Abstract**

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34 52 Switching weekly ALN or RIS to monthly MIN in patients with RA, of whom
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37 53 two-thirds were treated with low-dose PSL, significantly decreased bone turnover
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40 54 markers and increased BMD at 12 months, suggesting that monthly MIN may be an
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43 55 effective alternative treatment option of oral bisphosphonate treatment.
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48 49 50 57 **Introduction**

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53 58 **Increased risk of fractures in patients with rheumatoid arthritis (RA) compared to**
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56 59 **non-RA controls has been reported, with risk ratios (RR) varying from 2.0 to 3.0 at the**
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60 hip and 2.4 to 6.2 at the spine [1-3]. Pro-inflammatory cytokines, such as tumor necrosis
61 factor-alpha (TNF- α), interleukin (IL)-1, IL-6, and IL-17, are strongly involved in the
62 pathogenesis of RA, and also concerned with osteoclastogenesis and consequent bone
63 loss [4-7]. Indeed, high bone turnover and inflammation is associated with bone loss of
64 the femoral neck (FN) in postmenopausal RA patients [8]. Moreover, glucocorticoids
65 are often used to treat RA, which induce apoptosis of osteoblasts and osteocytes, and
66 result in increased fracture risk [9, 10]. Minodronate (MIN) is an oral
67 nitrogen-containing bisphosphonate (BP) developed in Japan which has a stronger
68 inhibitory effect on farnesyl pyrophosphate synthase in osteoclasts compared with
69 alendronate (ALN) or risedronate (RIS) [11]. It has been shown that switching daily or
70 weekly BP (mainly ALN and RIS) to monthly MIN increased bone mineral density
71 (BMD) of the lumbar spine (LS) and distal radius, and also decreased bone turnover
72 markers in patients with osteoporosis [12]. There are still considerable number of
73 patients who desire oral osteoporosis treatment, and we hypothesized that MIN can be a
74 convenient candidate of alternative oral BP treatment in patients with RA treated by
75 ALN and RIS, which may be more effective in decreasing bone turnover and increasing
76 BMD. The aim of this prospective study was to clarify the effect of switching weekly
77 ALN (35 mg) or RIS (17.5 mg) to monthly minodronate (50 mg) in patients with RA.

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79 **Materials and methods**

80 *Study design and subjects*

81 This twelve-month observational study was conducted based on a two-center,
82 prospective, open-label design. A total of 172 patients with RA who were treated with
83 oral weekly ALN or RIS in proportion to the Japanese guidelines for prevention and
84 treatment of osteoporosis 2011 [13] and the guidelines on the management and
85 treatment of glucocorticoid-induced osteoporosis of the Japanese Society for Bone and
86 Mineral Research 2004 [14], were enrolled in the study (Fig. 1). RA was diagnosed
87 based on the 1987 revised American College of Rheumatology (ACR) criteria [15].
88 C-reactive protein (CRP), matrix metalloproteinase-3 (MMP-3), and the Disease
89 Activity Score assessing 28 joints with CRP (DAS28-CRP) were evaluated as the
90 parameters reflecting inflammation as well as the disease activity of RA [16, 17].
91 Registered patients were asked their preference for a change to monthly oral BP
92 treatment and were allocated based on their preferences to either the “continue” group
93 (n=88), consisting of patients who wanted to continue their current therapies, or the
94 “switch-from-ALN” group (n=44) or “switch-from-RIS” group (n=40), consisting of
95 patients who were willing to switch over to MIN 50 mg from their current therapies.

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96 Other combined osteoporosis treatments, such as active vitamin D, vitamin K₂, and
97 **calcium** were continued during the study period. Patients' treatment persistence and
98 satisfaction levels with the therapies were assessed using a self-administered
99 questionnaire at 12 months (Table 1). **Patients were asked for their drug adherence every**
100 **time visiting outpatient clinic (every 1-3 months), and patients who didn't take their**
101 **medications more than twice of their interval (more than 2 weeks for weekly ALN or**
102 **RIS, and more than 2 months for monthly MIN) were considered as drop-out.**

103 This observational study was conducted in accordance with the ethical standards of the
104 Declaration of Helsinki and was approved by ethical review boards at the clinical center
105 (approval number 11273-2; Osaka University, Graduate School of Medicine). Written
106 informed consent was obtained from individual patients included in the study.

107
108 *BMD assessment*

109 Areal BMD in the LS (L2-L4), total hip (TH), and femoral neck (FN) were assessed by
110 dual-energy x-ray absorptiometry (Discovery A, Hologic, Inc., Waltham, MA, USA) at
111 baseline and after 6 and 12 months of treatment. Regions of severe scoliosis, vertebral
112 fracture, and operated sites were excluded from BMD measurements as previously
113 described [18].

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115 *Biochemical markers of bone turnover*

116 Bone turnover markers were measured in serum obtained from each patient at
117 approximately the same time in the morning after overnight fasting. The bone formation
118 marker, N-terminal type I procollagen propeptide (PINP); inter-assay coefficient of
119 variation (CV), 3.2%-5.2%, (Intact UniQ assay; Orion Diagnostica, Espoo, Finland),
120 and bone resorption marker, isoform 5b of tartrate-resistant acid phosphatase
121 (TRACP-5b); inter-assay CV, 5.0%-9.0%, (Immunodiagnostic Systems Ltd., Boldon,
122 UK) were measured by ELISA as previously described [19]. Levels of
123 undercarboxylated osteocalcin (ucOC) were measured by a solid-phase enzyme
124 immunoassay kit; inter-assay CV, 5.2%-8.3%, (Takara Bio, Shiga, Japan) with a
125 sensitivity of 0.25 ng/mL. **UcOC reflects not only vitamin K deficiency, but also total**
126 **bone turnover, as it is released from both osteoblasts and absorbed bone extracellular**
127 **matrix by osteoclast as previously described [20, 21].** Intact- parathyroid hormone
128 (PTH) was measured using a two-site immunoradiometric assay; inter-assay CV 8.4%,
129 (Nichols Institute Diagnostics, Valencia, USA).

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131 *Statistical analysis*

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132 The normal distributions of the data were examined by the Shapiro-Wilk test.
133 Differences between each study group were tested using analysis of variance for
134 normally distributed data and the nonparametric Kruskal-Wallis test was used for
135 non-normally distributed data. Changes in BMD and ranked bone turnover marker data
136 from baseline to specified time points within each study group were compared using the
137 nonparametric Wilcoxon signed-rank test. Results are expressed as the mean \pm standard
138 error. A *P* value < 0.05 indicated statistical significance. All tests were performed using
139 IBM SPSS Statistics version 22 software (IBM, Armonk, NY, USA).

140

141 **Results**

142 Baseline characteristics are shown in Table 2. Of the 172 study patients, 84 (48.8%)
143 were willing to switch to MIN 50 mg. No significant differences were observed in
144 baseline age, combined dose and prescription rate of active vitamin D or vitamin K₂ or
145 calcium or prednisolone (PSL), BMD, or disease activity of RA between the groups.

146 Duration of prior BP therapy at baseline was significantly longer in the
147 switch-from-ALN group (57.2 months) compared to the continue group (43.6 months; *P*
148 < 0.05) and the switch-from-RIS group (41.0 months; *P* < 0.05). Baseline serum
149 TRACP-5b levels in the switch-from-ALN group were significantly lower compared to

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150 the switch-from-RIS group (244.5 vs 309.8 mU/dL; $P < 0.05$). Eventually, 95.5%
151 (84/88) of patients in the continue group (2 patients were lost to follow up and 2
152 patients desired to change the medication) and 94.0% (79/84) of patients in the switch
153 group (3 patients were lost to follow up and 2 patients desired to change the medication)
154 completed the twelve-month trial (Fig. 1).

155

156 *Change in BMD*

157 BMD was monitored every 6 months (Fig. 2). Both the switch groups showed a
158 significant increase in LS and TH BMD from baseline to 6 and 12 months, while only
159 the switch-from-RIS group showed a significant increase in FN BMD from baseline to 6
160 and 12 months. Moreover, the switch-from-RIS group showed a significantly greater
161 increase compared to the continue group in the LS from 6 months (2.3 vs 0.6%; $P <$
162 0.05) to 12 months (4.1 vs 1.2%; $P < 0.001$), in the TH from 6 months (1.8 vs -0.5%; P
163 < 0.01) to 12 months (2.0 vs -0.7%; $P < 0.01$), and in the FN from 6 months (2.0 vs
164 -0.4%; $P < 0.05$) to 12 months (2.7 vs -0.5%; $P < 0.05$), respectively. On the other hand,
165 the switch-from-ALN group showed a significantly greater increase compared to the
166 continue group in LS BMD at 12 months (3.2 vs 1.2%; $P < 0.05$) and in the TH from 6
167 months (1.2 vs -0.5%; $P < 0.01$) to 12 months (1.5 vs -0.7%; $P < 0.01$). The

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168 switch-from-RIS group showed a significantly greater increase compared to the
169 switch-from-ALN group in the FN from 6 months (2.1 vs -0.3%; $P < 0.05$) to 12 months
170 (2.7 vs -0.6%; $P < 0.05$).

171

172 *Bone turnover markers*

173 Percent changes in bone turnover markers from baseline are shown in Fig. 3. The
174 switch-from-RIS group showed a significantly greater decrease compared to the
175 continue group in TRACP-5b levels from 6 months (-35.8 vs 1.3%; $P < 0.001$) to 12
176 months (-37.3 vs 2.5%; $P < 0.001$), in PINP levels from 6 months (-22.2 vs -3.3%; $P <$
177 0.05) to 12 months (-24.7 vs -6.2%; $P < 0.05$), and in ucOC levels from 6 months (-22.2
178 vs 12.4%; $P < 0.05$) to 12 months (-39.2 vs 13.0%; $P < 0.05$). On the other hand, the
179 switch-from-ALN group showed a significantly greater decrease compared to the
180 continue group only in TRACP-5b levels from 6 months (-14.6 vs 1.3%; $P < 0.01$) to 12
181 months (-12.5 vs 2.5%; $P < 0.05$). The switch-from-RIS group showed a significantly
182 greater decrease than the minimum significant change of serum TRACP-5b, PINP, and
183 ucOC levels, while the switch-from-ALN group showed only in the serum TRACP-5b
184 at 12 months. There were no greater changes than the minimum significant change of
185 serum TRACP-5b, PINP, and ucOC levels in the continue group. **The absolute value of**

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186 bone turnover markers are shown in Fig. 4. The average value of TRACP-5b, PINP, and
187 ucOC in all the groups were all within the reference value.

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189 *Rate of fragility fracture*

190 During the twelve-month period, the continue group patients experienced 3 vertebral
191 and 1 non-vertebral clinical fragility fractures (4.5%). The switch-from-ALN group
192 experienced 1 vertebral and 1 non-vertebral clinical fragility fractures (4.5%), and no
193 clinical fragility fracture was observed in the switch-from-RIS group (0.0%). No
194 statistically significant difference in the total clinical fragility fracture rate was observed
195 between the groups.

196

197 *Patient preference after switching to MIN 50 mg*

198 Patient preference after switching to monthly MIN 50 mg is shown in Fig. 5. The
199 questionnaire revealed that 80.8% of patients were satisfied with the switch to monthly
200 therapy and 88.7% preferred to continue the monthly treatment. The main reasons for
201 desiring continuation of monthly dosing was both the decreased frequency (69.8%) and
202 less worry about forgetting doses (47.2%), thus a perception of less overall burden.

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204 **Discussion**

205 In this study, we have demonstrated for the first time that in patients with RA, of whom
206 two-thirds were treated with low-dose PSL (< 10 mg/day), switching from weekly ALN
207 or RIS to monthly MIN was effective in increasing BMD and decreasing bone turnover
208 markers at 12 months. In addition, no previous studies have demonstrated the difference
209 of the effects of switching, by the difference of prior BP therapies.

210 In nitrogen-containing BP treatment, mineral binding affinities may influence their
211 distribution within bone and the period till anti-fracture effects are shown, and
212 inhibition of farnesyl diphosphate synthase (FPPS) may affect their anti-resorptive
213 effects by inducing apoptosis of osteoclasts [22].

214 It has been shown that ALN possesses a stronger binding affinity to hydroxyapatite
215 compared to RIS, while RIS possesses a stronger FPPS inhibition compared to ALN
216 [22]. Consequently, weekly ALN (70 mg) showed a greater increase in BMD and
217 decrease in bone turnover markers compared to weekly RIS (35 mg) in patients with
218 postmenopausal osteoporosis [23], while RIS showed lower rates of hip and
219 non-vertebral fractures than ALN during the first year of therapy [24].

220 Previous reports have demonstrated that MIN showed stronger FPPS inhibition [11] and
221 a weaker binding affinity to hydroxyapatite compared to ALN and RIS [25], which

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2 222 suggests that MIN inhibits bone resorption more strongly and is more quickly
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5 223 distributed within the bone compared to ALN and RIS. Indeed, MIN suppressed bone
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8 224 remodeling of cancellous and cortical bone more strongly than ALN in vitro [26], as
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11 225 well as in ovariectomized cynomolgus monkeys in vivo [27]. In the previous human
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14 226 study, switching ALN or RIS to monthly MIN for 6 months increased BMD +1.1% in
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18 227 LS, and the reduction rate of serum TRACP-5b was approximately 35% in the
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21 228 switching from RIS group at 6 months [12], which were consistent with our study.
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24 229 Finally, glucocorticoids have been shown to induce apoptosis of osteocytes, and BPs
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27 230 inhibit osteocyte apoptosis in vitro [28] as well as in glucocorticoid-treated animals [29].
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30 231 A systematic review and meta-analysis revealed that BPs can preserve bone mass and
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33 232 reduce the incidence of vertebral fractures in patients with rheumatic disease, mainly for
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36 233 those who are being treated with glucocorticoids [30], and both ALN and RIS strongly
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39 234 decreased the fracture risk associated with glucocorticoid-induced osteoporosis (GIO)
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42 235 [31, 32]. In this study, monthly MIN 50 mg resulted in a greater BMD increase and
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45 236 bone turnover decrease when patients were switched from ALN or RIS, which suggests
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48 237 its effectiveness not only in primary osteoporosis, but also in GIO.
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51 238 There are several limitations to this study. Due to the small number of subjects, fracture
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54 239 risk comparisons should be assessed in a randomized, larger cohort. As most of the
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240 patients showed remission or low disease activity in this study, the effects of switching
241 on high disease activity patients should be assessed in further study. Although most
242 patients were postmenopausal, some male patients were included in this study.

243 Concerning medication, the dose of ALN and RIS allowed in Japan is the half of
244 Caucasians, and the duration of prior BP therapy was significantly longer in switch-to
245 ALN group compared to other groups. In addition, only a small number of patients were
246 combined with calcium formulation, and total calcium intake couldn't be monitored.

247 In conclusion, switching weekly ALN or RIS to monthly MIN in patients with RA, of
248 whom two-thirds were treated with low-dose PSL, significantly decreased bone
249 turnover markers and increased BMD at 12 months, suggesting that monthly MIN may
250 be an effective alternative treatment option of oral BP treatment.

251

252 **Acknowledgments**

253 The authors thank Dr. Kenrin Shi for his excellent cooperation in conducting the study.

254

255 **Authors' roles**

256 Study design: KE, MH, JH, and HY. Study conduct: KE, TN, MH, and SK. Data
257 collection: KE, TN, SK, and MY. Data analysis: KE, TN, and MH. Data interpretation:

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258 KE and MH. Drafting the manuscript: KE and MH. Approving final version of the
259 manuscript: KE, TN, MH, JH, SK, and HY. KE takes responsibility for the integrity of
260 the data analysis.

261

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266 Kaneshiro, Masao Yukioka, and Hideki Yoshikawa declare that they have no conflict of
267 interest.

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371 **Figure legends**

372

373 Figure 1. Study design and schedule. Patients were asked for their willingness to switch
374 to monthly MIN 50 mg. Bone mineral density and bone turnover markers were
375 evaluated every 6 months in all the patients. The switch group patients were asked to
376 complete a patient preference questionnaire at 12 months.

377

378 Figure 2. Mean \pm standard error (SE) change from baseline in bone mineral density
379 (BMD) at the lumbar spine (panel a), total hip (panel b), and femoral neck (panel c). **P*
380 < 0.05 , ***P* < 0.01 , ****P* < 0.001 change from baseline within each treatment group. #*P*
381 < 0.05 , ##*P* < 0.01 , ###*P* < 0.001 continue group versus switch-from-RIS group. †*P* $<$
382 0.05 , switch-from-ALN group versus switch-from-RIS group.

383

384 Figure 3. Mean \pm standard error (SE) change from baseline in serum concentration of
385 bone turnover markers TRAP-5b (panel a), PINP (panel b), and ucOC (panel c).
386 TRAP-5b, isoform 5b of tartrate-resistant acid phosphatase; PINP, type I collagen
387 N-terminal propeptide; ucOC, undercarboxylated osteocalcin; #*P* < 0.05 , ##*P* < 0.01 ,
388 ###*P* < 0.001 continue group versus each switch group. **P* < 0.05 , ***P* < 0.01

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389 switch-from-ALN group versus switch-from-RIS group.

390

391 **Figure 4. Mean ± standard error (SE) absolute value of bone turnover markers TRAP-5b**
392 **(panel a), PINP (panel b), and ucOC (panel c). TRAP-5b, isoform 5b of tartrate-resistant**
393 **acid phosphatase; PINP, type I collagen N-terminal propeptide; ucOC,**
394 **undercarboxylated osteocalcin; #*P* < 0.05, ##*P* < 0.01, ###*P* < 0.001 continue group**
395 **versus each switch group. **P* < 0.05 switch-from-ALN group versus switch-from-RIS**
396 **group.**

397

398 **Figure 5. Patient satisfaction, preference, and reasons for preference after switching**
399 **weekly ALN or RIS to monthly MIN 50 mg treatment at 12 months.**

400

401

1 Table 1. Patient preference questionnaire

-
1. Rate your satisfaction with the current once-monthly dosing schedule ^{a)}
1 2 3 4 5
1-Low satisfaction 5-High satisfaction
 2. Which dosing schedule do you prefer?
a. Once weekly b. Once monthly c. No preference
 3. If you prefer once-monthly dosing schedule, check all the statements you agree with ^{b)}
 - a. This dosing schedule impose less burden of frequency
 - b. This dosing schedule has less worry to forget
 - c. I feel this dosing schedule is more effective
 - d. I expect less side effects with this dosing schedule
 - e. Others
-

2 a) Answer 4 and 5 are evaluated as satisfied, 3 as no preference, and 1 and 2 as not satisfied.

3 b) Multiple answers allowed.

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15 Table 2. Baseline clinical characteristics

Variable	Continue (n=88)	Switch-from-ALN (n=44)	Switch-from-RIS (n=40)
Age, (mean \pm SE years)	64.9 \pm 0.9	64.9 \pm 1.6	67.3 \pm 1.6
Gender, Females (%)	81/88 (92.0%)	40/44 (90.9%)	38/40 (95.0%)
Postmenopausal, n/N (%)	80/88 (90.9%)	38/44 (86.4%)	37/40 (92.5%)
Body mass index (kg/m ²)	21.9 \pm 0.4	21.2 \pm 0.6	22.2 \pm 0.6
Prior BP, ALN n/N(%)	58/88 (65.9%)		
Duration of prior BP therapy (months)	43.6 \pm 2.1	57.2 \pm 4.6*	41.0 \pm 5.5 [†]
Combined vitamin D, n/N(%)	46/88 (52.3%)	26/44 (59.1%)	25/40 (62.5%)
Combined vitamin K ₂ , n/N(%)	21/88 (23.9%)	12/44 (27.3%)	10/40 (25.0%)
Combined calcium, n/N(%)	5/88 (5.7%)	3/44 (6.8%)	3/40 (7.5%)
Prior vertebral fracture(s), n/N(%)	25/88 (28.4%)	9/44 (20.5%)	8/40 (20.0%)
Prior non-vertebral fracture(s), n/N(%)	22/88 (25.0%)	10/44 (22.7%)	7/40 (17.5%)
Bone mineral density (BMD)			
Lumbar spine BMD (g/cm ²)	0.856 \pm 0.017	0.861 \pm 0.028	0.858 \pm 0.019
Lumbar spine BMD (T-score)	-1.4 \pm 0.1	-1.3 \pm 0.2	-1.4 \pm 0.2
Femoral neck BMD (g/cm ²)	0.584 \pm 0.027	0.546 \pm 0.015	0.584 \pm 0.016
Femoral neck BMD (T-score)	-2.1 \pm 0.1	-2.3 \pm 0.1	-2.0 \pm 0.1
Total hip BMD (g/cm ²)	0.698 \pm 0.028	0.658 \pm 0.017	0.677 \pm 0.018
Total hip BMD (T-score)	-1.8 \pm 0.1	-1.9 \pm 0.1	-1.8 \pm 0.2
T-score < -2.5, n/N(%)	45/88 (51.1%)	22/44 (50.0%)	16/40 (40.0%)
PINP (μ g/l)	34.2 \pm 2.7	29.7 \pm 2.7	34.5 \pm 2.5
TRACP-5b (mU/dl)	258.1 \pm 11.2	244.5 \pm 17.6	309.8 \pm 22.7 [†]
ucOC (ng/ml)	2.7 \pm 0.3	3.6 \pm 0.9	3.7 \pm 0.6
Intact-PTH (pg/ml)	48.9 \pm 2.4	51.5 \pm 3.7	45.6 \pm 2.6
eGFR (ml/min/1.73m ²)	77.2 \pm 2.5	73.6 \pm 3.5	74.9 \pm 3.3
Duration of disease (years)	17.6 \pm 1.0	18.3 \pm 1.6	15.1 \pm 1.5
RF positivity, n/N (%)	73/88 (83.0%)	41/44(93.2%)	35/40(87.5%)
ACPA positivity, n/N (%)	75/88 (85.2%)	40/44(90.9%)	34/40(85.0%)
CRP (mg/dl)	0.7 \pm 0.1	0.6 \pm 0.1	0.5 \pm 0.1
MMP-3 (ng/ml)	158.4 \pm 16.2	118.1 \pm 16.4	118.2 \pm 30.1

DAS28-CRP	2.6±0.1	2.5±0.1	2.4±0.1
Remission (< 2.3), n/N (%)	41/88 (46.6%)	22/44 (50.0%)	22/40 (55.0%)
Low disease activity (< 2.7), n/N (%)	16/88 (18.2%)	11/44 (25.0%)	7/40 (17.5%)
Moderate disease activity (2.7 -4.1), n/N (%)	26/88 (29.5%)	10/44 (22.7%)	9/40 (22.5%)
High disease activity (> 4.1), n/N (%)	5/88 (5.7%)	1/44 (2.3%)	2/40 (5.0%)
MHAQ	0.5±0.1	0.4±0.1	0.6±0.1
Prednisolone dose (mg/day)	2.5±0.3	2.2±0.3	1.7±0.4
Prednisolone usage, n/N(%)	62/88 (70.5%)	32/44 (72.7%)	25/40 (62.5%)
MTX dose (mg/week)	5.0±0.4	5.6±0.6	4.7±0.6
MTX usage, n/N (%)	63/88 (71.6%)	35/44(79.5%)	28/40(70.0%)
Biologics usage, n/N (%)	20/88 (25.7%)	8/44(18.2%)	9/40(22.5%)

16 Mean ± Standard Error (SE), unless otherwise noted.

17 n/N (%) = number of patients with measurements / total number of patients (%)

18 ALN, Alendronate; RIS, Risedronate; BP, Bisphosphonate; PINP, Type I collagen N-terminal propeptide;

19 TRAP-5b, Isoform 5b of tartrate-resistant acid phosphatase; ucOC, Undercarboxylated osteocalcin; PTH,

20 parathyroid hormone; eGFR, Estimated glomerular filtration rate; RF, Rheumatoid factor; ACPA, Anti-

21 cyclic citrullinated peptide antibody; CRP, C-reactive protein; MMP-3, Matrix metalloproteinase-3;

22 DAS28-CRP, Disease activity score assessing 28 joints with CRP; MHAQ, Modified Health Assessment

23 Questionnaire; MTX, Methotrexate.

24 Differences between the groups were determined by ANOVA or chi-square test. *P<0.05 vs Continue

25 group. **P<0.01 vs Continue group. †P<0.05 vs Switch-from-ALN group. ††P<0.01 vs

26 Switch-from-ALN group.

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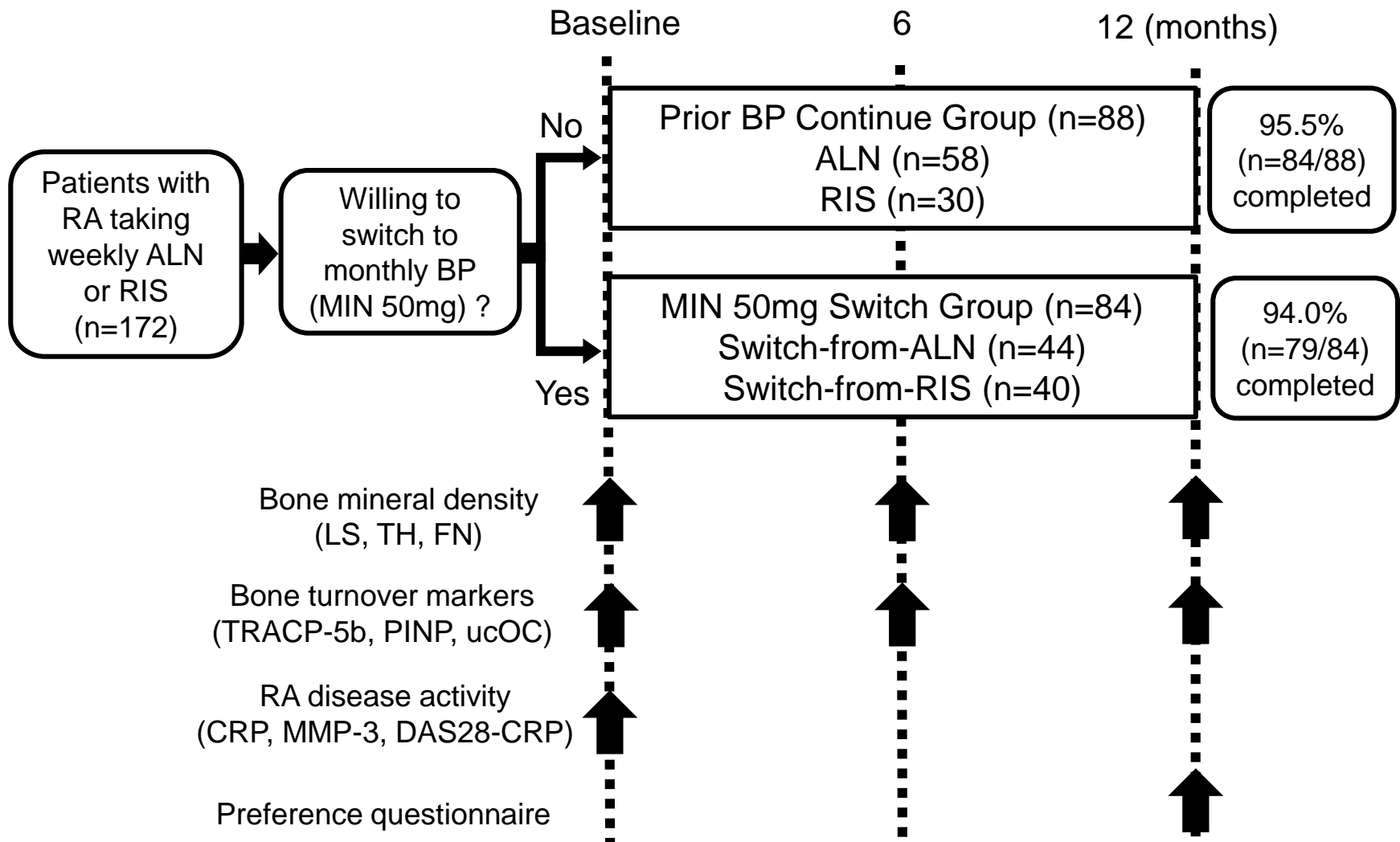
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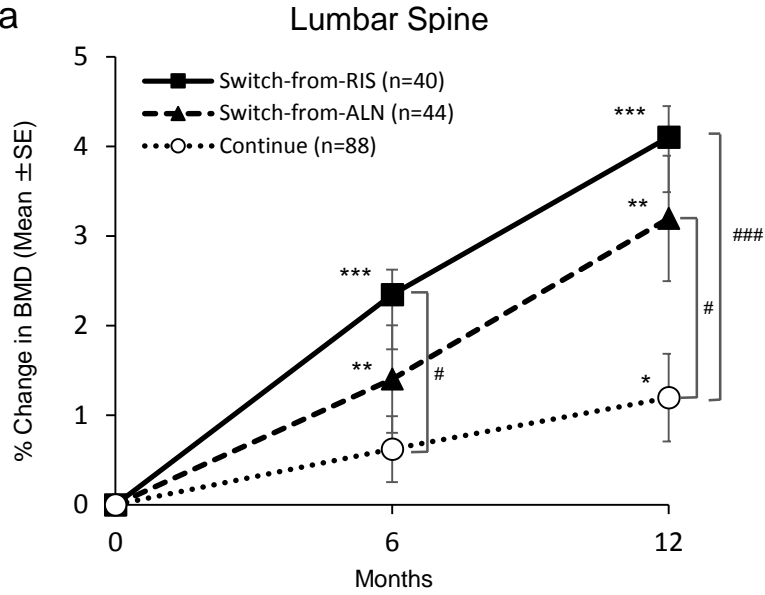
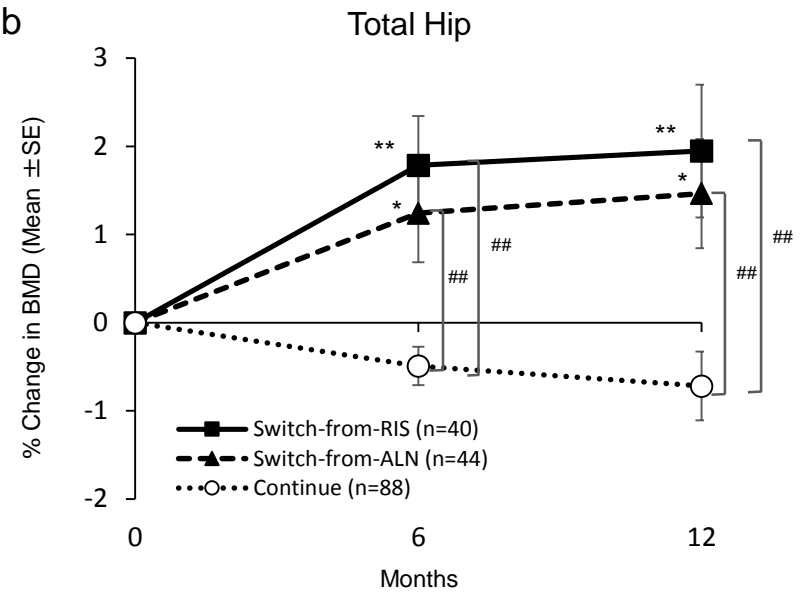
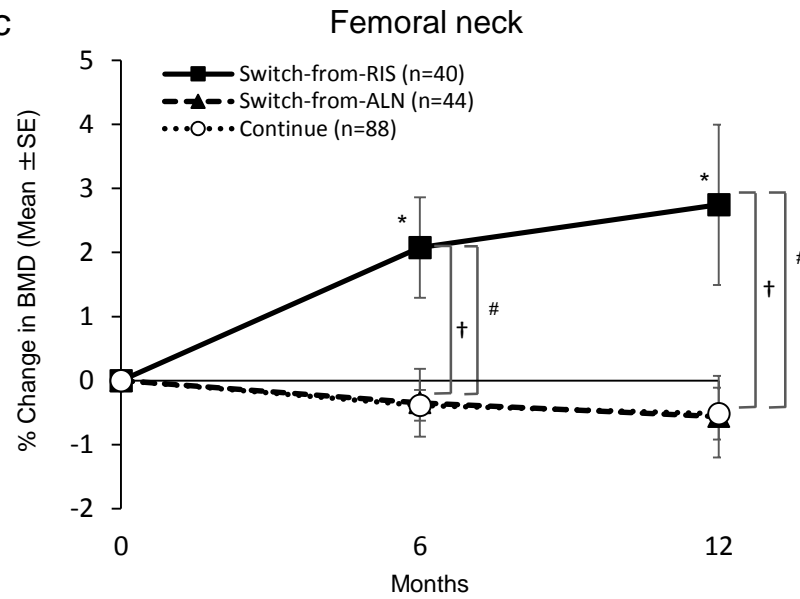
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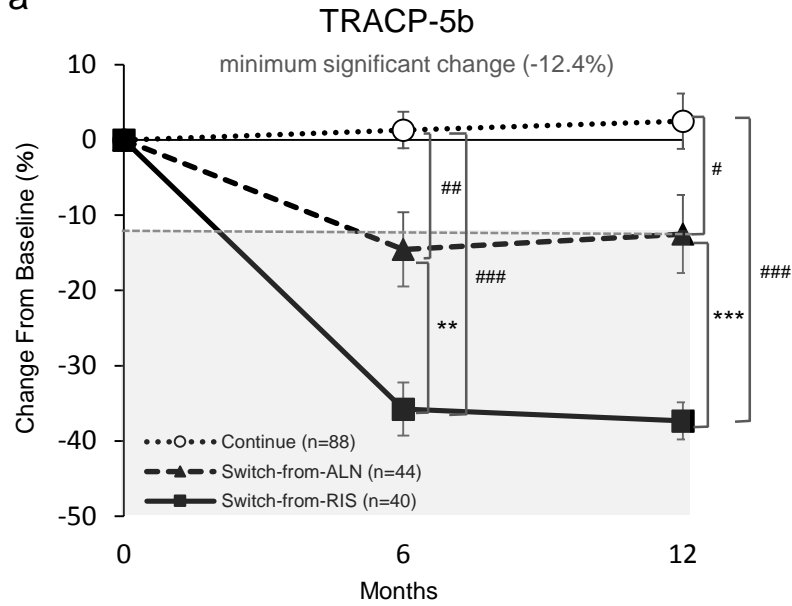
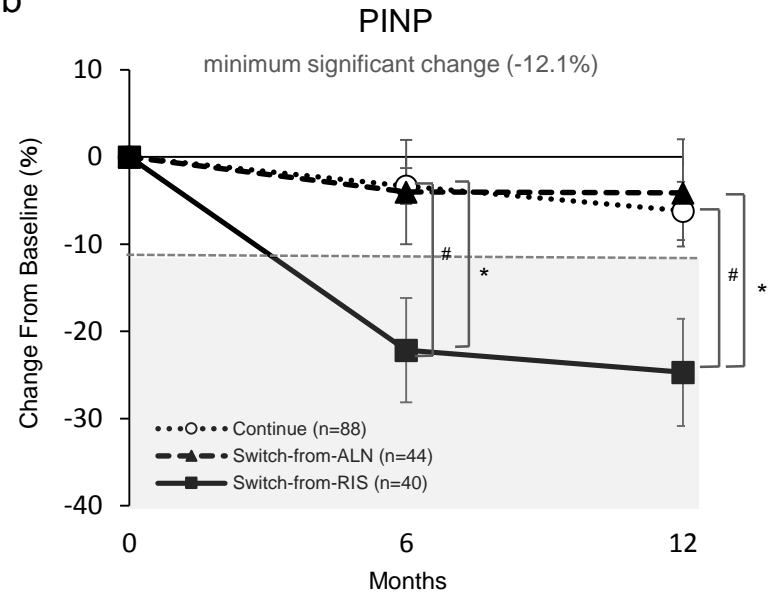
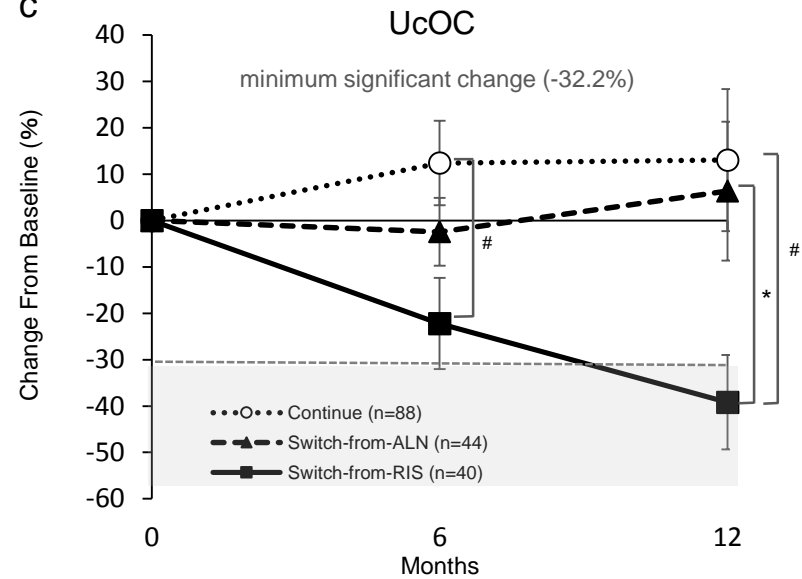
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Figure



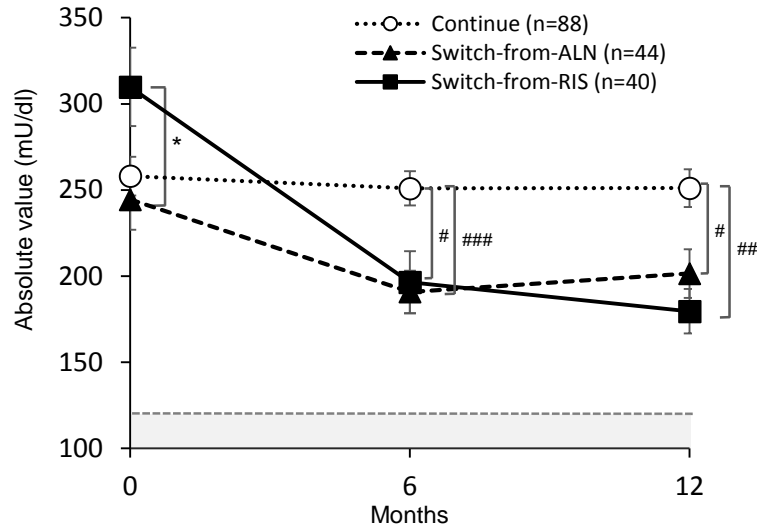
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TRACP-5b

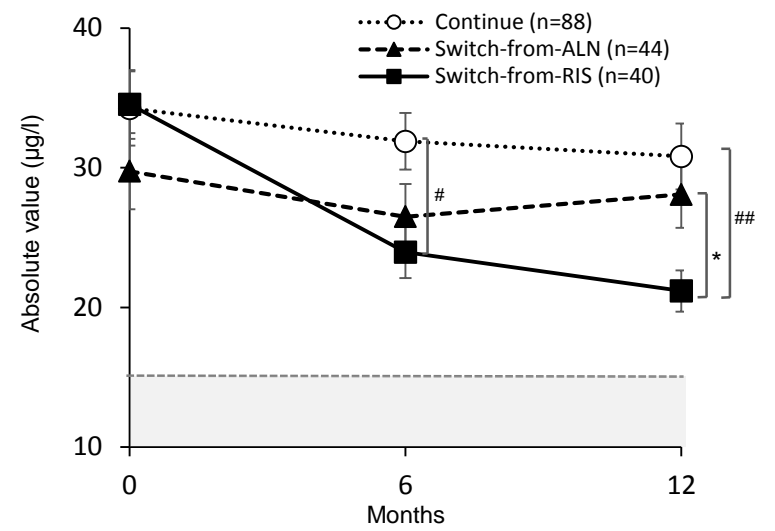
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PINP

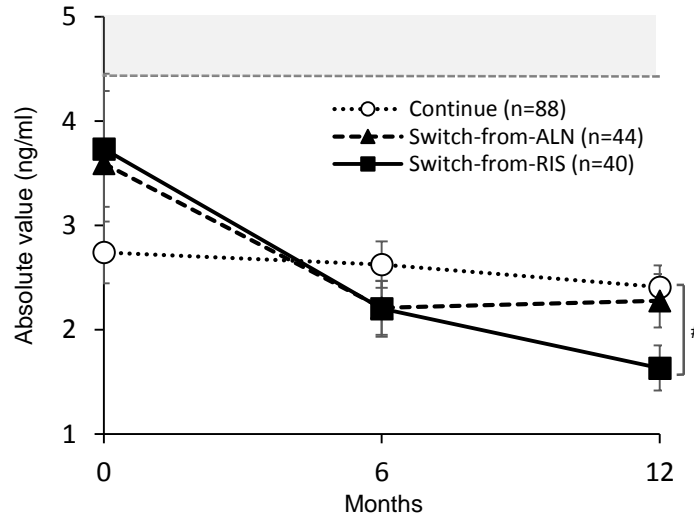
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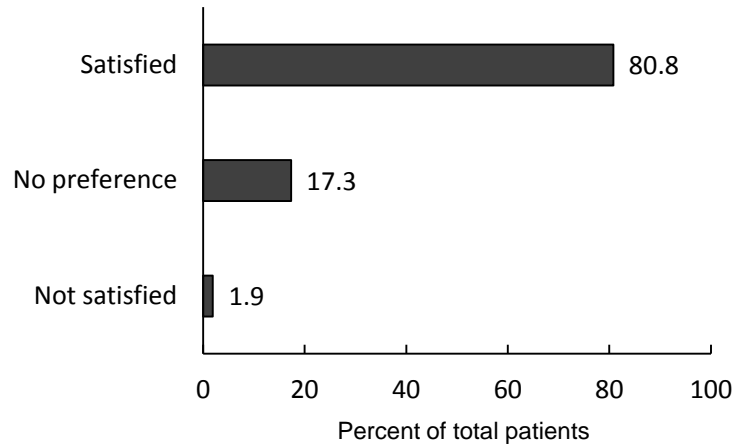
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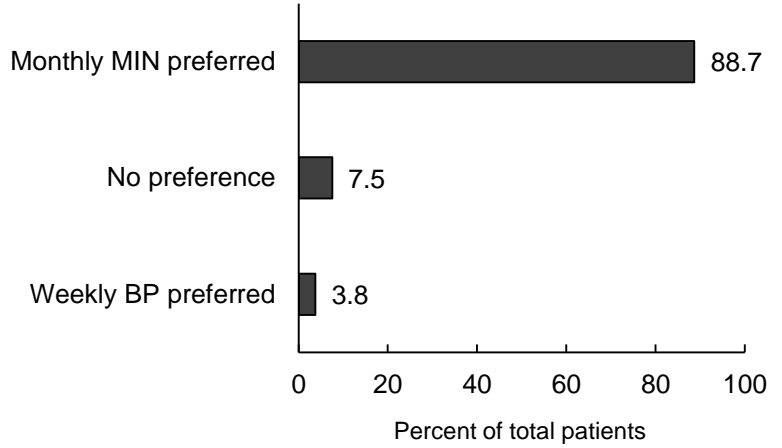
reference range: <4.5 ng/ml



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