WOMAC Meaningful Within-Patient Change: Results from Three Studies of Tanezumab in Patients with Moderate-To-Severe Osteoarthritis of the Hip or Knee

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**Running head:** WOMAC Within Patient Change.
ABSTRACT

Objective: To define meaningful within-patient change (MWPC) in the Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC).

Methods: Data were analyzed separately from 3 phase 3 clinical trials (ClinicalTrials.gov: NCT02697773, NCT02709486, NCT02528188) of tanezumab, a novel treatment intended for the relief of signs and symptoms of moderate-to-severe osteoarthritis, administered subcutaneously every 8 weeks. Patients with moderate-to-severe osteoarthritis of the hip or knee completed the WOMAC and Patient Global Assessment of Osteoarthritis (PGA-OA) at regular time points. A repeated measures longitudinal model with change in WOMAC Pain, Physical Function or Stiffness domain score as the outcome and change in PGA-OA as the anchor was used to establish MWPC for WOMAC domains.

Results: In the 3 studies there were 688, 844, and 2948 subjects available for analyses, respectively. Analysis showed that a linear relationship between changes in WOMAC domains and changes in PGA-OA was supported and justified. Moreover, the relationships between these changes were very similar for 2 trials and close for the third. The estimated MWPC for the 3 WOMAC domains were from 0.84–1.16 (0–10 numerical rating scale) and from 12.50–16.23%, depending on study and domain, that corresponded to a 1-category change on PGA-OA. For a 2-category change those values were from 1.68–2.31 and from 25.01–32.46%, respectively.

Conclusion: These results establish MWPCs for WOMAC domains, at the individual patient level, for patients with moderate-to-severe osteoarthritis of the hip or knee.
ClinicalTrials.gov identifiers: NCT02697773, NCT02709486 and NCT NCT02528188
INTRODUCTION

Clinical trial outcomes are typically determined by statistically significant mean differences between treatment and control groups; however, these “between-group” analyses do not indicate whether individual patients have experienced clinically meaningful benefits. To aid interpretation of the clinical meaningfulness of responses to treatment, it is important to determine thresholds for outcome measures. Such thresholds make it possible to assess group differences in the percentages of participants who have meaningful improvements after starting treatment.

Over thirty years ago, the minimal clinically important difference (MCID) was defined as “the smallest difference in score in the domain of interest which patients perceive as beneficial and would mandate, in the absence of troublesome side effects and excessive costs, a change in the patient’s management” (1). Although this definition of MCID refers to changes that patients perceive as meaningful for themselves, typically established using a global anchor rating of disease status or change, such MCIDs have been used in the interpretation of group differences observed in clinical trials (2,3). However, there are typically different considerations involved when interpreting what patients consider important improvements versus what should be considered an important group difference in a clinical trial.

The distinction between these two different concepts of clinical importance has been recognized for many years across a variety of therapeutic areas. It has typically been concluded that the clinical importance of group differences should not be based solely on the magnitude of the within-patient improvements that patients (or clinicians) consider important but should rather be based on the broader context of the disease and its available treatments along with a careful evaluation of risk...
versus benefit by relevant stakeholders(4,5). Indeed, in a landmark article, Guyatt and colleagues emphasized that “Clinicians and investigators tend to assume that if the mean difference between a treatment and a control is appreciably less than the smallest change that is important, then the treatment has a trivial effect. This may not be so. Let us assume that a randomized clinical trial shows a mean difference of 0.25 in a questionnaire in which the minimal important difference is 0.5. It might be concluded that the difference is unimportant and that the result does not support giving the treatment. This interpretation assumes that every patient treated scored 0.25 better than they would have done had they received the control and ignores the possibility that treatment might have a heterogeneous effect”(6).

Because of these considerations, use of the term MCID can be confusing when the definition is not very clearly specified and the difference between individual improvements and group differences is not clarified. Given this potential for misunderstanding, a recent US Food and Drug Administration (FDA) guidance on patient-focused drug development emphasized the difference between group- and individual-level changes, and highlighted the importance of defining what constitutes meaningful within-patient change (MWPC) in regulatory submissions(7). Just as thresholds for meaningful individual improvements should not necessarily be used to evaluate group differences, the FDA emphasized that “A treatment effect is different than a meaningful within-patient change. The terms minimally clinically important difference (MCID) and minimum important difference (MID) do not define meaningful within-patient change if derived from group level data and therefore should be avoided”(7).

The FDA guidance recommended the use of methods that utilize an external anchor measure to define the criteria for MWPC of an outcome measure at the
individual level(7). Anchor measures should be plainly understood and easier to interpret than the outcome measure. Recommended anchors include current-state global impression of severity scales. These scales may be preferable, as they avoid the recall bias associated with global impression of change scales and can also be used to assess change from baseline(7,8). Given a large range of thresholds reported in the literature, it is also important to evaluate the responder definition of an outcome for the target population of an intervention.

The Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC)\textsuperscript{1} is a disease-specific measure of osteoarthritis (OA) symptoms (pain and stiffness) and functional impairment that is regularly used in clinical trials. In order to facilitate the interpretation of OA clinical trial end points, it is important to determine thresholds for WOMAC that would indicate MWPC.

Previous attempts to define MWPC through meta-analysis of clinical trials in heterogeneous chronic pain conditions identified thresholds of a 10–20% reduction in pain as minimally important, ≥ 30% reductions as moderately important, and ≥ 50% reductions as substantial(4,9). A systematic literature review examined anchor-based clinically meaningful changes in WOMAC scores reported for patients with OA(10). Values for clinically important differences in WOMAC scores were between 0.3–1.3 for patients with OA of the hip and knee, between 0.5–3.4 for those with OA of the knee, and between 0.3–3.6 for those with OA of the hip, depending on the domain assessed (original standardized 0–100 scale converted to 0–10 scale).

The Patient Global Assessment of OA (PGA-OA) is a current-state global impression of disease severity scale regularly used in OA clinical trials in

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\textsuperscript{1} © 1996 Nicholas Bellamy. WOMAC\textsuperscript{®} is a registered trademark of Nicholas Bellamy (CDN, EU, USA).
combination with WOMAC(11,12). Given the large range of values reported for clinically meaningful changes in WOMAC, our aim was to estimate MWPC in WOMAC by examining the relationship between change in the outcomes of WOMAC domains and change in the anchor of PGA-OA using data from 3 trials that included patients with moderate-to-severe OA of the hip or knee.
MATERIALS AND METHODS

Statement of ethics and consent

The protocol for each clinical trial (ClinicalTrials.gov: NCT02697773, NCT02709486 and NCT02528188) was approved by an institutional review board or independent ethics committee at each participating investigational center. The studies were conducted in compliance with the ethical principles of the Declaration of Helsinki and Good Clinical Practice Guidelines. All patients provided written informed consent before entering the studies.

Patients

Data were analyzed separately from 3 phase 3 clinical trials of tanezumab, a novel treatment intended for the management of OA pain in adult patients for whom the use of other analgesics is ineffective or not appropriate. Study 1 (ClinicalTrials.gov: NCT02697773) was a 16-week randomized, double-blind, placebo-controlled trial(13). Patients in study 1 received subcutaneous (SC) placebo or tanezumab 2.5 mg at baseline and Week 8, or tanezumab 2.5 mg at baseline and 5 mg at Week 8. Study 2 (NCT02709486) was a 24-week randomized, double-blind, placebo-controlled trial(14). Patients in study 2 received SC placebo or tanezumab 2.5 mg or 5 mg at baseline and Weeks 8 and 16. Study 3 (NCT02528188) was a 56-week randomized, double-blind, double-dummy, active-controlled trial(15). Patients in study 3 received oral nonsteroidal anti-inflammatory drugs twice daily or SC tanezumab 2.5 mg or 5 mg every 8 weeks, with oral and SC study medication controls.

For all studies, patients aged ≥ 18 years with a body mass index of ≤ 39 kg/m² and a diagnosis of OA (Kellgren-Lawrence grade ≥ 2) of the hip or knee were eligible.
to enroll. Patients were required to have WOMAC Pain scores ≥ 5 in the index joint at baseline, defined as the most painful joint at baseline with a qualifying WOMAC Pain score and Kellgren-Lawrence grade, as confirmed by the Central Reader’s assessment. In all studies, patients were also required to have a WOMAC Physical Function score ≥ 5 in the index joint and a PGA-OA of “fair,” “poor,” or “very poor” at baseline.

**Measures**

WOMAC Index Version 3.1 numeric rating scale (NRS) consisted of 24 items assessed using a 0–10 NRS (0 = no pain/stiffness/difficulty to 10 = extreme pain/stiffness/difficulty), with higher scores indicating worse outcomes(16). The WOMAC Pain (5 items), Stiffness (2 items) and Physical Function (17 items) subscales measured the pain, stiffness or difficulty experienced in performing common tasks during a 48 hour recall period. Scores for total WOMAC and each subdomain were calculated from the average score of the component questions.

PGA-OA was a single question, “Considering all the ways your osteoarthritis in your hip/knee affects you, how are you doing today?” PGA-OA was measured on a 5-point Likert scale, with higher scores indicating worse symptoms (1 = “very good” [asymptomatic and no limitation of normal activities] to 5 = “very poor” [very severe symptoms that are intolerable and inability to carry out all normal activities]) (Supplementary Table 1). Patients were only asked to complete WOMAC and PGA-OA for their index hip or index knee, depending on the initial assessment for each subject.

WOMAC and PGA-OA were completed electronically on tablets at site visits. In study 1, all WOMAC subscales were completed at screening, baseline, and
Weeks 2, 4, 8, 12, 16, and 24; PGA-OA was completed at baseline and Weeks 2, 4, 8, 12, 16, and 24. In study 2, WOMAC Pain was completed at screening; all WOMAC subscales and PGA-OA were completed at baseline and Weeks 2, 4, 8, 12, 16, 24, and 32. In study 3, all WOMAC subscales were completed at screening, baseline, and Weeks 2, 4, 8, 16, 24, 32, 40, 48, 56, and 64; PGA-OA was completed at baseline and Weeks 2, 4, 8, 16, 24, 32, 40, 48, 56, and 64.

**Statistical analyses**

Changes from baseline in WOMAC and PGA-OA were calculated as the post-baseline value minus the baseline, with lower change score being more favorable. Negative changes therefore represent improvement in both measures.

A repeated measures longitudinal model with change in WOMAC domain score as the outcome and change in PGA-OA as the anchor measure was used to establish MWPC for WOMAC domains(17-21). In such a model, change from baseline in a WOMAC domain score was taken as a dependent variable using all available data from Weeks 2 to 24 in study 1, from Weeks 2 to 32 in study 2, and from Weeks 2 to 64 in study 3. The model contained data combined across treatment groups.

To study the appropriateness of the linear approximation of the relationship between predictor and outcome, we also studied the model when a predictor was used as a categorical variable. This version of the model does not impose any functional relationship, linear or otherwise, between an outcome and a predictor.

MWPC was evaluated using a 1-category change and, separately, a 2-category change on the anchor measure. The repeated measures longitudinal model calibrated the relationship between the outcome variable and anchor predictor by
taking the difference in mean change in outcome scores between adjacent
categories of the change in the anchor predictor (22). The theoretical range for
original unit changes in WOMAC scores is from –10 to 10. Conversion to percentage
change leads to a range from –100% to, theoretically, plus infinity for changes in
WOMAC domains scores.

In addition to producing estimates for MWPC for all patients within each study,
subgroups of patients based on index joint were separately analyzed to investigate
any potential differences in MWPC for patients with an index hip or knee.

The anchor-based repeated measures longitudinal model was supplemented
with empirical cumulative distribution function (eCDF) analyses, in accordance with
FDA recommendations(7). eCDFs were produced at the primary time points of Week
16 in studies 1 and 3 and of Week 24 in study 2.
RESULTS

The relationship between changes in PGA-OA and changes in WOMAC domains was analyzed in data collected from a total of 4480 patients, corresponding to 688, 844, and 2948 patients in studies 1, 2, and 3, respectively. Patient demographics and disease status were similar across the 3 studies (Table 1).

Using change in PGA-OA as a continuous or categorical variable revealed a robust relationship with change in all WOMAC domains across all 3 studies (Figure 1; Supplementary Figures 1 and 2). The correlations between changes in WOMAC Pain and Physical Function and changes in PGA-OA were approximately 0.6 for studies 1 and 3, and approximately 0.5 in study 2 (Table 2). Correlations between changes in WOMAC Stiffness and changes in PGA-OA were approximately 0.5 in studies 1 and 3, and approximately 0.4 in study 2. These data showed that a linear relationship between changes in WOMAC domains and changes in PGA-OA was supported and justified.

The relationship between changes in WOMAC domains and PGA-OA was examined across the 3 studies (Figure 2; Supplementary Tables 2a, 2b and 2c). The estimated MWPC for the 3 WOMAC domains were from 0.84–1.16 (original units) and from 12.50–16.23%, depending on study and domain, that corresponded to a 1-category change on PGA-OA (Tables 3a and 3b). For a 2-category change the corresponding values were from 1.68–2.31 (original units) and from 25.01–32.46%, respectively (Tables 3a and 3b). The estimated MWPC calculated separately for patients with an index hip or knee were not appreciably different from the pooled analyses or between subgroups of patients with an index hip or knee (Supplementary Tables 3 and 4).
To supplement the anchor-based method to estimate MWPC, eCDF curves for changes from baseline in WOMAC domains categorized by change in PGA-OA were generated. eCDFs for changes in WOMAC Pain in studies 1 and 3 showed a clear separation of curves for categories of PGA-OA changes with a sufficient number of available observations per category (Figure 3). Although not as clear, the curves for study 2 showed discernable separation, especially for the last 3 of the PGA-OA change categories. This pattern was consistent for all other domains across the 3 studies. In all studies and across all domains, the eCDF curves show that the majority of patients who reported no improvement in PGA-OA saw improvements in WOMAC scores.
DISCUSSION

These analyses of data from 3 randomized clinical trials of patients with moderate-to-severe OA of the hip or knee estimate that a 1-category change on PGA-OA corresponds to WOMAC score changes of 0.84–1.16 (original units) or 12.50–16.23%. A 2-category change on PGA-OA corresponds to an estimated 1.68–2.31 (original units) or 25.01–32.46% change in WOMAC, depending on domain and study. The same magnitude of MWPC is applied to meaningful within-person improvement and deterioration, the 2 differing only in sign. This is supported by evidence of a linear relationship between change in all WOMAC domains and change in PGA-OA.

Estimates for MWPC in WOMAC domains were produced using longitudinal data from all patients, regardless of treatment group or level of change from baseline in the current state anchor measure of PGA-OA. Consequently, these estimates are applicable at the individual patient level, in accordance with FDA guidance(7). This methodology contrasts with that used to calculate the MCID, which estimates improvement or worsening in an outcome at the individual or group level by comparing the difference in mean scores between categories of an anchor measure of change, which is subject to recall bias (1-3,8). When calculated at the group level, MCID estimates also need to take the wider disease context into consideration and cannot be used to determine meaningful change in outcomes at the individual patient level (23).

The estimated values for MWPC produced in this study are similar to and supportive of published data. The definitions of ≥ 10% reduction in pain as minimally important and ≥ 30% reduction as moderately important are consistent with the
MWPC in WOMAC for 1- and 2-category changes on PGA-OA, respectively (4,9). The definitions of an approximately 30% reduction in pain and an approximately 20% reduction in WOMAC Physical Function scores as clinically meaningful are similar to the data presented here(24). Clinically meaningful changes in patients who received rofecoxib, ibuprofen, or placebo were between 0.9–1.0 points on WOMAC (original 100 mm normalized visual analog scale converted to 0–10 scale), which is similar to the data presented here for a 1-category change on PGA-OA(25). Finally, the reported values for clinically meaningful changes in WOMAC domains in patients with OA of the hip or knee of between 0.3–1.3 (original standardized 0–100 scale converted to 0–10 scale), depending on domain, are consistent with data for a 1-category change on PGA-OA(10).

The relationships between change in WOMAC and change in PGA-OA were very similar across all domains and studies. Correlations between changes in WOMAC domains and PGA-OA were approximately 0.5–0.6 for studies 1 and 3, and 0.4–0.5 for study 2. Studies 1 and 3 were primarily based on data from US sites, while study 2 was primarily based on European data.

The eCDF curves for all studies show clear separation by changes in PGA-OA category, with the exceptions of –2 and –3 categories in study 2. When interpreting eCDFs and comparing with anchor-based modeling, it should be noted eCDFs are based on “completers” who had both outcomes collected. eCDFs simply visualize descriptive changes in outcome in a subgroup of subjects at a single time point in a study versus our anchor-based model, which uses all available data from all subjects and all time points.
Both anchor-based estimates of MWPC and eCDF curves show that patients who reported no change in PGA-OA experienced improvements in WOMAC scores. This could occur because WOMAC and PGA-OA measure similar but distinct concepts. Accounting for this, our model calibrates the relationship between WOMAC and PGA-OA by taking the difference in mean change in WOMAC scores between adjacent categories of the change in PGA-OA anchor; for example, by subtracting the mean change in WOMAC scores between the no-change and 1-category improvement categories on PGA-OA. Such a calibration is analogous to adjusting for placebo in an active intervention study; it is the relative or placebo-adjusted treatment effect that is important, rather than the unadjusted or absolute effect.

The model calibrates the relationship between change in WOMAC and change in PGA-OA over the entire span of their empirically predicted relationship. By contrast, a non-calibrated approach makes no such adjustment and therefore does not adjust for the no-change category on the PGA-OA by subtracting out its corresponding mean change score on WOMAC. This non-calibrated approach forces change on PGA-OA to correspond to no change on WOMAC and therefore assumes a perfect harmony exists in the relationship between the 2 measures, which is not the case in our data and in general.

In conclusion, these analyses of 3 tanezumab studies establish MWPCs for WOMAC domains in patients with moderate-to-severe OA of the hip or knee. A 2-category change on PGA-OA corresponds to an estimated 25.0–32.5% change in WOMAC domains, supporting previous studies describing an approximately 30% reduction in WOMAC as moderately clinically meaningful. An estimated 12.5–16.2%
change in WOMAC domains corresponded to a 1-category change on PGA-OA, suggesting that these improvements may be minimally clinically meaningful.

The large sample sizes used and the reproducibility of estimates for MWPC between each study demonstrate the robustness of the analyses performed. The estimates for MWPC were produced by combining data for both placebo and tanezumab treated patients for each study; however, the results are consistent with those of other studies of meaningful change in WOMAC, suggesting that they may be generalizable to this patient population. A possible limitation of this study is the use of only one anchor measure, as a credible alternative would have allowed sensitivity analyses to be performed.

These data may aid the interpretation of clinical trials in patients with OA of the hip or knee by defining the meaningful response to treatment in this patient population. Defining MWPC for patients with OA may also be valuable to healthcare professionals, as a means to assess the impact of their interventions. Further work should examine the most appropriate anchor measures and the best methods to determine MWPC.
Acknowledgements

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DATA SHARING

Upon request, and subject to review, Pfizer will provide the data that support the findings of this study. Subject to certain criteria, conditions and exceptions, Pfizer may also provide access to the related individual de-identified participant data. See https://www.pfizer.com/science/clinical-trials/trial-data-and-results for more information.
REFERENCES


nonsteroidal antiinflammatory drugs for hip or knee osteoarthritis: a randomized trial. Arthritis Rheumatol 2021; Feb 3 (in press).


FIGURE LEGENDS

Figure 1. Relationship between change in WOMAC domains and PGA-OA in study 3. Using PGA-OA as a continuous variable (blue lines), which implies a linear relationship with the outcome, or as a categorical variable (red lines), which does not impose any functional relationship, revealed an approximately linear relationship with all WOMAC domains in study 3. This relationship is observed when change in WOMAC is reported in original units (A) or as a percentage (B). A negative change represents an improvement in both measures. Pct: percent; PGA-OA: Patient Global Assessment of Osteoarthritis; WOMAC: Western Ontario and McMaster Universities Osteoarthritis Index.

Figure 2. Relationship between change in WOMAC domains and PGA-OA across all studies. The relationship between change in WOMAC domains and PGA-OA was observed across all domains and studies when reported in original units (A) or as a percentage (B). A negative change represents an improvement in both measures. Pct: percent; PGA-OA: Patient Global Assessment of Osteoarthritis; WOMAC: Western Ontario and McMaster Universities Osteoarthritis Index.

Figure 3. eCDF of change in WOMAC Pain score to support anchor-based models. eCDFs for the change in WOMAC Pain at the primary endpoint of each study, reported in original units (A) or as a percentage (B). eCDF: empirical cumulative distribution function; PGA-OA: Patient Global Assessment of Osteoarthritis; WOMAC: Western Ontario and McMaster Universities Osteoarthritis Index.
Figure 1. Relationship between change in WOMAC domains and PGA-OA in study 3.
Figure 2. Relationship between change in WOMAC domains and PGA-OA across all studies.

A

WOMAC Pain

WOMAC Physical Function

WOMAC Stiffness

WOMAC Total Score

Original Units

Mean change from baseline in WOMAC Pain

Mean change from baseline in WOMAC Physical Function

Mean change from baseline in WOMAC Stiffness

Mean change from baseline in WOMAC Total

Change from baseline in PGA-OA

Change from baseline in PGA-OA

Change from baseline in PGA-OA

Change from baseline in PGA-OA

B

Percentage

Mean Pt change from baseline in WOMAC Pain

Mean Pt change from baseline in WOMAC Physical Function

Mean Pt change from baseline in WOMAC Stiffness

Mean Pt change from baseline in WOMAC Total

Change from baseline in PGA-OA

Change from baseline in PGA-OA

Change from baseline in PGA-OA

Change from baseline in PGA-OA

Study 1

Study 2

Study 3
Figure 3. eCDF of change in WOMAC Pain score to support anchor-based models.

A

Study 1 – Week 16

Study 2 – Week 24

Study 3 – Week 16

B

Percentage

PGA-OA category change: -3, -2, -1, 0, 1

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Table 1. Baseline demographics and disease status.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Study 1 (n=696)</th>
<th>Study 2 (n=849)</th>
<th>Study 3 (n=2996)</th>
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<tr>
<td>Age (mean, [SD])</td>
<td>60.8 (9.6)</td>
<td>64.9 (9.4)</td>
<td>60.6 (9.4)</td>
</tr>
<tr>
<td>Female (n, [%])</td>
<td>453 (65.1)</td>
<td>587 (69.1)</td>
<td>1953 (65.2)</td>
</tr>
<tr>
<td>Index joint (n [%])</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hip</td>
<td>102 (14.7)</td>
<td>114 (17.0)</td>
<td>443 (14.8)</td>
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<tr>
<td>Knee</td>
<td>594 (85.3)</td>
<td>705 (83.0)</td>
<td>2553 (85.2)</td>
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<tr>
<td>Kellgren-Lawrence grade of index joint (n [%])</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>2 (0.2)</td>
<td>5 (0.2)</td>
</tr>
<tr>
<td>1</td>
<td>1 (0.1)</td>
<td>0</td>
<td>7 (0.2)</td>
</tr>
<tr>
<td>2</td>
<td>184 (26.5)</td>
<td>166 (19.6)</td>
<td>892 (29.8)</td>
</tr>
<tr>
<td>3</td>
<td>304 (43.7)</td>
<td>375 (44.2)</td>
<td>1425 (47.6)</td>
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<td>4</td>
<td>206 (29.6)</td>
<td>306 (36.0)</td>
<td>667 (22.3)</td>
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<tr>
<td>Disease duration (years) (mean [SD])</td>
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<td>7.5 (7.0)</td>
<td>8.8 (8.3)</td>
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<tr>
<td></td>
<td>Mean (SD) 1</td>
<td>Mean (SD) 2</td>
<td>Mean (SD) 3</td>
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<td>WOMAC Pain (mean [SD])</td>
<td>7.2 (1.2)</td>
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<td>WOMAC Physical Function (mean [SD])</td>
<td>7.3 (1.1)</td>
<td>6.7 (0.9)</td>
<td>7.1 (1.1)</td>
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<td>PGA-OA (mean [SD])</td>
<td>3.5 (0.6)</td>
<td>3.6 (0.6)</td>
<td>3.5 (0.6)</td>
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Baseline demographics and disease status for patients in each study. PGA-OA: Patient Global Assessment of Osteoarthritis; WOMAC: Western Ontario and McMaster Universities Osteoarthritis Index.
*Table 2. Correlations between change in WOMAC domains and change in PGA-OA.*

<table>
<thead>
<tr>
<th>WOMAC Domain</th>
<th>Study 1</th>
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<th>Study 3</th>
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<tr>
<td></td>
<td>Original Units</td>
<td>Percentage</td>
<td>Original Units</td>
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<td>0.613</td>
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<tr>
<td>Physical Function</td>
<td>0.624</td>
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<td>0.500</td>
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<tr>
<td>Stiffness</td>
<td>0.549</td>
<td>0.521</td>
<td>0.366</td>
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<tr>
<td>Total</td>
<td>0.615</td>
<td>0.578</td>
<td>0.479</td>
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Correlations between change in WOMAC domains and change in PGA-OA at the primary time points of week 16 in studies 1 and 3 and week 24 in study 2. PGA-OA: Patient Global Assessment of Osteoarthritis; WOMAC: Western Ontario and McMaster Universities Osteoarthritis Index.
Table 3a. Estimates of meaningful within-patient change by WOMAC domain and study in original units.

<table>
<thead>
<tr>
<th>WOMAC Domain</th>
<th>Study</th>
<th>1-Category Change in PGA-OA</th>
<th>2-Category Change in PGA-OA</th>
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<td></td>
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<td>(2.16-2.40)</td>
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<td>(2.16-2.39)</td>
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<th>2-Category Change in PGA-OA</th>
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<td>(1.63-1.82)</td>
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<tr>
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<td>2.25</td>
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<td>(1.10-1.15)</td>
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PGA-OA: Patient Global Assessment of Osteoarthritis; WOMAC: Western Ontario and McMaster Universities Osteoarthritis Index.

95% confidence intervals shown for each value.
**Table 3b.** Estimates of meaningful within-patient change by WOMAC domain and study as percentage change.

<table>
<thead>
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<td>31.02 (29.38-32.67)</td>
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<td>30.61 (29.02-32.21)</td>
<td>12.50 (11.81-13.20)</td>
</tr>
</tbody>
</table>

PGA-OA: Patient Global Assessment of Osteoarthritis; WOMAC: Western Ontario and McMaster Universities Osteoarthritis Index.

95% confidence intervals shown for each value.