Ixekizumab Improves Patient-Reported Outcomes in Non-Radiographic Axial Spondyloarthritis: Results from the Coast-X Trial

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ABSTRACT

Introduction: Ixekizumab, an interleukin-17A antibody, has shown efficacy in non-radiographic axial spondyloarthritis (nr-axSpA). The objectives of this analysis were (a) to measure improvement in ixekizumab-treated patients in Assessment of Spondyloarthritis International Society (ASAS) response domains and other patient-reported outcomes (PROs) and (b) to determine how ASAS responses were associated with changes in patient global disease activity (PtGA), spinal pain, function, stiffness, fatigue, and spinal pain at night.

Methods: COAST-X was a phase 3, 52-week multicenter, randomized, controlled trial investigating the efficacy and safety of 80-mg ixekizumab every 2 weeks (Q2W) and every 4 weeks (Q4W) in patients with active nr-axSpA. Changes from baseline in PROs were analyzed via mixed-effects models for repeated measures. Association analyses for ASAS responses used analysis of covariance with Scheffé’s method.

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**Results:** Patients treated with ixekizumab Q2W and Q4W reported significantly greater improvements in PtGA, spinal pain, function, and stiffness at week 1, when these measures were first assessed, compared with placebo ($p < 0.05$). ASAS40 responders, in comparison to ASAS20 non-responders, had the highest correlations with improvements in all response domains (PtGA, spinal pain, function, and stiffness) as well as fatigue and spinal pain at night ($p < 0.001$). ASAS40 responses were associated with 3.5- to 48.0-fold greater improvements in these PROs, with the highest values for PtGA and function, compared to ASAS20 non-achievement.

**Conclusions:** As early as week 1, patients with nr-axSpA treated with ixekizumab reported significant improvements in PtGA, spinal pain, function, and stiffness compared with those taking placebo. ASAS40 responders reported significantly greater improvements in all ASAS response domains (PtGA, spinal pain, function, and stiffness) as well as fatigue and spinal pain at night than ASAS20 non-responders. Improvements in PtGA and function appear to be major drivers in achieving ASAS40 response in patients with nr-axSpA.

**Trial Registration:** NCT02757352.

**Keywords:** Assessment of Spondyloarthritis International Society (ASAS) response; Biological therapy; Ixekizumab; Non-radiographic axial spondyloarthritis; Patient-reported outcomes

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**Key Summary Points**

**Why carry out this study?**

Patients with non-radiographic axial spondyloarthritis (nr-axSpA) experience negative impacts on health-related quality of life with pain and fatigue as notable symptoms.

This study measured improvement in ixekizumab-treated patients in patient-reported outcomes, including Assessment of Spondyloarthritis International Society (ASAS) treatment response domains to determine how ASAS responses were associated with changes in patient global disease activity, spinal pain, function, stiffness, fatigue, and spinal pain at night.

**What was learned from the study?**

Patients with nr-axSpA treated with ixekizumab reported greater improvements in patient-reported outcomes, and ASAS40 achievement had the highest correlation with improvements in patient global disease activity, spinal pain, function, stiffness, fatigue, and spinal pain at night.

Improvement in patient global disease activity and function may be drivers in achievement of ASAS40.

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**DIGITAL FEATURES**

This article is published with digital features, including a summary slide, to facilitate understanding of the article. To view digital features for this article, go to [https://doi.org/10.6084/m9.figshare.13187009](https://doi.org/10.6084/m9.figshare.13187009).
INTRODUCTION

Axial spondyloarthritis (axSpA), characterized by inflammation of the sacroiliac joints and axial skeleton, includes two subtypes: (1) ankylosing spondylitis (AS), also known as radiographic axSpA (r-axSpA), defined radiographically by substantial structural damage of the sacroiliac joints; and (2) non-radiographic axSpA (nr-axSpA), which lacks evidence of definite sacroiliitis on plain radiographs but may show evidence of inflammation on magnetic resonance imaging (MRI) in sacroiliac joints [1–4]. Of patients with axSpA, 40–60% may have nr-axSpA [5]. Patients with nr-axSpA experience similar negative impacts on health-related quality of life (HRQoL) to those in AS [1, 2, 6]. Spinal pain and fatigue particularly reflect the burden of disease in patients with axSpA [7].

Physical therapy and non-steroidal anti-inflammatory drugs (NSAIDs) are used as first-line treatments for nr-axSpA [8]. Biologic disease-modifying anti-rheumatic drugs (bDMARDs) are indicated for treatment of nr-axSpA patients with objective signs of inflammation who do not adequately respond to NSAID therapy [9]. While tumor necrosis factor inhibitors (TNFi) are efficacious in treating nr-axSpA, an unmet need remains since patients may not respond to TNFi or may only have partial responses [9–11].

Ixekizumab is a high-affinity monoclonal antibody that selectively targets interleukin (IL)-17A and has demonstrated efficacy in treating both AS and nr-axSpA [5, 11–13]. The Assessment of SpondyloArthritis International Society (ASAS) responses, ASAS20 and ASAS40 (defined by criteria including improvements of 20% and 40%, respectively) and partial remission, are primary endpoints in axSpA randomized controlled trials (RCTs). While ASAS responses are valuable in the context of RCTs, physicians in clinical practice focus on individual patient symptoms, such as patient global disease activity (PtGA), spinal pain, function, stiffness, fatigue, and spinal pain at night [14, 15]. The present study used data from the COAST-X trial to assess the impact of ixekizumab treatment over 52 weeks and explore how ASAS40 responses correlate with improvements in patient-reported outcomes (PROs).

METHODS

Study Design

COAST-X (NCT02757352) was a phase 3 multicenter, 52-week RCT assessing the efficacy and safety of 80-mg ixekizumab every 2 weeks (Q2W) and every 4 weeks (Q4W) compared with placebo in bDMARD-naive patients with active nr-axSpA.

Patients

Inclusion criteria for COAST-X have been detailed previously [5]. Eligible patients were adults with an established diagnosis of axSpA by a physician and fulfilled the Assessment of SpondyloArthritis International Society (ASAS) classification criteria for axSpA, with at least 12 weeks of previous therapy with NSAIDs and an inadequate response, as determined by the investigator, to two or more NSAIDs at the therapeutic dose range for a total duration of at least 4 weeks. Centrally read X-rays excluded patients with radiographic criteria of definite sacroiliitis according to modified New York criteria [4, 5]. Key inclusion criteria were active disease at screening and baseline (defined as a Bath Ankylosing Spondylitis Disease Activity Index [BASDAI] score of ≥ 4 and total back pain score of ≥ 4 on a 0–10 scale) and objective signs of inflammation, defined as evidence of elevated C-reactive protein (CRP > 5 mg/l) and/or presence of sacroiliitis on centrally read MRI, with sacroiliitis determined using the ASAS definition [16]. Enrolled patients could continue stable background medications including NSAIDs, conventional synthetic disease-modifying anti-rheumatic drugs (csDMARDs), corticosteroids, and analgesics. Ethical review boards approved COAST-X at each site before the trial began. Procedures involving human participants were performed within the ethical standards of the institutional and national research committees at all sites. COAST-X was conducted...
in accordance with the standards of the Declaration of Helsinki. All patients gave written informed consent before undergoing procedures related to the trials. The master ethics committee was Schulman Associates IRB, Cincinnati, OH, USA; complete listings of sites and investigators are available in the supplements of previously published results from COAST-X [5].

Randomization and Blinding

We randomized 303 patients 1:1:1 to receive placebo Q2W (N = 105), 80-mg of ixekizumab Q4W (N = 96), or 80-mg of ixekizumab Q2W (N = 102), via subcutaneous injection [5]. To achieve comparability between groups, randomization was stratified by country and screening MRI/CRP status (positive MRI and elevated CRP, positive MRI and non-elevated CRP, negative MRI and elevated CRP). The randomized, blinded treatment period ranged from weeks 0–52. From weeks 16–44, if patients’ disease activity required and at investigator discretion (with no specific, pre-defined switch criteria), adjustment of background medications for axSpA was allowed and/or patients were able to switch to open-label ixekizumab Q2W or subsequent TNFi treatment. Patients who switched to open-label treatment continued to be followed during the trial, and both investigators and patients remained blinded to original treatment assignment.

Outcome Measures

The primary endpoints for the COAST-X trial were the proportions of ASAS40 responders at weeks 16 and 52 and have already been reported [5]. Secondary endpoints included ASAS20 responders, changes from baseline in the response domains of ASAS criteria (PtGA, spinal pain, function, and stiffness), as well as BASDAI fatigue, Fatigue Numeric Rating Scale (NRS), and spinal pain at night.

PtGA, spinal pain, BASFI, and BASDAI responses were assessed at weeks 0 (baseline), 1, and 2, every 4 weeks from weeks 4–36, and then at weeks 44 and 52. Stiffness was measured as the average score based on responses to BASDAI questions 5 and 6: intensity and duration of morning stiffness [14]. Function was measured as the average score from responses to the Bath Ankylosing Spondylitis Functional Index (BASFI) [14]. Fatigue was measured with the Fatigue Severity NRS [17]. Fatigue NRS was recorded at weeks 0 (baseline), 8, 16, 36, and 52. Fatigue was also measured by BASDAI question 1 [14]. Spinal pain at night scores were recorded at weeks 0 (baseline), 1, and 2, every 4 weeks from weeks 4–36, and then at weeks 44 and 52.

Statistical Analyses

Statistical analyses were performed on data from the intent-to-treat population comparing ixekizumab dosing regimens to placebo during the double-blind treatment period (weeks 0–52) prior to biologic switches to ixekizumab 80-mg Q2W or TNFi. In patients treated with open-label ixekizumab 80-mg Q2W following biologic switch, only data up to the time of switch were included in the analysis. Mixed-effects models for repeated measures (MMRM) were used to compare changes from baseline in treatment groups at each visit for continuous outcomes for each PRO. Each model included treatment, geographic region, screening MRI/CRP status, baseline value, visit, baseline value-by-visit, and treatment-by-visit interactions as fixed factors. No prior imputation for missing data was performed for the MMRM modeling. P values were nominal.

Post hoc association analyses between ASAS responses and changes from baseline to weeks 16 and 52 in PROs were conducted by pooling patients with fully observed ASAS and outcome data at baseline and weeks 16 or 52, respectively, into three ASAS response groups: ASAS20 non-responders, ASAS20 responders (but not ASAS40 responders), and ASAS40 responders. Changes from baseline to weeks 16 and 52 in the above PROs were compared between ASAS responder groups using analysis of covariance (ANCOVA) models that included responses, baseline values, age, and gender as factors. Pairwise comparisons between response groups were conducted with Scheffé’s method for
No imputation was performed on missing values in ASAS responders or the PROs comprising the ASAS response criteria (PtGA, spinal pain, function, and stiffness); observed values were reported for these measures’ data. Modified baseline observation carried forward was used to impute missing fatigue and spinal pain at night data for ASAS correlations. SAS 9.4 was used for all analyses.
RESULTS

Baseline Characteristics

Baseline characteristics of patients enrolled in COAST-X are presented in Table 1 and were similar between treatment groups.

Changes from Baseline in Patient Global Disease Activity, Spinal Pain, Function, and Stiffness

Significant improvements in PtGA, spinal pain, function, and stiffness were reported as early as week 1 by patients treated with ixekizumab Q4W ($p = 0.002$, $p < 0.001$, $p = 0.034$, and $p = 0.008$, respectively) and ixekizumab Q2W ($p < 0.001$, $p = 0.002$, $p = 0.037$, and $p = 0.005$, respectively) compared with placebo (Fig. 1, Table 2). Significant improvements in PtGA were also reported at week 16 with ixekizumab Q4W and Q2W ($p = 0.01$ and $p = 0.001$, respectively) and week 52 with ixekizumab Q2W ($p = 0.027$) (Fig. 1b, Table 2). Significant improvements in function were also reported by patients treated with ixekizumab Q4W ($p = 0.040$) and Q2W ($p = 0.004$) at week 16 and sustained to week 52 ($p = 0.018$ and $p = 0.008$ for patients treated with ixekizumab Q4W and Q2W, respectively) (Fig. 1c, Table 2). Significant improvements in stiffness were reported by patients treated with ixekizumab Q4W ($p = 0.003$) and Q2W ($p < 0.001$) as early as week 16, and improvements were sustained to week 52 with ixekizumab Q4W ($p = 0.007$) and Q2W ($p < 0.001$) (Fig. 1d, Table 2).

Changes from Baseline in Fatigue and Spinal Pain at Night

At week 16, a significant improvement in Fatigue NRS was reported by patients receiving ixekizumab Q4W compared to placebo ($p = 0.024$; Fig. 2a, Table 2). Improvements in fatigue were not statistically significant with either ixekizumab dosing regimen at other time points (Fig. 2a, Table 2). Similarly, improvements in BASDAI fatigue were numerically greater but not statistically significant with both ixekizumab dosing regimens at weeks 16 and 52 (Fig. 2b, Table 2). Significant improvements in spinal pain at night were reported by patients treated with ixekizumab Q4W and Q2W as early as week 1 ($p = 0.029$ and $p = 0.013$, respectively) and at weeks 16 and 52 with ixekizumab Q2W as well ($p = 0.004$ and $p = 0.005$, respectively) (Fig. 2c, Table 2).

Association of ASAS Responses with Improvements in Patient Global Disease Activity, Spinal Pain, Function, and Stiffness

To examine how improvements in individual PROs relate to ASAS responses, improvement in the ASAS response domains PtGA, spinal pain,
Table 2  Changes from baseline in patient-reported outcomes at weeks 16 and 52

|                                | PBO  \((N = 105)\) | IXE Q4W \((N = 96)\) | IXE Q2W \((N = 102)\) |
|--------------------------------|---------------------|-----------------------|-----------------------|
| **Patient global disease activity** |                     |                       |                       |
| Week 16 Nx                      | 99                  | 96                    | 98                    |
| LSM (SE)                        | −1.3 (0.25)         | −2.3 (0.25)\(\dagger\) | −2.6 (0.25)\(\ddagger\) |
| Week 52 Nx                      | 34                  | 53                    | 52                    |
| LSM (SE)                        | −1.8 (0.38)         | −2.8 (0.32)           | −3.3 (0.32)\(\ddagger\) |
| **Spinal pain**                 |                     |                       |                       |
| Week 16 Nx                      | 99                  | 96                    | 98                    |
| LSM (SE)                        | −1.5 (0.24)         | −2.4 (0.25)*          | −2.6 (0.24)\(\ddagger\) |
| Week 52 Nx                      | 34                  | 53                    | 52                    |
| LSM (SE)                        | −2.3 (0.35)         | −2.9 (0.31)           | −3.3 (0.30)*          |
| **Function, BASFI**             |                     |                       |                       |
| Week 16 Nx                      | 99                  | 96                    | 98                    |
| LSM (SE)                        | −1.3 (0.2)          | −2.0 (0.2)*           | −2.3 (0.2)\(\ddagger\) |
| Week 52 Nx                      | 34                  | 53                    | 52                    |
| LSM (SE)                        | −1.6 (0.3)          | −2.6 (0.3)*           | −2.8 (0.3)\(\ddagger\) |
| **Stiffness (BASDAI questions 5 and 6)** |                   |                       |                       |
| Week 16 Nx                      | 99                  | 96                    | 98                    |
| LSM (SE)                        | −1.4 (0.24)         | −2.4 (0.25)\(\dagger\) | −2.9 (0.24)\(\ddagger\) |
| Week 52 Nx                      | 34                  | 53                    | 52                    |
| LSM (SE)                        | −1.9 (0.33)         | −3.2 (0.29)\(\ddagger\) | −3.5 (0.29)\(\ddagger\) |
| **Fatigue (NRS)**               |                     |                       |                       |
| Week 16 Nx                      | 99                  | 96                    | 98                    |
| LSM (SE)                        | −1.4 (0.24)         | −2.1 (0.24)*          | −1.9 (0.24)           |
| Week 52 Nx                      | 34                  | 53                    | 52                    |
| LSM (SE)                        | −2.1 (0.38)         | −2.6 (0.32)           | −2.7 (0.32)           |
| **Fatigue (BASDAI question 1)** |                     |                       |                       |
| Week 16 Nx                      | 99                  | 96                    | 98                    |
| LSM (SE)                        | −1.7 (0.23)         | −2.2 (0.24)           | −2.3 (0.24)           |
| Week 52 Nx                      | 34                  | 53                    | 52                    |
| LSM (SE)                        | −1.9 (0.34)         | −2.7 (0.29)           | −2.8 (0.29)           |

\(\dagger\) Adis
function, and stiffness (as well as the additional outcomes fatigue and spinal pain at night) were compared among three ASAS response levels pooled across treatment groups: ASAS20 non-responders, ASAS20 responders (but not ASAS40 responders), and ASAS40 responders. ASAS40 responders reported the greatest improvements in all PROs evaluated (Figs. 3 and S1). ASAS40 responders reported significantly greater improvements in PtGA, spinal pain, function, and stiffness at weeks 16 and 52 than ASAS20 non-responders. The magnitudes of improvement among ASAS40 responders compared to ASAS20 non-responders at week 52 were 6.4-fold in PtGA, 8.6-fold in spinal pain, 3.5-fold in function, and 3.5-fold in stiffness. ASAS20 responders also reported significant improvements in fatigue and spinal pain at night at weeks 16 and 52 compared with ASAS20 non-responders (Fig. S1). ASAS40 responders at week 16 reported an 11.7-fold improvement in fatigue and an 11.8-fold improvement in spinal pain at night compared to ASAS20 non-responders. The magnitudes of improvement among ASAS40 responders compared to ASAS20 non-responders at week 52 were 6.2-fold in fatigue and 3.5-fold in spinal pain at night. ASAS20 responders also reported significant improvements in fatigue and spinal pain at night at weeks 16 and 52 (Fig. S1).

**DISCUSSION**

ASAS40 responses at weeks 16 and 52 were the primary endpoints of the COAST-X clinical trial, a high bar that is challenging to meet since it necessitates considerable improvement in patients’ perceptions of symptoms [5, 15]. Because high placebo responses have been observed in AS studies, ASAS40 responses help better differentiate treatment effects [19]. At week 16 in the COAST-X trial, 35% of patients treated with ixekizumab Q4W and 40% of patients treated with ixekizumab Q2W were ASAS40 responders (compared with 19% receiving placebo). At week 52, 30% and 31% of patients receiving ixekizumab Q4W and ixekizumab Q2W, respectively, were ASAS40 responders.
responders (compared with 13% receiving placebo) [5]. ASAS40 responders reported significantly greater improvements in all ASAS components compared with ASAS20 responders and ASAS20 non-responders. In this study, ASAS40 responders reported a 48.0-fold improvement in PtGA and a 43.0-fold improvement in function compared to ASAS20 non-responders at week 16. With such notable improvements, PtGA and function appear to be major drivers in ASAS40 response for patients with nr-axSpA.

PtGA, spinal pain, stiffness, function, fatigue, and spinal pain at night reflect different burdens of disease that interact with each other and play an important part in overall health outcomes. In axSpA, severe spinal pain, impaired work productivity, and worse HRQoL have been linked [20]. Fatigue has been associated with greater disease activity and worse scores on the Ankylosing Spondylitis Quality of Life questionnaire [21]. It is now well established that the burden of disease is similar in both axSpA subtypes, nr-axSpA and AS [6, 22–24]. While widespread use of bDMARDs in nr-axSpA remains limited due to regulatory constraints, use of bDMARDs can improve patient outcomes [8, 11, 24].

In a previous study, patients with r-axSpA who were treated with ixekizumab and enrolled in the COAST-V and COAST-W trials reported greater improvements in patient-reported outcomes compared to placebo, and ASAS40 responders had greater improvements in PROs than ASAS20 non-responders [15]. In the current analysis of data from the COAST-X clinical trial of patients with nr-axSpA, we demonstrate that ixekizumab is effective at improving spinal PROs in patients with nr-axSpA. This analysis from COAST-X along with the previously reported results from COAST-V and COAST-W in r-axSpA indicate that treatment with ixekizumab is effective in improving the most impactful symptoms across the axSpA disease spectrum (r-axSpA and nr-axSpA) [15]. As previously reported in the r-axSpA trials assessing ixekizumab (COAST-V and COAST-W), here we also show that ASAS40 responders report greater improvements in PtGA, spinal pain, function, and stiffness, fatigue, and spinal pain at night compared with ASAS20 responders, and ASAS20 responders also report greater improvements in PROs than ASAS20 non-responders.

This study was limited due to the decrease in sample sizes after week 16 when patients who required changes in treatment were identified and switched either to ixekizumab Q2W or subsequent TNFi at the discretion of the investigator. The small sample sizes after week 16 affected analyses measuring significance in changes from baseline in PROs. The lack of specific pre-defined switch criteria to guide changes in study treatment may have impacted treatment effect size. Because of the decrease in sample size due to patients switched to ixekizumab Q2W after week 16, non-responder imputation was not applicable to the intent-to-
a Patient Global Disease Activity

b Spinal Pain

c Function (BASFI)

d Stiffness (BASDAI Questions 5 and 6)
treat sample. As a result, only observed data were used for post hoc association analysis between ASAS responses and PROs, potentially limiting the generalizability of results to the intent-to-treat population. The different recall periods of the Fatigue NRS and BASDAI questionnaires may have contributed to the significance of Fatigue NRS results and the non-significance of BASDAI fatigue results.

Strengths of this COAST-X RCT include a patient population representative of multiple global regions and inclusion criteria that required objective signs of inflammation (screening MRI/CRP status). Patients also had mean duration of symptoms over 10 years consistent with nr-axSpA and did not “convert” to r-axSpA. In addition, this analysis used recognized and validated instruments (ASAS response criteria, BASDAI, BASFI, and Fatigue NRS) to assess PROs.

CONCLUSIONS

Patients with nr-axSpA treated with ixekizumab reported significant improvements in PtGA, spinal pain, function, and stiffness compared with placebo at week 16, as early as week 1 and sustained in patients treated with ixekizumab Q2W through week 52. Greater improvements in fatigue were also reported by patients treated with either ixekizumab dosing regimen. ASAS40 responders reported significantly greater improvements in all response domains and in PROs than ASAS20 responders; ASAS20 responders also reported significantly greater PRO improvements than ASAS20 non-responders. ASAS40 responders reported 3.5- to 48.0-fold greater improvements in PtGA, spinal pain, function, stiffness, fatigue, and spinal pain at night compared to ASAS20 non-responders. Improvements in PtGA and function appear to be the major drivers in ASAS responses.

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**Compliance with Ethics Guidelines.** Ethical review boards approved COAST-X at each site before the trial began. Procedures involving human participants were performed within the ethical standards of the institutional and national research committees at all sites. COAST-X was conducted in accordance with the standards of the Declaration of Helsinki. All patients gave written informed consent before undergoing procedures related to the trials. The master ethics committee was Schulman Associates IRB, Cincinnati, OH, USA; complete listings of sites and investigators are available in the supplements of previously published results from COAST-X [5].

**Data Availability.** The datasets analyzed in this study are available from the corresponding author upon reasonable request.

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