Effect Of E-OA-07 On Improving Joint Health And Mobility In Individuals With Knee Osteoarthritis: A Randomized, Double-Blind, Placebo-Controlled, Parallel Group Study

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Aim: The aim of the present study was to evaluate the effect of E-OA-07 on individuals having osteoarthritis of the knee.

Background: Lanconone® (E-OA-07) is a widely marketed dietary supplement which has been previously studied in different clinical settings for managing chronic joint pain. This was a confirmatory study planned at a lowered dose regimen with the purpose of improving compliance and reducing consumer cost.

Methods: Male and female participants aged between 40 and 65 years, with history of joint pain for at least 3 years, were recruited. Knee joint dysfunction of grade II/III was radiographically characterized as per Kellgren-Lawrence system of classification. Enrolled participants were randomized to receive E-OA-07 at a dose of 1000 mg/day or placebo over a period of 8 weeks. The primary efficacy parameter was assessment of change in Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC) pain score. Whereas, the secondary parameters explored in the study included WOMAC subscales of stiffness and physical function, EQ-5D-5L questionnaire, systemic inflammatory marker (hs-CRP) and self-assessment of treatment satisfaction.

Results: At the end of 8 weeks, joint pain severity as per WOMAC was found to be significantly reduced in the E-OA-07 group as compared to placebo (p<0.001). Similar improvement was observed in the subscales of stiffness and physical function which corresponds to significant improvement in the quality-of-life standards of E-OA-07 participants (p<0.001), reporting higher treatment satisfaction (p<0.001).

Conclusion: E-OA-07 at a dose of 1000 mg/day was able to significantly reduce joint pain and thereby improve joint mobility in study participants. At the end of the study period, there was a clinically relevant change of 45.55%, 45.91% and 38.19% for pain, stiffness and physical function, respectively. Moving forward, studies could be planned for understanding the cartilage regenerative properties of E-OA-07.

Keywords: joint pain, joint inflammation, WOMAC, dietary supplement, osteoarthritis, Boswellia serrata

Introduction

In today’s society of unparalleled competition and stress, staying active is a hallmark of success mainly determined by strong bones and healthy joints. Age-related musculoskeletal disorders are one of the major causes for morbidity resulting in a worldwide socio-economic burden. There have been substantial advancements in
research as well as the number of therapies for degenerative joint disorders. However, a large extent of these are symptom oriented and are associated with unwanted adverse effects.

For instance, nonsteroidal anti-inflammatory drugs (NSAIDs), the cornerstone of pharmacotherapies, have been associated with life-threatening gastrointestinal adverse events such as bleeding and perforation.\(^2\) In 2015, the US Food and Drug Administration (FDA) took measures to strengthen the existing NSAIDs' label warnings, highlighting the increased risk of cardiovascular disorders with prolonged use of NSAIDs.\(^3\) Such events can significantly impact daily performance incurring increased health costs and physician visits.\(^4\) This emphasizes the need for therapeutic alternatives with proven safety profiles for chronic conditions of the joints.

The non-pharmacological treatments are viable therapeutic modalities for chronic conditions involving joint deterioration and inflammation.\(^5\) Physical activities are known to have a positive influence on patient’s overall quality of life (QoL) whilst improving an array of physical limitations.\(^6\) There is also a growing body of evidence for certain foods and nutrients that can significantly improve the symptoms of chronic joint conditions.\(^7,8\) Although research exploring their impact has grown significantly, it has been associated with varying findings and limitations. For instance, majority of these require strict patient adherence for an extended period of time, sometimes for life.\(^9\) As their impact on disease progression is currently unknown, their usage has been limited or in combination with other conventional therapies.

Individuals who do not respond well to conventional medical therapy are turning towards complementary and alternative medications, such as natural or herbal supplements. The efficacy and safety of joint health-related natural supplements such as *Zingiber officinale* (ginger)\(^10,11\) and *Curcuma longa* (turmeric)\(^12,13\) have already been established in several clinical trials and are known to have potent anti-inflammatory and analgesic effects. Even a recent meta-analysis concluded that standardized extracts of turmeric have the potential to alleviate joint pain and inflammation in osteoarthritic patients.\(^13\) With emerging evidence and clinical support promoting the use of natural alternatives, Enovate Biolife developed E-OA-07, a novel blend of 7 herbal ingredients for managing the symptoms of knee osteoarthritis (OA).

The previous studies of E-OA-07 have yielded promising and encouraging results, promoting its safe use in chronic joint conditions. An earlier proof of concept study for E-OA-07 (2700 mg/day) established its efficacy and safety profile, particularly for knee OA with no serious adverse events being reported.\(^14\) In another study, E-OA-07 (1000 mg/day) was compared to ibuprofen, to provide a safe and equipotent alternative for alleviating joint pain.\(^15\) Results demonstrated that E-OA-07 provided clinically meaningful relief within 3 hrs in activity-induced joint pain. Also, the quantum of relief experienced by participants was comparable to that of ibuprofen. With previous studies being indicative of the strong potency of E-OA-07, it encouraged us for conducting a confirmatory study in a different clinical setting. Therefore, the aim of the present study was to evaluate the efficacy and safety of E-OA-07 at a considerably lower dose with a view to improve treatment compliance and also for reducing cost borne by the consumer.

### Materials And Methods

#### Ethical Considerations

The study was approved by an independent ethics committee – Aditya, registered with the Office for Human Research Protections in the US Department of Health and Human Services (IRB00006475). Written informed consents were voluntarily obtained from all study participants, and the study was registered on public clinical trials registry of US National Library of Medicine (clinicaltrials.gov; NCT03658369). The study was performed in compliance with the Helsinki Declaration and ICH-GCP guidelines.

#### Participants

Participants of either sex, between the age group of 40 and 65 years suffering from OA of the knee were enrolled in the present study. The parameter of joint dysfunction of knee OA was characterized radiographically as grade II/III as per the “Kellgren-Lawrence” classification system.\(^16\) Volunteers with body mass index (BMI) of ≥25 and ≤29.9 kg/m\(^2\) and knee joint pain of ≥60 on a 100-point Pain-VAS (Visual Analogue Scale) score were included in the present study. Participants with prior or ongoing medical conditions (e.g., concomitant systemic illness, psychiatric condition, alcoholism, drug abuse, physical inability, electrocardiogram or laboratory abnormalities, a history of surgery or major trauma to the joints) were not included in the present study. Also, participants presenting any signs or having history of systemic infection during screening were excluded from the clinical trial. Urine test
for pregnancy was mandatory in females of childbearing potential. Individuals meeting the following criteria were included in the present 8 week randomized placebo-controlled study.

**Interventions**

The posology of E-OA-07 was determined through earlier conducted clinical trials. Based on the previous findings, the present study was designed to evaluate the efficacy of E-OA-07 at the lower dose of 1,000 mg in moderate-to-severe cases of knee OA. Lanconone® (Enovate Biolife, Wilmington DE, USA) is a buff-colored powder, rich in saponins, boswellic acids and other natural molecules purported to have analgesic and anti-inflammatory properties. It is a botanical compound based on 7 standardized herbal extracts, namely Shyonaka [Oroxyllum indicum], Ashwagandha [Withania somnifera], Shunthi [Zingiber officinale], Guggul [Commiphora mukul], Chochchini [Smilax chinensis], Rasana [Rasenlaceolata], Shallaki [Boswellia serrata] [Table 1]. The “0” size capsule used for the study contained 500 mg of E-OA-07 extract, and participants were instructed to take the 2 capsules after breakfast for the entire study period. Identical placebo capsules containing microcrystalline cellulose were manufactured and were matched for size, shape, color, texture, and packaging to preserve the blinding. The batch number of E-OA-07 used in this study was LN-AE-180801/02, and the control sample was retained with Enovate Biolife. Acetaminophen was allowed as a rescue medication at 500 mg dose in case of severe pain but prohibited 48 hours prior to each assessment. The products were manufactured in a “Good Manufacturing Practice” certified unit and were stored as per the manufacturer’s instructions throughout the study.

| S. No. | Ingredients   | Latin Name           | Quantity (mg) |
|-------|---------------|----------------------|---------------|
| 1     | Shyonaka      | Oroxyllum indicum    | 50            |
| 2     | Ashwagandha   | Withania somnifera   | 70            |
| 3     | Shunthi       | Zingiber officinale  | 30            |
| 4     | Guggul        | Commiphora mukul     | 140           |
| 5     | Chochchini    | Smilax chinensis     | 50            |
| 6     | Rasana        | Pluchea lanceolata   | 50            |
| 7     | Shallaki      | Boswellia serrata    | 110           |
|       |               |                      | Total: 500 mg |

**Study Conduct**

This was an 8 week randomized, double-blind, placebo-controlled, parallel group study in adults reporting knee joint pain. The study was conducted under the supervision of qualified orthopedicians. The first study participant was enrolled in August 2018, and final participant completed the protocol designated end of treatment visit in February 2019. Prior to randomization, a 7-day placebo run-in period was completed by all participants for identifying placebo responders and also for ascertaining treatment compliance. Participants with no placebo response were randomized in blocks of four using Stats Direct software (version 3.2) to either receive E-OA-07 or placebo. Once randomized on baseline (day 0), participants were followed-up on days 7, 28 and 56, respectively.

The master randomization chart was password protected and maintained in the electronic trial master file. Appropriate blinding was maintained throughout the study, and the blinding codes were secured in tamper-evident, sealed envelopes. Access was limited to authorized personnel on emergency basis as per Vedic Lifesciences standard operating procedures. The participants were also provided with a study diary to record study and non-study medications use during the course of the study. If any adverse events were experienced, the same was to be recorded by the participant in the study diary and communicated to the study investigator. Unused capsules were returned and checked by the trial coordinator for treatment compliance, who also checked the participant diary on scheduled visits to further verify IP compliance.

**Outcome Measures**

**Primary Outcome**

WOMAC is a self-administered validated instrument used widely in several clinical studies. The three WOMAC sub-scales comprise of 24 questions: Pain – 5, stiffness – 2, and physical function – 17, and each question is assessed on a Likert-based response rated from 0 to 4 points (0 indicates “no pain” and 4 “extreme pain”). Lower scores indicate improvement in participant’s QoL.

The primary outcome of the study was a reduction in WOMAC pain domain. The average response to 5 questions on a scale of 0–4 was calculated with a maximum score of 20. For evaluating the efficacy of E-OA-07, the change in the WOMAC pain values from day 0 to day 7, 28 and 56 was compared to that of the placebo.
Secondary Outcome
The secondary outcome for the study consisted of change in WOMAC subscales of stiffness (scale of 0–4; averaged response for 2 questions) and physical function (scale of 0–4; averaged response for 17 questions). Other measures performed included health-related quality of life (hr-QoL) as evaluated by a validated EQ-5D-5L questionnaire. It covers 5 domains – mobility, self-care, usual activities, pain/discomfort and anxiety/depression using 5 levels – no problems, slight problems, moderate problems, severe problems and extreme problems. Furthermore, treatment satisfaction was also assessed by a validated “Self-assessment of treatment (SAT)” questionnaire. The SAT questionnaire included five questions with 5-point response options for assessment of three areas, mainly pain relief, activity level and QoL.

Safety Assessments
Vital signs (blood pressure and pulse rate), laboratory parameters (complete blood count, liver – Serum glutamic oxaloacetic transaminase (SGOT) and Serum glutamic pyruvic transaminase (SGPT) and kidney profile – creatinine), frequency and occurrence of adverse events or serious adverse events, and use of rescue medication were considered for safety evaluations. Physical examination and vital signs were measured on days 0, 7, 28, and day 56. Blood samples were collected on days 0 and 56, and investigated using standard laboratory techniques by Suburban Diagnostics (Mumbai, India), accredited by “The College of American Pathologists” (CAP).

Quality Assurance
The study was conducted in compliance with the ICH-GCP guidelines laid down in E6 (R1) as per pre-approved monitoring and auditing plan by a Vedic Lifesciences team, independent of the clinical operational team.

Statistical Analysis
A study of continuous response variables from matched pairs of study participants in two groups of 1:1 ratio was planned. Based on the available data, the difference in the response of matched pairs is normally distributed with a standard deviation of 4.34. Taking into consideration the true difference in the mean response of matched pairs as 2.38, a sample size of 60 participants was needed to be studied for rejecting the null hypothesis. The Type I error probability associated with this test is 0.05 with power at 80%. To make up for some attrition during the trial period, a total of 72 participants were recruited and randomized in the ratio of 1:1. Data analysis was done for Per-Protocol (PP) population which was defined as the participants who completed the study without any major protocol deviations. Data normality was assessed using Shapiro–Wilk test. The baseline characteristics of the two groups were compared using independent t-tests for continuous variables and Chi-square tests for categorical variables. A “Mixed model Analysis of Variance (ANOVA)” was used to assess the effect of interventions across multiple time points. Furthermore, statistical significance between the study groups was also evaluated using “Mann–Whitney U-tests” (non-parametric) and “Wilcoxon signed rank tests” (parametric) for drawing best statistical conclusions. A value of p<0.05 was considered statistically significant, and data were expressed as mean ± standard deviation.

Data processing, tabulation of descriptive statistics, and calculation of inferential statistics were performed using Statistical Package for the Social Sciences (SPSS, IBM®) Python 3.0 (Armonk, New York, USA).

Results
A total of 78 potential participants were screened of which 72 were found eligible and were randomized in the ratio of 1:1. The enrolled population consisted of 36 participants each in E-OA-07 and placebo groups, respectively. As the study progressed, 5 participants from each group were lost to follow-up and withdrawn. The audit performed prior to data lock identified 18 participants, who were non-compliant to the study protocol and were hence excluded from
final statistical analysis. The end of the study population consisted of 44 participants with 21 in the study group and 23 in the placebo group, respectively. The participant disposition for the study has been presented in Figure 1.

Participant Demographics And Baseline Characteristics
At baseline, the two groups were comparable in terms of demographic characteristics. The mean (± SD) age of the participants in the E-OA-07 and placebo groups was 53.0 (± 6.35) and 52.43 (± 6.45) years, respectively. BMI values of the E-OA-07 and placebo groups were nearly similar (27.15 ± 1.46 and 27.60 ± 1.53). With respect to K&L classification, 16 participants were of grade II severity and 5 were of grade III in the E-OA-07 group, whereas 14 were grade II and 9 were of grade III severity in the placebo group. A summarized description of participant demographics and baseline characteristics is provided in Table 2.

Effect Of E-OA-07 On WOMAC Pain Score
At baseline, the mean ± SD pain score for E-OA-07 and placebo groups was 14.95 ± 1.72 and 14.87 ± 1.87, respectively (p=0.879). At the end of the study, E-OA-07 had a decreased mean score of 8.14 ± 3.83 (↓ 6.8 ± 3.6) and the placebo group score was 14.30 ± 3.81 (↓ 0.57 ± 3.03) [Tables 3 and 4]. The difference between the values of two groups clearly shows that E-OA-07 significantly reduced pain from the beginning
of the 4th week (p=0.033) and was further responsible for the decreased pain scores post 8 weeks of administration (p<0.001) [Figure 2A].

**Effect Of E-OA-07 On WOMAC Stiffness Score**

At baseline, the mean ± SD score for E-OA-07 and placebo groups was 5.38 ± 1.02 and 5.17 ± 1.61, respectively (p = 0.618). At the end of the study, stiffness score reduced significantly in the E-OA-07 to 2.91 ± 1.51, whereas the score increased to 5.44 ± 1.62 in the placebo group (p < 0.001) [Table 3 and Figure 2B]. At the end of 8 weeks, stiffness scores were significantly different in the E-OA-07 group as compared to placebo indicating significant reduction in stiffness subscale. Table 4 provides the absolute change in stiffness score for E-OA-07 and placebo groups, respectively.

**Effect Of E-OA-07 On WOMAC Physical Function Score**

A comparative analysis of WOMAC physical function at baseline showed a mean ± SD score of 63.71 ± 6.86 and 61.83 ± 8.70 (p = 0.431) for the E-OA-07 and placebo groups, respectively. At the end of the study, the physical function score was 39.38 ± 13.66 and 63.09 ± 11.64 (p < 0.001), indicating significant improvement in the physical functioning of E-OA-07 study participants as compared to placebo group [Table 3 and Figure 2C]. The absolute change in physical function score for E-OA-07 and placebo groups is provided in Table 4.

### Table 2 Participants Demographics And Baseline Characteristics

|                      | Placebo (n=23) |                      | E-OA-07 (n=21) |                      | p value |
|----------------------|----------------|----------------------|----------------|----------------------|---------|
|                      | Mean (SD)      | 95% CI               | Mean (SD)      | 95% CI               |         |
|                      | Min            | Max                  | Min            | Max                  |         |
| Age (years)          | 52.43 (6.45)   | 41-62                | 53 (6.35)      | 40-63                | 0.771   |
| Gender (%)           |                |                      |                |                      |         |
| Male                 | 4 (17.40)      |                      | 10 (47.62)     |                      | 0.032*  |
| Female               | 19 (82.61)     |                      | 11 (52.39)     |                      |         |
| BMI (kg/m²)          | 27.60 (1.53)   | 26.94-28.27          | 27.15 (1.46)   | 26.48-27.82          | 0.322   |
| Pulse Rate (Per Minute) | 85.26 (9.75) | 81.04-89.48          | 84.71 (10.93)  | 79.74-89.69          | 0.862   |
| BP Systolic (mmHg)   | 123.39 (6.77)  | 120.46-126.32        | 123.24 (7.01)  | 120.05-126.43        | 0.942   |
| BP Diastolic (mmHg)  | 82.43 (7.45)   | 79.21-85.66          | 80.43 (6.42)   | 77.50-83.35          | 0.346   |
| FBS (mg/dl)          | 103.26 (14.09) | 97.17-109.36         | 101.71 (15.11) | 94.83-108.59         | 0.727   |
| KL Grades of OA (n)  | Grade II – 14  |                      | Grade II – 16  |                      | 0.276*  |
|                      | Grade III – 9  |                      | Grade III – 5  |                      |         |
| VAS Pain Score       | 76.09 (9.88)   | 71.81-80.36          | 77.62 (11.79)  | 72.25-82.99          | 0.642   |
| Total EQ-5D Score    | 14.91 (2.92)   | 13.65-16.18          | 15.86 (2.69)   | 14.63-17.08          | 0.273   |

**Notes:** p, independent Student's t-test; *p, Chi-square test.

**Abbreviations:** n, number of participants; BMI, body mass index; BP, blood pressure; FBS, fasting blood sugar; K&L, Kellgren Lawrence; VAS, visual analogue scale; SD, standard deviation, CI, confidence interval; Min, minimum; Max, maximum.
Effect Of E-OA-07 On Total WOMAC Score

The total WOMAC score was significantly improved in the E-OA-07 group as compared to placebo. At baseline, total WOMAC score for E-OA-07 and placebo groups was reported to be 84.05 ± 8.65 and 81.87 ± 11.54, respectively (p = 0.486). At the end of study, the scores were decreased significantly in the E-OA-07 group to 50.43 ± 18.57, whereas the score was 82.83 ± 16.77 in the placebo group.
group; p < 0.001. The difference of scores as compared from baseline implies that there was a significant improvement in E-OA-07 participants at the end of 8 weeks [Table 3 and Figure 2D]. Table 4 provides the absolute change in total WOMAC score for E-OA-07 and placebo groups, respectively.

Table 4 Change In WOMAC Subscale Of Pain, Stiffness, Physical Function And Total WOMAC Scores

| Visit          | Placebo (n=23) | E-OA-07 (n=21) | p value* |
|----------------|----------------|----------------|----------|
|                | Mean (SD)      | 95% CI         | Mean (SD) | 95% CI     |
|                | Min            | Max            | Min       | Max        |
| Pain           |                |                |          |            |
| Change (Day 7) | 0.00 (1.09)    | −0.47          | 0.47     | −0.10 (1.41) | −0.737 | 0.547 | 0.802 |
| Change (Day 28)| −0.52 (2.35)   | −1.54          | 0.50     | −2.57 (2.66) | −3.781 | −1.362 | 0.010 |
| Change (Day 56)| −0.57 (3.03)   | −1.87          | 0.74     | −6.81 (3.60) | −8.448 | −5.171 | <0.001 |
| Stiffness      |                |                |          |            |
| Change (Day 7) | 0.22 (0.95)    | −0.19          | 0.63     | −0.05 (0.81) | −0.414 | 0.319 | 0.327 |
| Change (Day 28)| −0.13 (1.39)   | −0.73          | 0.47     | −0.86 (1.20) | −1.401 | −0.313 | 0.071 |
| Change (Day 56)| 0.26 (1.82)    | −0.52          | 1.05     | −2.48 (1.37) | −3.097 | −1.855 | <0.001 |
| Physical function |            |                |          |            |
| Change (Day 7) | 0.61 (5.48)    | −1.76          | 2.98     | −0.05 (3.54) | −1.66 | 1.57 | 0.643 |
| Change (Day 28)| −0.78 (8.45)   | −4.44          | 2.87     | −8.95 (8.87) | −12.99 | −4.91 | 0.003 |
| Change (Day 56)| 1.26 (12.10)   | −3.97          | 6.49     | −24.33 (13.80) | −30.62 | −18.05 | <0.001 |
| Total WOMAC   |                |                |          |            |
| Change (Day 7) | 0.83 (6.56)    | −2.009         | 3.661    | −0.19 (4.94) | −2.437 | 2.06 | 0.567 |
| Change (Day 28)| −1.44 (11.27)  | −6.306         | 3.436    | −12.38 (11.70) | −17.708 | −7.05 | 0.003 |
| Change (Day 56)| 0.96 (16.26)   | −6.073         | 7.986    | −33.62 (17.94) | −41.785 | −25.45 | <0.001 |

Note: *P, mixed analysis of variance.

Abbreviations: n, number of participants; SD, standard deviation; CI, confidence interval; Min, minimum; Max, maximum.

Effect Of E-OA-07 On European Quality Of Life-5 Dimension Score
At the end of 8 weeks, trend of reduction in the E-OA-07 group of participants was similar in each of the 5 domains of the EQ-5D questionnaire (mobility; self-care; daily activities; pain/discomfort; and depression – p < 0.001). At the baseline visit, the total EQ-5D score for E-OA-07 and placebo groups was 15.86 ± 2.69 and 14.91 ± 2.92, respectively. At the end of the study, total score for E-OA-07 group was significantly reduced to 8.57 ± 3.14 and was comparably different from the placebo group score of 15.96 ± 3.72; p < 0.001 [Table 5 and Figure 3].

Effect Of E-OA-07 On Treatment-Related Satisfaction
At the end of the study, overall SAT scores of E-OA-07 and placebo groups were 20.05 ± 4.60 and 11.30 ± 4.62, respectively, [Table 6]. The difference between the efficacy assessments of two groups was statistically significant (p < 0.001), and participants were to a greater extent satisfied in the E-OA-07 group as compared to placebo.

Effect Of E-OA-07 On High-Sensitivity C-Reactive Protein
At baseline, both groups had nearly similar levels of hs-CRP. The E-OA-07 value was 3.79 ± 3.43, and for placebo, it was 3.18 ± 2.87. The hs-CRP levels did not show any significant improvements in either of the groups as the hs-CRP values of the placebo group were 4.65 ± 6.09 and E-OA-07 were 3.86 ± 4.00, respectively. Though the values of placebo group were increased, the change between the groups was non-significant [Table 7].

Rescue Medication And Compliance
When compared from day 0 to 28, the acetaminophen usage in E-OA-07 was 9.38 ± 8.41 and placebo group was 10.35 ± 7.59, respectively. However, from day 28 to 56, it reduced to 4.14 ± 4.45 in the E-OA-07 group as compared to 8.83 ± 10.59 in the placebo group. Also, the compliance recorded in the E-OA-07 group of participants was more than 90% in the
analyzed population. This shows that participants in the E-OA-07 group experienced a greater degree of relief as compared to the comparator (placebo) group in the course of the study duration.

Safety Evaluations

Physical examination and vital signs were normal throughout the study, and there were no clinically significant abnormal findings in either of the study groups [Table 7]. The hemoglobin levels, RBC Count, WBC Count, and biochemical evaluations of SGOT and SGPT were in the range of normal laboratory reference values with no clinically significant observations in either of the groups. Overall, E-OA-07 was found to be safe and was well tolerated by the study participants [Table 8].

Adverse Events

During the course of the study, there were no serious/severe adverse events reported in either of the study groups.

Discussion

In the last few decades, drug discovery from plant-based extracts has been a primary area of research for scientists and academicians alike. The importance of nutraceuticals for the management of musculoskeletal disorders is now being recognized globally in terms of alleviating health-care standards and managing health-care costs. It is now evident from a number of clinical trials that nutraceuticals can be envisioned as one of the missing blocks for prevention and treatment of chronic disorders, such as OA. In the present study, we evaluated E-OA-07, a natural alternative for reducing joint pain and improving physical function, specifically in weight-bearing joints deteriorated by degenerative conditions.

Nutritional supplements are generally touted as safer alternatives to pharmacotherapies. However, due to their slow onset of action, they do not serve as a preferred choice for treating chronic disorders. A study by Majeed et al assessed the safety and efficacy of *Boswellia serrata* extract in the management of knee OA. The extract demonstrated a
19.6% reduction in the total WOMAC score as compared to the initial baseline values.\textsuperscript{27} Whereas, E-OA-07, during the same period, reduced total WOMAC score by 40% despite having higher baseline scores. Additionally, the perceived improvement in \textit{Boswellia serrata} group was evident only after a period of 120 days. Whereas, E-OA-07 achieved the same within 28 days of administration, thus demonstrating its strong analgesic action. In another study, Lugo and associates assessed the efficacy and tolerability of a collagen supplement in exercise-induced joint pain. Significant reduction in pain was observed after 6 months of IP administration, but with a number of side effects.\textsuperscript{28} In the present study, no adverse or serious adverse effects were reported during the 8 week study period. Overall, results of the present study support the effectiveness and safety of E-OA-07 in individuals suffering from symptoms of knee OA.

![EQ-5D - Quality of life Questionnaire](image)

Figure 3 Quality-of-life score response to placebo and E-OA-07 therapy. Significant improvement was observed in each of the domains: mobility, self-care, daily activities, pain/discomfort, and anxiety/depression. \textsuperscript{*}p <0.001.

| Table 5 Effect Of E-OA-07 And Placebo On EQ-5D-5L Scores |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Domains         | Visit           | Placebo (n=23)   | E-OA-07 (n=21)   | p value*        |
|                 | Mean (SD)       | 95% CI          | Mean (SD)       | 95% CI          |
|                 | Min Max         | Min Max         | Min Max         |
| Mobility        | Day 0           | 3.09 (0.67)     | 2.80 3.38       | 3.24 (0.70)     | 2.92 3.56       | 0.324 |
|                 | Day 56          | 3.52 (0.99)     | 3.09 3.95       | 1.67 (0.73)     | 1.33 2.00       | <0.001 |
|                 | p**             | 0.008           | <0.001          |
| Self-care       | Day 0           | 2.65 (1.03)     | 2.21 3.10       | 2.91 (0.77)     | 2.56 3.26       | 0.408 |
|                 | Day 56          | 2.96 (0.88)     | 2.58 3.34       | 1.62 (0.81)     | 1.25 1.99       | <0.001 |
|                 | p**             | 0.117           | <0.001          |
| Daily Activities| Day 0           | 3.48 (0.90)     | 3.09 3.87       | 3.48 (0.99)     | 3.03 3.92       | 0.786 |
|                 | Day 56          | 3.39 (1.03)     | 2.95 3.84       | 1.86 (0.66)     | 1.56 2.16       | <0.001 |
|                 | p**             | 0.691           | <0.001          |
| Pain/Discomfort | Day 0           | 3.35 (0.71)     | 3.04 3.66       | 3.67 (0.97)     | 3.23 4.11       | 0.213 |
|                 | Day 56          | 3.48 (1.04)     | 3.03 3.93       | 1.81 (0.75)     | 1.47 2.15       | <0.001 |
|                 | p**             | 0.490           | <0.001          |
| Anxiety/Depression | Day 0     | 2.35 (1.07)     | 1.89 2.81       | 2.57 (0.87)     | 2.18 2.97       | 0.459 |
|                 | Day 56          | 2.61 (0.78)     | 2.27 2.95       | 1.62 (0.74)     | 1.28 1.96       | <0.001 |
|                 | p**             | 0.238           | 0.001           |
| Total           | Day 0           | 14.91 (2.92)    | 13.7 16.2       | 15.86 (2.69)    | 14.63 17.08     | 0.265 |
|                 | Day 56          | 15.96 (3.72)    | 14.3 17.6       | 8.57 (3.14)     | 7.14 10.00      | <0.001 |
|                 | p**             | 0.141           | <0.001          |

Notes: *P, Mann–Whitney U-test; **Wilcoxon signed rank test.
Abbreviation: n, number of participants; SD, standard deviation; CI, confidence interval; Min, minimum; Max, maximum.
grade III chronic knee OA as per the K&L scale. The study showed an ~50% reduction in the WOMAC pain scale which is similar to the results observed in the present study, but at a significantly lower dose. This implies that the lower dose of 1000 mg/day is as effective as the previously studied dose of 2700 mg/day. Our second study by Girandola and associates recruited healthier individuals conforming to radiographic conditioning of K&L grade I/II. E-OA-07 was compared with ibuprofen for assessing the degree of relief experienced by study participants in activity-induced joint pain. Clinically meaningful relief was experienced by the participants within 3 hrs of E-OA-07 administration. This potent analgesic action of E-OA-07 was consistent across both studies even though the trial designs, study population, duration and objectives of Girandola’s research and ours were completely different. Also, by using healthier population and non-disease endpoints as efficacy measures, we believe that the analgesic action of E-OA-07 can also be applicable to younger or middle-aged adult population.

Table 6 SAT Scores Evaluated At The End Of Study Period (Day 56)

|Domains            | Placebo (n=23) Mean (SD) | 95% CI | E-OA-07 (n=21) Mean (SD) | 95% CI | p value* |
|--------------------|--------------------------|--------|--------------------------|--------|----------|
|Pain relief        | 2.17 (1.03) 1.73 (2.62)  |        | 3.91 (1.09) 3.41 (4.40)  | <0.001 |
|Activity levels    | 2.13 (0.97) 1.71 (2.55)  |        | 3.71 (1.15) 3.19 (4.24)  | <0.001 |
|Change in QOL      | 2.52 (0.95) 2.11 (2.93)  |        | 3.91 (1.14) 3.39 (4.42)  | <0.001 |
|Treatment satisfaction | 2.22 (1.17) 1.71 (2.72)  |        | 4.29 (0.90) 3.88 (4.70)  | <0.001 |
|Treatment preference| 2.26 (1.25) 1.72 (2.80)  |        | 4.24 (0.94) 3.81 (4.67)  | <0.001 |
|Total              | 11.30 (4.62) 9.31 (13.30) |        | 20.05 (4.60) 17.95 (22.14) | <0.001 |

Note: *P, Mann–Whitney U-test.
Abbreviations: n, number of participants; SD, standard deviation; CI, confidence interval; Min, minimum; Max, maximum.

Table 7 Vital Signs

|Visits            | Placebo (n=23) Mean (SD) | 95% CI | E-OA-07 (n=21) Mean (SD) | 95% CI | p value* |
|------------------|--------------------------|--------|--------------------------|--------|----------|
|Pulse Rate        |                          |        |                          |        |          |
|Day 0             | 87.57 (9.24) 68 (100)    |        | 82.38 (9.78) 61 (98)     |        | 0.003    |
|Day 56            | 80.04 (8.85) 65 (98)     |        | 80.29 (8.22) 62 (97)     |        |          |
|Systolic blood pressure |                |        |                          |        |          |
|Day 0             | 122.48 (7.52) 110 (134)  |        | 123.71 (7.46) 110 (130)  |        | 0.622    |
|Day 56            | 120.87 (7.33) 110 (130)  |        | 122.14 (6.81) 110 (130)  |        | 0.236    |
|Diastolic Blood Pressure |            |        |                          |        |          |
|Day 0             | 80.30 (6.04) 70 (90)     |        | 79.10 (7.22) 70 (90)     |        | 0.102    |
|Day 56            | 77.39 (7.52) 70 (90)     |        | 77.62 (7.68) 70 (90)     |        | 0.465    |

Note: *P, Wilcoxon Signed Rank test.
Abbreviations: n, number of participants; SD, standard deviation; CI, confidence interval; Min, minimum; Max, maximum.
### Table 8 Hematological And Biochemical Parameters

| Visits | Placebo (n=23) | E-OA-07 (n=21) |
|--------|----------------|----------------|
|        | Mean (SD)      | 95% CI         | Mean (SD)      | 95% CI         |
|        | Min | Max       | Min | Max       |
| Hemoglobin (g/dl) |         |         |         |         |
| Day 0  | 12.50 (0.92)  | 10.9 15.0   | 12.95 (1.66)  | 9.6 16.3      |
| Day 56 | 12.58 (1.03)  | 11.1 15.1   | 13.20 (1.66)  | 10.0 16.4     |
| p value | 0.385         | 0.321       |
| Red Blood Cells (RBC) Count (mil/cmm) | | | | |
| Day 0  | 4.50 (0.40)   | 3.44 5.20   | 4.66 (0.64)   | 3.31 5.97     |
| Day 56 | 4.57 (0.51)   | 2.98 5.60   | 4.75 (0.60)   | 3.88 6.21     |
| p value | 0.322         | 0.295       |
| White Blood Cells (WBC) Count (mil/cmm) | | | | |
| Day 0  | 8669.57 (2327.39) | 4900 12,000 | 7485.00 (1401.98) | 4600 10,300 |
| Day 56 | 9004.35 (2779.06) | 5400 16,100 | 8352.38 (1651.55) | 5000 11,200 |
| p value | 0.685         | 0.036       |
| Platelet Count (per cmm) | | | | |
| Day 0  | 291,086.96 (68,425.08) | 141,000 467,000 | 270,350.00 (72,955.12) | 112,000 455,000 |
| Day 56 | 294,521.74 (75,998.73) | 152,000 403,000 | 275,380.95 (63,242.77) | 128,000 354,000 |
| p value | 0.194         | 0.668       |
| Serum Glutamic-Oxaloacetic Transaminase (SGOT)(U/L) | | | | |
| Day 0  | 19.64 (6.87)  | 9.8 37.8    | 20.15 (7.26)  | 12.1 44.9     |
| Day 56 | 20.87 (7.52)  | 10.1 36.2   | 21.71 (7.74)  | 13.9 43.2     |
| p value | 0.354         | 0.651       |
| Serum Glutamic-Pyruvic Transaminase (SGPT) (U/L) | | | | |
| Day 0  | 19.02 (9.10)  | 8.9 45.2    | 19.12 (9.31)  | 5.5 35.8      |
| Day 56 | 19.35 (8.77)  | 8.9 42.1    | 20.47 (9.74)  | 9.1 40.7      |
| p value | 0.495         | 0.958       |
| High sensitivity C-Reactive Protein (hs-CRP) | | | | |
| Day 0  | 3.79 (3.43)   | 2.31 5.27   | 3.18 (2.86)   | 1.88 4.48     |
| Day 56 | 4.65 (6.09)   | 2.02 7.29   | 3.86 (4.00)   | 2.04 5.68     |
| p value | 0.438         | 0.455       |

Note: *P, Wilcoxon Signed Rank test.

Abbreviations: n, number of participants; SD, standard deviation; CI, confidence interval; Min, minimum and Max, maximum.
In the present study, we utilized EQ-5D questionnaire for measuring hr-QoL in participants with moderate-to-severe cases of knee OA. At the end of the study, majority of E-OA-07 participants had their QoL improved, and a clear pattern of improvement was observed in each of the EQ-5D domains. Additionally, these improvements led to better treatment satisfaction in E-OA-07 participants reporting higher SAT scores as compared to the placebo group. Therefore, results of the present study show that E-OA-07 bettered QoL by reducing pain intensity, which in turn raised the treatment satisfaction of study participants.

Elevated levels of hs-CRP have been correlated with OA disease progression and also with symptoms of pain and stiffness. However, the relationship between inflammatory markers and knee OA still remains controversial topic due to contradictory findings. In a recent meta-analysis, only fish oil was able to significantly reduce hs-CRP levels, whereas botanicals such as garlic, ginseng, saw palmetto, and pycnogenol-containing supplements failed to induce any changes. In the present study, there was no significant change observed in the hs-CRP levels of study participants. We believe this lack of effect can possibly be attributed to the presence of confounding factors such as obesity, asymptomatic viral or bacterial infection or age-related cardiovascular fatigue in the present study population. Longer duration studies may be required to ascertain the same.

Two of the most frequent causes of non-compliant behavior are misunderstanding of dosage regimens and the cost borne by the consumer. A lifelong compliance to a variety of regimens ultimately influences health-care costs. This is particularly of importance in individuals with repeated and overuse of joints, as they are at greatest risk for chronic conditions. If there was a way to manage this, not only the level of compliance would be improved but also the consumer costs would be lessened. In the present study, the efficacy of E-OA-07 on a reduced dose was approved and improvement in each of the efficacy variables was significantly superior to the placebo group. Also, results are further consolidated by the excellent safety and tolerability profile demonstrated by E-OA-07 in clinical as well as laboratory parameters. All these attributes can certainly have beneficial effects in terms of efficacy, compliance and cost-effectiveness. However, there is a potential limitation to the present study. There were no statistically significant changes observed in the hs-CRP marker between the groups or within the groups. Future studies can be attempted to explore the effect of E-OA-07 on hs-CRP, preconditioned to a larger but homogenized set of population with fewer confounding variables.

Conclusion
E-OA-07 at a dose of 1000 mg/day was able to significantly reduce joint pain, and thereby improve joint mobility in study participants. At the end of the study period, there was a clinically relevant change of 45.55%, 45.91% and 38.19% for pain, stiffness and physical function, respectively. Moving forward, studies could be planned for understanding the cartilage regenerative properties of E-OA-07.

Abbreviations
WOMAC, Western Ontario and McMaster Universities Osteoarthritis Index; EQ-5D-5L, European quality of life-5 dimensions-5 levels; NSAIDs, nonsteroidal anti-inflammatory drugs; OA, osteoarthritis; IEC, independent ethics committee; BMI, body mass index; VAS, visual analogue scale; K&L, Kellgren and Lawrence; SAT, self-assessment of treatment; hs-CRP, high-sensitivity C-reactive protein; ICH-GCP, International council of harmonisation – good clinical practice; ANOVA, analysis of variance; SPSS, statistical package for the social sciences; SD, standard deviation; CI, confidence interval; QoL, quality of life; RBC, red blood corpuscle; WBC, white blood corpuscle; SGOT, serum glutamic oxaloacetic transaminase; SGPT, serum glutamic pyruvic transaminase.

Data Sharing Statement
The datasets used and/or analyzed during the current study are available on a reasonable request from the authors.

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Disclosure
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