Clinical Study

The Safety and Efficacy of an Enzyme Combination in Managing Knee Osteoarthritis Pain in Adults: A Randomized, Double-Blind, Placebo-Controlled Trial

Wolfgang W. Bolten,1 Michael J. Glade,2 Sonja Raum,3 and Barry W. Ritz4

1Klaus-Miehlke Klinik, Leibnizstraße 23, 65191 Wiesbaden, Germany
2The Nutrition Doctor, 8612 Kedvale Avenue, Skokie, IL 60076, USA
3Mucos Pharma GmbH & Co. KG, Mirastraße 17, 13509 Berlin, Germany
4Atrium Innovations, 4 Hillman Drive, Suite 190, Chadds Ford, PA 19317, USA

Correspondence should be addressed to Barry W. Ritz; br@atrium-innovations.com

Received 17 September 2014; Revised 5 December 2014; Accepted 19 December 2014

Academic Editor: Burkhard Leeb

Copyright © 2015 Wolfgang W. Bolten et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

This randomized, double-blind, placebo-controlled, and comparator-controlled trial evaluated the safety and efficacy of an enzyme combination, as Wobenzym, in adults with moderate-to-severe osteoarthritis (OA) of the knee. Adults (n = 150) received Wobenzym, diclofenac (a nonsteroidal anti-inflammatory drug, NSAID), or placebo for 12 weeks. Improvement in pain scores (Lequesne Functional Index) did not differ between subjects treated with Wobenzym or diclofenac, and both treatment groups improved compared to placebo (P < 0.05). Reduction in total WOMAC scores (secondary outcome measure) did not differ between Wobenzym and diclofenac, although only diclofenac emerged as different from placebo (P < 0.05). The median number of rescue medication (paracetamol) tablets consumed was less in the Wobenzym group compared to placebo (P < 0.05), while there was no difference between diclofenac and placebo. Adverse events were similar in frequency in Wobenzym and placebo groups (7.2% and 9.1% of subjects, resp.) and higher in diclofenac group (15.6%). Wobenzym is comparable to the NSAID diclofenac in relieving pain and increasing function in adults with moderate-to-severe painful knee OA and reduces reliance on analgesic medication. Wobenzym is associated with fewer adverse events and, therefore, may be appropriate for long-term use.

1. Introduction

Osteoarthritis (OA), known as degenerative joint disease, is characterized by gradual loss of joint cartilage and intermittent or persistent local inflammatory processes. The hallmark signs of this progressive disorder are limitations in joint movement and pain. OA is the most common form of arthritis affecting nearly 14% of adults aged 25 to 65 and nearly 34% of adults over the age of 65 [1]. The most common site of osteoarthritis in individuals 40 years and older is the knee [2].

The signature pathologic feature of osteoarthritis is the progressive net loss of hyaline cartilage from within the joint, driven by the chondrocytic responses to aberrant loading of mechanical forces within the joint “mechanopathology,” with concomitant inflammatory responses to the degeneration of joint structures [3–5]. Nonsteroidal anti-inflammatory drugs (NSAIDs) are effective in reducing inflammation and pain but are associated with significant side effects that can interfere with therapeutic success, especially with long-term treatment [6].

Wobenzym is an orally administered combination of proteolytic enzymes and bioflavonoid. In combination, these active principles have produced beneficial outcomes in subjects with osteoarthritis of the knee and hip which were equivalent to those produced by the NSAID diclofenac [7–11]. However, these studies lacked a placebo control group.

In this current randomized, double-blind clinical trial we compared the effectiveness of an enzyme combination, as Wobenzym, with the effectiveness of the NSAID diclofenac and placebo. Outcome measures included amelioration of
pain, improvement of function, and type and frequency of side effects.

2. Methods

2.1. Data Integrity. The conduct of the experiment and the integrity of the collected data were monitored through oversight provided by Monitor Zentrale Deutschland (Sauerlach, Germany). The accuracy of the collected data was overseen by Clinical Research Facilities International (Schaik, Netherlands). The trial was reviewed and approved by an independent ethics committee and was performed according to the principles defined in the current edition of the Declaration of Helsinki, according to German Drug Law (AMG), and according to Good Clinical Practice (GCP). This trial was registered with ClinicalTrials.gov on March 13, 2014 (identifier NCT02088411).

2.2. Subjects. Middle-aged and older adult volunteers with moderate-to-severe knee osteoarthritis were enrolled in the study after satisfying all inclusion and exclusion criteria. Briefly, inclusion required the age of 40–80, a score on Lequesne Functional Index of 10–14 (the maximum possible index score is 24), WOMAC-A pain subscale score of ≥25 (scores range from 0 to 100), and arthritis of the knee confirmed by conventional radiography or tomography and swelling of the affected knee upon physical examination. Subjects were excluded based on a history of knee trauma, joint infection, joint surgery, or intra-articular injection (viscotherapy). Subjects with gastrointestinal diseases or with pharmacotherapy with corticosteroids, COX-II inhibitors, or glucosamine/chondroitin were excluded, as well as patients with known sensitivity to paracetamol, NSAIDs, or oral enzymes. Written informed consent was obtained from each patient prior to study enrollment following an oral and written explanation about the aim and the potential risks of the study.

2.3. Interventions. Subjects were recruited, screened, evaluated for eligibility, and enrolled into the study at clinical centers located throughout Germany (Bensheim, Berlin, Essen, Hamburg, Luneburg, Munchen, Norderstedt, Siegen, Stockach, Wiesbaden, and Wolfratshausen) and 2 centers in the Netherlands (Oos, Zwoll). Subjects were assigned randomly to receive three times daily either 1 tablet of diclofenac sodium (50 mg; Merckle GmbH, Blaubeuren, Germany) and 2 tablets of an indistinguishable placebo, or 2 tablets of Wobenzym (Mucos Pharma GmbH & Co. KG, Berlin, Germany) and 1 tablet of placebo, or 3 tablets of placebo. The resultant doses of active treatments were, therefore, Wobenzym 6 tablets (2 tablets, 3 times daily) or diclofenac 150 mg (1 tablet, 3 times daily). All interventions were identical in the total number of tablets and appearance. The random assignment of subjects to treatment groups was achieved through the use of a computer-generated randomization list in continuous recruitment order.

Wobenzym is an oral combination of natural compounds, including 288 mg trypsin (from porcine or bovine pancreas), 540 mg bromelain (from pineapples, Ananas comosus), and 600 mg rutoside trihydrate (rutin; from Japanese pagoda tree, Sophora japonica) per recommended daily dose. The tablets are enteric coated to prevent inactivation of the enzymes during gastric passage. The tablets must be consumed on an empty stomach, separate from meals.

2.4. Study Protocol. Each treatment was administered daily for 12 weeks. Compliance was determined by counting unconsumed tablets during examinations that were scheduled at 4, 8, and 12 weeks of the intervention. A minimum of 75% compliance was required. When necessary the subjects were allowed to self-medicate with the rescue medication paracetamol (acetaminophen, 500 mg per tablet) up to 2000 mg in a single 24-hour period. The use of rescue medication was not permitted 24 hours prior to each scheduled examination. The incidence of use and amount of paracetamol consumed were recorded in a patient diary. Other antirheumatic or analgesic therapies were prohibited, although continued use of daily low-dose (81 mg) aspirin for cardiovascular health was permitted. Self-administered exercise was allowed but the initiation of a new physical therapy regimen was prohibited.

2.5. Outcomes. The primary outcome was self-assessment of the affected knee joint by physician interview using the Lequesne Functional Index and reported as the total score. This index provides an estimate of the degree of pain associated with the affected joint, the maximum distance walked, and the activities of daily living [12]. Additional outcome criteria focused on self-assessments of pain and function utilizing the Western Ontario and McMaster Universities Index version 3.0 subscales for pain (WOMAC-A), joint stiffness (WOMAC-B), and physical joint function (WOMAC-C), as well as the total score after combining all three subscale scores [13]. In addition, selected indices of systemic inflammation (erythrocyte sedimentation rates at 1 and 2 hours and serum concentration of C-reactive protein) were evaluated [14]. These assessments were performed at baseline and 12 weeks.

The safety of each treatment was assessed by documenting the occurrence, nature, severity, and relevance of all adverse events, as well as potential adverse drug reactions that occurred during the course of the study, either recorded in patient diaries or communicated directly to study personnel. In addition, the following vital signs and blood chemistries were measured at baseline and week 12: resting systolic and diastolic blood pressure, heart rate, and body temperature; hematocrit; red and white blood cell count; thrombocyte count; fasting blood glucose concentration; fasting serum concentrations of total protein, albumin, urea-nitrogen, total bilirubin, creatinine, hemoglobin, calcium, sodium, potassium, and chloride; fasting serum activities of alkaline phosphatase, glutamic-oxaloacetic transaminase (SGOT; also known as aspartate aminotransferase; ASAT), glutamic/glutamine pyruvic transaminase (SGPT; also known as alanine aminotransferase; ALT), and γ-glutamyl transpeptidase (γ-GT; GGT); and prothrombin times and partial prothrombin times.
2.6. Data Analysis and Statistics. In order to account for baseline variability among subjects, the primary and secondary pain and joint function data were converted into the change and the percentage change for each subject during the 12 weeks of the study. Because the number of subjects that completed the study was too small for the meaningful application of tests of normality, all observations, 12-week changes, and 12-week percentage changes were expressed as medians (and their 95% confidence intervals) and were compared using nonparametric statistical procedures. Each combination of variable and treatment group, including the 12-week changes, 12-week percentage changes, and nontransformed blood data, was tested against the null hypotheses of "no change" or "no percent change" using the sign test ($\alpha = 0.05$) (InStat GraphPad Software, La Jolla, CA, USA). The change and percent change in each of the primary and secondary pain and joint function outcome variables and in each index of systemic inflammation were determined using the 2-sided Kruskal-Wallis procedure ($\alpha = 0.05$) [15]. When the null hypothesis was rejected for a particular outcome, three pairwise comparisons were performed using the 2-sided Mann-Whitney procedure for unpaired data ($\alpha = 0.05$). The median number of paracetamol tablets consumed by the subjects in the three treatment groups was compared using this same 2-step nonparametric approach. Within each treatment group, the percentage of subjects who consumed any paracetamol was compared to the percentage of subjects who consumed no paracetamol using the $z$-test for equality of proportions [15].

3. Results

A total of 295 subjects were enrolled and randomly assigned to treatment groups. Of these, 150 completed the full 12 weeks of the trial according to protocol; 46 received diclofenac, 52 received Wobenzym, and 52 received placebo (Figure 1). According to protocol, 59 subjects were determined ineligible at baseline because they no longer met inclusion criteria for Lequesne and/or WOMAC pain scores. Of the other 86 subjects who did not complete the study, 27 withdrew voluntarily, 16 were discontinued due to adverse reactions, 28 were enrolled in two study centers that did not follow protocol, and 15 subjects were noncompliant. Reasons for withdrawal included "patient request" (4 subjects), perceived lack of efficacy (4 Wobenzym subjects, 4 diclofenac subjects, and 7 placebo subjects), loss to follow-up (3 subjects), and other protocol interruptions (5 subjects). Efficacy data were analyzed based on 150 completers. All enrolled subjects were included in the safety analysis.

3.1. Baseline Characteristics. Gender, age, body weight, height, body mass index, the number of months since the initial medical diagnosis of knee osteoarthritis, and the duration (in months) of active symptoms present upon enrollment did not differ between the treatment groups (Table 1).

3.2. Primary Outcomes. The primary outcome of affected knee joint function and joint-associated pain quantified using the Lequesne Functional Index improved in all three treatment groups during the 12 weeks of the study (Table 2). In the subjects treated with diclofenac or Wobenzym, both the absolute reduction in total score (median change:
Table 3: WOMAC subset and total scores.

<table>
<thead>
<tr>
<th></th>
<th>Median score</th>
<th>Median change</th>
<th>Median % change</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>Week 12</td>
<td></td>
</tr>
<tr>
<td>WOMAC-A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diclofenac</td>
<td>29.0</td>
<td>15.0</td>
<td>−13.5*</td>
</tr>
<tr>
<td>Wobenzym</td>
<td>31.0</td>
<td>19.0</td>
<td>−12.5*</td>
</tr>
<tr>
<td>Placebo</td>
<td>30.0</td>
<td>21.0</td>
<td>−8.0*</td>
</tr>
<tr>
<td>WOMAC-B</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diclofenac</td>
<td>12.0</td>
<td>6.0</td>
<td>−4.0*</td>
</tr>
<tr>
<td>Wobenzym</td>
<td>11.0</td>
<td>8.0</td>
<td>−4.0*</td>
</tr>
<tr>
<td>Placebo</td>
<td>12.0</td>
<td>8.0</td>
<td>−3.0*</td>
</tr>
<tr>
<td>WOMAC-C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diclofenac</td>
<td>100.0</td>
<td>53.0</td>
<td>−32.0*</td>
</tr>
<tr>
<td>Wobenzym</td>
<td>101.5</td>
<td>70.0</td>
<td>−28.5*</td>
</tr>
<tr>
<td>Placebo</td>
<td>102.0</td>
<td>75.0</td>
<td>−19.0*</td>
</tr>
<tr>
<td>WOMAC total</td>
<td>136.5</td>
<td>74.0</td>
<td>−50.0*</td>
</tr>
</tbody>
</table>

All values presented as medians (95% confidence interval).
* Significantly different from "no change" (P < 0.05).
A,B Treatment groups with different superscript differ within the same column (P < 0.05).

−4.0 points and −3.3 points, resp.) and the relative reduction in total score (median % change: −32.7% and −27.4%, resp.) were nearly twice those of the corresponding improvements in the subjects treated with placebo (−2.0 points, −16.4%) (P < 0.05). Absolute and relative reductions in pain did not differ between the diclofenac and Wobenzym treatment groups.

3.3. Secondary Outcomes. All three treatment groups exhibited similar improvements in absolute and relative scores on the WOMAC-A (joint pain) and WOMAC-B (joint stiffness) subscales (P < 0.05), and there were no significant differences between the groups (Table 3). Scores on the WOMAC-C subscale (joint function) and WOMAC total score were also reduced in all three groups (P < 0.05), but the relative reductions reached significance compared to placebo only among subjects treated with diclofenac (P < 0.05). There were no differences in WOMAC-C or WOMAC total score between the Wobenzym and diclofenac groups.

3.4. Paracetamol Consumption. The median number of paracetamol tablets consumed during the study was significantly less (P < 0.05) in the group treated with Wobenzym (0.5 tablets per subject in 12 weeks) compared to the placebo group (10.0 tablets per subject in 12 weeks), and the median number of paracetamol tablets consumed by subjects treated with diclofenac (4.0 tablets per subject in 12 weeks) did not differ from the other two groups (Table 4). Within each of the three treatment groups, the percentage of subjects who consumed any paracetamol (users) was not different from the percentage of subjects who consumed no paracetamol during the study.
Table 4: Paracetamol (acetaminophen) consumption.

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>Median number of tablets consumed</th>
<th>Paracetamol users</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diclofenac</td>
<td>4.0^{AB}(0.0, 23.0)</td>
<td>26 56.5</td>
</tr>
<tr>
<td>Wobenzym</td>
<td>0.5^{A}(0.0, 7.0)</td>
<td>26 50.0</td>
</tr>
<tr>
<td>Placebo</td>
<td>10.0^{B}(2.0, 22.0)</td>
<td>33 63.5</td>
</tr>
</tbody>
</table>

^{A,B}Treatment groups with different superscripts differ within the same column (\(P < 0.05\)).

Table 5: Adverse events resulting in discontinuation of treatment.

<table>
<thead>
<tr>
<th>Treatment group</th>
<th>Gender</th>
<th>Adverse events</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diclofenac</td>
<td>F</td>
<td>Nausea</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>Diarrhea</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>Acute leukocytosis</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>Eczema</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>Stomach pain</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Edema</td>
</tr>
<tr>
<td>Wobenzym</td>
<td>F</td>
<td>Heartburn</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>Stomach pain</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>Back pain</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>Stomach pain accompanied by gastroesophageal reflux and nausea</td>
</tr>
<tr>
<td>Placebo</td>
<td>F</td>
<td>Nausea</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>Nausea</td>
</tr>
<tr>
<td></td>
<td>F</td>
<td>Edema of legs</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>Flatulence and diarrhea</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>Abnormal laboratory results</td>
</tr>
<tr>
<td></td>
<td>M</td>
<td>Stomach pain and episodes of vertigo</td>
</tr>
</tbody>
</table>

3.5. Indices of Systemic Inflammation. Erythrocyte sedimentation rates and serum C-reactive protein concentrations were not affected in any group during the 12 weeks of the study (data not shown).

3.6. Safety. There were no relevant changes in vital signs or blood chemistries in any group (see supplemental data in the Supplementary Material available online at http://dx.doi.org/10.1155/2015/251521). During the study, 16 subjects were discontinued due to adverse events (4 Wobenzym subjects, 6 diclofenac subjects, and 6 placebo subjects). These were reported as mostly mild-to-moderate gastrointestinal events, such as nausea, reflux, and stomach pain (Table 5). In the entire enrolled study population, there were 32 recorded adverse events, primarily gastrointestinal, which were considered possibly related to the intervention. Among these, 7 were in the Wobenzym group (7.2%), 16 in the diclofenac group (15.6%), and 9 in the placebo group (9.1%). None of these events included ulcers or ulcer complications.

4. Discussion

The enzyme combination, as Wobenzym (2 tablets, three times daily, for 12 weeks), produced significant improvements in joint pain and function as measured by the Lequesne Functional Index in middle-aged and older adults with moderate-to-severe knee osteoarthritis. Self-assessed reductions in the degree of pain associated with the affected joint were recorded according to physician interview. There were significant improvements in the ability to walk for a distance and affected knee joint flexibility. The benefits of Wobenzym were comparable to those experienced with diclofenac (150 mg) and were significantly greater than the responses to placebo. The adverse event profile of Wobenzym was similar to placebo.

In addition, the use of rescue medication (number of tablets consumed), potentially an indirect measure of treatment efficacy, was reduced by 95% in the Wobenzym group compared to placebo (0.5 versus 10 tablets, resp.), while there was no significant difference in the use of rescue medication between the diclofenac and placebo groups or between the Wobenzym and diclofenac groups. There was no difference in the percentage of “paracetamol users” between the three groups. The use of rescue medication was associated with less response to treatment in all three groups, although this reached significance only in the diclofenac group (data not shown). Because (1) paracetamol rescue was required only when self-assessed pain exceeded a self-determined threshold of discomfort, (2) the same percentage of subjects resorted to pharmacologic rescue at least once in each group, and (3) the median number of paracetamol tablets consumed per subject was reduced in the enzyme group, these data suggest that use of Wobenzym was accompanied by fewer episodes of suprathreshold discomfort, a clinically relevant outcome observed during the 12 weeks of study.

Both the enzyme combination and diclofenac have been reported to exhibit anti-inflammatory activity [16–18]. OA is not associated with a measurable systemic but local inflammatory reaction. Accordingly the data in this study do not support an effect on markers of systemic inflammation. A downregulation of local inflammation in OA is accompanied by an improvement in clinical symptoms (reduction of swelling and pain), but not in lab parameters. Clinical observations suggest that local anti-inflammatory activity accompanied diclofenac and Wobenzym use and produced improved functionality. However, self-assessment of pain by WOMAC, a secondary outcome measure, and function (WOMAC-C subscale) differed between diclofenac and placebo but did not differ between Wobenzym and either placebo or diclofenac. The trend for improving in the Wobenzym group as seen in the WOMAC analysis was similar to the Lequesne outcome but did not reach statistical significance.

There is a large body of literature that explores the effectiveness and safety of the enzyme combination in managing joint pain [7–11, 16, 19–23]. Several studies also have reported...
enhanced mobility and recovery following sports injuries, including a faster return to training or competition [24–27]. However, the vast majority of these studies were comparator trials that did not include a placebo control. The value of such studies is limited because it is not possible to discern the additional benefits of the enzyme combination beyond the well-established influence of placebo on subjective measures of joint pain [28, 29]. When a placebo group is omitted from an experimental design and in clinical trials assessing the effectiveness of oral agents in OA, the “placebo effect” can be both statistically and clinically significant [30]. The current study included both a diclofenac group, as an active treatment comparator, and a placebo control. Previous studies demonstrating the effectiveness of Wobenzym in managing joint pain were primarily comparative studies establishing noninferiority compared to NSAID as standard of care [7–11]. The randomized, double-blind, placebo-controlled, and comparator-controlled trial design is, therefore, a major strength of this study. The close monitoring of rescue medication use is another strength of this study, as it provided an additional clinically relevant outcome measure.

While the multisite design is a strength of the study, the data obtained from the subjects from two clinical study sites were excluded from analysis because those study centers did not follow study protocol. This resulted in a smaller patient population sample available for analysis than originally anticipated. The large number of subject exclusions and resultant smaller sample size is the major limitation of the study and severely limits the ability to interpret these results. As such, the current study must be considered a reanalysis of the data.

The current study did not evaluate efficacy in the management of acute knee pain; therefore, no conclusions can be drawn regarding acute efficacy, such as time to onset. However, because of the progressive nature of this condition, finding a medication or supplement that can be used safely and effectively for the long term is a prudent clinical objective. The moderate length of this study does not conclusively demonstrate whether Wobenzym would remain effective beyond 12 weeks or whether Wobenzym would remain safer than diclofenac for longer-term use. However, the data suggest comparable efficacy and a better safety profile, as compared to NSAID, with repeated use over the study period of 12 weeks. While some data support the safe long-term use of the enzyme combination, such as up to 1–5 years, these studies are limited and concern subject populations not included here (rheumatoid arthritis, multiple sclerosis) [31–33]. Therefore, further controlled trials are needed to confirm whether Wobenzym may safely support knee joint function in an adult population over an extended period of time.

5. Conclusions

The enzyme combination Wobenzym may be as effective as the NSAID diclofenac in the management of chronic OA of the knee when administered for 12 weeks, with a similar safety profile to placebo, although further study is needed. Use of Wobenzym reduced the reliance on paracetamol (acetaminophen) for additional pain relief. Wobenzym may be a safe and effective option for the daily management of long-term joint pain.

Conflict of Interests

Sonja Raum and Barry W. Ritz are employed by companies associated with the manufacture and sale of the investigational agent used in this study. There is no other conflict of interests.

Authors’ Contribution

Wolfgang W. Bolten conceived of the study and was the principle investigator in the acquisition of data. Michael J. Glade conducted the statistical analysis of data. Sonja Raum and Barry W. Ritz participated in the analysis and interpretation of the data, and Barry W. Ritz drafted the paper with the participation of all authors. All authors were involved in paper revisions and have given final approval of this paper version to be published. All authors agree to be accountable for all aspects of the work.

Acknowledgments

The authors wish to thank Karolyn Gazella and Kelly Gibson for their assistance in preparing the paper.

References

Arthritis


