Nucleotides Cytidine and Uridine Associated with Vitamin B12 vs B-Complex Vitamins in the Treatment of Low Back Pain: The NUBES Study

Purpose: We report the results of low back pain treatment using a combination of nucleotides, uridine (UTP), cytidine (CMP) and vitamin B₁₂, vs a combination of vitamins B₁, B₆, and B₁₂.

Patients and Methods: Randomized, double-blind, controlled trial, of a 60-day oral treatment: Group A (n=317) receiving nucleotides+B₁₂ and Group B (n=317) receiving B vitamins. The primary endpoint was the percentage of subjects in each group presenting adverse events (AEs). Secondary endpoints were visual analog scale (VAS) pain scores at Visit 2 (day 30) and Visit 3 (day 60) in relation to pretreatment values, Roland–Morris Questionnaire (RMQ) scores and finger-to-floor distance (FFD) (percentage of subjects per group presenting improvement ≥5 points and ≥3cm, respectively).

Results: Seventy-five (24%) and 105 (33%) subjects (P=0.21) presented 133 and 241 AEs, with 3159% of subjects presenting ≥2 AEs (P=0.019) in Group A and Group B, respectively. Twenty-four subjects in Group B were discontinued due to AEs, while no AE-related discontinuations occurred in Group A (P=0.001). VAS score reduction after 30 and 60 days of treatment was statistically significant (P=0.001) in both groups, with Group A showing greater reduction at Visit 2 (P=0.001). RMQ score improvement ≥5 points occurred in 99% of subjects from each group, and FFD improvement ≥3 cm occurred in all subjects.

Conclusion: Treatment with nucleotides+B₁₂ was associated with a lower number of total AEs, fewer AEs per subject, and no AE-related treatment discontinuation. Pain intensity (VAS) reduction was superior at 30 days of treatment in the nucleotides+B₁₂ group and equivalent between groups at 60 days of treatment. Improvements in efficacy measures RMQ and FFD were observed in both groups at treatment days 30 and 60.

Keywords: low back pain, uridine, cytidine, vitamin B₁, vitamin B₆, vitamin B₁₂

Introduction
Low back pain is a very common condition arising from strains or sprains to the muscles/ligaments of the lumbar region, described as pain, muscle tension, or stiffness in the region below the costal margin and above the inferior gluteal folds with or without accompanying leg pain.¹ Low back pain is often associated with trauma, sudden movements, or poor body mechanics, with an estimated 84% of the general population experiencing this condition at least once during adulthood.² In up to 90% of cases, low back pain is nonspecific, presenting in the absence of a detectable pathoanatomical cause.³,⁴ Low back pain is characterized by duration as acute (lasting up to
four weeks), subacute (lasting between four and 12 weeks), and chronic (persisting for 12 weeks or more).5

Contributing factors associated with low back pain include age—with the highest incidence in the third decade of life and increasing overall prevalence until the age of 60–65 years—educational status, psychosocial status, job satisfaction, occupational aspects, and obesity.6 Although often a self-limited condition not requiring active medical treatment, low back pain is one of the most common reasons for physician visits worldwide, leads the conditions associated with increased limitations to daily activities and missed work, and carries a significant direct and indirect economic burden at the individual, family, community, industry, and government levels.1,6,7

Treatment of low back pain aims to control pain and maintain function, and includes pharmacological and non-pharmacological therapies.4 Frequently used pharmacological therapies for low back pain include nonsteroidal anti-inflammatory drugs (NSAIDs), acetaminophen, tramadol, muscle relaxants, and antidepressants. However, those substances could have severe adverse effects.5

Evidence from previous exploratory clinical studies using either the combination of nucleotides and vitamin B12 or the B vitamins (B1, B6, and B12) has demonstrated symptomatic and functional improvement among patients treated for painful conditions including neuralgia, neuropathic pain manifestations, low back pain, and osteoarthritis-related pain.9–14 The objective of this study was to compare the clinical tolerability and efficacy of the combination of uridine triphosphate, cytidine monophosphate and hydroxocobalamin vs the combination of thiamine nitrate, pyridoxine hydrochloride, and cyanocobalamin in patients presenting low back pain, over a 60-day treatment period.

Written informed consent was obtained from each patient. The protocol was conducted in accordance with the Declaration of Helsinki and Good Clinical Practices and was approved by local independent ethics committee (Centro Universitário Serra dos Órgãos—UNIFESO—approval no. 1.295.147). This study is registered with the ClinicalTrials.gov identifier NCT 02640417.

Patient Population

Patients were included if they met the following criteria: patients of both genders, age ≥18 years, clinical presentation of acute low back pain—duration of ≤3 days from screening date, with and without functional alteration, with medium severity, defined as visual analog pain (VAS) scale score of 20–80 mm. Women of childbearing potential were to use adequate birth control methods throughout the duration of the study.

The main exclusion criteria were: known allergy or hypersensitivity to any component of the study drugs; need for surgical treatment; concomitant treatment with other analgesics; clinically significant laboratory alterations; gastric or duodenal ulcer; gastrointestinal, cerebrovascular or other bleeding; women who were pregnant, breastfeeding or of child-bearing potential not using acceptable birth control methods; and any disease or condition rendering the subject incompatible with participation in the trial.

Study Procedures

Subjects were treated on an ambulatory basis. The study included three visits: pretreatment, Visit 2 (after 30 days of treatment) and Visit 3 (after 60 days of treatment). Each study visit consisted of a physical examination and vital signs measurement, laboratory testing, and low back pain and functionality assessments. These variables included: a 100 mm VAS (ranging from 0=“no pain” to 100=“maximum pain intensity”); the 24-item Roland Morris Low Back Pain and Disability Questionnaire (RMQ) translated and validated in Brazilian Portuguese, consisting of 24 items assessing the impact of low back pain on functional activities, scored with 1 point for each item endorsed, for a total score ranging from 0 to 24 points, with higher scores representing higher levels of pain-related disability;15,16 as well as a finger-to-floor distance (FFD) evaluation in which the subject was asked to plant his/her feet firmly on the floor and lean forward as far as possible with arms outstretched to the front, and the distance between the subject’s middle finger and the floor was measured and

Materials and Methods

Study Design

This double-blind, randomized, parallel-group, prospective study was performed at UNIFESO Medical School facilities and hospital in Rio de Janeiro, Brazil from April 2016 to March 2017.

The primary objective of the study was to evaluate the tolerability of the interventions in the treatment of low back pain. The secondary objective was to evaluate the efficacy of the interventions in the treatment of low back pain.
recorded in centimeters. At each study visit, subjects were asked to rate their overall pain condition at the time of the visit on a scale of 1 to 10 points (1 point=worst evaluation and 10 points=best evaluation). The investigator also rated subject’s overall condition on a scale of 1–10 points (1 point=worst evaluation and 10 points=best evaluation) at each study visit.

At the final study visit, the investigator also evaluated overall treatment efficacy and tolerability using a four-level scale (poor, acceptable, good, and very good). Adverse event and compliance monitoring were performed throughout the study period.

The study was blinded both for subjects and investigators. There was a unique packaging and labelling for the treatment units with no identification of allocated study drug. Randomization to study drug groups was generated using random-allocation software. Randomization was performed sequentially for two groups, in blocks of 10, with a 1:1 ratio between the treatment groups.

**Study Drugs**

Subjects were randomized in order of arrival at the study center into one of two treatment groups. Group A received capsules containing active treatment: 1.5 mg uridine triphosphate trisodium (equivalent to 0.665 mg uridine)+2.5 mg cytidine monophosphate disodium (equivalent to 1.656 mg cytidine)+1000 µg hydroxocobalamin (Etna®, Laboratório Gross, Rio de Janeiro, RJ, Brazil). Group B received coated tablets containing active treatment: 100 mg thiamine+100 mg pyridoxine +5000 µg cyanocobalamin (Citoneurin®, Merck, Rio de Janeiro, RJ, Brazil). In order to maintain blinding, each subject also received dummy treatments identical to active treatment of the other group. Subjects in Group A received dummy coated tablets and subjects in Group B received dummy capsules. All subjects were instructed to take two capsules and one coated tablet, three times per day.

**Statistical Analysis**

The statistical analysis plan was approved and signed before the clinical database lock and treatment unblinding for the study. Two analysis populations were defined: the intent-to-treat (ITT) population included all randomized patients who received at least one dose of treatment; the safety population included all patients who received at least one dose of treatment. The safety population was the population evaluated for the primary (safety) endpoint, while the ITT population was the primary population for efficacy assessment.

The primary objective of this study was to evaluate the tolerability of the interventions in the treatment of low back pain. The primary endpoint was the percentage of subjects in each treatment group presenting with adverse events. The number of subjects included in this study was calculated taking into account treatment-related adverse event (AE) incidence reported from previous studies performed by our group with both combinations, with estimated AE rate difference of −0.12 between treatments (0.14 in nucleotides+B12 treatment vs 0.26 in B vitamin combination treatment groups), with the null hypothesis that the proportion of subjects presenting AEs would be identical in both populations. The criteria for significance (alpha) was determined at 0.050 and the test was two-tailed. With a sample size of 232 subjects per group, the study had a power of 90.1% to generate a statistically significant result, with a proportional between-group difference estimated at ±0.07 (95%CI: −0.19, −0.05). With an estimated rate of 37% major deviations, the total number of subjects to be included was 317 per group.

The secondary endpoints analyzed in the ITT population at each visit were RMQ scores (percentage of subjects in each treatment group presenting improvement ≥5 points), VAS pain scores at Visit 2 and Visit 3 in relation to pretreatment values, and FFD (percentage of subjects in each treatment group presenting improvement of ≥3cm in FFD in relation to pretreatment values).

Data analysis was performed using the software GraphPad Prism version 8.4.3 for Windows (GraphPad Software, San Diego, California USA, [www.graphpad.com](http://www.graphpad.com)). Baseline and demographic data were compared between groups using unpaired t-test or Fisher’s exact test for continuous and categorical variables, respectively. Primary endpoint analysis was performed using the Fisher’s exact test. Efficacy evaluations (VAS, FFD, and RMQ) were analyzed for the ITT population using mixed effects analysis followed by Tukey’s multiple comparisons test. For comparisons of categorical variables, we used the chi-squared or Fisher’s test, while continuous variables were analyzed using the Student’s t-test or mixed effects analysis for repeated measures.
Results
Disposition of Patients
A total of 1073 subjects were screened and 436 individuals not meeting criteria inclusion were excluded. Six hundred and thirty-four subjects were randomized: 317 in Group A and 317 in Group B (ITT population and safety population) (Figure 1). Early study discontinuations occurred for 28 subjects in Group A and 48 subjects in Group B, with 289 subjects in Group A and 269 subjects in Group B completing the study treatment.

Figure 1 Flow chart of subjects through the study.
Table 1  Characteristics of Patients at Baseline (n=634)

| Variables                              | Group A (n=317) | Group B (n=317) | Total (n=634) |
|----------------------------------------|----------------|----------------|--------------|
| Age (years), mean (SD)                 | 42.5 (11.0)    | 40.98 (9.8)    | 41.7 (10.5)  |
| Gender, n (%)                          |                |                |              |
| Male                                   | 191 (60.3)     | 178 (56.2)     | 369 (58.2)   |
| Female                                 | 126 (39.7)     | 139 (43.8)     | 265 (41.8)   |
| Ethnicity, n (%)                       |                |                |              |
| Asian                                  | 3 (0.9)        | 5 (1.6)        | 8 (1.3)      |
| Caucasian                              | 131 (41.3)     | 143 (45.1)     | 274 (43.2)   |
| Black                                  | 59 (18.6)      | 66 (20.8)      | 125 (19.7)   |
| Brown                                  | 124 (39.1)     | 103 (32.5)     | 227 (35.8)   |
| Body mass index (kg/m²), mean (SD)     | 25.7 (3.2)     | 25.7 (3)       | 25.7 (3.1)   |
| Height (cm), mean (SD)                 | 169.6 (9.1)    | 168.9 (8.5)    | 169.3 (8.8)  |
| Weight (cm), mean (SD)                 | 74.5 (13.7)    | 73.5 (12.4)    | 74.0 (13.1)  |
| Heart rate (bpm), mean (SD)            | 69.2 (5.6)     | 69.2 (6.2)     | 69.2 (5.9)   |
| Respiratory rate (ipm), mean (SD)      | 15.5 (2.04)    | 15.4 (2.2)     | 15.5 (2.1)   |
| Systolic blood pressure (mmHg), mean (SD) | 122.7 (8.0)   | 121.7 (7.8)    | 122.2 (7.9)  |
| Diastolic blood pressure (mmHg), mean (SD) | 78.4 (8.5)    | 77.0 (8.9)     | 77.7 (8.7)   |

Notes: Group A: subjects treated with uridine, cytidine, and vitamin B₁₂. Group B: subjects treated with vitamins B₁, B₆, and B₁₂.
Abbreviations: bpm, beats per minute; ipm, inspirations per minute, mmHg, millimeters of mercury.

Baseline Characteristics of Patients

Demographic and baseline characteristics of the study population are presented in Table 1. Demographic and clinical data were comparable at baseline between treatment groups (with the exception of mean diastolic blood pressure which was higher in Group A and a higher

Table 2  Frequency of Adverse Events Reported in Each Treatment Group by MedDRA System-Organ-Class (SOC)

| Adverse event (SOC)                                         | Group A     | Group B     |
|-------------------------------------------------------------|-------------|-------------|
|                                                             | Visit 2 (n=31) | Visit 3 (n=62) | Visit 2 (n=46) | Visit 3 (n=80) |
| Cardiac disorders                                           | 0           | 1           | 2            | 0           |
| Ear and labyrinth disorders                                 | 1           | 3           | 6            | 2           |
| Eye disorders                                               | 0           | 0           | 0            | 0           |
| Gastrointestinal disorders                                  | 0           | 0           | 7            | 15          |
| General disorders and administration site conditions        | 2           | 3           | 1            | 9           |
| Infections and infestations                                 | 0           | 2           | 0            | 0           |
| Investigations                                              | 1           | 3           | 8            | 12          |
| Metabolism and nutrition disorders                          | 1           | 2           | 2            | 2           |
| Musculoskeletal and connective tissue disorders             | 1           | 0           | 6            | 16          |
| Nervous system disorders                                    | 5           | 10          | 15           | 16          |
| Psychiatric disorders                                       | 3           | 7           | 5            | 3           |
| Renal and urinary disorders                                 | 16          | 59          | 23           | 71          |
| Respiratory, thoracic and mediastial disorders              | 3           | 2           | 0            | 0           |
| Skin and subcutaneous tissue disorders                      | 0           | 3           | 6            | 2           |
| Vascular disorders                                          | 1           | 1           | 5            | 7           |
| **Total number of AEs**                                     | **34**      | **99**      | **86**       | **155**     |

Notes: Group A: subjects treated with uridine, cytidine, and vitamin B₁₂. Group B: subjects treated with vitamins B₁, B₆, and B₁₂. *Number of subjects reporting AEs in each group at each study visit.
number of subjects reporting previous episodes of low back pain in Group A).

**Safety**
During the study treatment period, 75 (24%) subjects in Group A and 105 subjects (33%) in Group B reported AEs. While the percentage of subjects in the safety population reporting AEs was higher in Group B, the difference between groups did not reach statistical significance ($P=0.21$). The majority of AEs reported in both treatment groups affected the gastrointestinal tract (Table 2).

A total of 374 AEs was reported during the treatment period, of which 133 (35.6%) occurred in Group A and 241 (65.4%) in Group B. The distribution of number of AEs per subject during the study treatment period showed a larger percentage of subjects in Group B presenting two or more AEs ($q^2=17.04; df=4; P=0.002$). No serious AE was reported. Two severe AEs (vomiting, headache) were reported in Group A, and 13 severe AEs were reported in Group B: headache (n=3), hypokalemia (n=2), nausea (n=2), hot flushes (n=2), muscle cramps (n=1), muscle weakness (n=1), loss of appetite (n=1), and vomiting (n=1). At least one AE possibly or probably related to study drugs was reported in 17 (5.4%) subjects in Group A and 38 (12%) subjects in Group B; despite this numerical difference, the between-group difference was not statistically significant ($P=0.13$).

While there were no AEs leading to treatment discontinuation in Group A, 24 patients in Group B presented at least one AE leading to treatment discontinuation, a statistically significant between-group difference ($q^2=24.86; df=1; P<0.0001$). These AEs were: hypokalemia (n=5); nausea (n=3); muscle cramps, headache, hyperuricemia, loss of appetite, generalized pruritus, and vertigo (n=2); in addition to progressive numbness in the hands, tactile alterations in the hands, cramps, abdominal discomfort, gastric discomfort, numbness in face, numbness in hands, muscle spasms, tingling sensation in upper limbs, facial tingling sensation, muscle weakness, gout, hypertension, insomnia, palpitations, polyuria, paraesthesia in hands, pruritus and pruritus on upper limbs (n=1) (more than one AE could be reported per patient).

**VAS Pain**
Baseline VAS pain scores were 51.4 mm (±11.4; 95%CI: 50.2, 52.7) in Group A and 52.6 mm (±11.3; 95%CI: 51.4, 53.9) in Group B. Mean VAS scores at each study visit are displayed in Figure 2. Mean VAS pain scores at Visit 2 were 8.2 mm ±9.4 (95%CI: 7.1, 9.2) in Group A, with a mean difference from pretreatment of 43.3 (95%CI: 41.5, 45.1; $P<0.0001$) while in Group B, mean Visit 2 VAS pain score was 12.6 mm ±13.2 (95%CI: 11.2, 14.1), with a mean difference from pretreatment of 39.96 (95%CI: 37.8, 42.16; $P=0.0001$). At Visit 2, the mean difference in VAS scores was higher among patients in Group A compared to that in Group B (−4.5; 95%CI: −7.2, −1.8; $P<0.0001$).

At Visit 3, mean VAS pain scores in Group A were further reduced to 3.4 mm ±5.6 (95%CI: 2.7, 4.0) in Group A, mean difference in relation to pretreatment of 48.1 (95%CI: 46.3, 49.9) and 5.5 ±7.1 (95%CI: 4.7, 6.4) in Group B, reduction from pretreatment of 47.1 (95%CI: 45.2, 49.0). VAS score reduction from pretreatment at Visit 3 was statistically significant ($P<0.0001$) for both Group A and Group B. At Visit 3, mean difference when comparing VAS score reduction from pretreatment values was higher for Group A (−2.1; 95%CI= −3.7, −0.6; $P=0.002$).

**Finger-to-Floor Distance**
Mean pretreatment FD was 28.1 cm (±7.4; 95%CI: 27.3, 28.9) in Group A and 27.3 cm (±7.5; 95%CI: 26.5, 28.1) in Group B. All treated subjects had improvement of ≥3 cm in FFD at Visit 3 in relation to pretreatment values (Table 3). Change in FFD from pretreatment was statistically significant ($P<0.001$) in both groups at 30 days (Visit 2) and 60 days (Visit 3).
Table 3 Changes in Finger-to-Floor Distance During Treatment (ITT Population)

| Treatment Group/Study Visits | Mean FFD (SD) | Mean Difference a (95%CI) | P-value b |
|------------------------------|--------------|--------------------------|-----------|
| Group A (n=289) Pretreatment |              |                          |           |
| Visit 2                      | 28.1 (7.4)   | 10.7 (9.8, 11.7)         | <0.0001   |
| Visit 3                      | 17.4 (8.0)   | 14.6 (13.5, 15.8)        | <0.0001   |
| Group B (n=269) Pretreatment |              |                          |           |
| Visit 2                      | 27.3 (7.5)   | 12.8 (11.7, 13.9)        | <0.0001   |
| Visit 3                      | 14.5 (8.2)   | 17.7 (16.6, 18.9)        | <0.0001   |

Notes: Group A: subjects treated with uridine, cytidine, and vitamin B12; Group B: subjects treated with vitamins B6, uridine, and vitamin B12. aFrom pretreatment; bTukey’s multiple comparisons test.

Abbreviation: FFD, finger-to-floor distance (cm).

Roland–Morris Questionnaire

Pretreatment RMQ scores were 14.2 (±3.6; 95%CI: 13.8, 14.6) in Group A and 13.4 (±3.8; 95%CI: 13.0, 13.8). At Visit 3, 287 (98.9%) subjects in Group A and 265 (97.8%) subjects in Group B showed improvement of ≥5 points in RMQ scores. Mean scores at Visit 2 were 2.3 points (±3.0; 95%CI: 1.9, 2.6) in Group A mean difference of 11.9 points from pretreatment (95%CI: 11.3, 12.5) and 3.3 (±3.8; 95% CI: 2.9, 3.7); in Group B mean difference of 10.1 points from pretreatment (95%CI: 9.5, 10.8).

A further score improvement at Visit 3 was observed in both treatment groups, with Group A mean score of 0.9±1.8 (95%CI: 0.7, 1.1) and Group B mean score of 1.3±2.3 (95%CI: 1.1, 1.6). Visit 3 score mean differences in relation to pretreatment values were 12.8 points (95%CI: 12.1, 13.6) in Group A and 12.1 points (95%CI: 11.4, 12.7) in Group B.

Figure 3 Distribution of RMQ score changes from pretreatment at Visits 2 and 3.

Notes: Graph displaying percentage of subjects in each treatment group with worsening, no change, and improvement (categorized by <5, 5–10 points, 11–15 points, 16–20 points and >20 points).

Abbreviations: RMQ, Roland–Morris Questionnaire; V2, Visit 2 (after 30 days of treatment); V3, Visit 3 (after 60 days of treatment).
Score improvements within each treatment group were statistically significant ($P<0.001$) at Visit 2 and Visit 3 in relation to pretreatment scores in both treatment groups. When comparing scores between treatment groups, there was a greater RMQ score reduction in Group A at Visit 2 ($P=0.005$) but not at Visit 3 ($P=0.09$) in relation to Group B scores (Figure 3).

Compliance to Treatment
Drug compliance data was available for 589 subjects at Visit 2, with median compliance of 103.0% for tablets (95%CI: 103.5–104.1) and 103.0% for capsules (95%CI: 103.3–103.8). At Visit 3, median compliance (data from 558 subjects) was 100.0% (95%CI: 99.8–100.1) for tablets and 100.0% for capsules (95%CI: 99.7–100).

Patient Assessment of Overall Pain Condition
Pretreatment patient overall assessment scores were comparable between treatment groups ($\chi^2=2.9; df=3; P=0.96$). At Visits 2 and 3, the scores of this assessment improved within each treatment group ($P<0.0001$ for both groups), with no statistically significance between treatment groups in percentage of subjects presenting improvement ($P=0.06$ at Visit 2 and $P=0.54$ at Visit 3).

Physician Assessment of Overall Condition
Physician overall assessment scores were comparable between treatment groups at pretreatment ($P=0.72$). At Visits 2 and 3, the scores of this assessment improved within each treatment group ($P<0.0001$ for both groups), with no statistically significance between treatment groups in percentage of subjects presenting improvement ($P=0.06$ at Visit 2 and $P=0.33$ at Visit 3).

Investigators Assessment of Overall Treatment Efficacy and Tolerability
The global investigator evaluation of efficacy and tolerability was comparable for the two treatment groups. Overall, efficacy was rated as good-very good during the study by a large majority of investigators: 94% in Group A and 90% in Group B ($P=0.36$).

Discussion
We report here on the first comparative study of the combination of nucleotides plus vitamin B₁₂, vs the combination of vitamins B₁, B₆, and B₁₂ over a 60-day treatment period among patients with low back pain. The two treatments were comparable in the safety evaluation (primary study endpoint) as assessed by AE incidence among treated subjects. However, the number of AEs per subject was significantly higher among patients treated with the combination of vitamins B₁, B₆, and B₁₂, and only in the B vitamin combination group was there interruption of treatment due to AEs. No serious AEs were reported during the treatment period.

Previous studies in smaller populations assessing these combinations utilized shorter treatment periods of up to 30 days, however the longer treatment period employed in the present study does not appear to have resulted in a higher rate of AE occurrence. Comparing the results of the present study with those of the study by Goldberg et al, total AE rate in that study among 200 subjects who received the nucleotide+vitamin B₁₂ combination over 30 days of treatment was 19%, while total AE rate among subjects receiving the same treatment in the present study over 60 days was 24% ($P=0.49$).

The AEs observed during treatment were in keeping with the known safety profile of both treatments, with no previously unreported treatment-related AEs registered throughout the treatment period in either treatment group. Gastrointestinal AEs were the most frequently reported AEs in both treatment groups, in keeping with the reported AE profile of the previous studies employing these combinations. Known side effects associated with vitamin B₁₂ administration include allergic hypersensitivity reactions, with skin eruptions, rash and pruritus, hypokalemia, fever, chills, hot flushing, dizziness, malaise, and tremor has also been reported. Previously reported side effects of thiamine include skin alterations (discoloration, hives, pruritus), hyperhidrosis, restlesslessness, and gastrointestinal symptoms, especially nausea. Known side effects of pyridoxine include gastrointestinal symptoms, numbness and tingling, headache, sleepiness, and sensory nerve damage (following long-term use of high doses).

In our study, subjects in both treatment groups displayed significant improvement in all low back pain-related evaluation measures in relation to pretreatment status. VAS pain scores decreased significantly after 30 days of treatment within both treatment groups, with a greater score reduction among subjects in Group A compared to those in Group B, although this between-group difference was not observed after 60 days of treatment. The improvement in pain intensity as assessed by the VAS observed at Visit 2 was maintained at Visit 3, with further significant decrease in VAS scores at Visit 3. There was a significant improvement in functionality when assessed by...
the RMQ throughout the treatment period, with greater score reductions observed among subjects treated with the combination of nucleotides + vitamin B12. The positive effect on mobility (FFD) seen after 30 days of treatment was maintained or further improved at 60 days of treatment within both treatment groups. The same was observed for pain intensity, VAS and RMQ scores. The patient and investigator global assessments also improved significantly throughout the treatment period. These observations support the continued treatment for the period of 60 days among patients with low back pain.

The use of nucleotides and vitamin B12 employed to treat low back pain and other painful conditions has been the subject of continuing investigations, with encouraging results from previous studies. In patients with compression neuralgia associated with degenerative orthopedic alterations of the hip and spine, treatment with the combination of nucleotides and vitamin B12 resulted in significant pain improvement and increased functionality. In a study evaluating the use of the combination of nucleotides and vitamin B12 in patients with alcoholic polyneuropathy, pain reduction and improvement in motor coordination was reported following intramuscular injection of the combination for six days followed by 30-day oral treatment. In a previous study of peripheral neuropathies including low back pain, the same combination of nucleotides and vitamin B12 utilized in the present study was shown to be superior to vitamin B12 monotherapy in pain relief as assessed by a 100 mm VAS during a 30-day treatment period.

While the pyrimidine nucleotides cytidine and uridine lack direct analgesic or anti-inflammatory action, their contribution to symptomatic improvement in conditions involving the peripheral nervous system has been evidenced in preclinical studies following experimentally induced nerve damage. In animal models of induced crush injury, administration of nucleotides was shown to accelerate both axonal and myelin sheath repair of regenerating nerve and muscle fibers, with improvements in nerve fiber conduction velocity. Results of further preclinical investigations have indicated increased nerve conduction velocity, increased levels of neuritic protein neurofilaments, increased axon myelin surface area and thickness, and increased levels of neuronal cell membrane phospholipids phosphatidylcholine and phosphatidylethanolamine following nucleotide administration. More recently, Chavushyan et al reported increased excitatory and inhibitory responses in rats following sciatic nerve crush treated with the same combination of nucleotides and B12 employed in the present study, with recovery of motor and sensory function comparable to uninjured limbs after 30 days of treatment.

A salvage pathway is one way in which nucleotides (purine and pyrimidine) are synthesized from intermediates in the degradative pathway for nucleotides. Cytidine and uridine are absorbed by cells, and cytidine is deaminated to uridine. Uridine is phosphorylated to form uracil nucleotides by uridine kinase, a monophosphate kinase—specific for uridine monophosphate (UMP), CMP and dCMP—and an unspecific nucleoside diphosphokinase. Salvage pathway-synthesized UTP may preferably be used for RNA synthesis in cell nuclei and may contribute to a cytosolic free UTP input along with de novo pathway UTP. Free UTP can provide UTP to be stored in granules, or released to act on pyrimidine cell surface receptors.

The mechanisms underlying the therapeutic effects observed following uridine and cytidine administration appear to stem from the nucleotides’ impact on the purine P2Y receptors P2Y1 and P2Y2, which in turn have been shown to mediate nucleotide-induced sensory neuron excitation via inhibition of voltage-gated Kv7-family potassium channels via increased intracellular Ca\(^{2+}\), along with facilitation of TRPV1 vanilloid receptors in the dorsal root ganglia, via activation of protein kinase C. Additionally, purine nucleotides have been shown to induce activation of a1AR postsynaptic adrenergic receptors, which are involved in regulation of synaptic transmission, plasticity, and motor activity.

In both previous studies of the combination of nucleotides and vitamin B12 employing vitamin B12 monotherapy as control, improvement in pain and functionality was reported in the control group, which highlights the importance of vitamin B12 in the formulation. Use of intramuscular vitamin B12 monotherapy was evaluated by Mauro et al in the treatment of low back pain in patients presenting mechanical or irritative lumbago, reporting that after a two-week treatment period consisting of once-daily 1000 µg vitamin B12 injections, treated subjects experienced a significant decrease in pain and disability that was statistically superior to the placebo-treated control group. Vitamin B12 has also shown clinical efficacy in the treatment of diabetic neuropathy, demonstrating improvement in pain, paraesthesia, somatosensory and autonomic symptoms, as well as nerve conduction velocity.

In the nervous system, Vitamin B12 plays an essential role in myelin synthesis, nerve metabolism, and neuronal regeneration. In preclinical studies performed with rats, vitamin B12 has been demonstrated to exert an effect on nociception and pain response, with accelerated compound
The nucleotides+vitamin reduction continued the statistical once muscle action potential recovery and dose-dependent reduction of tactile allodynia reported. The proposed mechanisms of action of vitamin B₁₂ in pain relief include: promotion of nerve regeneration and/or remyelination by accumulation of exogenous B₁₂, selective blockade of sensory nerve conduction, promotion of protein synthesis necessary for nerve regeneration via upregulation of gene transcription.

Vitamins B₁, B₆, and B₁₂ belong to the family of neurotropic B vitamins, playing important roles in the maintenance of nervous system function, and deficiencies of which may result in peripheral nervous system disorders. Furthermore, evidence from both preclinical and clinical studies has shown that the combination of these B vitamins is effective in the treatment of neuropathic pain and symptoms of neuropathic conditions, even in the absence of deficiencies in these B vitamins.

The strong points of this study are mainly the 60-day study duration, the large number of patients and the assessments performed in the context of low back pain. The major study weakness is sponsorship by the manufacturer of the test product. Authors acknowledge potential for bias in sponsored studies. Given the more favorable safety with no subjects in the nucleotides+vitamin B₁₂ treatment group discontinuing treatment due to AEs, comparative clinical studies employing this combination in other types of peripheral neuropathies may also be of interest, as treatment periods of 60 days or more with therapeutic doses of B vitamins are often employed in these conditions.

**Conclusion**

In conclusion, while a larger percentage of subjects presented AEs in Group B, this difference did not reach statistical significance. However, >60% of total AEs occurred in Group B, a higher percentage of subjects in Group B presented two or more AEs during treatment, and AEs causing treatment discontinuation occurred only in the B vitamin combination group. Pain intensity (VAS) reduction was superior at 30 days of treatment in the nucleotides+vitamin B₁₂ group and equivalent between groups at 60 days of treatment. Improvements in efficacy measures RMQ and FFD were observed in both groups. The additional improvement in pain, mobility, and functionality observed at the final study visit support the continued treatment for the period of 60 days among patients with low back pain.

**Data Sharing Statement**

The authors agree to share study data upon request by e-mail to the corresponding author, via e-mail (maurogeller@gmail.com), to investigators whose proposed use of the data has been approved by an independent review committee or ethical committee and who agree to sign a data access agreement. Data to be shared includes individual deidentified participant data and study-related documents (ie, study protocol, informed consent text, or other study documentation requested), for a period of five years from the date of study publication.

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**Disclosure**

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