Objective: This study assessed the effects of 32 mg naltrexone sustained release (SR)/360 mg bupropion SR (NB) on body weight in adults with obesity, with comprehensive lifestyle intervention (CLI), for 78 weeks.

Methods: In this phase 3b, randomized, open-label, controlled study, subjects received NB + CLI or usual care (standard diet/exercise advice) for 26 weeks. NB subjects not achieving 5% weight loss at week 16 were discontinued, as indicated by product labeling. After week 26, usual care subjects began NB + CLI. Assessments continued through week 78. The primary end point was percent change in weight from baseline to week 26 in the per protocol population. Other end points included percentage of subjects achieving ≥5%, ≥10%, and ≥15% weight loss, percent change in weight at week 78, and adverse events (AEs) necessitating study medication discontinuation.

Results: NB + CLI subjects lost significantly more weight than usual care subjects at week 26 (8.52% difference; P < 0.0001). Weight loss persisted through 78 weeks. In total, 20.7% of subjects discontinued medication for AEs, including 7.0% for nausea.

Conclusions: Treatment with NB, used as indicated by prescribing information and with CLI, significantly improved weight loss over usual care alone. NB-facilitated weight loss was sustained for 78 weeks and was deemed safe and well tolerated.

Introduction

Obesity is a public health epidemic affecting more than one third of adults in the United States (U.S.); over two thirds of U.S. and half of European adults are overweight or at risk of developing obesity (1-4). Obesity-related complications such as type 2 diabetes, heart disease, stroke, and certain cancers are associated with decreased life expectancy and costs totaling up to $147 billion annually in the U.S. (5-7). Despite these health and economic consequences, many individuals are unable to combat obesity through diet and exercise alone, and few effective drugs are available to assist weight loss (5,8).

Appetite and eating behavior are controlled by several neural pathways in the brain (9). One key pathway involves pro-opiomelanocortin (POMC) neurons in the hypothalamus; activation of these neurons is thought to reduce hunger and therefore food intake. Bupropion has been demonstrated to increase POMC activation (10-13), while naltrexone lessens auto-inhibition of the POMC pathway by endogenous opioids, leading to a synergistic activation (14-16). Furthermore, both bupropion and naltrexone are thought to influence the mesolimbic dopaminergic reward pathway, which normally contributes to the reinforcement of behaviors, including feeding (9,17); the combination of naltrexone and bupropion (NB) may therefore also reduce reward-driven food intake.

Combination NB sustained release (SR), referred to as extended release (ER) in the U.S. and South Korea and prolonged release (PR) in Europe, has been evaluated for weight loss in four pivotal, placebo-controlled, double-blind, randomized phase 3 clinical trials (18-21). In each of these studies, NB was well tolerated and associated with significant weight loss (approximately 4%-5% greater mean weight loss compared with subjects receiving placebo and participating in the...
same lifestyle modification programs) and improvements in various secondary measures of cardiovascular risk. NB therapy was approved for the chronic treatment of obesity by the Food and Drug Administration in the U.S. in 2014 (Contrave®), by the European Medicines Agency in the European Union in 2015 (Mysimba®), and by the Ministry of Food and Drug Safety in South Korea in 2016 (Contrave) as an adjunct to lifestyle modification. Product labeling states that patients are to be evaluated after 16 weeks of treatment to determine whether they have lost at least 5% of initial body weight, and, if not, medication use should be discontinued. The present effectiveness study was designed to determine weight loss achieved when NB is used in a manner consistent with labeling in a real-world setting, including the use of a commercially available comprehensive lifestyle intervention (CLI) program. As the weight loss efficacy of NB compared with placebo in addition to lifestyle intervention programs of differing intensities was already demonstrated (19,21), in the current open-label placebo in addition to lifestyle intervention programs of differing intensities was already demonstrated (19,21), in the current open-label study the NB + CLI regimen was compared with general advice that patients might receive from their physician (usual care). While this design does not allow for individual assessment of the contribution of NB and CLI to weight loss, it does reflect the overall outcome that would be expected to occur when NB is prescribed according to product labeling and allows for comparison of those outcomes with what might be expected if a physician provides more general weight loss advice and support. Finally, this study examined the effects of NB for up to 78 weeks, longer than the previous phase 3 trials.

Methods
Study design
This was a phase 3b, multicenter, randomized, open-label, controlled trial designed to assess the effects of 32 mg naltrexone SR/360 mg bupropion SR (NB) in conjunction with a CLI program, compared with usual care (diet and exercise education and recommendations from the study site) on body weight for 26 weeks in subjects with obesity or overweight and dyslipidemia and/or controlled hypertension. Key exclusion criteria included: type 1 or 2 diabetes mellitus; myocardial infarction within 6 months before screening; angina pectoris grade III/IV (per the Canadian Cardiovascular Society grading scheme); clinical history of large vessel cortical strokes, including ischemic or hemorrhagic strokes (i.e., transient ischemic attack was not exclusionary); history (within 20 years before screening) of seizures, cranial trauma, bulimia, anorexia nervosa, or other conditions that predispose subjects to seizures; chronic use or positive screen for opioids; psychiatric conditions including mania, psychosis, acute depressive illness, or suicide risk; regular use of tobacco products.

Subjects and eligibility criteria
Adult male and female subjects, aged 18 to 60 years, had either obesity (body mass index [BMI] 30-45 kg/m²) or overweight (BMI 27-45 kg/m²) with dyslipidemia and/or controlled hypertension. Key exclusion criteria included: type 1 or 2 diabetes mellitus; myocardial infarction within 6 months before screening; angina pectoris grade III/IV (per the Canadian Cardiovascular Society grading scheme); clinical history of large vessel cortical strokes, including ischemic or hemorrhagic strokes (i.e., transient ischemic attack was not exclusionary); history (within 20 years before screening) of seizures, cranial trauma, bulimia, anorexia nervosa, or other conditions that predispose subjects to seizures; chronic use or positive screen for opioids; psychiatric conditions including mania, psychosis, acute depressive illness, or suicide risk; regular use of tobacco products.

Lifestyle intervention programs
CLI was a telephone- and Internet-based program that included a progressive nutrition and exercise program with individualized goal setting and tracking tools. Subjects were to receive up to 11 structured telephone calls from a coach/dietician during the first 26 weeks (controlled treatment period), with up to 12 additional calls offered over the remainder of the 78 weeks (uncontrolled treatment period). Usual care was a site-based lifestyle intervention program intended to mimic the weight loss support that might typically be provided to patients in a primary care setting. Usual care subjects were instructed at baseline and at week 10 to follow an exercise prescription and a hypocaloric diet, with a target daily caloric intake deficit of 500 kcal, and were given support tools such as a nutrition tracker, pedometer, and healthy weight literature.

Controlled treatment period (week 1-week 26)
Following a screening period, subjects were randomized in a 1.75:1 ratio to receive either open-label active study medication (NB) along with CLI (NB + CLI) or usual care for 26 weeks (Figure 1). In accordance with prescribing information, subjects randomized to NB + CLI initiated treatment with NB at a daily dose of 8 mg naltrexone/90 mg bupropion and escalated their dose over the subsequent 3 weeks. In addition, an evaluation of weight loss was performed in the NB + CLI group at week 16, with subjects discontinued from NB if they had not lost at least 5% of their baseline body weight. Additionally, subjects were to discontinue NB if they experienced increases in systolic or diastolic blood pressure of ≥10 mm Hg at both week 10 and week 16.

Uncontrolled treatment period (week 26-week 78)
After the controlled treatment period of 26 weeks, subjects in the NB + CLI group continued their open-label NB use and participation in the CLI program. Subjects in the usual care group were switched to open-label NB use and began participation in the CLI program. These subjects also underwent a weight loss and blood pressure evaluation 16 weeks after beginning NB use (at week 42), and treatment was discontinued, if necessary, based on the same criteria used for subjects originally assigned to NB + CLI.

Sample size determination
Based on the predicted difference in discontinuation rates and effect of a week 16 weight assessment, a randomization ratio of 1.75:1 was adopted. A total of 242 subjects were sufficient to provide ≥90% power to detect a significant difference (α = 0.05) between the treatment groups at week 26 for the per protocol (PP) population using a two-sample t-test assuming a true mean difference of 0.6 common standard deviation (SD). The assumptions are based on data from the NB phase 3 trials and publications pertaining to usual care for obesity (18-23).

Study end points
The primary end point was the percent change in body weight from baseline (day 1) to week 26, while the secondary end points were percentage of subjects achieving a loss of at least 5%, 10%, or 15% of baseline body weight at week 26. Additional study end points included changes in body weight, vital signs, and obesity-related risk factors from baseline to post-baseline visits (both before and after week 26). Patient-reported outcomes concerning weight-related quality of life, binge eating, and sexual function were assessed but are not reported in this article.
Statistical analysis

The primary analysis of the primary end point was based on the adjusted least-squares (LS) means estimated using analysis of covariance (ANCOVA), using the week 26 PP population (defined as modified intent-to-treat [mITT] subjects in compliance with the protocol in the controlled treatment period), with no imputation of missing data. As sensitivity analyses, the primary end point was also analyzed for the mITT population (defined as randomized subjects with at least one post-baseline body weight measurement while still on treatment) using mixed model repeated measures (MMRM) and last observation carried forward (LOCF). The week 78 analysis of percent change in body weight was conducted similarly with the primary analysis being ANCOVA in the week 78 PP population (subjects in compliance with the protocol throughout the study) and numerous sensitivity analyses.

Categorical analysis of number and percentage of subjects in the week 26 PP population achieving a loss of at least 5%, 10%, or 15% body weight change was conducted.
of baseline body weight at week 26 was assessed using a logistic regression model with a factor for treatment group and baseline BMI category with baseline body weight as a covariate. For treatment comparisons, odds ratios, 95% confidence intervals (CI) for the odds ratios, and P values are reported. Analyses were repeated at week 78 in the week 78 PP population. Other continuous variables were analyzed similarly to the primary end point. Data were only descriptively summarized beyond the controlled treatment period.

All statistical tests were performed two-sided at the 0.05 level of significance. Adverse events (AEs) were summarized in the ITT population and vital signs were collected for the week 78 PP population. All statistical analyses were performed using SAS® software (Version 9.3, SAS Institute Inc., Cary, NC).

Results

Baseline characteristics and subject disposition

A total of 242 CLI subjects were randomized and treated (ITT): 153 subjects to the NB + CLI group and 89 subjects to the usual care group (Figure 1). Subject demographics are summarized in Table 1. Of the 242 randomized subjects, 153 subjects (71 in the NB + CLI group and 82 in the usual care group) completed the 26-week controlled treatment period. The most common reason for discontinuation from the NB + CLI treatment regimen within the first 26 weeks was AEs (n = 35 of 153, 22.9%), followed by discontinuation due to the week 16 assessment (n = 32 of 153; 20.9% [n = 29 did not satisfy the weight criteria alone, n = 1 had blood pressure elevations that met the criteria for discontinuation alone, and n = 2 were discontinued due to both weight and blood pressure criteria]). The most common reason for discontinuation of usual care treatment was lost to follow-up (n = 5, 5.6%).

During the uncontrolled treatment period, the most common reason for discontinuation of the usual care/NB + CLI group was the week 42 assessment (n = 26 of 89, 29.2%). Of subjects from the original NB + CLI group, n = 16 discontinued after week 26. No subjects discontinued NB due to an AE with an onset after the first 26 weeks of NB use.

Treatment with NB + CLI led to greater weight loss than usual care

Subjects in the PP population treated with NB + CLI demonstrated a significant decrease in body weight compared with those in the usual care group after 26 weeks (9.46% reduction in the NB + CLI group; 0.94% reduction in the usual care group; 8.52% difference, P < 0.0001) (Figure 2A). When all NB + CLI subjects were included in the analysis using MMRM or LOCF methodology, including those who did not achieve weight loss of 5% at week 16, the reduction in the NB + CLI group relative to usual care at 26 weeks was 5.17% (MMRM) and 4.15% (mITT-LOCF) (P < 0.0001 in both sensitivity analyses). Changes in weight over the first 16 weeks were analyzed in subjects who achieved 5% weight loss at week 16 (responders) compared with those who did not achieve this threshold (nonresponders, including those who discontinued study medication before week 16) (Figure 2B). This figure demonstrates that the weight loss curves of these two populations separated at the first time point assessed (2 weeks).

Consistent with the greater percentage body weight reduction observed in the NB + CLI group, significantly more NB + CLI subjects achieved each weight loss threshold than usual care subjects at week 26 (84.5% NB + CLI vs. 12.2% usual care had ≥5% weight loss [P < 0.0001]; 42.3% NB + CLI vs. 3.7% usual care had ≥10% weight loss [P < 0.0001]; 12.7% NB + CLI vs. 0% usual care had ≥15% weight loss [P value cannot be determined]) (Table 2), indicating that while there are some subjects who are able to achieve these categorical weight loss values even with very basic advice, the combination of pharmacotherapy and more consistent lifestyle instruction dramatically increases a patient’s likelihood of success.

TABLE 1  Subject demographic and baseline characteristics

| Parameter                  | Usual care/ NB + CLI | NB + CLI |
|----------------------------|----------------------|----------|
| Age (y), mean (SD)         | 47.0 (9.98)          | 46.1 (9.66) |
| Sex, (% female)            | 86.5                 | 81.7     |
| Race (%)                   |                      |          |
| White                      | 71.9                 | 81.0     |
| Black or African American  | 27.0                 | 18.3     |
| Asian                      | 0.0                  | 0.7      |
| American Indian or Alaska Native | 1.1                 | 0.0      |
| Ethnicity (%)              |                      |          |
| Hispanic or Latino         | 5.6                  | 2.6      |
| Not Hispanic or Latino     | 94.4                 | 97.4     |
| Weight (kg), mean (SD)     | 100.2 (16.58)        | 101.4 (15.09) |
| BMI (kg/m²), mean (SD)     | 36.26 (4.36)         | 36.33 (4.20) |
| BMI category (%)           |                     |          |
| <30 kg/m²                  | 5.6                  | 2.0      |
| <35 kg/m²                  | 46.1                 | 45.8     |
| ≥35 to <40 kg/m²           | 31.5                 | 32.0     |
| ≥40 kg/m²                  | 22.5                 | 22.2     |
| Waist circumference (cm), mean (SD) | 111.9 (11.91) | 112.2 (11.23) |
| Triglycerides (mg/dL), mean (SD) | n = 86       | n = 147 |
| LDL cholesterol (mg/dL), mean (SD) | n = 84     | n = 147 |
| HDL cholesterol (mg/dL), mean (SD) | n = 84     | n = 147 |
| Glucose (mg/dL), mean (SD) | n = 86              | n = 147 |
| Insulin (uIU/mL), mean (SD) | n = 84            | n = 147 |
| HOMA-IR, mean (SD)         | 17.3 (10.68)         | 19.5 (19.65) |
| Systolic blood pressure (mm Hg), mean (SD) | 120.6 (11.41) | 123.7 (9.51) |
| Diastolic blood pressure (mm Hg), mean (SD) | 78.8 (7.60)  | 80.4 (7.26) |
| Heart rate (bpm), mean (SD) | 69.8 (6.68)       | 70.9 (8.66) |

BMI, body mass index; CLI, comprehensive lifestyle intervention; HDL, high-density lipoprotein; HOMA-IR, homeostasis model assessment of insulin resistance; LDL, low-density lipoprotein; NB, naltrexone/bupropion; SD, standard deviation.
After week 26, when the subjects in the usual care group began treatment with NB + CLI, a similar pattern of weight loss was observed as previously observed in those initially randomized to NB + CLI (Figure 3A), with approximately 90% of the maximum weight loss in each cohort observed over the first 6 months of NB treatment. Changes in weight at week 78 were similar between those who received 52 and 78 weeks of NB + CLI treatment. The change in body weight from baseline to week 78 for each subject who remained in the study is also presented graphically in Figure 3B, highlighting the range of weight changes observed in the population, even when the week 16 criteria was used to select for responders.

### Obesity-related risk factors

In addition to the greater reductions in weight observed with NB + CLI compared with usual care at week 26, subjects in the NB + CLI group had significantly greater reductions in triglycerides, waist circumference, glucose, insulin, and a measure of insulin resistance, as well as a significantly greater increase in high density lipoprotein cholesterol (Table 3). At week 78, these changes were generally maintained in subjects who received long-term treatment with NB + CLI, as shown by the pooled data from all subjects.

### Safety results

Safety and tolerability of NB was thoroughly studied in the phase 3 trials, therefore only AEs leading to discontinuation of study medication and serious AEs (SAEs) were collected. AEs leading to NB discontinuation were similar between subjects who initially were randomized to NB + CLI and to subjects originally randomized to usual care who began treatment with NB after week 26; data are therefore presented for the two groups combined in Table 4. Gastrointestinal and neuropsychiatric side effects, known to be associated with NB pharmacology, contributed to 82% of AEs in both treatment groups and tended to occur within the first 4 weeks of treatment. The most frequent AEs that led to discontinuation of NB for the two groups combined included nausea (7.0%), anxiety (2.1%), headache (1.7%), dizziness (1.2%), and insomnia (1.2%). All AEs leading to discontinuation from the NB + CLI group had onset dates before week 26. Two subjects experienced SAEs that were deemed not related to NB by the investigator (breast cancer, one case diagnosed 3 months into NB treatment and the other 1 year after NB treatment had been discontinued).

Systolic and diastolic blood pressure and heart rate were collected over time to further evaluate the safety of NB treatment. Mean (standard error, SE) systolic blood pressure decreased by 2.2 (2.0) and 2.7 (1.4) mm Hg from baselines of 120.9 and 126.0 mm Hg in the usual care (N = 28) and NB + CLI (N = 55) groups, respectively, at week 78. Mean (SE) diastolic blood pressure decreased by 0.3 (1.6) and 1.7 (1.0) mm Hg from baselines of 78.5 and 80.9 mm Hg,

### Table 2 Subjects who achieved a weight loss of at least 5%, 10%, or 15% of body weight at week 26

| Weight Loss | Usual care (N = 82) | NB + CLI (N = 71) |
|-------------|---------------------|------------------|
| ≥5% weight loss at week 26, n (%) | 10 (12.2) | 60 (84.5) |
| Odds ratio (95% CI) | 44.0 (16.6 to 116.3) | 0.43 (0.14 to 1.28) |
| P | <0.0001 | 0.07 |
| ≥10% weight loss at week 26, n (%) | 3 (3.7) | 30 (42.3) |
| Odds ratio (95% CI) | 21.4 (6.0 to 76.7) | 2.7 (1.2 to 6.1) |
| P | <0.0001 | 0.025 |
| ≥15% weight loss at week 26, n (%) | 0 (0.0) | 9 (12.7) |
| Odds ratio (95% CI) | N/D | N/D |
| P | N/D | N/D |

P value for testing the null hypothesis that the odds ratio equals 1 from a logistic regression model with a factor for treatment group and for baseline body mass index category and baseline body weight as a covariate.

CI, confidence interval; CLI, comprehensive lifestyle intervention; NB, naltrexone/bupropion; N/D, not determined.
Discussion

In this randomized, open-label study, treatment with 32 mg naltrexone SR/360 mg bupropion SR (NB + CLI) as indicated by its prescribing information was shown to result in approximately 10% weight loss from baseline over a 26-week period, which was approximately 8.5% more than was observed with usual care, a diet and exercise advice program designed to reflect what a patient might receive in a primary care setting. In this study, subjects who did not achieve at least 5% body weight reduction by 16 weeks of use (i.e., those who did not respond to treatment) or had a sustained elevation in blood pressure were discontinued from study medication. In those who remained on medication through 78 weeks, longer than the observation periods of previous NB studies (56 weeks) (18-21), the initial weight loss observed at 26 weeks was sustained.

This study, conducted to approximate the real-world experience, is consistent with and builds on the results of the previous NB phase 3 trials, all of which demonstrated that NB used as an adjunct to lifestyle modification is associated with weight loss of approximately 4% to 5% greater than seen with placebo subjects receiving the same lifestyle intervention in subjects with overweight or obesity (18-21). The previous studies did not prospectively utilize an efficacy assessment at week 16 of NB treatment and were limited to a 56-week duration. This study assessed the effects of NB in the population of subjects who responded to the medication by losing at least 5% of their initial body weight by 16 weeks, and demonstrated that NB + CLI associated weight loss were maintained over 78 weeks. The safety profile was also consistent with previous studies: most subjects tolerated NB well, and those who developed AEs did so early in the treatment protocol. The most common AE leading to NB discontinuation was nausea (7.0% of all subjects), which is consistent with the rate in the phase 3 trials (6.3%). While the mechanism by which NB causes nausea is unknown, it was demonstrated in the phase 3 trials that nausea tends to occur early in treatment, is primarily mild to moderate in severity, and is not a major contributor to weight loss (24). Only two AEs led to discontinuation in the NB + CLI group after week 26 (both with AE onset before week 26), and no AEs necessitating discontinuation had an onset date during the extended time period (weeks 52-78). These results strengthen the body of evidence suggesting combination therapy of NB along with lifestyle to promote weight loss is a promising approach to lowering the prevalence of obesity.

| TABLE 3 Changes in obesity-related risk factors |
|-----------------------------------------------|
| Usual care (N = 82)                           |
| NB + CLI (N = 71)                             |
| P                                             |
| LS mean change (SE) from baseline at week 26  |
| Waist (cm)                                    |
| −1.6 (0.66)                                   |
| −7.0 (0.71) < 0.0001                          |
| Triglycerides (mg/dL)                         |
| 2.8 (4.63)                                    |
| −13.6 (4.96) 0.0019                           |
| HDL cholesterol (mg/dL)                       |
| 0.1 (0.73)                                    |
| 4.1 (0.77) 0.0001                             |
| LDL cholesterol (mg/dL)                       |
| −1.9 (2.11)                                   |
| −2.0 (2.20) 0.9686                            |
| Fasting plasma glucose (mg/dL)                |
| 1.6 (0.96)                                    |
| −2.9 (1.04) 0.0016                            |
| Fasting insulin (μIU/mL)                      |
| −3.4 (0.76)                                   |
| −7.5 (0.79) 0.0004                            |
| HOMA-IR                                       |
| −0.8 (0.19)                                   |
| −2.0 (0.20) 0.0003                            |
| Mean change (SE) from baseline at week 78     |
| (all subjects with week 78 data) (N = 83)      |
| Waist (cm)                                    |
| −7.5 (0.81)                                   |
| Triglycerides (mg/dL)                         |
| −12.4 (8.17)                                  |
| HDL cholesterol (mg/dL)                       |
| 7.0 (0.94)                                    |
| LDL cholesterol (mg/dL)                       |
| 0.3 (2.32)                                    |
| Fasting plasma glucose (mg/dL)                |
| −0.8 (1.10)                                   |
| −7.9 (2.51)                                  |
| Fasting insulin (μIU/mL)                      |
| −1.9 (0.67)                                   |
| P value for ANCOVA LS mean difference between treatment groups. CLI, comprehensive lifestyle intervention; HDL, high-density lipoprotein; HOMA-IR, homeostasis model assessment of insulin resistance; LDL, low-density lipoprotein; LS, least squares; SE, standard error.
The evaluation for body weight reduction of 5% at week 16 (after 12 weeks at full dose NB) is in line with product labeling. The recommendation to discontinue due to sustained blood pressure increases of 10 mm Hg at week 16 in the present study was more specific than product labeling, which recommends monitoring at regular intervals and discontinuing NB if concerns about increased blood pressure are present. In this real-world setting, 67% (72 of 107) of NB subjects who received study medication for 16 weeks responded to the medication per the evaluation criteria used in the present study. Similarly, 57% (37 of 65) of usual care/NB + CLI subjects who received study medication for 16 weeks met the early evaluation threshold. The overwhelming majority of discontinuations at week 16 were due to insufficient weight loss relative to baseline (more than 90% of the discontinuations in each treatment group) rather than adverse blood pressure effects.

An advantage of various sensitivity analyses (MMRM and LOCF) is to demonstrate the robustness of the conclusion from the primary analysis in the PP population. In the present manuscript, the planned sensitivity analyses confirmed the positive findings in efficacy, while also allowing contribution of the earlier data from subjects who were discontinued at the week 16 weight assessment.

A major caveat of this study is inherent to its design. In order to test NB effects in a manner consistent with the label, continued study participation in the NB treatment group was limited to those who responded to the drug, whereas no such limitation applied to the control group. Other limitations of the present study include its open-label nature, the uncontrolled nature of the extension time period, and the high number of dropouts. The NB phase 3 trials were specifically designed to test the effects of NB, and thus utilized a common lifestyle intervention program in both NB and placebo groups. This study design compared lifestyle intervention that can be considered standard in clinical care (i.e., encouragement to lose weight, educational materials, and tools designed to enable weight loss, but without regular, consistent reinforcement of the behavior change), against a more supportive lifestyle intervention combined with NB treatment that followed the prescribing information whereby treatment could only be continued if at least 5% weight loss was observed after 12 weeks of treatment with the full dose (16 weeks after start of treatment). The lack of a CLI group without study medication could be viewed as a limitation, as one cannot estimate the contribution from each component of the weight loss assistance provided (NB vs. CLI), but based on the phase 3 trials, the effect of NB above that of a placebo plus lifestyle intervention ranged from 3.68% (20) in patients with type 2 diabetes to 6.24% in a more general obesity population (19).

**Conclusion**

Treatment with 32 mg naltrexone SR/360 mg bupropion SR (NB) as indicated by the prescribing information and along with a CLI significantly improved weight loss relative to usual care alone. The efficacy was maintained for at least 78 weeks, a longer duration than has been previously studied. The results of this study reflect what may be expected to occur in real-world clinical practice.

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