Efficacy of Guanfacine Extended Release Assessed During the Morning, Afternoon, and Evening Using a Modified Conners’ Parent Rating Scale–Revised: Short Form

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Abstract

**Objective:** The purpose of this study was to evaluate the efficacy of once-daily guanfacine extended release (GXR) monotherapy administered either in the morning or evening, using a modified Conners’ Parent Rating Scale–Revised: Short Form (CPRS–R:S) assessed three times/day in children with attention-deficit/hyperactivity disorder (ADHD).

**Methods:** This multicenter, double-blind, placebo-controlled study randomized children 6–12 years of age with ADHD into three groups: GXR a.m. (GXR in the morning and placebo in the evening), GXR p.m. (placebo in the morning and GXR in the evening), or twice-daily placebo. The CPRS–R:S, administered in the morning, afternoon, and evening prior to each study visit, was a secondary measure of efficacy.

**Results:** A total of 333 subjects were included in the analysis population (GXR a.m., n = 107; GXR p.m., n = 114; placebo, n = 112). At visit 10, last observation carried forward (LOCF), subjects receiving GXR demonstrated significantly greater improvement from baseline in the daily mean CPRS–R:S total score, as well as in each of the morning, afternoon, and evening CPRS–R:S assessments, compared with placebo, regardless of the time of GXR administration (p < 0.001 vs. placebo for GXR a.m. and GXR p.m.). In addition, subjects receiving GXR showed significantly greater improvements from baseline in each subscale score (oppositional, cognitive problems/inattention, hyperactivity, and ADHD index) compared with those receiving placebo, regardless of time of administration (p < 0.003 vs. placebo across all subscales for GXR a.m. and GXR p.m.).

**Conclusions:** These results provide further support for the demonstrated efficacy of once-daily GXR in reducing ADHD symptoms, and demonstrate that response is consistent throughout the day regardless of the time of administration, with improvement seen in ratings of oppositional as well as of ADHD symptoms.

Introduction

**Attention-deficit/hyperactivity disorder (ADHD)** is a disorder of hyperactivity, impulsivity, and inattention estimated to affect ~9.5% of children and adolescents 4–17 years of age in the United States (Biederman and Faraone 2005; Centers for Disease Control and Prevention 2010). In ADHD, school/work functioning, psychological functioning, and/or social functioning are impaired, and ADHD often persists into adulthood (Biederman and Faraone 2005; Faraone et al. 2009). Stimulant medications are the most widely used pharmacological treatments for ADHD; however, in some patients, these medications have been associated with suboptimal efficacy and poor tolerability (Banaschewski et al. 2004; Faraone et al. 2009).

Nonstimulants, such as α2A-adrenoceptor agonists (e.g., guanfacine extended release [GXR] and clonidine XR), have also demonstrated efficacy in the treatment of ADHD (Biederman et al. 2008; Sallee et al. 2009; Jain et al. 2011; Wolraich et al. 2011). GXR is indicated in the United States for the treatment of ADHD in children and adolescents 6–17 years of age, both as monotherapy

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including any severe comorbid Axis II or Axis I disorders that could psychiatric disorders (except oppositional defiant disorder), in-
they had a current diagnosis of controlled or uncontrolled comorbid expressions–Severity score
have an ADHD-RS-IV total score (K-SADS-PL), were enrolled in this study. Subjects were required to
by the ADHD Rating Scale IV (ADHD-RS-IV), the study’s primary efficacy outcome. In addition, the tolerability profile of GXR administered either in the morning or evening was similar to that of previous clinical trials of GXR (Biederman et al. 2008; Sallee et al. 2009).

The objective of the current exploratory analyses was to examine additional potential efficacy differences between once-daily GXR (1–4 mg/day) monotherapy, administered either in the morning or evening, compared with placebo in children 6–12 years of age—specifically, consistency of response throughout the day and effect on oppositional as well as ADHD symptoms. The Conners’ Parent Rating Scale–Revised: Short Form (CPRS–R:S), a parent-report measure designed to assess children’s problem behaviors over the previous month, has become a useful tool for assessing treatment outcomes in children with ADHD; the CPRS–R:S can provide ADHD-related, behavior-specific outcome measures (e.g., hyperactivity/impulsivity) of treatment over time (Conners et al. 1998). In this study, the CPRS–R:S administration schedule was modified to assess ADHD symptoms at three time points (morning, afternoon, and evening) throughout the day, and analyzed as a secondary measure of efficacy. A modified CPRS has been utilized as an efficacy measure at similar time points throughout the day in previous studies of stimulants for the treatment of ADHD (Lopez et al. 2008; Coghill et al. 2013; VYVANSE 2013). The CPRS–R:S has widespread clinical utility and provides information on op-
positional symptoms (Conners et al. 1998), a parameter not examined by the ADHD-RS-IV. Because z agonists have been associated with sedative adverse events (AEs), including somnolence, sedation, and hypersomnia (Jain et al. 2011; Kollins et al. 2011), the Pediatric Daytime Sleepiness Scale (PDSS) was also administered as an exploratory secondary measure of safety and tolerability, to determine the extent of daytime sleepiness reported by children treated with GXR monotherapy (and whether this varied as a function of time of administration).

Methods  
Subjects

Children 6–12 years of age with a primary diagnosis of ADHD with combined subtype or hyperactive/impulsive subtype, as defined by the Diagnostic and Statistical Manual of Mental Disorders, 4th ed., Text Revision (DSM-IV-TR) (American Psychiatric Association 2000) based on psychiatric assessment using the Kiddie-Schedule for Affective Disorders and Schizophrenia–Present and Lifetime version (K-SADS-PL), were enrolled in this study. Subjects were required to have an ADHD-RS-IV total score ≥28 and a Clinical Global Impressions–Severity score ≥4 at baseline. Subjects were excluded if they had a current diagnosis of controlled or uncontrolled comorbid psychiatric disorders (except oppositional defiant disorder), including any severe comorbid Axis II or Axis I disorders that could potentially confound efficacy and safety measures or for which GXR is contraindicated, a previous or present risk for suicide, or a history or active presence of cardiac abnormalities or a primary sleep disorder.

Study design

This was an 8 week, multicenter, double-blind, randomized, placebo-controlled, dose optimization, phase 3 study. A screening visit (visit 1) to determine eligibility occurred in the month prior to randomization. Eligible subjects were randomized in a 1:1:1 ratio to the following treatment arms: GXR A.M. (GXR administered in the morning and placebo administered in the evening), GXR P.M. (placebo administered in the morning and GXR administered in the evening), or placebo (placebo administered in the morning and evening). The study consisted of a 5 week dose-optimization period, a 3 week dose-maintenance period, and a 9 day dose-taper period.

During dose optimization, a starting dose of 1 mg/day was titrated upward in 1 mg increments after a minimum of 1 week at the previous dose, based on clinical response and tolerability, up to a maximum of 4 mg/day (visits 2–7). The objective of dose optimization was to determine the dose required to achieve an adequate response (i.e., optimal dose), defined as a clinically significant reduction in ADHD symptoms (≥30% reduction in ADHD-RS-IV total score from baseline) with an acceptable level of side effects. If a ≥30% reduction in ADHD-RS-IV scores was achieved, the op-
timal dose was well tolerated, and the clinician felt that the subject could potentially achieve additional symptom reduction, the dose could be increased to the next dosage strength. The investigator could decrease the dose by 1 mg based on tolerability; however, only one dose reduction was allowed per subject during the study.

Subjects were maintained on their optimal dose for an additional 3 weeks, during which efficacy and safety were assessed weekly (visits 8–10). During the maintenance period, the subject’s dose could be reduced by 1 mg based on tolerability, provided that the dose had not been decreased during the optimization period. Final on-treatment assessment was defined as the last non-missing postbaseline value while the subject was on treatment (before dose tapering) with study drug (analogous to visit 10, last observation carried forward [LOCF]).

Assessments

Results of the primary efficacy measure of change from baseline in the clinician-rated ADHD-RS-IV score have been reported previously (Newcorn et al. 2013). Secondary efficacy endpoints included the change from baseline in the parent/guardian-rated CPRS–R:S score; these results are reported herein. The CPRS–R:S consists of 27 questions grouped into four subscales: oppositional (6 items), cognitive problems/inattention (6 items), hyperactivity (6 items), and ADHD index (12 items). Each item is rated on a four point scale; from 0 (not true at all) to 3 (very much true); total score ranged from 0–81 (higher scores represent greater severity). In the current study, the administration of the CPRS–R:S administration was modified from the validated guidelines in order to evaluate the duration of efficacy response throughout the day (Lopez et al. 2008; VYVANSE 2013). The CPRS–R:S was administered upon awakening (6:00 A.M.), at midday (2:00 P.M.), and during the evening (8:00 P.M.) prior to each of the following visits: 2 (baseline), 7, 8, 9, and 10, and assessments were to be performed within 1 hour of the specified times, rating the child’s behavior immediately before the assessment time.

Safety evaluations included assessments of AEs, vital signs, laboratory test results, physical examination findings, and ratings
on the PDSS. The PDSS, an eight question self-report scale suitable for children and adolescents 11–15 years of age (Drake et al. 2003), was administered to subjects (with parental assistance) at visits 1 (screening), 2 (baseline), and all subsequent scheduled visits through visit 10 (week 8 of treatment). Each item on the PDSS is scored using Likert scale ratings from 0 (never) to 4 (always) for a maximum total score of 32, with higher scores reflecting greater levels of daytime sleepiness.

**Data analyses**

The analysis population included all subjects who were randomized and had taken one or more doses of the study drug. Mean CPRS–R:S scores and mean change from baseline in CPRS–R:S scores were summarized at each time point (morning, afternoon, and evening), and mean scores across the three time points were calculated by treatment group at visits 7, 8, 9, and 10; if scores at any time point from a given visit were missing, the mean was taken from available time points at that visit. Changes in CPRS–R:S total scores were examined at each time point, and a daily mean score was calculated across all time points combined. Changes in CPRS–R:S subscale scores were only examined across all time points combined (daily mean scores).

PDSS total score and mean change from baseline were summarized at each visit through visit 10, using LOCF. A post-hoc item analysis of the PDSS was conducted to further evaluate the effects of treatment on daytime sleepiness. Response distributions to individual PDSS questions were evaluated using the Cochran–Mantel–Haenszel test. Strengths of association were evaluated between PDSS total score change from baseline to visit 10, LOCF, and weight (Pearson correlation), age (Spearman rank correlation), and sex (Spearman rank correlation). Changes from baseline in CPRS–R:S and PDSS scores were analyzed using analysis of covariance (ANCOVA) models at each applicable visit, using LOCF method-

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**Table 1. Demographic and Baseline Characteristics**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>GXR a.m. (n = 107)</th>
<th>GXR p.m. (n = 114)</th>
<th>Placebo (n = 112)</th>
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<tr>
<td>Mean (SD) age, y</td>
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<td>9.3 (1.76)</td>
<td>8.9 (1.78)</td>
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<td>Sex, n (%)</td>
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<td></td>
<td></td>
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<td>Male</td>
<td>72 (67.3)</td>
<td>78 (68.4)</td>
<td>85 (75.9)</td>
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<tr>
<td>Female</td>
<td>35 (32.7)</td>
<td>36 (31.6)</td>
<td>27 (24.1)</td>
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<tr>
<td>Race, n (%)</td>
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<td></td>
<td></td>
</tr>
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<td>White</td>
<td>66 (61.7)</td>
<td>68 (59.6)</td>
<td>56 (50.0)</td>
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<tr>
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<td>47 (42.0)</td>
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<td>0</td>
<td>1 (0.9)</td>
</tr>
<tr>
<td>American Indian or Alaska Native</td>
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<td>1 (0.9)</td>
<td>0</td>
</tr>
<tr>
<td>Other</td>
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<td>10 (8.8)</td>
<td>8 (7.1)</td>
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<td>Mean (SD) body weight, lb</td>
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<td>80.38 (20.91)</td>
<td>75.79 (17.57)</td>
</tr>
<tr>
<td>Mean (SD) BMI, kg/m²</td>
<td>17.92 (2.42)</td>
<td>18.25 (2.35)</td>
<td>17.96 (2.33)</td>
</tr>
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<td>ADHD subtype, n (%)</td>
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<td>Predominately inattentivea</td>
<td>3 (2.8)</td>
<td>3 (2.6)</td>
<td>1 (0.9)</td>
</tr>
<tr>
<td>Predominately hyperactive-impulsive</td>
<td>3 (2.8)</td>
<td>2 (1.8)</td>
<td>1 (0.9)</td>
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<tr>
<td>Combined subtype</td>
<td>101 (94.4)</td>
<td>109 (95.6)</td>
<td>110 (98.2)</td>
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<td>Mean (SD) time since ADHD diagnosis, y</td>
<td>1.5 (2.12)</td>
<td>2.0 (2.24)</td>
<td>1.6 (2.13)</td>
</tr>
</tbody>
</table>

*aPredominately inattentive subtype was exclusionary.

ADHD, attention-deficit/hyperactivity disorder; BMI, body mass index; GXR, guanfacine extended release; SD, standard deviation.

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(morning, afternoon, and evening). At visit 10, LOCF, subjects receiving GXR had a significantly greater improvement from baseline in the daily mean CPRS–R:S total score than did those receiving placebo, regardless of the time of GXR administration ($p < 0.001$ for GXR a.m. and GXR p.m.; Fig. 1A). Effect sizes were similar for both morning and evening GXR administration (0.71 and 0.62, respectively). Subjects on GXR also showed significant improvement compared with the placebo group starting at visit 7 (i.e., the first CPRS–R:S assessment postbaseline, 5 weeks on treatment or LOCF; $p < 0.001$ for GXR a.m. and GXR p.m.).

At each of the morning, afternoon, and evening CPRS–R:S assessments at visit 10, LOCF, subjects receiving GXR showed significantly greater improvements from baseline than did those receiving placebo, regardless of the time of GXR administration ($p < 0.001$ for morning, afternoon, and evening CPRS–R:S assessments for GXR a.m. and GXR p.m.; Fig. 1B–D). At each of the three daily CPRS–R:S assessments, significant improvement from baseline in the total score was evident in subjects receiving GXR compared with those receiving placebo, starting at the first visit at which a postbaseline measurement was obtained (visit 7 [5 weeks on treatment or LOCF]; $p \leq 0.002$ for morning, afternoon, and evening assessments).

CPRS–R:S subscale scores

At baseline, all mean CPRS–R:S subscale scores were similar among treatment groups. At visit 10, LOCF, subjects receiving GXR showed significantly greater improvements from baseline in each subscale score (oppositional, cognitive problems/inattention, hyperactivity, and ADHD index) than did those receiving placebo, regardless of time of administration ($p < 0.003$ across all subscales for GXR a.m. and GXR p.m.; Fig. 2). Effect sizes were similar within each subscale across the different times of GXR administration (effect size range, 0.55–0.85 for GXR a.m. and 0.47–0.68 for GXR p.m.). Subjects on GXR also showed significant improvement on each subscale compared with those receiving placebo starting at visit 7 (5 weeks on treatment or LOCF; $p < 0.008$ across all subscales for GXR a.m. and GXR p.m. vs. placebo).

Safety

Treatment-emergent adverse events (TEAEs) from this study have been previously discussed in detail (Newcorn et al. 2013). In brief, TEAEs occurred in 81.4% of subjects receiving GXR (79.4% for GXR a.m. and 83.3% for GXR p.m.) compared with 57.1% of subjects receiving placebo. The most frequently reported AEs (reported in >10% of subjects) in the GXR groups were somnolence, headache, sedation, upper abdominal pain, and fatigue. The majority of AEs were mild or moderate, and 7.2% of those on GXR (eight subjects each in the a.m. and p.m. groups) discontinued the study because of AEs. Three subjects reported serious AEs (SAEs): one subject in the GXR a.m. group (syncope) and two subjects in the GXR p.m. group (syncope and self-injurious ideation/suicidal ideation). All SAEs were determined by the investigators to be of mild/moderate intensity and to be related to study drug; all resolved by each subject’s final study visit.

FIG. 1. Placebo-adjusted least squares (LS) mean (95% CI) change from baseline in Conners’ Parent Rating Scale–Revised: Short Form (CPRS–R:S) total scores at visit 10, last observation carried forward (LOCF): (A) across all three time points; (B) at morning assessment; (C) at afternoon assessment; and (D) at evening assessment. *$p < 0.001$. GXR, guanfacine extended release. LS means and $p$ values are based on type III sum of squares from an analysis of covariance (ANCOVA) model for the change from baseline, with treatment group as a fixed effect and baseline value as a covariate.
The impact of GXR lowering blood pressure and pulse was consistent with the known safety profile of GXR. For the GXR a.m. and p.m. groups, respectively, mean decreases in supine pulse rate were -3.7 and -3.8 bpm; decreases in systolic blood pressure were -1.6 and -2.1 mm Hg; and decreases in diastolic blood pressure were -0.8 and -2.1 mm Hg. At visit 10, LOCF, subjects receiving GXR demonstrated a mean decrease from baseline in supine pulse rate, systolic blood pressure, and diastolic blood pressure (-3.8 bpm, -1.9 mm Hg, and -1.5 mm Hg) compared with subjects receiving placebo (+1.0 bpm, -0.5 mm Hg, and -0.3 mm Hg).

**PDSS scores**

At baseline, mean (SD) PDSS total scores were similar among treatment groups: 13.9 (5.73) for the GXR a.m. group, 14.9 (5.93) for the GXR p.m. group, and 14.8 (5.75) for the placebo group. By comparison, the normative sample of 450 students from 6th to 8th grades had a mean (SD) PDSS total score of 15.3 (6.2) (Drake et al. 2003), indicating that sleep was not especially disrupted in this population. No significant correlations were found between change from baseline to visit 10, LOCF, in PDSS total scores by treatment group and age, weight, or gender (p ≥ 0.260 among groups for all coefficients). In addition, there were no consistent effects of treatment on PDSS total scores based on whether subjects experienced a sedative TEAE, including somnolence, sedation, and hypersomnia.

**Discussion**

There is increasing awareness that children with ADHD can experience symptoms throughout the day: In school, at home, and during evening activities (Pelham et al. 2001). Therefore, children with ADHD may have a need for long-lasting therapeutic options for the treatment of ADHD symptoms. The current analysis demonstrates that morning or evening administration of once-daily GXR reduces ADHD-related symptoms compared with placebo, as measured by a modified parent/guardian-rated CPRS–R:S. Importantly, symptom reductions were observed at three intervals during the day, at morning, afternoon, and evening CPRS–R:S assessments, regardless of morning or evening GXR administration, supporting once-daily dosing of GXR.

Several other studies have used the CPRS–R:S as an outcome measure. Clinical studies with other nonstimulant therapies such as atomoxetine and clonidine have also demonstrated improvements in CPRS–R:S scores (Michelson et al. 2001, 2002; Spencer et al. 2002). These studies presumably administered unmodified versions of the CPRS–R:S, with baseline and endpoint assessments at least 1 month apart. The current study examined a modified CPRS–R:S at three time points across the day, as was done previously with the stimulant lisdexamfetamine dimesylate (Lopez et al. 2008; VY-VANSE 2013). Results not only demonstrated that once-daily GXR is effective in reducing the CPRS–R:S total score, but these improvements were observed consistently throughout the day.
However, as this study used a modified version of the CPRS–R:S, direct comparisons cannot be made with the previous studies of nonstimulants.

In addition to improvements in the CPRS–R:S total score, morning or evening administration of GXR was more effective than placebo across all symptom subscales (oppositional, cognitive problems/inattention, hyperactivity, and ADHD index). Effect sizes for morning or evening administration of GXR were similar across all subscales; the overall improvement in CPRS–R:S total score was not predominantly driven by any particular subscale. Importantly, GXR showed improvements in the oppositional symptomatology as assessed by the oppositional subscale of the CPRS, symptomatology that is not assessed by the ADHD-RS-IV.

Exploratory secondary safety analyses were conducted using the PDSS to evaluate the effects of GXR (morning or evening administration) on overall daytime sleepiness in children with ADHD, as nonstimulant agents including GXR have been associated with somnolence, sedation, and hypersomnia (Jain et al. 2011; Kollins et al. 2011). Baseline PDSS total scores across treatment groups were similar to the normative sample, and there were no consistent effects of GXR treatment (morning or evening administration) on PDSS total scores based on whether subjects experienced a sedative TEAE. This result is noteworthy, given the sedative side effects typically associated with α agonists, and the fact that sedation was a frequently occurring AE in this study, as previously indicated.

There are several limitations to the methodology of this study that should be considered. First, as previously mentioned, the CPRS–R:S was modified to evaluate ADHD-related symptoms at several time points throughout the day, rather than completing the measure while considering symptoms during the previous month (as it is validated for use) (Conners 1997), which may limit the interpretation of these results. Second, although the study drug was administered at two different time points (morning and evening), this study was not powered to formally assess differences between the A.M. and P.M. cohorts. Last, the PDSS is a subjective assessment and has not demonstrated correlation with objective measures of sleep (e.g., polysomnography) (Patil 2010). As a self-report measure, the PDSS results may have been skewed by inherent rater bias; there appears to be a trend toward inaccuracy for self-appraisal by children and adults with ADHD, although controversy exists regarding this topic (Knouse et al. 2005; Owens et al. 2007; Rizzo et al. 2010; Manor et al. 2012). In addition, although parental assistance was permitted, it is important to note that the PDSS was developed for children and adolescents 11–15 years of age (Drake et al. 2003), but was administered in a younger population (6–12 years) in this study. Furthermore, it is possible that the PDSS scores did not correlate with sedative TEAE incidence because the event may have resolved prior to being captured by the PDSS. For these reasons, the PDSS, as employed in this study, may not have been the most appropriate measure to examine daytime sleepiness.

Conclusions

Once-daily GXR monotherapy has demonstrated efficacy in reducing ADHD symptoms in children, as measured previously by the clinician-rated ADHD-RS-IV, and in the current study by using a modified parent/guardian-rated CPRS–R:S. Once-daily GXR was effective in reducing ADHD symptoms consistently throughout the day as assessed by the CPRS–R:S, regardless of whether the medication was administered in the morning or evening. These improvements were observed in oppositional symptoms and consistently across all other subscales of the CPRS–R:S. Furthermore, the long-lasting effects of GXR support once-daily dosing.

Clinical Significance

These data suggest that once-daily GXR effectively reduces ADHD symptoms and oppositional symptoms in children with ADHD throughout the day. GXR demonstrated similar efficacy whether administered in the morning or evening, consistent with its long half-life and extended time to maximum blood concentrations, thus providing the convenience of either morning or evening administration, as preferred by clinicians and/or patients.

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