Brief report

Improvement in hematological, visceral, and skeletal manifestations of Gaucher disease type 1 with oral eliglustat tartrate (Genz-112638) treatment: 2-year results of a phase 2 study

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Eliglustat tartrate is an investigational oral substrate reduction therapy for Gaucher disease type 1 that is pharmacologically distinct from intravenous enzyme replacement therapy. Eliglustat tartrate improved clinical manifestations in patients who received 50 or 100 mg twice daily for 1 year during an open-label phase 2 study (Blood. 2010;116(6):893-899). We report further improvements after 2 years of treatment in 20 patients (11 females, 9 males; mean age, 33 years) with baseline splenomegaly and thrombocytopenia and/or anemia. Statistically significant (P < .001) percentage improvements from baseline occurred in platelet count (mean ± SD, 81% ± 56%), hemoglobin level (20% ± 15%), spleen volume (~52% ± 11%), and liver volume (~24% ± 13%). Mean platelet count increased ~50 000/mm². Mean hemoglobin level increased 2.1 g/dL overall and 3.1 g/dL in 10 patients with baseline anemia. Organ volume reductions were greatest in patients with severe baseline organomegaly. Seventeen (85%) patients met established therapeutic goals for ≥3 of the 4 parameters. Lumbar spine bone mineral density increased 7.8% ± 10.6% (P = .01) and T-score 0.6 ± 0.8 (P = .012), with major gains in osteoporotic and osteopenic patients. Magnetic resonance imaging assessment showed that bone marrow infiltration by Gaucher cells was decreased (8/18 patients) or stable (10/18 patients). No safety-related trends emerged during 2 years of treatment. This multisite, open-label, single-arm phase 2 study is registered at www.clinicaltrials.gov as NCT00358150. (Blood. 2010;116(20):4095-4098)

Introduction

Eliglustat tartrate (formerly Genz-112638) is an investigational oral substrate reduction therapy (SRT) for Gaucher disease type 1 (GD1), an autosomal recessive lysosomal storage disorder caused by mutations of acid β-glucosidase (glucocerebrosidase; glucosylceramidase; EC 3.2.1.45). In GD1, normal catabolism of glycosphin-golipids (predominantly glucosylceramide) is impaired, and degraded substrates accumulate in “Gaucher cells” of the monocyte/macrophage system, leading to multiorgan disease pathology.1 Thrombocytopenia, anemia, hepatosplenomegaly, and skeletal complications are the most common clinical manifestations, although the pattern and extent of organ involvement vary widely among individuals.

Eliglustat tartrate partially inhibits glucosylceramide synthase, thereby reducing endogenous production of glucosylceramide.2 We recently reported the pharmacokinetics and maximum tolerated doses of eliglustat tartrate from phase 1 studies in healthy normal subjects3 as well as promising results from the 1-year primary analysis period of a phase 2 trial in GD1 patients.4 Here we present longer-term efficacy and safety data obtained after 2 years of treatment in the phase 2 study and assess patient responses to eliglustat tartrate with reference to evidence-based therapeutic goals established for intravenous enzyme replacement therapy (ERT) with imiglucerase.5

Methods

This multisite, open-label, single-arm phase 2 study is registered as NCT00358150 at www.clinicaltrials.gov. Each site’s Ethics Committee or Institutional Review Board approved the protocol. Patients provided written informed consent in accordance with the Declaration of Helsinki. As previously described,4 26 patients with GD1 enrolled, and 6 (13%) withdrew before or at 1 year. The remaining 20 patients continued in the study extension and completed 2 years of treatment. Eliglustat tartrate capsules were self-administered; the dosage was 50 mg (n = 5) or 100 mg (n = 15) twice daily, based on day 10 plasma drug concentrations.

Hemoglobin level, platelet count and plasma biomarkers were analyzed at central laboratories. Spleen and liver volumes (reported as multiples of normal [MN]) were derived from magnetic resonance imaging (MRI) or computed tomography (CT) scans, and skeletal assessments were based on x-ray, dual-energy x-ray absorptiometry (DEXA), and MRI. Images were obtained at the study sites but evaluated by central reviewers who were blinded for organ volume determinations and unblinded for skeletal assessments. Safety assessments, which were periodically reviewed by an


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Figure 1. Improvements in hematological, visceral, and skeletal parameters: overall and stratified by baseline disease severity. (A-F) Data are reported as means ± SD. N = 20, except n = 16 for lumbar spine BMD (excluding 4 patients for receipt of zolendronate during the study [n = 1], assessment with different DEXA machines at different time points [n = 1], lack of 1-year data [n = 1], or presence of radiographic abnormalities preventing BMD assessment [n = 1]). BMD indicates bone mineral density; MN, multiples of normal (volume); F, female; and M, male. Baseline lumbar spine BMD T-scores were categorized as normal, osteopenic, or osteoporotic, as previously described.11
Results and discussion

The 20 patients who completed 2 years of eliglustat tartrate treatment included 11 females and 9 males, aged 18 to 55 years (mean, 33 years) at first treatment. Acid β-glucosidase activity ranged from nondetectable levels to 12% of normal.

Figure 1A shows that statistically significant ($P < .001$) mean percent improvements from baseline occurred for platelet count (+81%), hemoglobin level (+20%), spleen volume (−52%), and liver volume (−24%) at 2 years. Stratification of each disease parameter by baseline severity (Figure 1B–E) revealed that platelet count recovered steadily, on average by ~50 000/mm$^3$, regardless of the severity of baseline thrombocytopenia. Hemoglobin level rose by a mean of 2.1 g/dL for all patients and by 3.1 g/dL for those with more severe baseline anemia. Similarly, organ-volume reductions were relatively greater among patients with more extensive baseline organomegaly. The therapeutic goals established for the 4 disease parameters during 2 years of ERT with imiglucerase$^5$ were met by 90%-95% of patients for spleen, liver, and hemoglobin and by 60% for platelets (Table 1). A slower platelet response also were met by 90%-95% of patients for spleen, liver, and hemoglobin.

<table>
<thead>
<tr>
<th>Clinical parameter</th>
<th>Baseline disease severity</th>
<th>Subgroup</th>
<th>Overall</th>
</tr>
</thead>
<tbody>
<tr>
<td>Platelet count per mm$^3$</td>
<td>Moderate thrombocytopenia</td>
<td>61 000-105 500</td>
<td>64% (7/11)</td>
</tr>
<tr>
<td></td>
<td>Severe thrombocytopenia</td>
<td>39 000-58 000</td>
<td>56% (5/9)</td>
</tr>
<tr>
<td>Hemoglobin level, g/dL</td>
<td>Normal hemoglobin to mild anemia</td>
<td>Females, 11.2-12.5; males, 12.0-14.6</td>
<td>100% (10/10)</td>
</tr>
<tr>
<td></td>
<td>Mild to moderate anemia</td>
<td>Females, 8.7-9.8; males, 9.7-11.9</td>
<td>90% (9/10)</td>
</tr>
<tr>
<td>Spleen volume, MN</td>
<td>Mild to moderate splenomegaly</td>
<td>8.2-14.6</td>
<td>92% (12/13)</td>
</tr>
<tr>
<td></td>
<td>Moderate to severe splenomegaly</td>
<td>15.1-49.2</td>
<td>86% (6/7)</td>
</tr>
<tr>
<td>Liver volume, MN</td>
<td>Mild to moderate hepatomegaly</td>
<td>0.81-1.50</td>
<td>100% (8/8)</td>
</tr>
<tr>
<td></td>
<td>Moderate to severe hepatomegaly</td>
<td>1.53-2.47</td>
<td>92% (11/12)</td>
</tr>
</tbody>
</table>

MN indicates multiples of normal (volume).

That is, while statistically significant increases in mean (SD) T-score [0.6 (0.8); $P = .012$] and Z-score [0.6 (0.7); $P = .003$] were observed for the cohort, the major gains occurred among osteoporotic and osteopenic patients with little change among normal patients (see Figure 1F). For 14 patients with femur BMD data, baseline T-score and Z-score means (SD) of −0.2 (0.9) and 0.1 (0.8), respectively, were normal and did not change at 2 years (0.0 [0.3] and −0.1 [0.4], respectively). Of 18 patients with baseline evidence of Gaucher cell infiltration of bone marrow (dark marrow) by MRI,$^{12}$ 8 showed reductions and 10 were stable. Lytic lesions, evident in 9 patients at baseline, remained stable with no new lesions detected. Among 7 patients with bone infarcts at baseline, 1 improved and 6 remained stable. No bone crises or pathologic fractures were reported.

GD1 biomarkers were elevated in nearly all patients at baseline. Plasma chitotriosidase activity (n = 18) and chemokine CCL18 level (n = 19) each decreased by a median of ~50% at 1 year and 75% at 2 years. Median plasma concentrations of glucosylceramide (n = 19) and ganglioside GM3 (n = 20), exploratory markers of glucosylceramide synthase inhibition, normalized by 6 months and remained normal through 2 years.

Overall, during 2 years of treatment, 23 of 26 patients experienced 126 AEs, with 76% assessed as mild in intensity and 94% considered unrelated to treatment. AEs reported in >2 patients were viral infection (6 patients), urinary tract infection, increased blood pressure, and abdominal pain (3 patients each). The high frequency of diarrhea occurring early in treatment with miglustat, an SRT restricted to adults with mild-to-moderate GD1 who are unable/unwilling to receive ERT,$^{13}$ has not been observed during the phase 2 study of eliglustat tartrate. Safety data for the first year of eliglustat tartrate treatment have been published, including details of patient withdrawals due to pregnancy (n = 3), asymptomatic nonsustained ventricular tachycardia (n = 2), and progression of preexisting osteonecrosis (n = 1).$^4$ During the second year, no patient withdrew, and, of 47 AEs reported for 14 of the 20 patients, none were serious. All AEs were mild (n = 36) or moderate (n = 10) in intensity except for a severe hiatal hernia. Holter monitoring performed at the beginning of the second year revealed no abnormalities, and no treatment-related cardiac events were subsequently reported. The only AE considered possibly treatment-related during the second year was based on results of NCV testing at 18 months that showed decreased sural nerve...
amplitude (bilateral) and velocity (unilateral) in an asymptomatic 41-year-old male. This patient had a prior history of symptomatic peripheral neuropathy that emerged during miglustat treatment and resolved upon its termination. Mild sural nerve abnormalities were also detected by NCV in a 50-year-old male during the first year of eliglustat treatment, but these were not considered treatment-related. No worsening has been evident for either patient on subsequent NCV tests, and both patients have remained asymptomatic with normal neurologic examinations after 3 years of treatment. Peripheral neuropathy, often subclinical, has been reported as a possible manifestation of GD1, as discussed by Cherin et al.14 Miglustat has been associated with symptomatic peripheral neuropathies and tremors that usually arise within the first few months and may lead to cessation of treatment.13

In summary, the treatment responses observed during the first year of eliglustat tartrate therapy were confirmed and extended during the second year, as hematological, visceral, and skeletal manifestations of GD1 continued to improve. This phase 2 study is small and uncontrolled, but spontaneous reversal of disease is not expected and treatment responses for the heterogeneous cohort are consistent for all monitored clinical parameters. The frequency and severity of AEs reported during the second year were remarkably low, and no trends emerged from review of all 2-year safety data. The phase 2 study extension is ongoing, and controlled phase 3 trials of eliglustat tartrate have been initiated.

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Authorship

Contribution: M.J.P. designed the study; E.L., E.A.A., M.I., N.W., H.R., M.P., M.D., and G.M.P. recruited patients and conducted the study research; M.K. performed the statistical analyses; D.I.R. and R.S.K. evaluated the X-ray, MRI, and DEXA images of bone; and M.J.P., A.C.P., and T.S. analyzed and interpreted the results and wrote the manuscript. All authors reviewed an early and final draft of the manuscript; the authors were fully responsible for the content and editorial decisions related to this manuscript.

Conflict-of-interest disclosure: M.J.P., M.K., A.C.P., and T.S. are employees and stockholders of Genzyme Corporation; E.L. received honoraria for travel and speaking from Genzyme Corporation; G.M.P. is the recipient of research grants from Amicus, Actelion, Biomarin, Genzyme, Shire HGT, and Protalix; and D.I.R. and R.S.K. were contracted by Genzyme to review bone images. N.W., E.A.A., M.D., M.I., H.R., and M.P. declare no competing financial interests.

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References

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