Twelve years of experience with miglustat in the treatment of type 1 Gaucher disease: The Spanish ZAGAL project


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Abstract

We report data from a prospective, observational study (ZAGAL) evaluating miglustat 100 mg three times daily orally, in treatment-naïve patients and patients with type 1 Gaucher Disease (GD1) switched from previous enzyme replacement therapy (ERT). Clinical evolution, changes in organ size, blood counts, disease biomarkers, bone marrow infiltration (S-MRI), bone mineral density by broadband ultrasound densitometry (BMD), safety and tolerability annual reports were analysed. Between May 2004 and April 2016, 63 patients received miglustat therapy; 20 (32%) untreated and 43 (68%) switched. At the time of this report 39 patients (14 [36%] treatment-naïve; 25 [64%] switch) remain on miglustat. With over 12-year follow-up, hematologic counts, liver and spleen volumes remained stable. In total, 80% of patients achieved current GD1 therapeutic goals. Plasma chitotriosidase activity and CCL-18/PARC concentration showed a trend towards a slight increase. Reductions on S-MRI (p = 0.042) with an increase in BMD (p < 0.01) were registered. Gastrointestinal disturbances were reported in 25/63 (40%), causing miglustat suspension in 11/63 (17.5%) cases. Thirty-eight patients (60%) experienced a fine hand tremor and two a reversible peripheral neuropathy. Overall, miglustat was effective as a long-term treatment. Efficacy can be maintained in the long term with miglustat. Although gastrointestinal disturbances may appear in the long term, these are reversible and manageable.
1. Background

Type 1 Gaucher disease (GD1) (OMIM 230800) is an autosomal-recessive inborn disorder of lysosomal metabolism characterized by the accumulation of glucocerebroside in different organs and systems as a consequence of the deficient activity of the enzyme, glucocerebrosidase (acid β-glucosidase). GD1 is a multisystemic disease associated with a considerable degree of heterogeneity in terms of symptoms type and severity. It is also one of the most common lysosomal storage disorders, and the first for which multiple approved therapies were available [1,2].

Enzyme replacement therapy (ERT) has been used widely to treat GD1 since 1990 and, in general, studies have demonstrated good efficacy and favourable safety/tolerability [3,4,5]. The achievement and maintenance of GD1 therapeutic goals is considered a useful way to monitor clinical efficacy, and good rates of goal achievement have been reported in a number of clinical studies assessing ERT [6,7]. However, some GD1 patients on long-term ERT can develop a plateau in their response to ERT in terms of haematological and visceral disease parameters. The objective of continued treatment in such patients is to maintain symptom control and ensure adequate quality of life using the most cost-effective long-term treatment [6,7,8,9,10].

Miglustat (Zavesca®; Actelion Pharmaceuticals) is an orally active iminosugar that reversibly inhibits UDP-glucosylceramide synthase – the enzyme that catalyses the rate-limiting stage in the glycosphingolipid biosynthesis pathway [11]. Miglustat was approved in Europe in March 2003 and in the USA in 2004 for use as a substrate reduction therapy (SRT) in adult patients with mild-to-moderate GD1 for whom ERT is either unsuitable or is not a therapeutic option [12,13,14].

There are a number of published data sets on clinical experience with miglustat in GD1, the majority of which report findings from clinical trials, multicenter retrospective cohorts and case series [15,16,17,18,19]. In Spain the prospective, observational ZAGAL study was initiated by the Spanish Gaucher Disease Foundation after miglustat was approved by the European Medicines Agency in 2004. We have previously reported ZAGAL study data regarding the effects of both short-term (6-month) and long-term (5-year) miglustat therapy during everyday clinical use in treatment-naïve patients and those previously treated with ERT [15,16]. Here, we report the long-term efficacy, safety and tolerability of miglustat in ZAGAL study patients with mild-to-moderate GD1 treated in clinical practice settings during a follow-up period of 12 years.

2. Methods

2.1. Study design

The ZAGAL study is a prospective observational study that was initiated to establish a set of recommendations for the collection of efficacy, safety and quality-of-life data in a structured longitudinal manner, and to coordinate the use of miglustat for the treatment of GD1 in real-life settings. All consecutive patients with confirmed GD1 who fulfilled criteria to receive miglustat therapy are included. The study was designed in accordance with recommendations from the European Working Group on GD Advisory Council [20], and was subsequently approved by the Ethics Committee for Clinical Investigation of Aragon (CEICA) and was conducted in accordance with the Declaration of Helsinki of 1975, as revised in 2008.

2.2. Study treatment

Miglustat therapy was administered according to guidelines devised by the Spanish Foundation for the Study and Therapy of Gaucher Disease (FEETEG) in order to optimize and standardize miglustat use. All treated patients received miglustat 100 mg three times daily orally, with dietary recommendations to exclude carbohydrates during the first weeks of starting therapy.

2.3. Efficacy and safety assessments

Specific assessment methods have been described in detail in previous publications [15,16]. Effects on core GD disease parameters were considered in relation to established therapeutic goals for anaemia, thrombocytopenia, hepatomegaly and splenomegaly according to reports published by Pastores et al. [6,7]. Specific assessment methods have been described in detail in previous publications [15,16].

Briefly, organ volumes were evaluated annually at local sites, and were corroborated in a subset of patients using magnetic resonance imaging (MRI). Bone marrow infiltration was evaluated every 12 months based on the published S-MRI scoring protocol for Gaucher cell infiltration patterns in the spine, pelvis and femur [21]. Bone mineral density (BMD) was evaluated using ultrasound measurements in both heels (calcaneus bone) using a Cuba Clinical Bone Densitometer (Norland Medical Systems). Broadband ultrasound attenuation (BUA), speed of sound (SOS), and estimated BMD were all determined at baseline and after 2 and 4 years of therapy. Plasma chitotriosidase (CHT) activity and CLL18/PARC concentration were analysed in the FEETEG laboratory using a 4-methylumbelliferyl-b-D-N,N′-triacylchitotrioside substrate and enzyme-linked immunosorbent assay, respectively. Safety and tolerability assessments were based on patient- and treating-physician reports.

2.4. Data analysis

Descriptive statistics for continuous variables, sample size (n), mean and its standard error (SE), standard deviation (SD), median, and range were calculated for haemoglobin (Hb) concentration, platelet count, spleen volume, liver volume, and changes in biomarker activity. Spleen and liver volumes were calculated as multiples of normal (MN; where normal spleen volume is 2 mL/kg of body weight and normal liver volume is 25 mL/kg of body weight). Patient numbers and percentages were calculated for categorical variables based on patients with available data per time point.

Bone parameters were analysed by study cohort quartiles. Kaplan-Meier analysis estimate of the probability of drug discontinuation or death. Changes between pre- and post-therapy S-MRI scores were assessed using the paired t-test. For all statistical tests, differences were considered significant if p < 0.05.

3. Results

3.1. Patients and disposition

To date, a total of 351 GD1 patients have been diagnosed in Spain [22]. Between 2004 and May 2016, 63 patients from 43 hospitals were treated with miglustat, among whom 20 (32%) received miglustat as first-line therapy and 43 (68%) switched from previous ERT (mean [range] ERT duration 6.9 [1.2–13.5] years).
Baseline patient and disease characteristics for all treated patients, grouped according to treatment path (i.e., treatment naive or switched), are summarized in Table 1. Overall, the mean (range) age at the time of GD1 diagnosis was 34.8 (2–79) years and the mean (range) age at initiation of miglustat therapy was 43.7 (18–79) years.

Of 63 patients exposed to miglustat, a total of 24 discontinued treatment during the study observation period: 11 due to uncontrollable gastrointestinal disturbances, six due to an increase in bone infiltration (two of whom had a bone crisis), one patient due to weight loss >10%, one due to persistent headache (during the early stages of treatment), and two due to planned pregnancy.

Five patients died during the study observation period due to reasons that were not considered related to treatment, including two cases of cancer in treatment-naïve patients (melanoma after 2 years and endometrial adenocarcinoma after 3 years of follow up), and one case each of myocardial infarction, liver failure and progressive general deterioration in switch patients.

Currently, 39 (62%) patients continue to receive miglustat therapy (mean [range] age 44.2 [20–68] years; 41.0% female), among whom 14 (36%) were treatment-naïve at commencement of miglustat and 25 (64%) had previously received ERT for ≤5 years. The median previous ERT dose among switch patients was 30 U/kg every other week; once every other week: nine were receiving ERT ≥30 U/kg every other week at switching.

Among all patients who currently remain on miglustat, the mean (range) overall duration of miglustat treatment is 89.2 (6–120) months. Thirty-eight patients (60%) have received miglustat for ≥5 years, and 15 (38%) have been on miglustat for 12 years. All currently treated patients have undergone at least one post-baseline follow-up visit, providing a median (range) efficacy follow-up period of 99 (9–120) months.

3.2. Core disease parameters

Absolute values of Hb concentration, platelet count, spleen and liver volume, and plasma CHT and CCL18/PARC levels in the entire cohort (N = 63) at baseline and annual follow-up are summarized in Fig. 1. Haemoglobin concentration increased between baseline and Year 4, and appeared stable thereafter (Fig. 1a). Overall, there were no statistically significant differences in mean Hb versus baseline at 2, 5 or 10 years of follow up. Among patients with available data and anaemia at baseline (n = 6), the mean (SD) change in Hb concentration versus baseline was +1.3 g/dL (0.74) at 6-month follow up, and Hb remained stable thereafter. In switch patients with available data (n = 3), Hb concentration was <12 g/dL at the time of switching and improved by +1.8 g/dL (0.39) versus baseline at subsequent follow up.

Platelet counts appeared stable over the 12-year course of follow up, and there was no statistically significant overall change from baseline in mean platelet counts at 2-, 5- or 10-year follow-up (Fig. 1b). Among 20 patients with available data at 10-year follow up, absolute values had decreased slightly, with a mean (range) change from baseline of −35 (10–86) × 10^9/L. Four patients with available data had a platelet count <100 × 10^9/L (70, 77, 90 and 95 × 10^9/L) at final follow up.

Table 1

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Treatment-naïve (N = 20)</th>
<th>Switch (N = 43)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age in years, median (range)</td>
<td>55.7 (18–79)</td>
<td>48.4 (21–70)</td>
</tr>
<tr>
<td>Male: female, n (%)</td>
<td>11 (55.0)</td>
<td>9 (45.0)</td>
</tr>
<tr>
<td>Genotype, n (%):</td>
<td>N370S/N370S 6 (30.0)</td>
<td>3 (69)</td>
</tr>
<tr>
<td></td>
<td>N370S/L444P 6 (30.2)</td>
<td>13 (30.2)</td>
</tr>
<tr>
<td></td>
<td>N370S/other 8 (40.0)</td>
<td>19 (44.1)</td>
</tr>
<tr>
<td></td>
<td>Other/other 0</td>
<td>8 (18.6)</td>
</tr>
<tr>
<td>SSI, median (range)</td>
<td>6.6 (3.0–7.5)</td>
<td>6.6 (2.8–7.6)</td>
</tr>
<tr>
<td>Time on ERT (years)</td>
<td>0</td>
<td>6.9 (1.2–13.5)</td>
</tr>
<tr>
<td>Spleenectomy (%)</td>
<td>3 (15.0)</td>
<td>7 (16.3)</td>
</tr>
</tbody>
</table>

ERT, enzyme replacement therapy; SSI, severity score index.

Spleen and liver volumes did not show statistically significant variations over the 10-year course of follow-up (Figs. 1c and d).

3.3. Biomarkers

At 10-year follow up, we observed wide variability in CHT activity with increased mean in 24/42 (57%) patients with available data (+2448 nmol/mL·h; range 135–13,687), and CCL18/PARC was also increased in 18/42 (43%) patients (mean 250 ng/mL; range 19–1016). These increases were not statistically significant versus baseline values (p = 0.41 and p = 0.08, respectively) and were not associated with clinical manifestations (Figs. 1e and f).

3.4. Therapeutic goals analysis

Over 80% of patients achieved or maintained accepted therapeutic goals for GD1 in the switched on miglustat therapy during the study observation period. Haemoglobin concentration was normalised (to >11.0 g/dL for women; >12.0 g/dL for men) after 12–24 months on miglustat, independent of transfusions. No symptoms of anaemic syndrome were identified over the initial 2 years of follow up. Overall, Hb values were stable at 2-year follow up, and remained so up to the end of the 10-year observation period (Fig. 1) Platelet counts increased in treatment-naïve patients with moderate thrombocytopenia (40–99 × 10^9/L) during the first 12 months of miglustat therapy, and continued improvement up to 60 months. Among switch patients, platelet counts were maintained at ≥100 × 10^9/L throughout 10-year follow-up. The accepted goal for reducing spleen volume is a 30–50% decrease in the first year and 50–60% at 5 years, and the target reduction in liver volume is 20–30% over 12–24 months and 30–40% after 3–5 years. Both of these goals were achieved during miglustat therapy in this cohort.

3.5. Bone parameters

The majority of patients with available bone data (32/43 [74.4%]) reported an overall reduction in chronic bone pain. Mean (range) overall S-MRI scores at baseline and 2-year follow up were 9.6 points (0–25) and 7.2 points (0–21), respectively (p = 0.042). The mean S-MRI score in the treatment-naïve group was 8.8 points (0–14) and 10.0 (0–25) in the switch group at start therapy. Ultrasound examinations of calcaneus bone among a total of 24 patients who received miglustat for ≥2 years (seven treatment-naïve and 17 switch patients) indicated bilateral statistically significant increases in BMD (Table 2).

3.6. Tolerability and safety

The most commonly recorded adverse events in the whole study cohort (N = 63) were mild-to-moderate gastrointestinal disturbances, recorded in 25 (40%) patients. Gastrointestinal adverse events were reversible and were improved by adherence to a controlled diet or disaccharidase supplementation in most cases. An exploratory comparative subgroup analysis to identify differences in general characteristics between patients who showed good gastrointestinal tolerability (n = 38) and those in whom miglustat was not well tolerated (n = 25) did not reveal any significant differences (data not shown). Gastrointestinal disturbances were recorded as the reason for miglustat discontinuation in a total of 11 patients.

Thirty-eight patients (60%) experienced a reversible fine-hand tremor without functional consequences during the first month of therapy, but only 20 patients reported persistence of this effect after 4 years on therapy. The tremor was reversible upon discontinuation of therapy. Electrophysiological signs of subclinical peripheral neuropathy were observed in 8% of patients before miglustat initiation. Periodic neurological tests and electromyography (EMG) in these patients during follow-
Fig. 1. Hb concentration (g/dL), platelet count (×10^9/L), liver and spleen volume (MN), CHT activity (nM/mL-h) and CCL18/PARC concentration (ng/mL) during long-term miglustat therapy in all miglustat-treated patients (N = 63). No significant difference in mean Hb concentration at baseline compared with values after 2 years (p = 0.570), 5 years (p = 0.500) or 10 years (p = 0.778) of miglustat therapy; no significant difference in mean platelet counts at baseline compared with values after 2 years (p = 0.577), 5 years (p = 0.793) or 10 years (p = 0.657) of miglustat therapy. In spite of wide variability in CHT activity and CCL18/PARC concentration there are not significant differences at baseline compared with values after 2, 5 and 10 years (CHT Baseline vs 10 years (p = 0.41)), (CCL18/PARC Baseline vs 10 years (p = 0.08)).

Table 2

<table>
<thead>
<tr>
<th>Limb</th>
<th>BUAa, mean (range)</th>
<th>Pb</th>
<th>Score</th>
<th>Z-T score, mean (SD)</th>
<th>Change</th>
<th>Pf</th>
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<tr>
<td></td>
<td>Baseline Month 24</td>
<td></td>
<td></td>
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<tr>
<td>Right</td>
<td>82.4 (47–101)</td>
<td>0.04</td>
<td>Z</td>
<td>-1.3 (0.9)</td>
<td>-0.09, 0.42</td>
<td>-0.01</td>
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<tr>
<td></td>
<td>84.2 (66–100)</td>
<td></td>
<td>T</td>
<td>-1.3 (0.9)</td>
<td>0.09, 0.48</td>
<td>-0.01</td>
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<td>-1.1 (0.9)</td>
<td>0.09, 0.30</td>
<td>-0.01</td>
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<td></td>
<td>-1.5 (1.1)</td>
<td>0.08, 0.45</td>
<td>-0.01</td>
</tr>
<tr>
<td>Left</td>
<td>74.2 (32–100)</td>
<td>0.06</td>
<td>Z</td>
<td>-1.1 (0.9)</td>
<td>0.09, 0.30</td>
<td>-0.01</td>
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<td></td>
<td>75.6 (48–101)</td>
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<td>T</td>
<td>-1.1 (0.9)</td>
<td>0.09, 0.48</td>
<td>-0.01</td>
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<td>-1.1 (0.4)</td>
<td>0.08, 0.45</td>
<td>-0.01</td>
</tr>
</tbody>
</table>

a Broadband ultrasound attenuation.
b Data based on seven treatment-naïve patients and 17 previous ERT patients with available data.
c P-values related to a comparison of month 24 values with baseline values, as assessed based on the paired t-test.

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up did not reveal any evidence of progressive neuropathy. However
during follow-up two patients developed a peripheral neuropathy.

No patients in the study appeared to develop cognitive deterioration
during miglustat therapy for up to 12 years. Weight loss >10% was re-
ported for three patients (4.7%). Among patients who reported gastroin-
testinal disturbances as the reason for miglustat discontinuation, six
also showed increased bone infiltration, two had a bone crisis, and
one had >10% weight loss. One patient also had Gaucher cell infiltration
in the myocardium (at biopsy), and one had persistent headache that
led to discontinuation in the early stages of treatment.

Kaplan-Meier estimate of the probability of drug discontinuation or
death in the entire cohort (n = 63) indicated a tendency to lower over-
all rate of miglustat discontinuation among treatment-naive patients
compared with the switch group, but the comparison by Log-Rank
test is not significant. (Fig. 2).

4. Discussion

We present long-term data on changes in haematological and
visceral indices and biomarkers in both treatment-naive and ERT-
stabilized adult GD1 patients treated with miglustat for up to 12 years
in clinical practice settings. To our knowledge, this report represents
the longest real-world assessment of the effectiveness and safety of
miglustat in GD1 to date, and builds on previous findings from this co-
hort and on data reported in other open-label studies and case series
[15,17,18,19,23,24].

Throughout up to 12 years of therapy with miglustat in this Spanish
cohort, Hb concentration remained largely unchanged and platelet
count remained stable, with moderate fluctuations. Liver and spleen
volumes were also stable during follow up. These findings are consistent
with accepted therapeutic goals for GD1 therapy in treatment-naive pa-
tients reported by Pastores et al. in 2004 [6,7]. The data from this cohort
add further support for the use of miglustat in patients who refuse ERT
or who are otherwise unsuited to regular intravenous infusions. The ef-
cacy of miglustat as a maintenance monotherapy for GD1 over 24
months has previously been reported in 36 patients previously stabi-

dized with ERT [25]. In the observational, retrospective multicentre
study in 115 miglustat-treated patients reported by Kuter et al. [19],
Hb and platelet counts tended to increase in treatment-naive patients
during miglustat therapy, and remained stable or decreased slightly in
ERT pre-treated patients.

The efficacy of miglustat may result from two mechanisms of action.
Besides inhibiting substrate formation, miglustat may also accelerate
destruction of the glycolipid complex by increasing glucocerebrosidase
(GC) activity through a chaperoning function. The chaperoning effect
of miglustat was shown to be specific for glucocerebrosidase with mu-
tations in the catalytic domain (domain III) [25,26]. In vitro studies dem-
strated that glucocerebrosidase activity was increased 2.5-fold in
cells transfected with the N370S mutation when cultured in a medium
containing miglustat [26]. It is therefore noteworthy that the majority
of the patients in our series had at least one glucocerebrosidase gene al-

lele with the N370S mutation.

Disease biomarkers during miglustat therapy showed a tendency to
increase during follow up in this study without any appearance of clinical
manifestations. It is therefore possible that differences in patient compli-
ance or inter-individual variations affected the observed CHT values.
Overall our data suggest that plasma CHT and CCL18/PARC might not
fully reflect systemic disease burden in GD1, and cast doubt on the strict
validity of these biomarkers in clinical decision making [25].

Bone disease is the most frequent and debilitating manifestation of
GD1 [27]. A prospective pooled analysis of data from three open-label
studies with miglustat as monotherapy for GD1 demonstrated that
miglustat reduces the incidence of bone pain and improves BMD [28]
– an effect thought to stem from the fact that this agent has a wide dis-
tribution throughout body tissues, including bone [29]. Early and
sustained increases in lumbar spine and femoral neck BMD were report-
ed, with significant increases from baseline evident at 6, 12 and
24 months [28]. For these reasons, miglustat might be a valuable treat-
ment option for the improvement of bone disease in GD1.

Based on the current cohort we assessed long-term changes in BMD
and bone elasticity using ultrasound methodology in the calcaneus
(heel) bone. Data from heel bone ultrasound analyses have been shewn to correlate well with DXA measurements, although ultrasound
data indicate greater degrees of change over time compared with DXA
findings [30]. Evaluations in our patients were performed by the same
radiologist using the same machine, and according to the same protocol,
at baseline and follow up, and showed statistically significant increases
of both ultrasound-derived BUA and DXA-derived Z- and T-scores after
2 years on miglustat. The majority of patients in the whole cohort also
showed improvements in chronic bone pain and Gaucher cell infiltr-
ation of bone (as measured by S-MRI) during 2 years on therapy.

Avascular necrosis was observed in two patients (3.1%), both of
whom were non-splenectomized switch patients, and one of which
did not demonstrate good adherence to miglustat therapy. GD1 is a
chronic disorder and requires an effective and continuous treatment
regimen on a life-long basis. For patients on prolonged oral therapy
the correct adherence to every day therapy is a widespread problem
[31].

Gastrointestinal tolerability showed a high degree of individual vari-
ability in this study, which was possibly related to differences in food
habits and levels of disaccharidase inhibition (Medrano B et al. in
press). Similar to therapies for other chronic diseases such as diabetes,
arterial hypertension and dyslipidaemia, gastrointestinal disturbances
such as diarrhoea, flatulence and vomiting are known, common effects
seen during miglustat treatment [12,13]: it has been estimated that ap-
proximately 56% of drug therapies have an influence on gastrointestinal
function. Although uncomfortable if they occur, gastrointestinal adverse
events related to miglustat therapy are reversible and do not require in-
tensive therapy in most cases [31]. While generally moderate in intensi-

ty and not related to severe complications, gastrointestinal effects were
the most frequent reason for miglustat discontinuation in this cohort,
which is in line with previous published data [31].

Tremor was the main neurological side effect observed during the
study, particularly at commencement of treatment. Two new cases of
peripheral neuropathy with clinical symptoms that were attributable
to miglustat were observed in this study. Previous published data dem-
strating that peripheral neuropathy should be considered part of the
natural history of GD1 [32,33,34], and data from up to 9 years of post-
marketing clinical safety surveillance for miglustat that showed no

Fig. 2. Estimated probability to discontinuation of miglustat therapy (all miglustat-treated
patients; N = 63). Data censored to account for discontinuation of miglustat therapy or
death from any cause (Log-Rank test 0.482).
association between miglustat and peripheral neuropathy [17,23], however treating physicians should be aware of this association to improve early detection.

No other adverse long-term effects of miglustat treatment have been documented [17,23].

5. Conclusions

The first trials of oral SRT with miglustat in GD1 began >16 years ago [14,25,35,36]. As a result there is now considerable, widespread experience of its use in clinical practice. The general opinion regarding oral miglustat therapy is that it is satisfactory both in terms of controlling disease in treatment-naive patients and in maintaining disease stability in patients previously stabilized on ERT. Miglustat is not a curative therapy, but it does prevent excess lipid storage in the spleen, liver and bone marrow, and has been shown to maintain disease stability in line with established therapeutic goals in the majority of patients. Overall, long-term treatment with miglustat appeared associated with only mild-to-moderate adverse events in this Spanish cohort. Importantly, miglustat also provided additional benefits on bone parameters, with statistically significant decreases in bone marrow infiltration (as measured by S-MRI) and increases in BMD (as measured by BUA and DXA). Plasma CHT activity and CCL-18/PARC concentration showed a trend to increase [14,25,35,36]. As a result there is now considerable, widespread experience of its use in clinical practice. The general opinion regarding oral miglustat therapy is that it is satisfactory both in terms of controlling disease in treatment-naive patients and in maintaining disease stability in patients previously stabilized on ERT. Miglustat is not a curative therapy, but it does prevent excess lipid storage in the spleen, liver and bone marrow, and has been shown to maintain disease stability in line with established therapeutic goals in the majority of patients. Overall, long-term treatment with miglustat appeared associated with only mild-to-moderate adverse events in this Spanish cohort. Importantly, miglustat also provided additional benefits on bone parameters, with statistically significant decreases in bone marrow infiltration (as measured by S-MRI) and increases in BMD (as measured by BUA and DXA). Plasma CHT activity and CCL-18/PARC concentration showed a trend to increase over time, but did not reach initial values. As an aside, biomarker alterations did not correlate with clinical.

As we have commented in previous reports [15,16], and reinforce now with the benefit of accumulated experience, miglustat appears to be a valid alternative to ERT as a maintenance therapy in GD1. This agent also has the added advantage of convenient oral administration, which serves as a useful therapeutic option for maintaining efficacy in patients who discontinue ERT due to preference, holidays, travel, work or other commitments.

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References


