A comprehensive monocentric ophthalmic study with Gaucher disease type 3 patients: vitreoretinal lesions, retinal atrophy and characterization of abnormal saccades

Susanne Hopf1*, Norbert Pfeiffer1, Matthias Liesenfeld2, Karl-Eugen Mengel3, Julia B. Hennermann3, Irene Schmidtmann4 and Susanne Pitz5

Summary

Background: The differentiation between Gaucher disease type 3 (GD3) and type 1 is challenging because pathognomonic neurologic symptoms may be subtle and develop at late stages. The ophthalmologist plays a crucial role in identifying the typical impairment of horizontal saccadic eye movements, followed by vertical ones. Little is known about further ocular involvement. The aim of this monocentric cohort study is to comprehensively describe the ophthalmological features of Gaucher disease type 3. We suggest recommendations for a set of useful ophthalmologic investigations for diagnosis and follow up and for saccadometry parameters enabling a correlation to disease severity.

Methods: Sixteen patients with biochemically and genetically diagnosed GD3 completed ophthalmologic examination including optical coherence tomography (OCT), clinical oculomotor assessment and saccadometry by infrared based video-oculography. Saccadic peak velocity, gain and latency were compared to 100 healthy controls, using parametric tests. Correlations between saccadic assessment and clinical parameters were calculated.

Results: Peripapillary subretinal drusen-like deposits with retinal atrophy (2/16), preretinal opacities of the vitreous (4/16) and increased retinal vessel tortuosity (3/16) were found. Oculomotor pathology with clinically slowed saccades was more frequent horizontally (15/16) than vertically (12/16). Saccadometry revealed slowed peak velocity compared to 100 controls (most evident horizontally and downwards). Saccades were delayed and hypometric. Best correlating with SARA (scale for the assessment and rating of ataxia), disease duration, mSST (modified Severity Scoring Tool) and reduced IQ was peak velocity (both up- and downwards). Motility restriction occurred in 8/16 patients affecting horizontal eye movements, while vertical motility restriction was seen less frequently. Impaired abduction presented with esophoria or esotropia, the latter in combination with reduced stereopsis.

Conclusions: Vitreoretinal lesions may occur in 25% of Gaucher type 3 patients, while we additionally observed subretinal lesions with retinal atrophy in advanced disease stages. Vertical saccadic peak velocity seems the most promising “biomarker” for neuropathic manifestation for future longitudinal studies, as it correlates best with other neurologic symptoms. Apart from the well documented abduction deficit in Gaucher type 3 we were able to demonstrate motility impairment in all directions of gaze.

Keywords: Saccadometry, Video-oculography, Saccades, Gaucher’s disease, Gaucher disease type III, Retina, Vitreous opacity, Ophthalmology, Neuro-ophthalmology, Lysosomal storage disease

* Correspondence: hopf101@gmail.com
1Department of Ophthalmology, University Medical Center Mainz, Langenbeckstr.1, 55131 Mainz, Germany
Full list of author information is available at the end of the article

© The Author(s). 2019 Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
Background

Context and summary of the literature

Gaucher disease is the most common sphingolipidosis and is defined by the autosomal recessively inherited deficiency of lysosomal acid β-glucocerebrosidase. The gold standard for diagnosing Gaucher disease is the confirmation of deficient enzyme activity. Mutation detection in the affected GBA-gene can enhance the diagnostic accuracy [1]. However, there is a strong genetic heterogeneity of patients with similar symptoms and vice versa [2, 3]. Few genotypes are known to have a prognostic impact. The homozygote L444P allele is associated with Gaucher disease type 3 (GD3), which is the rare chronic neuronopathic form [4–7]. The diagnosis of Gaucher disease type 3 (GD3) is challenging as specific neurologic symptoms develop at various ages [8]. Reclassification of type 1 (visceral) to type 3 (chronic neuronopathic) in later stages occurs [9]. The most common early neurologic symptoms are oculomotor disturbances [10], such as slowed and hypometric saccades [8, 10] and saccadic initiation errors [8], and esotropia [8, 11] and/or abduction deficits [8, 12–14]. Horizontal saccades decelerate earlier, while vertical saccades follow [15–17]. Vertical downwards saccades are known to be impaired earlier than the less affected upwards saccades [8, 15, 16, 18]. As the clinical examination of oculomotor abnormalities has a limited reproducibility, depending on patient’s cooperation as well as the investigator’s evaluation, an objective oculomotor testing has the potential to result in a more objective assessment and a better quality of data for follow-up investigation [8].

Beside neuro-ophthalmologic abnormalities, Gaucher type 3 may present posterior segment abnormalities of the eye. Specific vitreous opacities are better known from Gaucher type 1, with a prevalence of 3% [19]. The prevalence of posterior segment abnormalities in Gaucher type 3 is not yet established. Recent case reports describe preretinal accumulations of matter [20–24]. In fact, they even may promote severe tractive retinal detachment in Gaucher type 3 [25, 26].

Purpose of this study

The purpose of this study is to propose recommendations for a set of useful ophthalmologic investigations for diagnosis and follow up as well as saccadometry parameters of Gaucher disease type 3 patients enabling a correlation to disease severity.

Methods

This monocentric cohort study was conducted in 2015 in the University Medical Center Mainz in Germany. The study was approved by the Medical Ethical Committee of the State Chamber of Medicine of Rhineland Palatinate in Mainz, Germany (reference number 837.374.14 (9613)).

The patients or their parents/guardians gave a written consent to publication of their anonymized clinical data.

Characteristics of participants

Inclusion criteria for participants were a biochemically and genetically diagnosed Gaucher disease type 3 and the cognitive ability to perform the investigations. Children under 6 years were excluded, because cooperation was regarded not to be sufficient for the examinations at that age. A visual acuity of better than 1.3 logMAR (at far distance) was required.

Sixteen patients were included in this study. A cohort of 100 healthy individuals at a stratified age distribution served as control.

Ophthalmologic and neuro-ophthalmologic examination procedure

The patients underwent an ophthalmologic and neuro-ophthalmologic examination according to a standardized investigation form. Ophthalmologic examination included: refraction, best corrected visual acuity, anterior and posterior segment examination, Spectral Domain (SD)-OCT (Spectralis OCT, Heidelberg Engineering GmbH, Heidelberg, Germany) of the optic nerve head and the macula. Only OCT scans with a quality index above 20 were analyzed. Neuro-ophthalmologic examination included: orthoptic status (stereopsis, cover test, motility, optokinetic nystagmus and clinical saccade testing). A technical saccadometry was performed using an infrared-videooculography device (see below). Saccadometry was done by a specifically trained investigator. Orthoptic examination was carried out by an experienced orthoptist in 12 cases.

Quantification of neuropathic Gaucher symptoms

The neuropathic Gaucher symptoms were assessed quantitatively by the disease specific “modified severity scoring tool” (mSST) using 12 categories [27]. Additionally, the “scale for the assessment and rating of ataxia” (SARA) including 8 items was assessed. Herein, the items ataxia and dysarthria [28] are characteristic in GD3 [29–31] and are also part of the mSST [27]. The IQ, one of the categories of the mSST, represents the cognitive function and in Gaucher patients serves to evaluate the neurologic involvement of the disease. The wide range of cognitive impairment in Gaucher patients has been reported by several authors [11, 15, 32].

Video-oculographic saccadometry using EyeSeeCam in Gaucher patients

Saccades were recorded using the infrared video-oculography (VOG) device EyeSeeCam HIT (Eyeseetec,
Fürstenfeldbruck), sampling at 220 Hz (every 3.6 ms) which was linked to a MacBook Pro 13” (OS X Version 10.9.5, Apple Inc.) equipped with the EyeSeeCam software (EyeSeeCam VOG HIT System Reversion r3429 and r3444, EyeSeeTec Fürstenfeldbruck, Germany). The study participants were seated on a height-adjustable chair, facing the center of a computer monitor (Dell Technologies Inc., 19” screen BQR-1908FPb, 1280 x 1024 resolution, 60 Hz refresh rate, 300 CD/m² display luminance, and 5 ms images building time) at a distance of 60 cm from the glabella to the monitor. The linear visual range was at 5°/15°/30° leftwards and rightwards and 5°/10°/20° upwards and downwards. Each target was a smiley icon. Prior to the measurement, a qualitative calibration was performed. Movements of the left eye were measured. To avoid alternating fixation in heterotropia, the right eye was covered in these cases.

Saccades were counted and calculated at a threshold of 100°/s. The beginning and ending of a saccade was determined at a threshold of 5°/s. Further criteria were an occurrence 0.5 s after stimulus at latest and an amplitude of more than 0.5 of the relevant stimulus. In case that standard calculation did not detect enough saccades due to pathologically decelerated saccades, a “special saccades calculation” with more sensible criteria was used (threshold of 5°/s, beginning and end at 2°/s, latest detection 0.7 s after stimulus and detection of saccades even if they measure less than 0.5 of the stimulus amplitude).

The evaluation by “special saccades calculation” was necessary in 13/14 patients horizontally and in 8/14 patients horizontally to allow the detection of a sufficient number of saccades. Thereby a minimum of 4 saccades could be included in 82.4% of the 4 direction and 3 degrees of 12 different eccentricities) in the 14 patients.

Statistical analysis, power calculation
With 16 patients and 100 controls, it was possible to establish a difference of 0.67 standard deviation between groups at 5% level with a power of 80% using a two sample t-test. This required that the parameter of interest followed a normal distribution with common variance. This study did not need an age matching as the main measurement parameters have been shown not to be age dependent in the targeted population (as described in a parallel paper) [33].

Statistical Package for Social Sciences (SPSS 23) was used for the analysis. To evaluate the distribution of data, Shapiro-Wilk-test and diagnostic plots were performed for each parameter to assess symmetry and peaks. Between the patient and the study group, normally distributed parameters were compared using t-tests. Within the patient group paired t-test was used. As this is an explorative study, we consider p-values ≤0.05 as indication for statistically significant difference. No adjustment for multiple testing has been applied, and thus only the error rate per comparison is controlled.

To assess correlations between the saccade parameters (peak velocity, amplitude, latency) and the mSST, SARA and IQ, Spearman correlation was performed. Spearman’s rank correlation coefficient of ≥0.8 was considered a strong correlation; ≥ 0.5 was considered a moderate correlation; ≥ 0.2 a weak correlation; and < 0.2 no correlation.

Results
A total of 16 patients with biochemically and genetically diagnosed Gaucher disease type 3 was examined, of which 14 performed sufficient quality data in saccadometry. An ophthalmologic summary of the patient series is shown in Table 1, while Table 2 contains a detailed description of all patients.

In the anterior segment, a diffuse corneal opacity was found in one patient (Fig. 1) besides a slight retraction of the eye lids. The patient was heterozygous G202R/D409H. The corneal haze did not affect visual acuity (0.05 as indication for statistically significant difference. No adjustment for multiple testing has been applied, and thus only the error rate per comparison is controlled.

In the posterior segment, vitreous lesions were seen by fundoscopy or OCT. These mostly large opacities stood out as white or hyperreflective, often round, dot or cloud like opacities without disturbing patient’s subjective vision or visual acuity (Fig. 2).

In another three cases we found epiretinal particles on the ILM and the posterior hyaloid membrane by OCT. These lesions escaped fundoscopy.

While the previous ocular findings were located preretinally, two patients showed peculiarities, which were located subretinally. The lesions arose in the peripapillary region in a regular pattern in the shape of whitish drusen-like deposits, potentially reaching the macula area (Fig. 3). The deposits’ size was about 1/6 of the optic nerve head diameter which itself was slightly enlarged. Additional small lesions were found in the perifoveal area of the macula (Fig. 3). In the OCT, the deposits were identified in the subretinal region as hyperreflective dome shaped structures (Fig. 4). Importantly, the retina adjacent to these deposits was atrophic, showing photoreceptor and retinal pigment epithelium (RPE) loss in an extended area towards the optic nerve. The more affected case was a patient with a severe phenotype (Fig. 3), while the other one had intermediate phenotype and presented few deposits and retinal atrophy close to the optic nerve head in addition to extensive vitreous opacities (Figs.5 and 6). Both patients had a comparable disease duration of about 19 years.
Oculomotor segments
Posterior segments
Visual Acuity normal Median 0
Refraction 5 x myopic (OD + OS)
Refractive error in type 3 GA was used to describe the eyes as either hypertonic (OD + OS)
Corneal opacity + slight retraction of the eye lids
Anterior segments
Conical opacity + slight retraction of the eye lids
Posterior segments
Vitreous opacities
Epiretinal particles
Retinal deposits + atrophy
Oculomotor results
Abduction deficits
Up−/Downgaze deficits
Saccadometry
Horizontal saccades
Vertical saccades

Tortuosity or caliper changes of retinal vessels were detected in four patients (Additional file 1).

Optic nerve head appearance was normal; two patients exhibited tilted discs due to moderate/high myopia. All optic discs had normal retinal nerve fiber layer (RNFL) thickness in OCT.

Oculomotor investigation
Bilateral abduction deficits were found in 8/16 patients, while 6 patients additionally presented downgaze or and upgaze restriction. While patients with mild phenotype show free motility until the middle of the 3rd decade, restricted motility already occurs in the 2nd decade in patients with a severe phenotype (Table 2). The motility deficit was of marked severity in two of the patients precluding saccadometric measurements due to poor quality. Strabismus was present in 8 of 16 patients (most common strabismus convergens, followed by strabismus convergens alternans and decompensated esophoria). Half of the patients had full stereopsis in Titmus ring test, however stereo resolution was significantly worse than in the control group (mean degree of disparity in patients 130 ± 149 arc seconds vs. 54 ± 45 arc seconds in the controls, p = 0.041). Limited triggerable optokinetic nystagmus was seen in 14/16 patients (stronger impairment horizontally in 8/16). Binocular clinical assessment revealed horizontal saccades to be more often decelerated (15/16) than vertical ones (12/16) (Table 3). The patient presenting normal saccades horizontally suffered epilepsy and was genetically diagnosed as type 3 as both mutations N188S and L395P are known in type 3 GD. No type 1 mutation/ no neuroprotective mutation was present.

Saccadometric measurements
A minimum of 4 recorded saccades (as response to 6–8 stimuli per direction and target displacement) in each of the 16 patients was detected in 82.4% of the 12 eccentricities in 16 patients (98.5% in the 100 control subjects). Missing values were few. The saccade parameters had an approximately normal distribution except for latency.

Saccade parameters, especially peak velocity, showed to be impaired in a characteristic pattern of damage: horizontally (rightwards/leftwards) followed by downwards and last upwards (Fig. 7).

Peak velocity was significantly reduced (exemplarily peak velocity 15° saccades horizontally: 69°/s rightwards and 83°/s leftwards in GD3 patients vs. 339°/s rightwards and 364° leftwards in the controls, p < 0.001), while main sequence was preserved. The difference to the control group was more evident horizontally and downwards (p ≤ 0.001), followed by upward saccades (p = 0.001–0.003). Moreover, paired t-test showed downwards saccades in GD3 patient to be slower (144°/s ± 87°/s) compared to upwards saccades (192°/s ± 94°/s).

Further saccadometric data are provided in the Additional file 2 displayed as boxplots [see Additional file 2]. Gain or saccadic accuracy as defined per amplitude divided by target eccentricity proved to have a larger range (in terms of interquartile distance) in GD3 patients. The saccades were of similar accuracy vertically, while being less accurate horizontally. Both study groups showed hypometric saccades more often in horizontal than in vertical direction (see Additional file 2A).

The latency of Gaucher patients was not normally distributed. The non-parametric Wilcoxon rank sum test revealed that latency was significantly longer in Gaucher patients compared to controls except for upward direction and large target displacements (20° vertically and 30° horizontally) (Table 4) (see Additional file 2B).
| ID | Phenotype severity | Genotype       | Age | Disease duration | SARA IQ | Adjusted mSST | Motility (smooth pursuit) | Cover Test | Ocular findings | Horizontal velocity (e.g. 30° rightwards) | Vertical velocity (e.g. 20° upwards) | logMAR visual acuity (right eye) | logMAR visual acuity (left eye) |
|----|--------------------|----------------|-----|------------------|---------|---------------|---------------------------|------------|----------------|------------------------------------------|-------------------------------------|-------------------------------|-------------------------------|
| 1  | Mild               | L444P/D409H    | 8   | 7                | 0       | 100 0         | Full, synkinetic blinking, head thrust | Normal     | Normal         | 22                                        | 337                                 | 0                             | 0                             |
| 2  | Mild               | L444P/D409H    | 21  | 12               | 0       | 100 0.5       | Full, synkinetic blinking, head thrust | Normal     | Epiretinal particles on the ILM and posterior hyaloid, partially detached -> normal | 46                                        | 293                                 | 0                             | 0                             |
| 3  | Mild               | L444P/D409H    | 22  | 15               | 2       | 85 0.5        | Full                       | Normal     | Normal         | 57                                        | 266                                 | 0                             | 0                             |
| 4  | Mild               | N188S/L385P    | 24  | 3                | 0       | 100 1.5       | Full                       | Normal     | Partial vitreous detachment- > normal | 332                                       | 323                                 | 0                             | 0                             |
| 5  | Mild               | –               | 27  | 21               | 0       | 100 0         | OD: minimal adduction defect, OS: minimal abduction deficit, OU: elevation 25°, depression normal. Synkinetic blinking. | Decompressing esophoria | OD: partial vitreous detachment, slight peripapillary atrophy, OD/OS Lamina cribrosa- > normal | 114                                       | 242                                 | 0                             | 0                             |
| 6  | Mild               | G202R/D409H (*1) | 44  | 38               | 39      | 71 18         | *2: abduction 10°, adduction 20°, elevation 20°, depression 10° | Decompressing esophoria | Sight retraction of the eyelids, diffuse corneal opacity, tilted disc, prominent posterior hyaloid membrane, OD: large optic nerve head and excavation | 25                                        | 39                                  | 0                             | 0                             |
| 7  | Intermediate       | L444P/L444P     | 6   | 5                | 1       | 100 1         | Full                       | Esotropia OD | Tortuous vessels | 119                                       | 405                                 | −0.1                          | −0.1                          |
| 8  | Intermediate       | L444P/L444P     | 7   | 6                | 0       | 104 1         | Full                       | Normal     | Tortuous vessels | 119                                       | 452                                 | −0.1                          | −0.1                          |
| 9  | Intermediate       | L444P/L444P     | 8   | 6                | 0       | 79 1          | Full                       | Normal     | Normal          | 88                                        | 333                                 | 0                             | 0                             |
| 10 | Intermediate       | L444P/L444P     | 10  | 10               | 16      | 60 10         | OD/OS: abduction 5° before primary position (p.p./p.p; adduction 30°/25°, minimal elevation and depression deficit) | Esotropia OD (*4) | Normal (cave: bad image quality in OCT) | –                                         | –                                  | 0                             | 0                             |
| 11 | Intermediate       | L444P/L444P     | 12  | 12               | 10      | 68 6          | Adduction 35–40°            | Alternating esotropia | Large vitreous opacities, drusen-like deposits, retinal atrophy, partial vitreous detachment | 98                                        | 134                                 | 0                             | 0                             |
| 12 | Intermediate       | L444P/L444P     | 19  | 18               | 18      | 60 15         | Full, synkinetic blinking | Esotropia OD (*4) | Large vitreous opacities | –                                         | –                                  | 0.1                          | 0.1                          |
| 13 | Intermediate       | L444P/L444P     | 21  | 19               | 4       | 75 5.5        | Abduction 45°/adduction 40°, adduction 35°, elevation 15° (*3), depression normal. Synkinetic blinking. | Alternating esotropia (*5) | Large vitreous opacities, tortuous vessels, epiretinal particles on an irregular ILM and the posterior hyaloid membrane | 76                                        | 233                                 | 0                             | 0                             |
| ID | Phenotype severity | Genotype | Age | Disease duration | SARA IQ | Adjusted mSST | Motility (smooth pursuit) | Cover Test | Ocular findings                                                                 |
|----|--------------------|----------|-----|------------------|---------|---------------|--------------------------|------------|--------------------------------------------------------------------------------|
| 14 | Intermediate       | L444P/L444P | 23  | 0                | 100 2.5 | Normal         | Peripapillary atrophy due to high myopia -> normal | 95 396 0 0 |                                                                              |
| 15 | Intermediate       | L444P/L444P | 33  | 9                | 68 18.5 | Normal         | Partial vitreous detachment, caliper changes, fine granular background, thin retina and thin vessels | 91 173 0 0.1 |                                                                              |
| 16 | Severe             |          | 19  | 7                | 79 12   | Secondary alternating esotropia (*6) | Vitreous opacities, druse-like deposits, retinal atrophy (RPE and photoreceptor loss), enlarged optic nerve diameter (5.5 mm) | 107 105 0 0.1 |                                                                              |

*1: G202R mutation known from GD2 with slightly reduced enzyme activity, while neuronopathic involvement is severe [34]. *2: only elicitable by VOR. *3: elevation by VOR 25°. *4: patient had undergone once strabismus surgery OD. *5: patient had undergone 4 times strabismus surgery. *6: initially esotropia. Patients with ID 4, 6, 15, 16 were under anticonvulsive medication.
Correlation analysis between saccade parameters and other neurologic parameters

Clinical scores correlated with subjective and objective saccadic peak velocity and with saccade hypometria. Phenotype severity showed mild correlation with longer latency. High mSST (adjusted for saccades), high SARA, long duration of disease and low IQ correlate with slow and hypometric saccades. The correlation was found to be greater in vertical than in horizontal saccades. Phenotype severity correlated with longer latency, while its correlation with other saccade parameters was not universal. The underlying analyses data are provided in the Additional file 3, and the neurologic items are listed in the Additional file 4 [see Additional files 3 and 4].

Discussion

We report ocular involvement and saccadometric parameters in this to date largest monocentric ophthalmologic cohort of GD3.

Ophthalmologist’s findings

Diffuse corneal opacities have been infrequently reported in GD 3 patients [24]. They occurred in D409H homozygous patients [35–38] in association with cardiac valve calcification, while we found corneal haze in a heterozygous patient as did Inui et al. in toddler Gaucher type 2 [39]. Our heterozygous patient with corneal opacity was significantly older (44 years) than other patients. If the corneal opacity is due to the presence of the D409H mutation (albeit in the heterozygous compound state), it could be hypothesized that the absence of this finding in the other carriers of the same mutation (L444P/D409H) might be due to their still young age. In another case of Gaucher type 1 disease (F216Y/L444P), corneal abnormality preceded the diagnosis almost 15 years [40].

First descriptions on posterior segment abnormalities in Gaucher patients, before enzyme replacement therapy was established, focused on vitreous opacities [41, 42]. Recent case reports confirmed vitreous opacities, condensations, preretinal hyperreflective dots and posterior vitreous detachment by OCT [21, 23] [24]. Sheck et al. [23] located a hyperreflective preretinal accumulation between the posterior hyaloid interface and the nerve fiber layer of the retina in the mid peripheral and perimacular area (as also shown in Fig. 3). An additional sign in this 14-year-old girl was partial posterior vitreous detachment [23]. One year later in 2013, Coussa showed comparable OCT recordings in a 13-year-old girl – preretinal deposits - with slight progression after a follow up period of 5 years [21]. These are in accordance with our experiences in 16 GD3 patients, in which we found epiretinal particles in the macula, epiretinal and vitreous opacities in the mid periphery, and partial detachment of the posterior vitreous. However, in the present study, we did not only see vitreoretinal lesions (at the
interface), but also subretinal lesions. Their occurrence seems more typical with severe phenotype and longer disease duration. The OCT characteristics seem similar to drusen but unlike drusen, the color was whitish instead of yellowish, similar to the pre-retinal presumed Gaucher cell bodies. One could speculate that the drusen-like deposits are Gaucher cell collections. These lesions located at the peripapillary region, reaching the macula were accompanied by retinal atrophy per loss of photoreceptors and retinal pigment epithelium. We were unable to find comparable findings in a computerized literature search. While chorioretinal atrophy is described in other storage diseases [43], their appearance in these diseases is considerably less marked and without deposits.

Likewise, tortuosity of the retinal vessels is known from other lysosomal storage diseases such as Fabry disease and alpha-mannosidosis. Unlike the isolated finding of tortuosity in our cohort, a previously reported case of GD showed severe tortuosity with extensive opacities in the vitreous body accompanied by visual impairment [44]. Red cherry spot is no specific sign for Gaucher disease [24], which we could confirm. In our cohort, we could not detect any nerve fiber layer deposits as described by Sawicka-Gutaj et al. [22].
At the time being, we cannot answer the crucial question whether the occurrence of subretinal (or vitreoretinal) affection indicates a neuronopathic (or visceral) manifestation.

Watanabe [25] and Zhao [26] described tractional retinal detachment due to strong vitreoretinal adhesions and massive vitreous opacities in GD3. In teenagers, the underlying mechanism might include early development of liquefied cavity that exercises traction by the vitreous [25]. If retinal detachment develops, surgical treatment will be essential to preserve vision [25], otherwise permanent vision loss might occur at a young age [26].

Therefore, from the ophthalmologist’s point of view, Gaucher patients whether they are diagnosed type 1 or 3, should undergo the following diagnostic procedures:

- Macula and papillary dense optical coherence tomography (the resolution of OCT is 7–25 μm, which makes it a sensitive technique) helps to localize subretinal and vitreoretinal lesions.
- Fundoscopy (with dilated pupils) and fundus pictures enable to document the extent of the lesions in selected cases for follow-up. Differential diagnosis of these lesions – especially the distinction from intraocular lymphoma – is challenging, as GD
patients are known to have an increased risk of haematological malignancies [24]. Ideally, fundus photography should be performed by obtaining coloured and red-free recordings, as the case may be.

- Electroretinogram and visual field investigation might be informative to detect retinal involvement and optic disc anomalies or in case of otherwise unexplained visual impairment.

### Neuroophthalmologic aspects

GD3 patients develop a progressive horizontal supranuclear gaze palsy [15, 45]. Motility restriction, namely bilateral abduction deficits, indicate abducens motoneuron/nucleus affection [12], esotropia being the likely result of this N. abducens involvement. We found esotropia or decompensated esophoria in 8/18 patients (44%), which is in a comparable range as described earlier [11], and considerably more frequent than its prevalence in healthy subjects (2–3.5%). We could additionally demonstrate motility restriction involving all directions of gaze in 6/16 patients (38%). Oculo-motor disturbances follow a typical pattern in Gaucher type 3, in accordance to the topo-anatomical areas (PPRF (paramedian pontine reticular formation), rMLF (rostral interstitial nuclei of the medial longitudinal fasciculus), motoneurons of the abducens nucleus, flocculus/cerebellum and vestibular system [12, 18]. Besides slowed saccadic velocity, the initiation of saccades is delayed and [10] and saccadic gain (accuracy) is reduced, horizontally more severely than vertically [15]. Horizontal gaze may be affected so severely that technical measurement is impossible. Therefore, being a more sensitive measurement, the investigation of the less impaired vertical saccades has gained much more attention in saccadometric studies or possible effects of medical treatment [15, 17]. 2/16 of our patients exhibited saccades that were not reliably measurable due to motility restriction.

### Saccade parameters: peak velocity, latency and gain

**Peak velocity:** The relation between increasing peak velocity by increasing amplitude, termed main sequence in healthy subjects [46], is preserved in GD3 patients despite abnormal saccades. The overall saccadometric results of decelerated horizontal and secondary vertical
saccades confirms previous data on slowed saccades in GD3 patients [8, 10, 18].

Latency: As in previous reports, latency was found to be prolonged [10, 12, 15, 18], except for upwards saccades, which impair lastly. The results show enlarged interquartile distances in the boxplots, which in our view rather reflects the susceptibility for confounders such as age than for disease severity.

Gain: Saccadic accuracy (gain) received relatively little attention in studies on oculomotor involvement in GD3 patients. A study on Norrbottian GD3 has recently reported normal accuracy [18], while a previous report described it to be reduced [15]. In our cohort, horizontal saccades were less precise and rather hypometric. The same held true for vertical ones, although to a lesser extent. One of the reasons for low gain values in general is motility restriction, especially regarding the maximal target eccentricity. Anticonvulsive medication might be a confounder resulting in slowed and hypometric saccades [45]. We figured out the patients without anticonvulsive medication (12/16) to have slowed saccades (just like the patients with anticonvulsive medication (4/16)) with one exception corresponding to a mild genotype (N188/−).

Saccadometry is useful

Saccadometry was feasible in all Gaucher type 3 patients (14/16 performed saccadometry with sufficient data quality). We also recommend performing OCT as it is a widely used, expeditious, and informative investigation method feasible in less than 10 min in Gaucher patients, while saccadometry rather takes 10 to 30 min depending on the protocol used and the necessity to retake a test. The additional, typical signs of oculomotor abnormalities such as synkinetic blinking [8] are assessable by clinical evaluation. Another technique is an upwards curvature to horizontal saccades likely a Bell’s phenomenon (but keeping the lids open). Both are felt to inhibit the omnipause region allowing the saccade to begin. The latter phenomenon is noticeable during saccadometry and may disturb measurements by producing artefacts. Clinical assessment is difficult and less sensitive: While clinical assessment could not reveal differences between upwards and downward saccades, saccadometry was able to measure significant differences regarding this measurement. This is why we recommend to screen Gaucher type 1 patients and to monitor Gaucher type 3 patients for their neurologic manifestation by meticulous motility evaluation, ideally by video-oculography. We suggest that GD1 patients should be assessed for individual follow-up and that a normative GD1 cohort data, which does not exist to the best of our knowledge, should be established.

Infrared video-oculography is non-invasive, not stressful and easy to understand. It is well tolerated by children, however case-by-case-decision is surely advocated. The measurements are time consuming and editing of the raw data is needed to ensure optimum data acquisition. The quality of measurements depends primarily on the patient’s cooperation of the investigator’s experience.

Correlation between peak velocity and neurologic status (adjusted SARA, disease duration, IQ and to a slightly less extend the mSST) was greatest vertically. We were unable to reproduce the strongest correlation for downwards saccades as reported by Bremova (2018) [12]. However, the better preserved vertical spectrum of velocity allows stronger correlations. This fact has been explained by the horizontal floor effect [15] or ceiling effect [12]. Vertical saccadic peak velocity therefore functions as indicator for neurologic manifestation [12, 15, 17]. Saccadometry has the potential to qualify for those diagnostic procedures allowing quantification of neurologic manifestation such as the mSST. It might replace the less objective clinical saccade assessment and optimize the scoring system. Although correlation between clinical assessment of saccadic impairment and adjusted mSST, SARA and IQ were demonstrated, saccadometry is definitely more objective.

Strengths and limitations of this study

Despite the low prevalence of all GD-phenotypes of 0.7 to 1.75 per 100,000 inhabitants [47] and an amount of 5% GD3 [48] within these, we were able to investigate a group of 16 GD3 patients. Most studies reporting on oculomotor disorders in GD3 are based on smaller samples [8, 10, 14, 15, 18] or display a multicenter study design [12]. Schiffmann et al. performed saccadometry on 30 GD3 patients in two study centers as part of a randomized controlled study on miglustat [17]. One older

### Table 4 Latency of Gaucher patients vs. controls computing Wilcoxon rank sum test

| Target eccentricity | 5° R  | 15° R  | 30° R  | 5° L  | 15° L  | 30° L  |
|---------------------|------|-------|-------|------|-------|-------|
| P-value (t-test)    | *0,013 | *0,066 | 0,768 | *0,004 | *0,003 | 0,147 |

| Target eccentricity | 5° UP  | 10° UP  | 20° UP  | 5° DOWN  | 10° DOWN  | 20° DOWN  |
|---------------------|-------|--------|--------|----------|----------|----------|
| P-value (t-test)    | 0,128 | 0,584  | 0,855  | *0,048   | *0,071   | 0,817    |
report exists on clinical oculomotor and ophthalmological findings in 22 Norbottanian GD3 patients [11]. Statistical analysis of the limited samples encountered in rare diseases is challenging. In this explorative study, the significance level has not been adjusted for multiple testing, which may result in an excess of false positive results. Confirmatory studies are required to verify the results.

Another problem is a possible selection bias as the included patients mostly presented mild or intermediate phenotype severity. For patients with severely affected oculomotor disturbances, a favorable atmosphere for the measurement was created and the eyelid was fixed to minimize blinking artefacts.

A potential flaw of our investigation is the fact that in presbyopic study participants, no near correction was used. Quite interestingly, up to now there is no data on how near correction influences saccadometric precision [49]. Myopes and emmetropes show similar saccadic eye movements [50].

Regarding the pre-set program of the video oculography system we used, it would be useful to provide a shorter protocol to maintain attention and thus to reduce artefacts during saccadometry in patients with abnormal eye movements. As patients with restricted motility are not able to reach large target eccentricities, smaller maximal target displacements should be considered in those cases. Longitudinal saccadometry might be appropriate for monitoring and seems to be more sensitive than the clinical testing used in the scoring system.

**Conclusions**

Characteristic ocular manifestations of Gaucher disease type 3 may present as vitreoretinal lesions (vitreous opacities) and subretinal lesions (deposits with retinal atrophy in advanced disease stage). We recommend an ophthalmologic assessment including fundoscopy with dilated pupils and optical coherence tomography (optic nerve head and macula). The pattern of saccadic impairment in Gaucher type 3 affects predominantly horizontal, later downwards and finally upwards eye movements. Peak velocity vertically as a biomarker for neuropathic manifestation is valuable for future longitudinal studies, as it correlates best with other neurologic symptoms. We could additionally demonstrate motility restriction to involve all gaze directions, not only abduction deficits which cause esotropia in Gaucher patients. This study is valuable for ophthalmologists because they can contribute to diagnosis of Gaucher type 3 by neuro-ophthalmologic examination and may alert in case ocular manifestations are present indicating severe disease stage.

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s13023-019-1244-9.

- **Additional file 1.** Retinal vessels in Gaucher type 3 patients (Figure).
- **Additional file 2.** Saccadometric data including gain (A) and latency (B) in Gaucher type 3 patients compared to 100 controls split by target eccentricity (Boxplots).
- **Additional file 3.** Correlation analysis between saccade parameters and other neurologic items (Table).
- **Additional file 4.** Neurologic items in Gaucher type 3 patients (Table).

**Abbreviations**

GD3: Gaucher disease type 3; mSST: Modified severity scoring tool; mSST: Modified severity scoring tool; OCT: Optical coherence tomography; PPRF: Paramedian pontine reticular formation; riMLF: Rostral interstitial nuclei of the medial longitudinal fasciculus; RNFL: Retinal nerve fiber layer; RPE: Retinal pigment epithelium; SARA: Scale for the assessment and rating of ataxia.

**Acknowledgements**

We thank Dr. med. Katharina Bell of the Department of Ophthalmology (University Medical Center Mainz) for language-editing and Dr. med. Shahrzad Ashayer for her support in the study design.

**Authors’ contributions**

SH and ML investigated the participants for ophthalmologic and neuro-ophthalmologic findings (acquisition of data), recruited the participants, analyzed and interpreted the data (data management) and drafted the manuscript; EM recruited the patients and investigated the patients for neurologic symptoms, contributed the genetic and general patient data and interpreted the data; IS supported the statistical analyses, study design and power calculation; JBH contributed general patient data and conception of the work; SPF made enabled data collection, revised and supervised critically the work; SP made substantial contributions to conception and interpretation of the data and revising critically the manuscript for important intellectual content. All authors read and approved the final manuscript.

**Funding**

This research received no specific grant.

**Availability of data and materials**

Data generated or analyzed during this study are included in this published article.

**Ethics approval and consent to participate**

The study was approved by the Medical Ethical Committee of the State Chamber of Medicine of Rhineland Palatinate in Mainz, Germany (reference number 837.374.14 (9613)).

**Consent for publication**

The patients or their parents/guardians gave a written consent to publication of their anonymized clinical data.

**Competing interests**

The authors declare that they have no competing interests.

**Author details**

1Department of Ophthalmology, University Medical Center Mainz, Langenbecksstr.1, 55131 Mainz, Germany. 2Clinic for Anaesthesia, Surgical Intensive Care, Emergency Medicine and Pain Therapy, Klinikum Frankfurt Höchst, Frankfurt, Germany. 3Villa Metabolica, Center for Pediatric and Adolescent Medicine, University Medical Center Mainz, Mainz, Germany. 4Institute of Medical Biostatistics, Epidemiology and Informatics (IMBEI), University Medical Center Mainz, Mainz, Germany. 5Orbital Center, Ophthalmic Clinic, Bürgerhospital Frankfurt, Frankfurt, Germany.
References

1. Grabowski GA. Phenotype, diagnosis, and treatment of Gaucher’s disease. Lancet. 2008;372:1263–71.

2. Sidransky E. Gaucher disease: complexity in a “simple” disorder. Mol Genet Metab. 2004;83:6–15.

3. Sidransky E, Tzigi S, Martin BM, Stubblefield B, Ginns EI. DNA mutation analysis of Gaucher patients. Ann J Med Genet. 1992;42:331–6.

4. Beutler E. Discrepancies between genotype and phenotype in hematology: an important frontier. Blood. 2001;98:2597–602.

5. Goker-Alpan O, Hruska KS, Orvisky E, Kishnani PS, Stubblefield BK, Schiffermann R, et al. Divergent phenotypes in Gaucher disease implicate the role of modifiers. J Med Genet. 2005;42:e39.

6. Ida H, Rennert OM, Iwasawa K, Kobayashi M, Eto Y. Clinical and genetic studies of Japanese homozygotes for the Gaucher disease L444P mutation. Hum Genet. 1999;105:120–6.

7. Kopriwa V, Stone DL, Park JK, Callahan M, Frisch A, Cohen IJ, et al. Analysis of neurological involvement in Gaucher disease. Neuropediatrics. 1999;30:289–93.

8. Capablo J, Saenz de Cabezon A, Fraile J, Alfonso P, Pocovi M, Giraldo P, et al. Neurological evaluation of patients with Gaucher disease diagnosed as type 1. J Neurol Neurosurg Psychiatry. 2008;79:219–22.

9. Accardo AP, Perssino S, Perissiutti P. Saccadic analysis for early identification of neurological involvement in Gaucher disease. Ann N Y Acad Sci. 2005;1039:503–7.

10. Dreborg S, Erikson A, Hagberg B. Gaucher disease—Norrbottnian type. I. General clinical description. Eur J Pediatr. 1980;133:107–18.

11. Bremova-Ett T, Schiffermann R, Patterson MC, Belmatou N, Billiette de Veulemer T, Bardin S, et al. Oculomotor and Vestibular Findings in Gaucher Disease Type 3 and Their Correlation with Neurological Findings. Front Neurol. 2017;8:711.

12. Patterson MC, Horowitz M, Abel RB, Currie JN, Yu KT, Kanesi C, et al. Isolated horizontal supranuclear gaze palsy as a marker of severe systemic involvement in Gaucher’s disease. Neurology. 1993;43:1993–7.

13. Perssino S, Accardo A, Pitts MG, Ciana G, Bembali B, Perissiutti P. Saccade testing in the diagnosis and treatment of type 3 Gaucher disease. Neurology. 2005;65:1837.

14. Benko W, Ries M, Waggas GA, Brady RO, Schiffermann R, Fitzgibbon EJ. The saccadic and neurological deficits in type 3 Gaucher disease. PLoS One. 2011;6:e22410.

15. Garbutt S, Hanwood NR, Harris CM. Comparison of the main sequence of reflexive saccades and the quick phases of optokinetic nystagmus. Br J Ophthalmol. 2001;85:1477–83.

16. Schiffermann R, Fitzgibbon EJ, Harris C, De Ville C, Davies EH, Abel L, et al. Randomized, controlled trial of miglustat in Gaucher disease type 3. Ann Neurol. 2008;64:514–22.

17. Blume J, Benaminov S, Kampe Björkvik C, Machaczka M, Svenningson P. Saccadic impairments in patients with the Norrbottinian form of Gaucher’s disease type 3. Front Neurol. 2017;8:295.

18. Wolfstein G, Eitzen D, Strossman I, Seekenfreund M, Zylbermann R, Zimran A. Pretermal white dots in adult-type Gaucher disease. Retina Phila Pa. 1999;19:570–6.

19. Anand S, Kidd D, Hughes D. Photo essay: retinal changes in type 3 Gaucher disease. Am J Ophthalmol. 2019;103:315–26.

20. Watanabe A, Gelka T, Arii K, Tsunoeoka H. A case of traction retinal detachment in a patient with Gaucher disease. Ophthalmic Genet. 2017;38:273–6.

21. Sawto T-T, Li H-L, Guo X-J, Luo J, Liu X, Fang F. Retinal detachment in a boy with Gaucher disease. Int J Ophthalmol. 2018;11:1566–8.

22. Davies EH, Mengel E, Tylko-Szymanska A, Kleintoniene G, Reinke J, Vellodi A. Four-year follow-up of chronic neuropathic Gaucher disease in Europeans using a modified severity scoring tool. J Inherit Metab Dis. 2011;34:1053–9.

23. Schimpf-Hübch T, de Montcolot ST, Baliok I, Berciano J, Boeschen S, Depontd C, et al. Scale for the assessment and rating of ataxia: development of a new clinical scale. Neurology. 2006;66:1717–20.

24. Ericson A, Forsberg H, Nilsson M, Åström M, Månsson J-E. Ten years’ experience of enzyme infusion therapy of Norrbottinian (type 3) Gaucher disease. Acta Paediatr. 2006;95:312–7.

25. Machaczka M, Paucar M, Björkvik CK, NIC S, Cox TM, Forsgren L, et al. Novel hyperkinetic dystonia-like manifestation and neurological disease course of Swedish Gaucher patients. Blood Cells Mol Dis. 2018;68:86–92.

26. Park JK, Orvisky E, Tayeib N, Kanesi C, Lammers ME, Stubblefield BK, et al. Myoclonic epilepsy in Gaucher disease genotype-phenotype insights from a rare patient subgroup. Pediatr Res. 2003;53:387–95.

27. Davies EH, Surtees R, De Ville C, Schoon I, Vellodi A. A severity scoring tool to assess the neurological features of neuropathic Gaucher disease. J Inherit Metab Dis. 2007;30:768–82.

28. Hopf S, Liesenfeld M, Schmidtmand I, Ashayer S, Pitts S. Age dependent normative data of vertical and horizontal reflexive saccades. PLoS One. 2018;13:e0204008.

29. Zimmerman KP, le Coutre P, Aerts HM, Harzer K, Fukuda M, O’Brien JS, et al. Intracerebral transport of acid beta-glucosidase and lysosome-associated membrane proteins is affected in Gaucher’s disease (G202R mutation). J Pathol. 1999;188:407–14.

30. Abrahamov A, Elstein D, Gross-Tsur V, Farber B, Glaser Y, Hadash-Halpern I, et al. Gaucher’s disease variant characterised by progressive calcification of heart valves and unique genotype. Lancet. 1995;346:1000–3.

31. Chabas A, Combard B, Grignon D, Burguera JM, Balcells S, Mentino J, et al. Unusual expression of Gaucher’s disease cardiovascular calcifications in three sibs homozygous for the D409H mutation. J Med Genet. 1995;32:740–7.

32. Cindik N, Ozfay F, Sümen D, Akbıyık I, Gökdemir M, Varan B, et al. Gaucher disease with communicating hydrocephalus and cardiac involvement. Clin Cardiol. 2010;33:E26–30.

33. Guemes A, Kosmorsky GS, Moodie DS, Clark B, Meisler D, Traboulsi EI. Corneal opacities in Gaucher disease. Am J Ophthalmol. 1998;126:833–5.

34. Inui K, Yanagihara K, Otaki K, Suzuki Y, Akagi M, Nakayama M, et al. A new variant neuropathic type of Gaucher’s disease characterized by hydrocephalus, corneal opacities, deformed toes, and fibrous thickening of spleen and liver capsules. J Pediatr. 2001;138:137–9.

35. Geens S, Kestelyn P, Claes H. Corneal manifestations and in vivo confocal microscopy of Gaucher disease. Cornea. 2013;32:e169–72.

36. Cogan DG, Chu FC, Gittinger J, Tychsen L. Fundal abnormalities of Gaucher disease. Ophthalmic Genet. 2017;38:273–6.

Publisher’s Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.