Electrophysiological Study of Visual Pathways in Nevoid Basal Cell Carcinoma Syndrome Patients

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Introduction: Gorlin-Goltz syndrome (GGS) also known as nevoid basal cell carcinoma syndrome (NBCCS) is a complex rare genetic disorder characterized by a wide range of clinical and radiological manifestations. Ophthalmological alterations have always been reported, but no study on the eventual pattern visual evoked potentials (pVEPs) abnormalities has yet been published.

Purpose: The purpose of the study was to evaluate the functionality of the optic pathways in a group of NBCCS patients through pattern reversal VEPs, after a thorough exclusion of subjects with preexisting ocular and optic pathways pathologies.

Methods: Nineteen NBCCS patients (31 eyes) and 20 healthy controls (40 eyes) have been recruited for this study. All subjects underwent an evaluation of the functionality of the optic pathways through pVEPs with small (120′), medium (60′), and large (15′) check size stimulation.

Results: NBCCS patients showed a statistically significant alteration in the transmission of the macular pathway function when compared to controls. PVEPs analysis confirmed a reduced amplitude and an increased latency of the P100 component, suggesting an involvement of the visual pathway even in the absence of ocular clinical manifestations.

Conclusion: Visual pathways may have been affected both by a subclinical myelination deficit, determined directly by the genetic alteration, as well as by neurological abnormalities typical of this syndrome. Further studies are warranted.

Keywords: Gorlin-Goltz syndrome, nevoid basal cell carcinoma syndrome, pattern visual evoked potentials, rare diseases, electrophysiology, visual pathway, genodermatosis, PATCH1

Introduction

In 1960 Robert Gorlin and Robert Goltz first described a syndrome characterized by nevoid basal cell epithelioma, jaw cysts, and bifid ribs, nowadays known as Gorlin-Goltz syndrome (GGS) or nevoid basal cell carcinoma syndrome (NBCCS).\(^1\) NBCCS (OMIM 109400) is an autosomal dominant disease with near complete penetrance and variable expressivity. NBCCS is considered a rare disease and its prevalence, similar in men and women, is estimated between 1/57,000 to 1/256,000 with significant differences between races and ethnicity.\(^2,3\) This genodermatosis is determined by a mutation of the PATCH1 gene located on chromosome 9q22.3-q31. PATCH1 acts as a tumor suppressor gene and codes for the transmembrane glycoprotein Patched 1. Patched 1 plays a cardinal role in numerous pathways as it binds transcription factors such as Sonic Hedgehog (SHH). The most frequent mutations are insertion or deletion, but nonsense, missense, and splicing mutations
have been reported as well.\(^4\) Mutations of the PATCH1 gene are responsible for about 50–85% of the cases of NBCCS with higher estimates closer to 90% in more recent studies.\(^5\) The loss of function of PATCH1 determines the unchecked activation of cellular proliferative pathways.\(^6\) Recently two other genes have been recognized responsible for the development of NBCCS. PTCH2, highly homologous to PTCH1, located on chromosome 1p32.1-p32.3 and suppressor of the fused gene (SUFU) located on chromosome 10q.\(^7,8\) Interestingly, patients with SUFU mutation show an increased incidence of medulloblastoma when compared to patients carrying PATCH1 mutation.\(^9\)

NBCCS is a complex syndrome with more than 100 recognized clinical manifestation. Cardinal features are multiple basal cells carcinoma (BCC) that usually develop around the second or third decade but have reported in children as well;\(^10\) keratocystic odontogenic tumor, usually multiple lesions can be found in 90% of the affected patients above the age of 40;\(^11\) ribs or spine abnormalities such as bifid ribs and early calcification of the falx cerebri.\(^12,13\)

Major diagnostic criteria include multiple (>2) BCCs or 1 BCC by ≤30 years of age, or more than 10 basal cell naevi, any odontogenic keratocysts of the jaw proven by histology, palmar or plantar pitting, lamellar calcification of the falx cerebri and first-degree relative with NBCCS. Minor diagnostic criteria include congenital skeletal anomalies such as bifid, fused or missing ribs, medulloblastoma, ovarian and cardiac fibromas, lymphomesenteric cysts, increased circumference of the head, and other congenital malformations (frontal bossing, coarse facies, cleft lip or palate, moderate or severe hypertelorism, etc.).\(^10\) Diagnosis requires the concomitant presence of 2 major criteria and 1 minor criterion or 1 major criterion and 3 minor criteria. Oftentimes though, in developed countries diagnosis is achieved by genetic analysis on the PTHC1 gene or the other involved genes.\(^13\)

Numerous ocular manifestations have been reported in patients with NBCCS with a higher prevalence compared to the general populations. Most of these reports are sporadic and confined to case reports as only a few works have considered ocular manifestations of NBCCS.

Black et al demonstrated in 30 NBCCS patients the tendency to develop fibrogial epiretinal membrane (ERM), abnormal ganglion-cell axon myelination due to the inactivating mutation of the PATCH1 gene.\(^14\) Other common findings in NBCCS patients are hypertelorism, congenital cataract, strabismus, myelinated fibers, retinal hamartoma, optic nerve colobomas, nystagmus and palpebral ptosis.\(^15-18\)

The objective of the present study is to evaluate the integrity of the macular pathways and functionality in NBCCS patients by means of pattern reversal visual evoked potentials (pVEPs). The results obtained were compared with a control group (CG) of healthy subjects. In literature, there is only a single case report that studied pattern VEP changes in a patient with GGS, which found reduced amplitude in both eyes and delayed latency in one eye.\(^19\)

Materials and Methods

We recruited 19 patients affected by NBCCS for a total of 37 eyes (1 patient suffered from congenital unilateral anophthalmia) and 20 age-matched healthy controls for a total of 40 eyes. The two groups also matched for ethnicity (Caucasian) and sex. The NBCCS group consisted of 19 subjects 13 females and 6 males with a mean age of 45.5 years ±20.2 standard deviation (SD). NBCCS diagnosis was performed according to the Kimonis’s diagnostic criteria (two major criteria or one major and two minor criteria) during the dermatological exam that preceded the recruitment in the present study.\(^20\) Moreover, the molecular testing showed the variant of the PTCH1 gene in every patient included in the NBCCS group.

CG consisted of 20 subjects, 15 females, and 5 males (mean age 46.6 ±13.6 SD). Controls showed no manifest ocular pathology, nor systemic pathologies that could have altered the outcome of the study.

The enrollment took place between May 2019 and February 2020 at the Umberto I Policlinico. The study protocol was approved by the Ethics Committee of the Sapienza University of Rome. The research followed the tenets of the Declaration of Helsinki and informed consent was obtained by all the participants before the enrollment.

Both groups underwent complete ophthalmological examination that included: monocular best-corrected visual acuity (BCVA) measured to 5 meters with Snellen chart; intraocular pressure measurement with Goldman applanation tonometer; anterior segment biomicroscopy and mydriatic indirect fundus biomicroscopy. Furthermore, all patients and controls were examined with spectral-domain optical coherence tomography.
SD-OCT; Spectralis Family Acquisition Module, V5.1.3.0; Heidelberg Engineering with Heidelberg Eye Explorer V1.6.2.0).

In the NBCCS group, we excluded from the study eyes with visual acuity less than 10/10 (logMAR >0.00), refractive defects greater than ±4 diop ters (spherical equivalent), and simultaneous ocular pathologies such as glaucoma, epiretinal membrane, macular pucker, optic nerve coloboma, or cataract. This led to the exclusion of 6 eyes, therefore only 31 eyes in the NBCCS group were considered for both the statistical analysis and pVEPs recording, while all 40 eyes in the control group underwent pVEPs examination.

To record the pVEPs we placed 1 midline active skin-electrode on the scalp over the visual cortex at O2 with the reference electrode at Fz on the forehead, and 1 neutral electrode on the earlobe of the subjects. Each eye was examined separately at 100 cm with the optoelectronic stimulator Vision Monitor MonPack 120 by Metrovision (Pérenchies, France): stimulus field size 23.6 arc deg centered on the fovea, contrast 100%, mean luminance 50 cd/m2, and 2 reversals/s. The VEP were elicited by checkerboard stimuli with large (120`) (106 min arc), medium (60`) (59 min arc), and small (15`) (15.4 min arc) checks. The waveform of the transient examination was obtained at low temporal frequencies of the stimuli; then the values of the amplitude and latency (or time-to-peak) of P100 peaks were studied. A minimum of two recordings of each VEP condition were acquired, measured and displayed. After the recording, we manually analyzed 3 peaks N75, P100 and N135, each characterized by an amplitude and latency. The following results refer to the amplitude and latency of mean peak P100. The whole recording of pVEPs was performed adhering to ISCEV standards.21

Statistical analysis was performed with Excel for Windows. Descriptive statistical analysis was performed for all collected data. For each quantitative variable, mean, standard deviation and median were calculated. An ANOVA analysis was performed to test the hypothesis of equal means between the two groups, for each variable separately. The correlation matrix was calculated and, due to a high correlation between the measures, a p-value <0.01 was considered statistically significant. The assumption of equal variance in the two groups is not always evident, also from the inspection of the boxplots for the 6 variables. Hence, a t-test with not-equal variances (heteroscedastic) was applied in order to confirm ANOVA results (data not shown). The number of observations in each group is large enough to ensure that the t-test is robust even if some deviations from the normality assumption is observed.

### Results

The present study collected data from a total of 19 patients and 20 controls. The 2 groups were homogeneous for gender and age. Six eyes of 6 NBCCS patients had interfering conditions and were excluded from the statistical analysis. The ophthalmological examination highlighted several ocular abnormalities described in Table 1. The most common conditions were hypertelorism, strabismus, and epiretinal membranes. However, the ocular abnormalities (Figure 1) of the patients examined did not cause subjective disturbances in visual acuity. Visual acuity of all subjects included in this study was ≥10/10 (logMAR ≤0.00) with refractive error not exceeding 4 diop ters. The results obtained after the recording of pVEPs are shown in Table 2. For each check size to 120‘, 60‘ and 15‘ were evaluated mean, median, and SD for amplitude and latency of the P100 component. The boxplots comparing the distribution for each studied parameter are available in Supplementary Figure S1. We reported a statistically significant reduction in amplitude and an increase in latency of the P100 component in the NBCCS group when compared to the CG with both ANOVA and t-test. (Table 2)

### Discussion

The objective of the present study was to examine the macular pathway function in NBCCS patients without any significant ocular impairment confronting the results with a CG. To the best of our knowledge, this is the first VEP study to involve a relatively large number of patients (19 patients) in this rare condition. It is known that physiological factors such as age, gender, macular status, pupil

| Ocular Abnormalities                  | Frequency in GGS Group |
|---------------------------------------|------------------------|
| Hypertelorism                         | 7/19                   |
| Strabism                              | 8/19                   |
| Epiretinal membranes                  | 6/19                   |
| Myelinated fibers                     | 5/19                   |
| Congenital cataract                   | 3/19                   |
| Optic nerve coloboma                  | 1/19                   |
| Ptosis                                | 1/19                   |
| Palpebral cysts                       | 2/19                   |
| Anophthalmia                          | 1/19                   |
diameter, refractive errors, type of stimulus, electrode position and anatomical factors may affect pVEPs. To try to minimise the interference of these factors, the two groups were matched by age and sex, and strict exclusion criteria were applied.\(^{21-23}\)

In this research, the analysis of VEP demonstrated an increase in latency as well as a reduction in amplitude in the observed patient, indicative of a qualitative and quantitative disorder of the optic pathway conduction (Figure 2).\(^{10,24}\) The significant results, throughout all frequencies of spatial stimulation, suggest a dysfunction of the macular pathway in NBCCS patients compared to the CG.

It is known that myelin involvement in numerous diseases determines an alteration in VEP P100 wave latency and amplitude. The ongoing VEP analysis of optic neuritis in patients suffering from multiple sclerosis (MS) showed an increase latency of P100 wave with normal amplitude.\(^{25,26}\)
Interestingly, the amplitude reduction and the significant increase of the latency in NBCCS group in the transmission time of the optical impulse may correlate to a subclinical myelin alteration in NBCCS.\textsuperscript{14,19} Besides, we know that from a genetic point of view the activation of the SHH pathway is required for the development of the visual system. In fact, SHH expression is required for the induction of the optic nerve as well as photoreceptor differentiation.\textsuperscript{27}

It has been demonstrated that the inactivation of the PATCH gene may be responsible for the development of fibroglial epiretinal membrane and abnormal ganglion-cell axon myelination.\textsuperscript{18}

This is also shown by the common recurrence in our NBCCS group of epiretinal membranes and myelinated fibers. PATCH deficient murine models showed dysplastic foci in the retina and abnormal gliotic response further confirming the key role of SHH/PATCHED in glial cells

Table 2 Mean, Median and Standard Deviation (SD) for Each Check Size of Stimulation

| Gorlin-Goltz (n = 31 Eyes) | Controls (n = 40 Eyes) | ANOVA |
|---------------------------|------------------------|-------|
|                           |                       |       |
| pVEP120 ampl              | 11.6                   | 15.8  | <0.01 |
| pVEP120 lat               | 114.1                  | 102.1 | <0.01 |
| pVEP60 ampl               | 11.1                   | 16.6  | <0.01 |
| pVEP60 lat                | 114.5                  | 102.2 | <0.01 |
| pVEP15 ampl               | 11.5                   | 19.0  | <0.01 |
| pVEP15 lat                | 127.2                  | 111.2 | <0.01 |

Notes: Amplitude is expressed in microvolts = μV, and latency in milliseconds = ms. Significant results in bold.
Abbreviations: Ampl, amplitude; Lat, latency; SD, standard deviation.

Figure 2 Transient pattern visual evoked potentials (p-VEPs) in Gorlin-Goltz syndrome (GGS) patient. Values with increased latency and reduced amplitude of the P100 components, obtained for each spatial stimulation frequency (120°, 60°, and 15°). Diagram of normal transient VEP illustrating how measurements of negative and positive waves were made (bottom right).
homeostasis. Moreover, SHH is responsible for the development of oligodendrocytes progenitors and their differentiation into mature oligodendrocytes. These evidences suggest that the results obtained through the pVEPs may be caused by altered myelination at a subclinical level during the organogenesis.

The observed electrophysiological alterations could be determined also by other common manifestations of GGS. Dysgenesis/agenesis of the corpus callosum has been reported in around 10% of GGS patients. Corpus callosum is responsible for the interhemispheric connection in the brain including the occipital lobe where V1 is located. Moreover, the integrity of corpus callosum is necessary for the development and plasticity of the visual system.

Rugg et al studied the callosal function through VEPs elicited by lateralized stimuli in two patients affected by callosal agenesis. They found that VEPs from ipsilateral (to the visual field of stimulus) hemisphere were aberrant compared to contralateral, suggesting that VEPs for lateralized stimuli could be used to detect suspected callosal disease. Therefore, subclinical alterations in the development of this anatomic structure may be partially responsible for the observed results.

Other relatively common alterations in GGS such as falx cerebri calcification, bridging of sella turcica or cavum septum pellucidum could be also participate to the development of altered central conduction of the macular pathway function. Unfortunately, we were not able to find pre-existing literature on the role of these conditions. The association remains, therefore, purely speculative as magnetic resonance imaging (MRI) has not been yet performed on these patients; in the future this study will be a further object of evaluation for our research.

One consideration that we would still to discuss in patients with GGS is addressed to the dysmorphism linked to the congenital malformations of the skull which often presents with greater head circumference. This defect is associated with characteristic facies with frontal bossing, coarse facies, cleft lip, or palate, and moderate or severe hypertelorism.

Such anomalies could cause impaired functioning of the visual apparatus, as they have been described in the literature for other genetic conditions linked to cranial and cerebral malformations.

The cause could also be traced to an altered path and/or an extension of the optical pathways beyond the norm. Or the craniofacial malformations could be the primary cause responsible for abnormal ganglion-cell axon myelination, or the lack of integrity of corpus callosum necessary for the development and plasticity of the visual system. Moreover, also the falx cerebri calcifications and the alterations of the bridging of sella turcica or cavum septum pellucidum could be caused by the craniofacial malformations. The discussion becomes complex as the cranial and brain morphological anomalies could both coincide and cause each other. Many studies will have to investigate in detail these morphogenetic and functional aspects related to the hereditary pathologies of the Gorlin-Goltz syndrome. In addition, another parameter to be considered is whether such craniofacial malformations are associated with an increase in Intracranial Pressure (ICP), a factor capable of determining abnormal VEPs. None of our patients showed evidence of papilledema, but that alone does not rule out the possibility of increased ICP. Some of these concepts will be evaluated in future studies on patients examined by us through careful evaluation together with colleagues neuroradiologists.

In conclusion, due to the lack of preexisting literature on the specific topic, only hypotheses can be made on the obtained results. Further morphological, functional, and laboratory studies will be needed to correlate the present results with the common brain affections found in GGS patients.

Abbreviations
GGS, Gorlin-Goltz syndrome; NBCCS, nevoid basal cell carcinoma syndrome; pVEPs, pattern visual evoked potentials; SHH, Sonic Hedgehog; SUFU, Suppressor of the fused gene; BCC, basal cells carcinoma; ERM, epiretinal membrane.

Ethical Approval
All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki Declaration and its later amendments or comparable ethical standards.

Informed Consent
Informed consent was obtained from all individual participants included in the study.

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