Motor and functional evaluation of patients with spastic paraplegia, optic atrophy, and neuropathy (SPOAN)

Zodja Graciani1, Silvana Santos2, Lucia Inês Macedo-Souza3, Carlos Bandeira de Mello Monteiro4, Maria Isabel Veras5, Simone Amorim6, Mayana Zatz7, Fernando Kok8

ABSTRACT
Spastic paraplegia, optic atrophy, and neuropathy (SPOAN) is an autosomal recessive complicated form of hereditary spastic paraplegia, which is clinically defined by congenital optic atrophy, infancy-onset progressive spastic paraplegia and peripheral neuropathy. In this study, which included 61 individuals (age 5–72 years, 42 females) affected by SPOAN, a comprehensive motor and functional evaluation was performed, using modified Barthel index, modified Ashworth scale, hand grip strength measured with a hydraulic dynamometer and two hereditary spastic paraplegia scales. Modified Barthel index, which evaluate several functional aspects, was more sensitive to disclose disease progression than the spastic paraplegia scales. Spasticity showed a bimodal distribution, with both grades 1 (minimum) and 4 (maximum). Hand grip strength showed a moderate inverse correlation with age. Combination of early onset spastic paraplegia and progressive polyneuropathy make SPOAN disability overwhelming.

Key words: hereditary spastic paraplegia, peripheral nervous system disorder, optic atrophy, scales, psychomotor performance.

Avaliação motora e funcional de pacientes com paraplegia espástica, atrofia óptica e neuropatia (SPOAN)

RESUMO
A paraplegia espástica, atrofia óptica e neuropatia (SPOAN) é uma forma complicada de paraplegia espástica de herança autossômica recessiva, caracterizada por atrofia óptica congênita, paraplegia espástica progressiva de início na infância e neuropatia periférica. Este estudo avaliou o desempenho motor e funcional de 61 indivíduos com SPOAN (5 a 72 anos), por meio do índice de Barthel modificado, a escala modificada de Ashworth, da avaliação da força de preensão das mãos com dinamômetro hidráulico de Jamar e escalas de paraplegia espástica hereditária. O índice de Barthel modificado, que investiga aspectos funcionais, mostrou-se mais sensível para avaliar a progressão da doença do que as escalas de paraplegia espástica. A espasticidade apresentou distribuição bimodal, com o grau 1 (mínimo) e 4 (máximo). A força de preensão mostrou correlação inversa moderada com a idade. A combinação de paraplegia espástica de início precoce com polineuropatia progressiva faz da SPOAN uma condição incapacitante.

Palavras-chave: paraplegia espástica hereditária, doença do sistema nervoso periférico, atrofia óptica, escalas, performance psicomotora.
Spastic paraplegia, optic atrophy and neuropathy (SPOAN) (OMIM # 609641) is a neurodegenerative disorder recently described in Brazil by our group in individuals from Rio Grande do Norte State. This condition is characterized by: (a) congenital and non progressive optic atrophy; (b) progressive spastic paraplegia, with onset early in life, leading to loss of autonomous ambulation before adolescence; (c) axonal neuropathy, with clinical onset after the first decade of life and causing progressive loss of function of upper limbs; (d) startle response with unexpected sounds; (e) dysarthria; and (f) joint retractions and spine deformities.

Up to now, we are aware of 71 patients with SPOAN, belonging to 44 nuclear families. 41 of each are known to be consanguineous. Most of them are living in SW Rio Grande do Norte state, concentrated in Serrinha dos Pintos and São Miguel municipalities. It is estimated that 1 in every 170 inhabitants of Serrinha dos Pintos (population in 2000: 4,250) is affected by SPOAN. SPOAN is linked to chromosome 11q13, but its responsible gene remains unknown. No other form of hereditary spastic paraplegia, optic atrophy or neuropathy maps to the same chromosomal region of SPOAN.

The purpose of this study is to perform a quantitative motor and functional evaluation in a series of patients with SPOAN, in order to better understand disease progression and overall performance. As most of those individuals are living in small communities, with scarce healthcare resources, as a rule they previously had no access to any kind of structured rehabilitation program.

**METHOD**

**Patients**

Sixty-one individuals (42 females) with clinical diagnosis of SPOAN and in which genetic study demonstrates linkage to chromosome 11q13 participate of this study. Age at ascertainment ranged from 5 to 72 years [mean age of 34 (±13) years]. Clinical data of those patients have already been presented elsewhere. Information used in this study was collected through direct observation and with patients interview, usually conducted close to their hometown.

This study was approved by Biosciences Institute Ethics Committee and an informed consent for participation was obtained from patients or their caregivers.

**Procedures**

The following scales and quantitative instruments were used in this study:

1. The Spastic Paraplegia Rating Scale (SPRS), which evaluate 13 variables graded from 0 (normal) to 4 (severe). The maximum obtainable score is 52.

2. The Functional Hereditary Spastic Paraplegia Rating Scale (FHSPS), which is a semi-quantitative instrument to measure gait dysfunction level. It is graded from 0 (normal) to 5 (maximum).

3. The Modified Barthel Index (MBI), to evaluate functional performance in 10 functional categories (personal hygiene, bath, feeding, toilet, stairs climbing, dressing, gait, transfer ability, bowel and bladder control). Dependence level was ranked according to achieved performance in total (0-24 points), severe (25-50), average (51-75), and minimum (76-99).

4. Sural triceps, quadriceps and hip adductors spasticity was evaluated using Modified Ashworth Scale (MAS).

5. Left and right hand grip strength was measured using a Jamar hydraulic dynamometer (Asimow Eng., Trenton, Canada), following established standards. Mean value of three attempts with one minute interval was calculated for each hand. Reference values were supplied by the manufacturer.

**RESULTS**

For data analysis, sample was divided according to age in five groups: under 20 years, 20 to 29 years ([20, 30]), 30 to 39 years ([30, 40]), 40 to 49 years ([40, 50]) and above 50 years.

**Table.** Mean score, standard deviation, maximum and minimum values obtained in the five age groups with the implementation of Spastic Paraplegia Rating Scale.

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![Fig 1. Mean score obtained in five age groups in the Modified Barthel Index.](image-url)
SPRS was determined in 60 individuals (the 5-year-old subject was not included), and an average score of 40.7 points (±5.5) was obtained. Mean scores in five age ranges is presented in Table. Lowest individually achieved score was of 27 and highest of 52 points; 60% of individuals had score above 40 points. Mean score was not significantly different across 5 different ranges of age, and remained stable after 20 years of age, with average of 41 to 43 points. Therefore no correlation was seen between SPRS score and age.

Applying FHSPS to 61 individuals with SPOAN, it was determined that 44 (72.1%) had level 5, or maximum, of dysfunction, which is characterized by severe loss of gait and wheelchair use for more than 50% of time. 14 (22.9%) were at level 4, one (1.6%) at level 3 and one (1.6%) at level 2 of dysfunction.

BMI, used for functional assessment, disclosed maximum level of dependency in 28 individuals (45.9%), total dependency in 16 (26.2%), average in 14 (22.9%) and minimum in 2 (3.3%) subjects. Pearson linear correlation coefficient demonstrate a moderate correlation between scores and age, demonstrating that individual score decreases with age, which means that a functional deterioration occurs in SPOAN (Fig 1).

According to MAS, lower limbs spasticity had a bi-modal distribution, with subjects concentration in levels 1 (minimum) and 4 (maximum), as can be appreciated in Figure 2.

Hand grip measurement showed that strength decreases with age and that right hand grip was not different form left one (Fig 3).

**DISCUSSION**

SPOAN is a complicated form of HSP which leads to an early and severe handicap. Nevertheless, SPRS and FHSPS, the two available scales for HSP evaluation, were not useful indicators of disease progression, as most patients achieve early in life high scores in both scales. In this sense, MBI was more sensitive to appreciate disease progression, as it evaluates other functions, some of them highly dependable of upper limbs functionality.
As in SPOAN occurs a combination of upper and lower neuron compromise, which progress at different rates and have opposite effect on tonus, disease pattern can change overtime. Nevertheless, independently of age, in some individuals symptoms secondary to lower motoneurons compromise prevail and in other subjects, upper motoneuron clinical manifestations are dominant. Bimodal distribution of lower limbs tone, as seen with MAS, is probably secondary to this mutually opposed effect of spinal cord and motor neuropathy on tonus.

Hand grip evaluation allows quantification of age related strength deterioration, secondary to relentless progression of sensory and motor axonal neuropathy. Loss of hands strength correlates positively with neuropathy progression.

This study did not evaluate visual function, which might be contributory to dependency. Nevertheless, there is no evidence that in SPOAN there is a deterioration of visual function and therefore it is not expected to have a functional change. For lack of an adequate instrument, it was not analyzed the influence on overall performance of spine and joints deformities, present in variable degree in most patients.

In summary, this investigation allows quantify the motor and functional performance of 61 individuals diagnosed with SPOAN and suggests that this complicated form of HSP has a lifelong progression. Specific scales for HSP did not appreciate disease progression, as functional deterioration of lower limbs is an early event and achieves a maximum before 20 years of age. By the other hand, functional scales as BMI, with a more comprehensive inventory of evaluated items, perform better for investigation of disease progression and overall individual performance.

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REFERENCES