

Psicothema 2017, Vol. 29, No. 4, 446-452 doi: 10.7334/psicothema2016.298 ISSN 0214 - 9915 CODEN PSOTEG Copyright © 2017 Psicothema www.psicothema.com

brought to you by

Pre-symptomatic testing for neurodegenerative disorders: Middle- to long-term psychopathological impact

Susana Lêdo¹, Ângela Leite⁴, Teresa Souto², Maria Alzira Pimenta Dinis³ and Jorge Sequeiros¹ ¹ Universidade do Porto (Portugal), ² Lusophone University of Oporto (Portugal), ³ UFP Energy, Environment and Health Research Unit (FP-ENAS), University Fernando Pessoa (UFP) (Portugal) and ⁴ Universidade Europeia

Abstract

Background: Over the past 20 years, studies have revealed that the communication of a pre-symptomatic test (PST) result for late-onset diseases, such as Huntington's disease (HD), doesn't cause psychological disturbance. This cross-sectional study investigated the middle- (4 years) to long-term (7 and 10 years) psychological impact of PST for 3 autosomal dominant late-onset diseases: HD, Machado-Joseph disease (DMJ) and familial amyloid polyneuropathy (FAP). Method: The study included 203 subjects: 170 (84%) agreed to make the PST for FAP, 29 (14%) for HD and 4 (2%) for MJD. They were mostly women (58%) and married (67%). It was considered the cutoffs points: 4 years (middle-term) and 7 and 10 years (long-term) indicating the time after receiving the TPS results. Results: women and widows (oldest) presented the highest mean values for almost all BSI dimensions and the highest values correspond to the obsessivecompulsive dimension. MJD participants presented the highest mean values. No differences were found concerning the PST test results while participants are still asymptomatic. Psychopathology was only present in symptomatic carriers. Conclusions: The onset of the disease seems to assume the trigger for psychological disturbance, regardless the time that has elapsed since the PST result communication or the individual carrier/ non-carrier condition.

Keywords: Pre-symptomatic testing, psychological impact, late-onset genetic diseases.

Resumen

Pruebas pre-sintomáticas de enfermedades neurodegenerativas: el impacto psicopatológico a largo plazo. Antecedentes: el presente estudio transversal investigó el impacto psicopatológico a medio (4 años) y a largo plazo (7 y 10 años) de la prueba pre-sintomática (PPS) para tres enfermedades autosómicas dominantes de aparición tardía: enfermedad de Huntington EH, la enfermedad de Machado-Joseph (EMJ) y la polineuropatía amiloide familiar (PAF). Método: participaron 203 sujetos: 170 (84 %) realizaron el PPS para PAF, 29 (14 %) para EH y 4 (2 %) para EMJ. La muestra, en su mayoría, estuvo compuesta por mujeres (58 %) y por personas casadas (67 %). Fueron considerados como puntos de corte los 4, 7 y 10 años después de haber recibido el resultado de la PPS. Resultados: las mujeres y los viudos presentan las medias más altas. Los participantes con EMJ presentaron las medias más elevadas. No se encontraron diferencias significativas en lo concerniente a los resultados de PPS. La perturbación psicológica fue escasamente observada en los sujetos portadores que ya evidenciaban síntomas. Conclusiones: la aparición de los primeros síntomas parece constituir el detonante para la existencia de perturbaciones psicológicas, independientemente del intervalo de tiempo sucedido desde la comunicación de los resultados de la PPS o de la condición genética (portador/no portador).

Palabras clave: prueba pre-sintomática, impacto psicopatológico, enfermedades genéticas de aparición tardía.

The predictive testing (PST) model for late-onset neurodegenerative diseases such as Huntington's disease (HD), a rare disorder with a prevalence of ~1-7 in 100,000 individuals of European ancestry (Ramos et al., 2015), has been implemented and adapted for other late-onset diseases around the world (Hawkins, Ho, & Hayden, 2011), namely, Machado-Joseph disease (MJD) and Familial Amyloidotic Polyneuropathy (FAP), two Portuguese monogenic, autosomal and dominant diseases (Sequeiros et al., 2006), with a severe neurodegenerative evolution and no effective cure. In Continental Portugal, MJD has a prevalence of 1: 100,000, and is considered a rare disease, except for the area of the Tejo Valley (1: 1000) (Bettencourt & Lima, 2011); signs of cerebellar ataxia, progressive external ophthalmoplegia and pyramidal signs are reported (Coutinho, 1996; Sequeiros et al., 2006). Although Amyloidosis is very rare (less than 1 case in 100,000 people worldwide) it is more frequent in some countries such as Portugal, wherein the Val30Met mutation occurs in 1 in 1,000 people in areas of higher incidence like Póvoa Varzim/Vila do Conde, the most likely focus of origin of the disease (with 1/3 of total Portuguese patients). Sousa (1995) described a disease prevalence of 90.3/100,000 and an average age of onset of 33.5 years. For FAP, an abnormal amyloid protein (TTR) is deposited in various organs leading patients to experience progressive limitations (Saraiva, 1986).

There are several studies about the PST psychosocial short-term impact (one year) that did not demonstrate a severe negative impact (Lêdo, Leite, & Sequeiros, 2013; Lêdo, Paneque, Rocha, Leite,

Received: September 28, 2016 • Accepted: April 20, 2017 Corresponding author: Susana Lêdo Instituto de Investigação e Inovação em Saúde Universidade do Porto 4200-1 Porto (Portugal) e-mail: susanaledo@gmail.com

& Sequeiros, 2013; Rolim et al., 2006; Tibben, 2007). However, there are few studies investigating the PST psychosocial impact in the mid- to long-term (Almqvist et al., 2003; Decruyenaere et al., 2003; Decruyenaere et al., 2004; Gargiulo et al., 2009; Gonzalez et al., 2012; Timman, Roos, Maat-Kievit, & Tibben, 2004).

At the Center for Predictive and Preventive Genetics (CGPP). Institute of Molecular and Cell Biology (IBMC), University of Porto (Portugal), a national reference model was developed for one year of genetic counseling protocol for individuals at-risk for HD, MJD and FAP (Rolim et al., 2006). Lêdo and colleagues (2013) studied the psychopathological impact on this population a year after the PST protocol and noticed that values decreased significantly one year after the implementation of the PST, regardless of the disease studied or the test result; however, for all Brief Symptom Inventory (BSI) dimensions and global indexes, significantly higher values were found than those of control groups. Therefore, it became a priority to study the mid- to long-term PST psychological impact as a result of communicating the genetic status to subjects that underwent the PST, and compare the results obtained with those of the few studies for HD (Decruyenaere et al., 2003; Decruyenaere et al., 2004; Gargiulo et al., 2009; Timman et al., 2004). The main aim of this research is to increase the knowledge about followup studies investigating the long-term consequences of PST, as suggested by Timman and colleagues (2004) so that the adjustment of psychological support for this population may be possible in the context of the Portuguese reality.

Method

This research is a descriptive cross-sectional study, resulting from the compilation of the medical records of the subjects who completed the one year PST protocol at CGPP (including the molecular study) and discovered their genetic status at least three years ago, for three autosomal dominant late-onset conditions: HD, MJD and FAP.

Participants

Fifty eight percent of subjects were female and the majority of the responses correspond to age ranging up to 30 years [21-77]. The majority of subjects had mainly professions involving some responsibilities (1st Graffar Index) and a high level of education. Most of the subjects underwent the PST for FAP (84%) and 37% were identified as carriers; of these, 15% had become symptomatic and 5% having had a liver transplant (Table 1).

Out of the 203 subjects, 32% had been informed about their genetic status 4 years ago (middle-term), 47% 7 years ago and 21% 10 years ago (the long-term). Concerning the three different cutoff points, data were similar to data observed in the total sample, with the exception of those subjects who completed the protocol 10 years ago, where the age increases (31-40 years) and, consequently, raising the number of pensioners. Those subjects, who underwent the PST protocol 7 years ago, present the highest mean age and many of them are already retired.

Instruments

The socio-demographic data - gender, age, profession, and current marital status - were collected from a questionnaire sent to all participants. The questionnaire sent to carriers also included the following questions (clinical variables) "Current clinical

status", "*Still without symptoms*?", and "*Had a significant change in your life in recent years*?". No clinical variables were included in the questionnaire to non-carriers.

The dependent variable *psychopathology* was assessed using the BSI (Derogatis, 1993) adapted for the Portuguese population by Canavarro (2007). This instrument is composed of 53 items, rated on a Likert scale of five grades (0 "rarely" to 4 "very often"), nine dimensions and three global indexes which express psychometric ratings of emotional distress: global severity index (GSI), positive symptoms total (PSTI) and positive symptom distress index (PSDI).

Procedure

This study was accepted by the IBMC ethics committee. Information about the researcher, the nature and objectives of the study and the principle of confidentiality was displayed when the subjects were originally registered in the PST protocol. Participants were contacted by mail in order to answer the questionnaire that included sociodemographic and clinical data and the BSI.

Data analysis

The statistical analysis was performed using the PASW Statistics Program, version 22.0. Descriptive [frequencies (N

Table 1 Sample description					
		Frequencies (N = 203)	Percentage (100%)		
Gender	Female	118	58.1		
o en der	Male	85	41.9		
	≤ 30 years	88	43.3		
	31 - 40 years	62	30.5		
	41- 50 years	20	9.9		
Age	51 - 60 years	22	10.8		
	61 - 70 years	8	3.9		
	≥ 71 years	3	1.5		
	Single	53	26.6		
M 1 10.	Married	132	66.5		
Marital Status	Divorced	10	5.0		
	Widow	3	1.5		
	Retired	40	19.7		
	Unemployed	19	9.4		
	Student	15	7.4		
Destaution	1 St Graffar Index	59	29.1		
Profession	2 nd Graffar Index	11	5.4		
	3rd Graffar Index	4	2.0		
	4th Graffar Index	22	10.8		
	5th Graffar Index	33	16.3		
	HD	29	14.3		
Type of Disease	MJD	4	2.0		
	FAP	170	83.7		
DCT Docult	Non-carrier	91	44.8		
PS1 Result	Carrier	112	55.2		
	Non-carrier	89	44.5		
	Asymptomatic Carrier	73	36.5		
Clinical Status	Symptomatic Carrier	20	14.5		
	Liver Transplanted (FAP carriers)	9	4.5		

and n, mean (M) and standard deviation (SD)] and inferential [bi-variate statistical (ANOVA, chi-square test and bi-varying correlation)] analyses were carried out.

Results

BSI descriptive analysis

When considering the three cutoff points of 4, 7 and 10 years, mean and standard deviation were very similar to the ones relating to BSI scores (Table 2). For the three cutoff points, α was similarly high, pointing to a good scale reliability. Comparisons of means were performed but no statistically significant differences were found.

Comparison between the BSI means regarding independent variables

Analyzing the mean values for the BSI variables, regarding socio-demographic and clinical variables, some statistically significant results were found:

BSI total, nine dimensi	ons and t of	<i>Table</i> hree index 4, 7 and 1	2 kes (<i>M</i> , <i>SD</i> 0 years	and α) at	the cutoff j	points	
	4 years (<i>n</i> = 65)		7 years (<i>n</i> = 95)		10 years (<i>n</i> = 42)		
	М	SD	М	SD	М	SD	
	(1	α)	(1	α)	(4	α)	
BSI Total	36.28	33.72	36.65	28.24	31.32	24.41	
DSI Iotal	(.9	97)	.9	(.97)		96)	
Competing time	4.15	5.28	4.39	4.67	3.01	3.51	
Somauzation	3.)	38)	3.)	37)	(.80)		
01 . 1.	5.42	4.71	5.85	4.49	5.13	3.79	
Obsessive-compulsive	8.)	86)	3.)	37)	(.1	17)	
Texterna and the state	2.72	2.82	2.97	2.61	2.55	2.45	
Interpersonal sensitivity	8.)	30)	(.7	79)	(.1	/8)	
Deserve	4.58	4.97	4.53	4.49	3.25	3.00	
Depression	8.)	38)	3.)	39)	(.1	17)	
A 1.	3.75	4.19	3.96	3.37	3.61	3.27	
Anxiety	(.84)		(.7	(.78)		(.84)	
T T	3.97	3.85	4.15	3.47	3.30	2.70	
Hostility	(.85)		3.)	(.86)		(.75)	
DI 11 1	1.68	3.31	1.62	2.55	2.00	2.75	
Phobic anxiety	(.85)		(.7	(.77)		(.67)	
D	4.49	4.38	4.54	3.76	4.40	3.41	
Paranoid ideation	3.)	34)	3.)	33)	(.1	/6)	
D	2.72	3.42	2.51	3.21	1.90	2.48	
Psychoticism	(.75)		(.82)		(.77)		
GSI	0.68	0.64	0.69	0.53	0.59	0.46	
PSTI	21.02	14.19	23.00	14.87	20.10	12.43	
PSDI	1.55	0.53	1.42	0.44	1.41	0.37	

Note: BSI = Brief Symptom Inventory; GSI = Global Severity Index; PSTI = Positive Symptoms Total Index; PSDI = Positive Symptom Distress Index; M = Mean; SD = Standard deviation; α = Cronbach's alpha

Socio-demographic variables

Gender variable. Female presented higher mean values than men for the BSI total scores and for BSI somatization, interpersonal sensitivity, depression and phobic anxiety dimensions. The mean values for GSI and PSTI revealed statistically significant differences (Table 3).

Age variable. Significant results were found in the obsessive-compulsive dimension - F (5, 189) = 2.325, p = .045; η^2 = .058 - and in the PSTI – F (5, 189) = 2.551, p = .029; η^2 = .066 -, meaning that mean age and values increase in a similar direction. The exception occurred with older subjects that presented the lower averages for the same obsessive-compulsive dimension, $M_{\leq 30}$ (n = 88) = 4.71; M_{31-40} (n = 62) = 5.71; M_{41-50} (n = 20) = 6.22; M_{51-60} (n = 22) = 7.00; M_{61-70} (n = 8) = 9.29; $M_{\geq 71}$ (n = 3) = 4.33. The same trend is verified for the PSTI - $M_{\leq 30}$ (n = 88) = 21.24; M_{31-40} (n = 62) = 22.37; M_{41-50} (n = 22) = 23.73; M_{61-70} (n = 8) = 21.75; $M_{\geq 71}$ (n = 3) = 13.67.

Marital status variable. Widows presented significantly lower mean values for almost all dimensions than single, married and divorced subjects; and divorced subjects the highest for all the BSI dimensions (Table 4).

Clinical variables

Type of disease. Phobic anxiety [*F* (2, 192) = 9.434, *p* = .000; η^2 = .091], *psychoticism* [*F* (2, 192) = 3.958, *p* = .021; η^2 = .040] and PSDI values [*F* (2, 192) = 5.170, *p* = .007; η^2 = .054] presented significant differences. MJD subjects showed higher mean values than FAP and HD subjects, regarding *phobic anxiety* [*M*_{HD} (*n* = 29) = 1.15; *M*_{MID} (*n* = 4) = 7.50; *M*_{FAP} (*n* = 170) = 1.70], *psychoticism* [*M*_{HD} (*n* = 29) = 2.35; *M*_{MID} (*n* = 4) = 6.75; *M*_{FAP} (*n* = 170) = 2.34] and PSDI [*M*_{HD} (*n* = 29) = 1.47; *M*_{MID} (*n* = 4) = 2.27; *M*_{FAP} (*n* = 170) = 1.44].

PST results variable. Significant differences were found in *somatization* [*F* (1, 193) = 6.035, *p* = .015; η^2 = .029] and PSDI [*F* (1, 193) = 4.569, *p* = .034; η^2 = .021], where carriers (c) showed higher mean values than non-carriers (nc): *somatization:* M_c (*n* = 112) = 4.78 and M_{nc} (*n* = 91) = 3.14; PSDI: M_c (*n* = 112) = 1.52 and M_{nc} (*n* = 91) = 1.37. *Current clinical status variable.* Statistical significant values were found in *somatization* [*F* (3, 189) = 7.451, *p* = .000; η^2 = .104] and PSTI [*F* (3, 189) = 3.269, *p* = .023; η^2 = .048]. Non-carriers [*M* (*n* = 89) = 3.14] and asymptomatic carriers [*M* (*n* = 73) = 3.58] had lower mean values than symptomatic carriers [*M* (*n* = 9) = 5.43]. For PSTI, non-carriers [*M* (*n* = 89) = 21.72], asymptomatic carriers [*M* (*n* = 9) = 17.44] presented lower mean values than symptomatic carriers values than symptomatic carriers [*M* (*n* = 9) = 27.00].

"Still without symptoms?" variable. In the carriers group, significant differences were found for all BSI dimensions and GSI, PSTI and PSDI, except for the *phobic anxiety* dimension (Table 5): subjects who still had no symptoms had lower mean values than those who already had symptoms; subjects that answered "perhaps" were those with the highest mean values. Carriers that answered "no" (n = 30) or "perhaps" (n = 9) presented significant differences for *somatization* [F (3, 189) = 3.966, p = .016; $\eta 2 = .218$].

Subjects that considered having severe [M (n = 11) = 11.33]and moderate [M (n = 7) = 10.86] symptoms had higher means in *somatization* than those subjects that present minimal symptoms

<i>Table 3</i> Comparison of means for the gender variable: BSI total, four dimensions and two indexes $(M, F, df, p \text{ and } \eta^2)$							
	п	M (SD)	F	df	р	η^2	
BSI Total	Female (118) Male (85)	39.48 (33.01) 29.87 (22.37)	5.007	1,193	.026	.026	
Somatization	Female (118) Male (85)	4.81 (5.24) 2.99 (3.48)	7.397	1,193	.007	.037	
Interpersonal sensitivity	Female (118) Male (85)	3.13 (2.77) 2.33 (2.36)	4.480	1,193	.036	.023	
Depression	Female (118) Male (85)	4.91 (4.70) 3.40 (3.80)	5.701	1,193	.018	.029	
Phobic anxiety	Female (118) Male (85)	2.17 (3.31) 1.11 (1.89)	6.596	1,193	.011	.033	
GSI	Female (118) Male (85)	0.75 (0.62) 0.56 (0.42)	5.007	1,193	.026	.026	
PSTI	Female (118) Male (85)	23.55 (14.70) 19.51 (13.19)	4.070	1,193	.045	.020	

Note: BSI = Brief Symptom Inventory; GSI = Global Severity Index; PSTI = Positive Symptoms Total Index; M = Mean; F = Snedecor's F Distribution; df = degrees of freedom; p = p-value; η^2 = effect size

-			I total, six dimensions and one index $(M, F, df, p and \eta^2)$						
	п	M (SD)	F	df	р	η^2			
	Single (104)	29.36 (25.44)			010				
BSI Total	Married (89)	36.85 (28.71)	3 800	2		060			
DSI Iotai	Divorced (5)	62.22 (50.43)	5.077	5	.010	.000			
	Widow (2)	15.67 (2.89)							
	Single (104)	2.60 (3.55)							
Comptingation	Married (89)	4.51 (4.54)	4 160	2	007	.066			
Somauzation	Divorced (5)	7.33 (9.30)	4.109	2	.007				
	Widow (2)	1.00 (1.73)							
	Single (104)	4.66 (3.14)		3 .006 3 .007					
01	Married (89)	5.60 (4.46)	4.025	2	007	064			
Obsessive-compulsive	Divorced (5)	10.22 (7.98)	4.235	3	.006	.004			
	Widow (2)	5.67 (0.58)							
	Single (104)	4.11 (4.41)	4.121	3	.007	.062			
	Married (89)	4.09 (4.13)							
Depression	Divorced (5)	9.11 (6.72)							
	Widow (2)	2.00 (0.00)							
	Single (104)	3.38 (3.17)	4.121 3 2.762 3						
Apprinter	Married (89)	3.90 (3.55)		0.42	042				
Anxiety	$ \begin{array}{ccccc} \text{binded (5)} & \text{if } 0.22 (1.50) \\ & \text{Widow (2)} & 5.67 (0.58) \\ & \text{Single (104)} & 4.11 (4.1) \\ & \text{Married (89)} & 4.09 (4.13) \\ & \text{Divorced (5)} & 9.11 (6.72) \\ & \text{Widow (2)} & 2.00 (0.00) \\ & \text{Single (104)} & 3.38 (3.17) \\ & \text{Married (89)} & 3.90 (3.55) \\ & \text{Divorced (5)} & 6.78 (6.57) \\ & \text{Widow (2)} & 1.33 (0.58) \\ & \text{Widow (2)} & 1.33 (0.58) \\ \end{array} $.045	.045						
	Widow (2)	1.33 (0.58)							
	Single (104)	0.68 (1.17)		 3.007 3.006 3.007 3.007 3.043 3.003 3.011 					
Dhahia anviatu	Married (89)	2.10 (3.12)	4 790		002	072			
sessive-compulsive pression xiety pbic anxiety	Divorced (5)	3.56 (4.53)	4.789		.005	.073			
	Widow (2)	0.33 (0.58)							
	Single (104)	3.49 (3.21)	2.820	3	011				
Demonsid idention	Married (89)	4.80 (3.92)				059			
Paranoid ideation	Divorced (5)	7.33 (5.92)	5.820		.011	.058			
	Widow (2)	1.33 (1.15)							
	Single (104)	0.55 (0.48)	2,000	2	010				
CSI	Married (89)	0.69 (0.54)				0.00			
031	Divorced (5)	1.17 (0.95)	3.899	5	.010	.060			
	Widow (2)	0.29 (0.54)							

Table 5 Comparison of means for the variable "Still without symptoms?": BSI total, eight dimensions and three indexes $(M, F, df, p \text{ and } \eta^2)$						
	n	M (SD)	F	df	р	η^2
BSI Total	Yes (67) Perhaps (9) No (30)	29.62 (25.76) 67.22 (35.09) 44.92 (33.97)	8.042	2	.001	.142
Somatization	Yes (67) Perhaps (9) No (30)	3.06 (4.22) 10.44 (6.60) 7.00 (5.20)	13.925	2	.000	.218
Obsessive-compulsive	Yes (67) Perhaps (9) No (30)	4.88 (4.20) 9.89 (5.03) 6.48 (5.24)	5.264	2	.007	.095
Interpersonal sensitivity	Yes (67) Perhaps (9) No (30)	2.43 (2.26) 5.44 (2.65) 3.19 (3.10)	5.823	2	.004	.104
Depression	Yes (67) Perhaps (9) No (30)	3.40 (3.75) 7.56 (4.59) 5.44 (4.88)	5.377	2	.006	.097
Anxiety	Yes (67) Perhaps (9) No (30)	3.24 (3.30) 6.33 (3.81) 5.11 (4.14)	4.682	2	.011	.086
Hostility	Yes (67) Perhaps (9) No (30)	3.51 (3.56) 7.56 (5.66) 5.07 (4.28)	4.888	2	.009	.089
Paranoid ideation	Yes (67) Perhaps (9) No (30)	3.66 (3.35) 6.56 (5.68) 5.48 (4.40)	3.675	2	.029	.068
Psychoticism	Yes (67) Perhaps (9) No (30)	1.75 (2.45) 4.89 (4.57) 3.31 (4.13)	5.293	2	.007	.097
GSI	Yes (67) Perhaps (9) No (30)	0.56 (0.49) 1.27 (0.66) 0.85 (0.64)	8.042	2	.001	.142
PSTI	Yes (67) Perhaps (9) No (30)	19.61 (12.81) 35.89 (10.09) 23.90 (15.99)	6.002	2	.003	.104
PSDI	Yes (67) Perhaps (9) No (30)	1.41 (0.45) 1.81 (0.65) 1.67 (0.51)	4.453	2	.014	.085
Note: BSI = Brief Symptom Inventory; C	GSI = Global Severity Index; PSTI = Positi	ve Symptoms Total Index; PSDI =	Positive Symptom Di	stress Index; $M = N$	Mean; $F = $ Snedecor	's F Distribution;

Note: BSI = Brief Symptom Inventory; GSI = Global Severity Index; PSTI = Positive Symptoms Total Index; PSDI = Positive Symptom Distress Index; M = Mean; F = Snedecor's F Distribution; df = degrees of freedom; p = p-value; η^2 = effect size

[M (n = 16) = 5.73] and those who didn't specified their symptoms in the questionnaire (n = 5).

"Has there been a significant change in your life (marriage, divorce, death of loved one, illness, job change, earth, etc.) in recent years?" variable. Significant differences were found concerning somatization, depression, anxiety and hostility dimensions and GSI and PSTI, emphasizing that carriers who experienced meaningful life changes presented higher mean values than those that did not (Table 6).

Discussion

This study does not corroborate previous studies suggesting the absence of negative psychological impact resulting from the PST long-term outcome (Timman et al., 2004), because the BSI dimensions values obtained in this sample were higher than impact (Lêdo et al., 2013b) and when compared with the standard values reached for the Portuguese population. Nevertheless, and regarding the GSI, PSTI and PSDI, the obtained values did not reflect the presence of clinically psychological disturbance since the PSDI was always inferior to 1.7 (Canavarro, 2007). The lower GSI, PSTI and PSDI scores may be justified by the existence of a self-selection prior to PST of those subjects who were psychologically more prepared (Codori et al., 1994; Paneque et al., 2007; Rolim et al., 2006; Tibben, 2007) and that could be the same who responded to the present study. Subjects who were less psychological disturbed prior to the PST were those who did not drop out of the follow-ups or did not avoid the reality of the disease (Timman et al., 2004).

those obtained in previous studies about short-term psychological

Age variable and the obsessive-compulsive dimension presented a positive correlation. This may be explained due to

Comparison of means for the variable "Has there been a significant change in your life in recent years?": BSI total, four dimensions and two indexes $(M, F, df, p \text{ and } \eta^2)$								
	п	M (SD)	F	df	р	η^2		
BSI Total	No (54) Yes (56)	29.27 (27.43) 45.77 (33.09)	7.631	1	.007	.070		
Somatization	No (54) Yes (56)	3.58 (4.39) 5.94 (5.76)	5.666	1	.019	.051		
Depression	No (54) Yes (56)	3.34 (3.97) 5.61 (4.87)	6.991	1	.009	.062		
Anxiety	No (54) Yes (56)	3.00 (3.25) 5.21 (3.98)	9.798	1	.002	.086		
Hostility	No (54) Yes (56)	3.49 (3.68) 5.15 (4.34)	4.528	1	.036	.041		
GSI	No (54) Yes (56)	0.55 (0.52) 0.86 (0.62)	7.631	1	.007	.070		
PSTI	No (54) Yes (56)	19.41 (13.13) 25.66 (15.15)	5.334	1	.023	.047		

Note: BSI = Brief Symptom Inventory; GSI = Global Severity Index; PSTI = Positive Symptoms Total Index; M = Mean; F = Snedecor's F Distribution; df = degrees of freedom; $p = p - \eta^2$ = effect size

a progressive concern with the outbreak of the first symptoms, as suggested by Licklederer, Wolff and Barth (2008). Divorced subjects were those who presented the highest values in almost all BSI dimensions and widows the lowest: that points to the importance of the real/imaginary experiences of rejection/ abandonment as realities that might be interfering with these results, instead of the real experiences of loss or feelings of loneliness that might be present with the widowhood (Lêdo et al., 2013b).

Subjects who underwent PST for MJD presented the highest values in *psychoticism* and *phobic anxiety* maybe because MJD patients have shown some emotional changes related to disruption of frontal-subcortical systems (Zawacki, Grace, Friedman, & Sudarsky, 2002) and cognitive disorders (Rezende et al., 2015); although the MJD sample dimension is very small and a previous study conducted with subjects at-risk for MJD showed no psychological disturbance at post-test and after they knew their carriers/non-carriers status (Gonzalez et al., 2004). Additionally, FAP group is aware of a therapeutic solution that prevents progression of the disease to an advanced state (Coelho, Maia, Martins da Silva, Waddington, Planté-Bordeneuve, Lozeron et al., 2012).

All carriers had the highest values in *somatization* and PSDI, what is understandable since the carrier condition leads them to be more focused on their physical and body sensations, suggesting that higher levels of *somatization* are associated with real symptoms. The presence of real symptoms appeared to increase the tendency for these individuals to report somatic reactions (Lêdo, Leite, Souto, Dinis, & Sequeiros, 2016; Licklederer et al., 2008), although the perspective that they may probably already have symptoms let them more disturbed than the certainty of having symptoms (Licklederer et al., 2008).

Subjects who had significant changes in their lives were those who presented higher values in *somatization*, *anxiety* and *hostility* dimensions. Being one of the mentioned changes reported the "loss or illness of a close relative" item, it explains the high values on the referred dimensions (Lêdo et al., 2016).

Although the results did not present differences between the psychological impact in the mid- to long-term, they suggest that this impact exists but without being possible to differentiate it regarding the time resulting from the completion and notification of the PST result. So, the age of symptoms onset was not recognized as being determinant to the level of psychological disturbance (Lêdo et al., 2016; Licklederer et al., 2008).

References

- Almqvist, E. W., Bloch, M., & Hayden, M. (1999). A worldwide assessment of the frequency of suicide, suicide attempts, or psychiatric hospitalization after predictive testing for Huntington Disease. *American Journal of Human Genetics*, 64, 1293-1304.
- Almqvist, E. W., Brinkman, R. R., Wiggins, S., & Hayden, M.R. (2003). Canadian collaborative study of predictive testing. Psychological consequences and predictors of adverse events in the first 5 years after predictive testing for Huntington's disease. *Clinical Genetics*, 64, 300-309.
- Bettencourt, C., & Lima, M. (2011). Machado-Joseph disease: From first descriptions to new perspectives. Orphanet Journal of Rare Diseases, 2, 35-35.
- Bloch, M., Fahy, M., Fox, S., & Hayden, M. (1989). Presymptomatic testing for Huntington disease: II. Demographic characteristics, life-style patterns, attitudes, and psychosocial assessments of the first fifty-one test candidates. *American Journal of Medical Genetics*, 32, 217-224.
- Canavarro, C. (2007). Inventário de Sintomas Psicopatológicos (BSI). Uma revisão crítica dos estudos realizados em Portugal [Brief Symptom Inventory (BSI). A critical review of the studies carried out in Portugal]. In M. Simões, C. Machado, M. Gonçalves, L. Almeida (Eds), Avaliação Psicológica. Instrumentos validados para a população portuguesa [Psychological evaluation. Instruments validated for the Portuguese population] (vol. III, pp. 305-331). Portugal: Quarteto Editora.

- Codori, A. M., Hanson, R., & Brandt, J. (1994). Self-selection in predictive testing for Huntington's disease. *American Journal of Medical Genetics*, 54, 167-173.
- Codori, A., Slavney, P. R., & Brandt, J. (1997). Predictors of psychological adjustment to genetic testing of Huntington's Disease. *Health Psychology*, 16, 36-50.
- Coelho, T., Maia, L. F., Martins da Silva, A., Waddington, M., Planté-Bordeneuve, V., Lozeron, ..., Grogan, D. R. (2012). Tafamidis for transthyretin familial amyloid polyneuropathy: A randomized, controlled trial. *Neurology*, 79, 785-792.
- Decruyenaere, M., Evers-Kiebooms, G., & Van Den Berghe, H. (1997). Non-participation in predictive testing for Huntington's Disease: Individual decision-making, personality and avoidant behaviour in the family. *European Journal of Human Genetics*, 5, 351-363.
- Decruyenaere, M., Evers-Kiebooms, G., Cloostermans, T., Boogaerts, A., Demyttenaere, K., Dom, R., & Fryns, J. P. (2003). Psychological distress in the 5-year period after predictive testing for Huntington's disease. *European Journal of Human Genetics*, 11, 30-38.
- Decruyenaere, M., Evers-Kiebooms, G., Cloostermans, T., Boogaerts, A., Demyttenaere, K., Dom, R., & Fryns, J. P. (2004). Predictive testing for Huntington's disease: Relationship with partners after testing. *Clinical Genetics*, 65, 24-31.
- Derogatis, L. R. (1993). BSI: Brief Symptom Inventory. Minneapolis: Nacional Computers Systems.
- DudokdeWit, A. C., Tibben, A., Duivenvoorden, H. J., Niermeijer, M. F., & Passchier, J. (1998). Predicting adaptation to presymptomatic DNA testing for late onset disorders: who will experience distress? Rotterdam Leiden Genetics Workgroup. *Journal of Medical Genetics*, 35, 745-754.
- Gargiulo, M., Lejeune, S., Tanguy, M., Lahlou-Laforet, K., Faudet, A., Cohen, D., Feingold, J., & Durr, A. (2009). Long-term outcome of presymptomatic testing in Huntington disease. *European Journal of Human Genetics*, 17, 165-171.
- Gonzalez, C., Lima, M., Kay, T., Silva, C., Santos, C., & Santos, J. (2004). Short-term psychological impact of predictive testing for Machado-Joseph disease: Depression and anxiety levels in individuals at risk from the Azores (Portugal). *Community Genetics*, 7, 196-201.
- Gonzalez, C., Gomes, E., Kazachkova, N., Bettencourt, C., Raposo, M., Taylor, T., MacLeod, P., Vasconcelos, J., & Lima, M. (2012).
 Psychological well-being and family satisfaction levels five years after being confirmed as a carrier of the Machado-Joseph disease mutation. *Genetic Testing and Molecular Biomarkers*, 16, 1-6.
- Hawkins, A., Ho, A., & Hayden, M. (2011). Lessons from predictive testing for Huntington disease: 25 years on. *Journal of Medical Genetics*, 48, 649-650.
- International Huntington Association and World Federation of Neurology Research Group on Huntington's Disease (1994). Guidelines for the molecular genetics predictive test in Huntington's disease. *Journal of Medical Genetics*, 31, 555-559.
- Lêdo, S. (2002). O primeiro dia do resto de suas vidas. Alguns aspectos psicológicos da Paramiloidose [The first day of the rest of their lives. Some psychological aspects of Paramyloidosis]. MSc Dissertation, ISPA - Lisbon, Portugal.
- Lêdo, S., Leite, A., & Sequeiros, J. (2013a). Anxiety and pre-symptomatic testing for neurodegenerative disorders. *Open Journal of Genetics*, 3, 14-26.
- Lêdo, S., Leite, A., Souto, T., Dinis, M. A., & Sequeiros, J. (2016). Middleand long-term anxiety levels resulting from presymptomatic testing of HD, MJD and FAP neurodegenerative diseases. *Revista Brasileira de Psiquiatria*, 38, 113-120.
- Lêdo, S., Paneque, M., Rocha, J., Leite, A., & Sequeiros, J. (2013b). Predictive testing for two neurodegenerative disorders (FAP and HD): A psychological point of view. *Open Journal of Genetics*, 3, 270-279.
- Leite, A. (2006). Determinantes Psicossociais da Adesão ao Teste Pré-Sintomático em Doenças Neurológicas Hereditárias de Aparecimento

Tardio [Psychosocial Determinants of Adhesion to Pre-Symptomatic Testing in Late-Onset Hereditary Neurological Diseases]. PhD Dissertation, University of Porto - Porto, Portugal.

- Lerman, C. (1997). Psychological aspects of genetic testing: Introduction to the special issue. *Health Psychology*, *16*, 3-7.
- Licklederer, C., Wolff, G., & Barth, J. (2008). Mental health and quality of life after genetic testing for Huntington disease: A long-term effect study in Germany. *American Journal of Medical Genetics*, 146A, 2078-2085.
- Lopes, A., & Fleming, M. (1996). Doença somática e organização psíquica: reflexões a partir da Polineuropatia Amiloidótica Familiar [Somatic illness and psychic organization: reflections from Familial Amyloid Polyneuropathy]. *Revista Portuguesa de Psicanálise*, 15, 93-100.
- Lopes, A., & Fleming, M. (1998). Aspectos psicológicos da Polineuropatia Amiloidótica Familiar: a trama subterrânea intergeracional [Psychological aspects of Familial Amyloid Polyneuropathy: The intergenerational underground]. *Brotéria Genética*, XIX, 183-192.
- Paneque, H. M., Prieto, A. L., Reynaldo, R. R., Cruz, M. T., Santos, F. N., Almaguer, M. L., et al. (2007). Psychological aspects of presymptomatic diagnosis of spinocerebellar Ataxia type 2 in Cuba. *Community Genetics*, 10, 132-139.
- Paneque, M., Lemos, C., Sousa, A., Velázquez, P. L., Fleming, M., & Sequeiros, J. (2009). Role of the disease in the psychological impact of pre-symptomatic testing for SCA2 and FAP ATTRV30M: Experience with the disease, kinship and gender of the transmitting parent. *Journal* of Genetic Counseling, 18, 483-493.
- Rezende, T. J., D'Abreu, A., Guimarães, R. P., Lopes, T. M., Lopes-Cendes, I., Cendes, F., Castellano, G., França, & M. C. Jr. (2014). Cerebral cortex involvement in Machado-Joseph disease. *European Journal of Neurology*, 22, 277-283.
- Rolim, L., Leite, A., Lêdo, S., Paneque, M., Sequeiros, J., & Fleming, M. (2006). Psychological aspects of pre-symptomatic testing for Machado-Joseph disease and familial amyloid polyneuropathy type I. *Journal of Clinical Genetics*, 69, 297-305.
- Saraiva, M. J., & Costa, P. (1986). Familial Amyloidotic Polyneuropathy, Portuguese type: Phenotype and genotype. In M. L. Sales-Luís (Eds.), Symposium on Peripheral Neuropathies (pp. 207-212). Lisboa.
- Sequeiros, J. (1998). Prenatal diagnosis of late-onset diseases. Progresos en Diagnostico Prenatal, 10, 218-220.
- Sequeiros, J., Pinto-Basto, J., Rocha, J., Lêdo, S., Leite, A., Rolim, L., ..., Fleming, M. (2006). Ten years of a programme for presymptomatic testing (PST) and prenatal diagnosis (PND) in late-onset neurological diseases in Portugal: Machado-Joseph disease (MJD), Huntington disease (HD) and familial amyloid neuropathy type I - ATTRV30M (FAP-I). European journal of Human Genetics, 14, 1-1.
- Skirton, H., Goldsmith, L., Jackson, L., & Tibben, A. (2013). Quality in genetic counselling for presymptomatic testing - clinical guidelines for practice across the range of genetic conditions. *European Journal* of Human Genetics, 21, 256-260.
- Sousa, A. (2006). Epidemiologia Genética da Polineuropatia Amiloidótica Familiar [Genetic Epidemiology of Familial Amyloid Polyneuropathy]. *Sinapse*, 6, 74-79.
- Tibben, A., Timman, R., Bannink, E., & Duivenvoorden, H. (1997). Three years follow-up after presymptomatic testing for Huntington's Disease in tested individuals and partners. *Health Psychology*, *6*, 20-35.
- Tibben, A. (2007). Predictive testing for Huntington's disease. *Brain Research Bulletin*, 72, 165-171.
- Timman, R., Roos, R., Maat-Kievit, A., & Tibben, A. (2004). Adverse effects of predictive testing for Huntington Disease underestimated: Long-term effects 7-10 years after the test. *Health Psychology*, 23, 189-197.
- Weil, J. (2003). Psychosocial genetic counseling in the post-nondirective era: A point of view. Journal of Genetic Counseling, 12, 199-211.
- Zawacki, T. M., Grace, J., Friedman, J. H., & Sudarsky, L. (2002). Executive and emotional dysfunction in Machado-Joseph disease. *Journal of Movement Disorders*, 17, 1004-1010.