Refine penetrance estimates in the main pathogenic variants of transthyretin hereditary (familial) amyloid polyneuropathy (TTR-FAP) using a new non-parametric approach (NPSE)

Farida Gorram, Flora Alarcon, Hervé Perdry, Bérénice Hébrard, Thibaud Damy, Pascale Fanen, Benoît Funalot, Gregory Nuel, Violaine Planté-Bordeneuve

To cite this version:


HAL Id: hal-01539752
https://hal.archives-ouvertes.fr/hal-01539752
Submitted on 28 May 2018

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Refine penetrance estimates in the main pathogenic variants of Transthyretin Familial Amyloid Polyneuropathy (TTR-FAP) by use of a new non-parametric approach (NPSE)

Farida Gorram1,2,3, Flora Alarcon4, Hervé Perdry5, Bérénice Hébrard6, Thibaud Damy2,3,7, Pascale Fanen6, Benoît Funalot5,6,8, Gregory Nuel9 and Violaine Planté-Bordeneuve1,2,3

1 Department of Neurology, Henri Mondor University Hospital, AP-HP, Créteil, France. 2 Amyloid research Institute, Amyloid Network, Créteil, France. 3 Paris-Est Créteil University, Créteil, France. 4 Laboratory MAP5 UMR CNRS 8145 Paris Descartes University, France. 5 University Paris-Sud, UMR-S 669 - Inserm, U669, Villejuif, France. 6 Department of genetics, Henri Mondor University Hospital, AP-HP, Créteil, France. 7 Department of Cardiology, Henri Mondor University Hospital, AP-HP, Créteil, France. 8 Inserm U.955, Institut Mondor de Recherche Biomédicale, Créteil, France. 9 Institute of Mathematics, National Center for French Research, Laboratory of Probability, University Pierre et Marie Curie, Sorbonne University, France.

Address for correspondence: Violaine Planté-Bordeneuve of Department of Neurology, Henri Mondor University Hospital, AP-HP, Créteil, France. E-mail: Violaine.plante@aphp.fr

Background: Significant variability of phenotype and age of onset are well known in TTR-FAP associated to a wide spectrum of pathogenic TTR variants, among which Val30Met is the most frequent [1,2]. Recently, new therapeutic options became available in TTR-FAP that should be administered from the very onset of symptoms. In this context, the knowledge of the risk of being symptomatic for mutation carriers (penetrance) is essential to adjust the follow-up of carriers and for patient management [3,4]. This study aims to refine penetrance estimates in the main pathogenic TTR variants encountered in our TTR-FAP population using a newly developed non-parametric approach named NPSE for Non-Parametric Survival Estimate.

Materials and methods: A systematic genealogical enquiry was carried out in each family assessed in our center. Relevant data were collected with special attention to the genotype, phenotype, age at onset (AO) for affected individuals and age at last news for asymptomatic carriers and relatives. Portuguese families were not included in this study.

In previous works, we estimated penetrance as function of age, using a parametric survival method the PEL (Proband’s Exclusion Likelihood) [6] in which the AO is modeled by a Weibull distribution (WD). However, such method does not fit properly when penetrance function is far from WD. To avoid this bias, we have developed a non-parametric approach NPSE [5] based on a survival analysis (Kaplan-Meier estimator) that fits any penetrance shape. Also, NPSE can take into account covariates, here the TTR variant, through a Cox Model.

Results: We obtained genealogical data from 71 unrelated kindreds, including Val30Met (35 families, 90 patients), Ser77Tyr (15 families, 47 patients), Ile107Val (12 families, 21 patients), Ser77Phe (9 families, 30 patients) with information on 1654 subjects (188 affected /115 asymptomatic carriers).

Mean age at onset (± Standard Deviation ) was 54.6 years (± 14.9) for Val30Met, 55.6 years (± 9.1) for Ser77Tyr, 58.7 years (± 6.4) for Ser77Phe and 62.3 years (± 6.8) for Ile107Val.

Penetrance estimates were significantly different between the 4 TTR variants tested (p= 0.003) and remained incomplete in elderly. Hence, at 80 years of age, penetrance (95% Confidence Interval) varies from 46.9%, (16.6-66.2) in Ile107Val up to 71.3% (51.1-83.1) for Ser77Tyr kindred.

The disease risk is virtually null up to the age of 50 years for all variants except in the Val30Met group where it increases progressively from 3.1% (95% CI, 0.3-5.7) at 30 years to 16.5% (9.8-22.7) at 50 and 64.4% (49.4-74.9) at 80 years old. In contrast, in non Val30Met variants, the risk increased abruptly after 50 years old. In Ser77Phe and Ser77Tyr carriers, it raised from 7% (0.8-
15.8) and 12.8% (3.3-21.4) respectively up to 65.8% (37.1-81.4) and 71.3% (51.1-83.1). Risk estimate was found the lowest in Ile107Val families where it increases progressively from 50 years to 37.5% (14.2-54.4) at 70 y-o.

**Discussion and Conclusions:** This study showed significant differences of penetrance profiles in TTR-FAP with various pathogenic mutations of TTR. Our results should be replicated and implemented on larger samples of families to refine the management of carriers. NPSE method is more accurate and powerful than PEL previously used to unravel penetrance estimates. In future studies, it will allow testing the effect of other covariates, like gender, origin, and parent of origin on disease expression.

**Declaration of interest:** The authors declare no conflict of interest.