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


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BRIEF COMMUNICATION

Harmonizing results of ataxia rating scales: mFARS, SARA, and ICARS

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Abstract

The ever-increasing body of ataxia research provides opportunities for large-scale meta-analyses, systematic reviews, and data aggregation. Because multiple standardized scales are used to quantify ataxia severity, harmonization of these measures is necessary for quantitative data pooling. We applied the modified Friedreich Ataxia Rating Scale (mFARS), the Scale for the Assessment and Rating of Ataxia (SARA), and the International Cooperative Ataxia Rating Scale (ICARS) to a large cohort of people with Friedreich's ataxia. We provide regression coefficients for scale interconversion and discuss the reliability of this approach, together with insights into the differential sensitivities of mFARS and SARA to disease progression.

Introduction

Friedreich's ataxia (FRDA) is a progressive neurological disease characterized by gait and limb ataxia, dysarthria, loss of reflexes, proprioceptive dysfunction, and muscle weakness, as well as non-neurological features including cardiomyopathy, diabetes, and scoliosis.¹ Quantifying severity and tracking changes in the clinical symptom profile is challenging, and usually focuses on the predominant movement features of FRDA, which largely reflect underlying cerebellar pathology.

Clinical rating scales have been developed to quantify disease severity based on a codified neurological exam. Two such scales have evolved as the cornerstones for assessing disease progression in FRDA: the modified Friedreich Ataxia Rating Scale (mFARS)² and the Scale for the Assessment and Rating of Ataxia (SARA).³ Besides common and unavoidable issues of such instruments (e.g., ceiling effects at longer disease duration), the earlier developed ICARS score⁴ has shown additional problems (e.g., item redundancies and a questionable subscale grouping)^{5–7} and is not used anymore in large studies. While SARA (8 items and 4

bilateral) emphasizes fast assessment, mFARS is more granular in scale with 18 items (7 bilateral) grouped into 4 sub-components. Regardless, SARA and mFARS have shown similar psychometric properties.^{6,8–11}

A growing amount of data is available from research studies in FRDA and other ataxias that utilize these scales (e.g., spinocerebellar ataxias, SCAs). This has seeded the establishment of worldwide consortia that retrospectively combine datasets to allow for greater statistical sensitivity, reliability, and generalizability of outcomes (e.g., the ENIGMA-Ataxia consortium for neuroimaging).¹² However, a major barrier to quantitatively compare existing studies in meta-analyses and systematic reviews is the use of different clinical rating scales.

While the overall approaches to assess clinical features inherent to cerebellar ataxia are similar between the scales, the relative weights for axial (balance, gait), appendicular (upper and lower limb function) and speech differ. Specifically, these components comprise 39%/56%/5% of the mFARS, 45%/40%/15% of the SARA, and 34%/52%/8% of the ICARS, respectively, and only ICARS includes oculomotor disorders (6%; see Table S1).

This has implications for the comparability of the results and for the sensitivity to symptom progression at different disease stages. For instance, it has been shown in a large natural history study that ambulant individuals progress mostly in axial items,¹¹ whereas appendicular function declines largely after people transition into a wheelchair. Bulbar and speech as measured by these clinical scales, play a minor role, and progress largely independent of the disease phase. Scale behavior and comparability may therefore be dependent on the disease stage.

Here, we examine the interchangeability of these scales, and provide empirically driven guidance regarding the homogenization and conversion of scores for use in retrospective and meta-analytic research, building on previous correlation studies between the FARS, SARA, and ICARS.¹³

Material and Methods

Data collection

The dataset was collected at a single center (Murdoch Children's Research Institute, Melbourne) during yearly study visits of the Friedreich Ataxia Clinical Outcome Measures study (FACOMS, NCT03090789) between January 2011 and May 2021. The subset included here encompasses all individuals participating at this center with available scores on all three scales. This study had approval from the Monash Health Human Research Ethics Committee (Project Number 02114A). All participants gave their informed, written consent in accordance with the 1964 Declaration of Helsinki.

mFARS scores were collected according to the FACOMS case report form¹⁴; afterward common items in the SARA/ICARS were rescored, followed by administering outstanding items from SARA and ICARS with the goal to minimize assessment burden and to ensure all items are assessed at approximately the same time of day. On average administering, all scales concurrently increased the overall assessment time by ~10 min. In addition, Functional Disease Staging (FDS)¹⁵ was recorded. Ratings correspond to no (1), minimal (2, symptoms present), mild (3, symptoms overt and significant), and moderate (4, requires a walker or other aids) disability. Reaching FDS 5 defines loss of ambulation (LoA).¹⁶

Statistical analysis

All results were divided by their maximum possible score and multiplied by 100, that is, the resulting %-scales range from 0 (no disability) to 100 (maximum disability). Percentiles were chosen over other normalization metrics (such as z-scores) due to better external comparability

and descriptiveness. The %-scores were then compared at each FDS stage using boxplots, followed by correlation analyses using linear regression functions and analyzed by Pearson's R^2 statistics. Data from all available visits were used; regression coefficients derived from cross-sectional data (first visit only, last visit only) were only minimally different (generally <1%, data not shown). Distributions were compared using density plots.¹⁷ Such plots are not affected by the number of bins used in histograms, facilitating the comparison of distribution shapes.¹⁷

Results

Data comprised 605 visits of 166 individuals, whose demographics are representative of the complete FACOMS cohort¹¹ (Table S2); importantly, nearly the full range of clinical and genetic severity is represented in our sample.

All scales and components showed significant overlap across FDS stages, with appendicular function progressing most similarly between all scales (Fig. 1A). Conversely, axial %-scores differed between mFARS and SARA/ICARS: At FDS 1 (least severe), median mFARS-AX was almost 50%, but median SARA-AX was below 25%, with similar results through to FDS 4 (Fig. 1B). Total score results matched the axial pattern across FDS 1 to 4, with mFARS scores roughly similar to SARA and ICARS at one later (more severe) disease stage (Fig. 1C). ICARS total scores were close to SARA results.

Correlation analyses (Fig. 2) showed high Pearson's R^2 values: >0.93 for total scores, >0.86 for axial scores, and >0.85 for the appendicular components. Appendicular functions were highly similar across scales (Fig. 2A–C). For axial components, linear correlation coefficients showed large intercepts vs. FARS-AX (39% and 29% from SARA-AX and ICARS-AX, respectively, translating to 14 and 10 absolute points) and low slopes (0.61 and 0.70, respectively). Second-order polynomials provided improved r^2 values and model fits (Fig. 2D–F; Table S3 for model statistics). Again, total scores followed the axial component pattern, with smaller distinctions between the linear and polynomial models (Fig. 2G–I. For a full list of coefficients, see Table 1).

To evaluate the extent of information retained during conversion with different polynomial functions, density plots of axial components of mFARS-AX and SARA-AX were created (Fig. 3). The use of second-order polynomial models lead to an improved replication of the measured results.

Discussion

In an attempt to harmonize ataxia rating scales, we provide inter-conversion procedures between the mFARS,

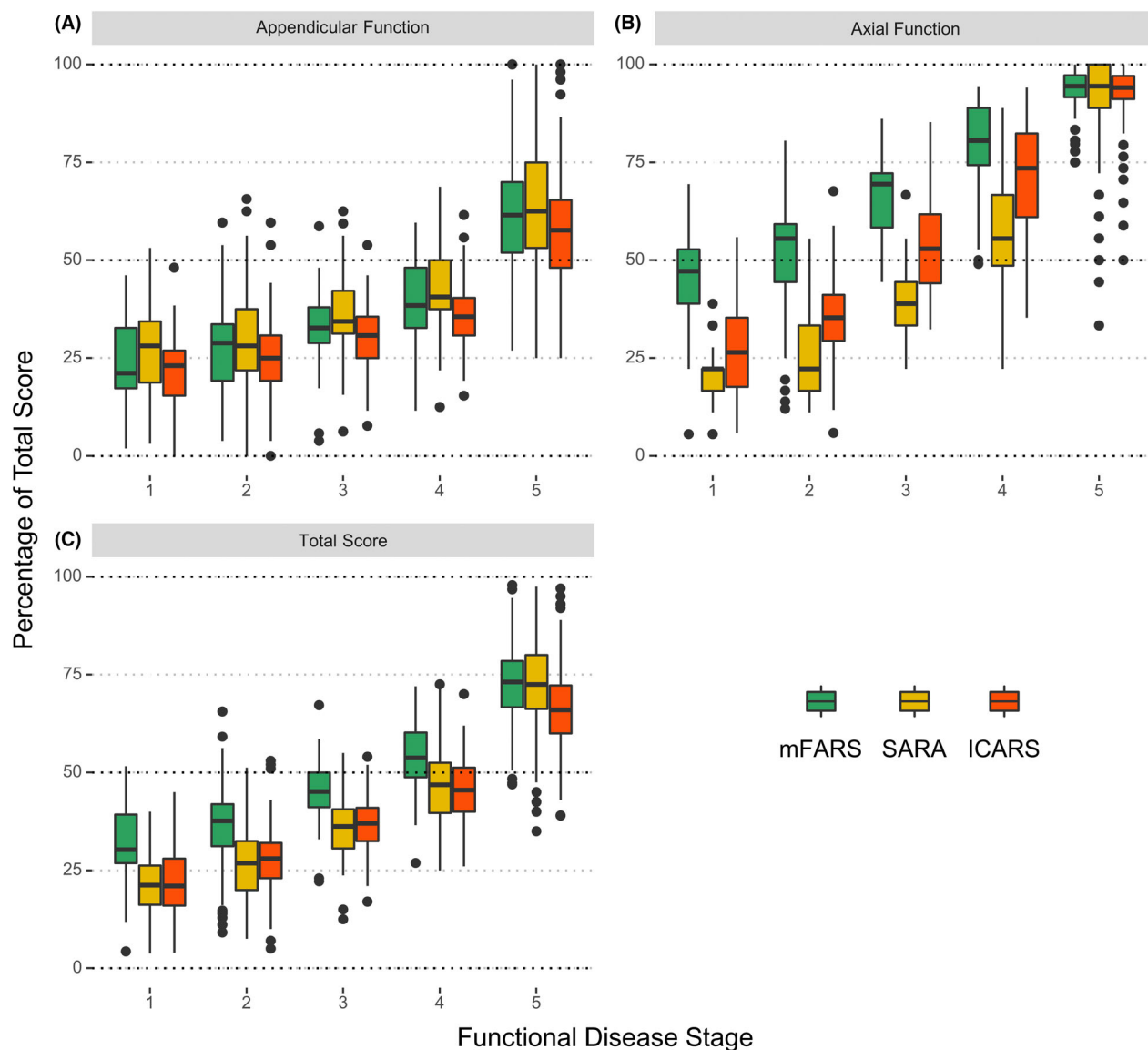


Figure 1. Appendicular (A), axial (B), and total scores (C), percentage of total scores by functional disease stage.

SARA, and ICARS. This facilitates the comparison of scores at different disease stages in FRDA cohorts scored with differential rating scales. We show that linear conversion functions are mostly sufficient, although in ambulatory cohorts, the use of second-order polynomials and/or focusing on axial function alone might offer advantages (if sub-scores are available). Appendicular functions do not require conversion factors.

A comparison of results by FDS revealed that a percentage-scale is insufficient for harmonization, due to differential progression characteristics of each scale. Scale scores also provide a poor proxy for identifying specific disease milestones that are captured more directly and with greater face validity by the FDS itself.

Interestingly, the systematic offsets we observed between SARA/ICARS and mFARS scores were not detected in earlier analyses¹³ which used the full FARS-neuro score (FARS_n, maximum of 125 points). However, the FARS_n behaved very similar to the mFARS in our results (Table 1), and the reason for the difference may be due to the larger sample and the broader scale coverage in our study.

Of particular relevance are differences in stance function testing. The mFARS assesses stance using 1 min test (in triplicate, if necessary) and the relative absence of intermittent scores in these items¹⁶ indicates that once a stance trial is missed, the incentive and motivation to retry might be low. Also, as shown previously,¹⁶ in most patients three out of six stance functions in the mFARS are already lost at the time of

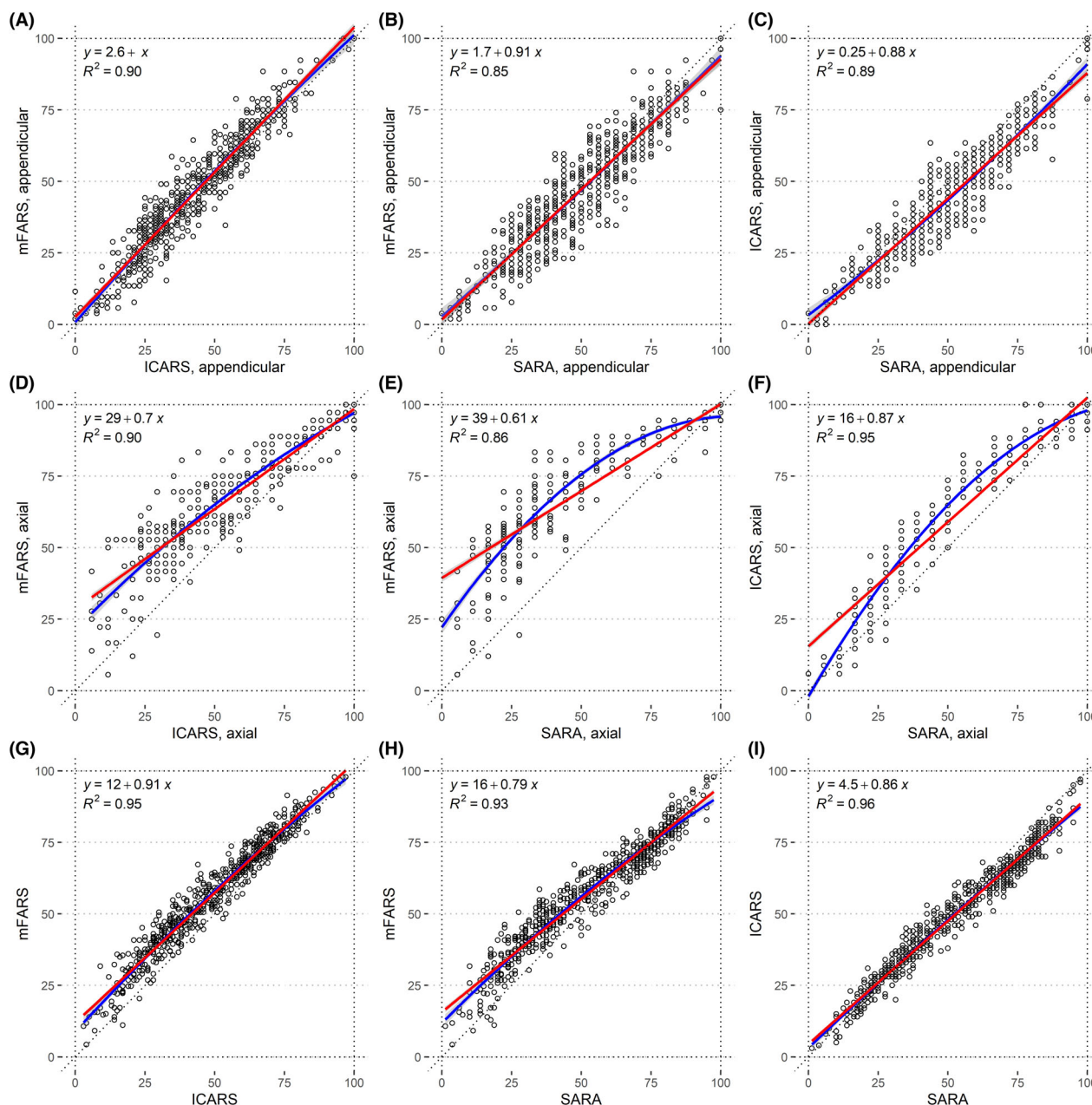


Figure 2. Correlation plots of percentages of maximum score for the appendicular components of mFARS, SARA, and ICARS (A–C), their axial components (D–F), and the full scores (G–I). Linear correlation is depicted in red, and second-order polynomial regression is in blue.

FRDA diagnosis. This early loss is in part responsible for the large intercepts when converting between the SARA/ICARS and the mFARS. More importantly, however, these items should offer high sensitivity in early or prodromal stages of the disease, making the instrument a promising candidate for other, more slowly progressive ataxias where axial function loss is potentially less aggressive and diagnosis occurs at lower disability or before onset.¹⁸ SARA and ICARS on the other hand use shorter stance test intervals (10 sec) and show a later ceiling. SARA and ICARS may therefore better

distinguish disease burden in patients who are approaching loss of ambulation. Of note, it remains unclear if this difference translates into meaningful improvements in the measures, that is, a better signal/noise ratio.

Taken together, there are differences in the capacity of mFARS, ICARS, and SARA to capture cerebellar impairment at different disease stages. Since both have comparable psychometric quality, the choice between the two should depend on the population studied, rather than the region and history of the center.

Table 1. Conversion coefficients for total scores and axial components (the use of second-order polynomial equations is only recommended for ambulatory populations). For example, $mFARS = 14.59 + (SARA \times 1.84)$, or $mFARS(axial) = 22.20 + (SARA \times 1.40) + (SARA^2 \times -0.0066)$. The association between mFARS and FARSn is included for legacy comparison (see "Discussion" section).

	Score	Model Units Predicted from	Linear model				Second-order polynomial model			
			Points		Percentages		Percentages			
			Inter.	Slope	Int.	Slope	Inter.	x	x ²	
Total scores	mFARS	FARSn	-0.10	0.80	-0.10	1.07				
	mFARS	SARA	14.57	1.84	15.67	0.79				
	mFARS	ICARS	11.06	0.85	11.89	0.91				
	SARA	mFARS	-5.84	0.51	-14.61	1.17				
	SARA	ICARS	-1.27	0.45	-3.18	1.12				
	ICARS	mFARS	-9.67	1.12	-9.67	1.04				
Axial component	ICARS	SARA	4.51	2.15	4.51	0.86				
	mFARS	SARA	14.19	1.22	39.41	0.61	22.20	1.40	-0.0066	
	mFARS	ICARS	10.27	0.74	28.53	0.70	21.39	0.99	-0.0023	
	SARA	mFARS	-8.49	0.71	-47.15	1.42	34.93	-1.28	0.0197	
	SARA	ICARS	-2.48	0.58	-13.79	1.09	12.87	0.00	0.0087	
	ICARS	mFARS	-10.27	1.22	-30.19	1.29	11.07	-0.06	0.0099	
		ICARS	SARA	5.29	1.64	15.56	0.87	-1.97	1.68	-0.0068

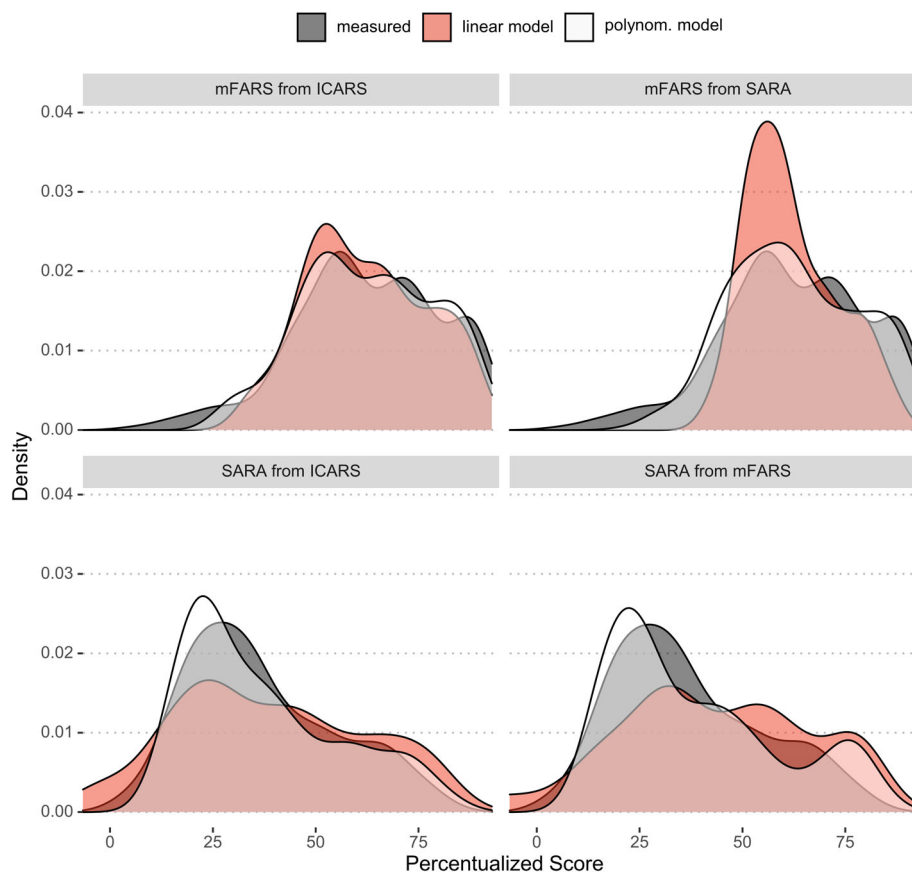


Figure 3. Density plots of axial component scores. Measured results (dark gray), compared to predictions derived from second-order polynomial conversion (white) versus from simpler linear regression models (red).

In conclusion, we provide conversion coefficients for mFARS, SARA, and ICARS, as well as their axial and appendicular components, useful for the harmonization and comparison of disease stages within cohorts from diverse origins. Comparing results using percentages is not recommended. Caution is advised when generalizing to other diseases (e.g., SCAs), where progression profiles between, for example, axial and appendicular function might differ from FRDA.

Conflict of Interest

The authors report no relevant disclosures.

Author Contributions

Christian Rummey: Conception or design of the work. Data analysis and interpretation. Drafting the article. Ian H. Harding: Conception or design of the work. Critical revision of the article. Martin B. Delatycki: Critical revision of the article. Data Collection. Genevieve Tai: Critical revision of the article. Data Collection. Thiago Rezende: Critical revision of the article. Louise A. Corben: Conception or design of the work. Critical revision of the article. Data Collection.

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Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Table S1. Items and functionality (abbreviated) across mFARS, SARA, and ICARS, including the sequence of administration. Numbers in brackets show the maximum possible scores (highest disability). For full details, please refer to the primary references.

Table S2. Demographic characteristics and amount of follow-up, grouped by age of onset*. Data are percentage or median [IQR].

Table S3. Summary statistics for simple linear and polynomial models.