Exonic Deletions of FXN and Early-Onset Friedreich Ataxia

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Background: Friedreich ataxia (FA) is the most frequent type of autosomal recessive cerebellar ataxia, occurring at a mean age of 16 years. Nearly 98% of patients with FA present with homozygous GAA expansions in the FXN gene. The remaining patients are compound heterozygous for an exonic expansion and a point mutation. Patients who are compound heterozygous for an exonic deletion and an expansion are exquisitely rare.

Objectives: To describe 6 patients affected with FA due to an exonic deletion mutation (FAexdel) and to compare these 6 patients with FAexdel with 46 patients consecutively diagnosed with typical FA due to homozygous GAA expansion and whose small expansions were within the same range as that of the expansions of the patients with FAexdel.

Design: Description of a series.

Setting: Academic research.

Patients: Six patients with FAexdel and 46 patients with typical FA.

Intervention: FXN gene analysis, including assessment of GAA expansion and exon sequencing and determination of exonic copy numbers using multiplex ligation-dependent probe amplification.

Results: We identified 6 patients with FA who presented with the combination of 1 GAA expansion and 1 FXN exonic deletion. The mean (SD) age at onset of the disease was earlier for patients with FAexdel (7 [4] years [range, 3-12 years]) than for patients with typical FA (15 [5] years [range, 6-30 years]) (P = .001), and the median time to confinement to wheelchair was shorter for patients with FAexdel (20 years) than for patients with typical FA (28 years) (P = .002). There was no difference between the mean (SD) size of the expansion for the patients with FAexdel (780 [256] GAA triplet repeat sequences [range, 340-1070 GAA triplet repeat sequences]) and the mean (SD) size of the short expansion for the patients with typical FA (634 [163] GAA triplet repeat sequences [range, 367-1000 GAA triplet repeat sequences]) (P = .10). The mean disease duration before becoming wheelchair bound was shorter for patients with FAexdel (9 years) than for patients with typical FA (13 years), and the incidence of cardiomyopathy was higher for patients with FAexdel (84%) than for patients with typical FA (68%). However, these differences were not significant, probably owing to the small size of the FAexdel group. The other extraneurological signs, such as scoliosis or diabetes mellitus, were particularly frequently observed in the FAexdel group. One patient presented at 9 years of age with severe angina and marked cardiomyopathy that confined her to a wheelchair. Three patients had disabling autonomic disturbances. It appears that exonic deletion significantly contributes to the clinical picture of patients with FA.

Conclusions: Friedreich ataxia due to an exonic deletion mutation corresponds to an early onset and severe variant of FA. FXN should be investigated for exonic deletion in patients with early-onset FA in which only 1 GAA expansion without a point mutation is found. Patients with FAexdel have to be carefully observed using cardiological, orthopaedic, endocrinological, gastroenterological, and ophthalmological data. Friedreich ataxia due to an exonic deletion mutation should be suspected in young patients presenting with severe scoliosis.

mozygous GAA expansions in the FXN gene encoding the frataxin protein. About 2% of these patients have both 1 GAA expansion and 1 point mutation in FXN. Patients compound heterozygous for an exonic deletion and an expansion are exquisitely rare. Although the clinical phenotype of FA due to GAA expansion or point mutations has been well described, the clinical spectrum of patients with FA due to an exonic deletion mutation (FAexdel) remains unknown. We report 6 patients with FAexdel who presented with the combination of 1 GAA expansion and 1 FXN exonic deletion responsible for early-onset FA with precocious neurological disability and cardiomyopathy.

**METHODS**

Approximately 600 patients, recruited from French centers during the period from 1988 to 2010, received a diagnosis of FA, which was determined by use of a genetic test. The DNA samples of 13 ataxic patients whose clinical phenotypes were compatible with FA but who were heterozygous for a GAA expansion and had no point mutation were assessed for large FXN molecular analysis.
deletion or duplication using a multiplex ligation-dependent probe amplification kit (SALSA MLPA kit P316-A1 Recessive Ataxias; MRC-Holland) according to the manufacturer’s recommendations. We retrospectively reviewed the clinical, molecular, electrophysiological, and morphological data of the patients with FAexdel and of 46 patients consecutively diagnosed with typical FA due to homozygous GAA expansion and whose small expansions were within the same range as that of the expansions of the patients with FAexdel.

Data, reported as mean (SD) values along with minimum and maximum values (ie, range), were analyzed by use of the Mann-Whitney test. When percentages were compared, the Fisher exact test was used. For the median time to confinement to a wheelchair and for the median disease duration before confinement to a wheelchair, the Kaplan-Meier method was performed.

RESULTS

Seven patients had a normal FXN exonic copy number dosage, excluding the diagnosis of FA. These patients, whose molecular diagnosis remains unknown, presented with late-onset ataxia (ranging from 30 to 58 years of age) and had large expansions (from 700 to 1100 GAA triplet repeat sequences). Some had atypical presentation because they had cerebellar atrophy and/or a lack of sensory neuropathy. They were assumed to be heterozygous carriers of the expansion by chance, given the high frequency of the GAA expansion in the European population (1 in 90 individuals). The remaining 6 patients were found to be affected with FA due to heterozygous FXN exonic deletion. Three of them (patients 1, 2, and 3) were previously reported in a series of patients with an FA-like disease who presented with only 1 GAA repeat expansion and no point mutation. The main findings are summarized in Table 1 and Figure 1, and the comparison between patients with FAexdel and patients with typical FA are presented in Table 2 and Figure 2. Cardiomyopathy was diagnosed at 4 years of age in patient 1 but was particularly stable until the patients turned 24 years of age. Patient 2 presented with severe angina and cardiomyopathy with a left ventricular ejection fraction of 52% at 9 years of age that confined her to a wheelchair. Interestingly, patient 3 was the grandaunt of patient 2 (Figure 1); therefore, autosomal dominant cerebellar ataxia was first suspected in this family. The high frequency of GAA expansion in heterozygous carriers led to a pseudodominant inheritance of FA in this family because the expansion of the 2 patients was transmitted by 2 unrelated individuals (Figure 1). The 6

![Figure 1. Pedigree showing 4 generations of a family that included patient 2 (IV1) and patient 3 (II3) with pseudodominant inheritance of Friedreich ataxia. The squares represent male individuals; circles, female individuals; filled circles, affected female individuals; diagonal line, deceased individual; arrow, proband. Genotypes are indicated as follows: E, GAA expansion; exdel, exonic deletion; WT, wild type. ND indicates analysis was not done.](https://jamanetwork.com/)

![Figure 2. Data on patients with Friedreich ataxia due to an exonic deletion mutation (FAexdel) and patients with typical FA who were not wheelchair bound (determined by use of the Kaplan-Meier method).](https://jamanetwork.com/)

| Table 2. Comparison Between Patients With FAexdel and Patients With Typical FA |
|-------------------------------------------------|-----------------|-----------------|--------|
| Parameter                                      | Typical FA      | FA exdel        | P Value |
| Age at onset, mean (SD) [range], y             | 15.4 (5) [6-30] | 7 (4) [3-12]    | .001   |
| Time to confinement to wheelchair, median, y   | 28              | 20              | .02    |
| DD before confinement to wheelchair, median, y | 13              | 9               | .42    |
| Age at last follow-up, mean (SD) [range], y    | 31 (10) [14-57] | 33 (22) [9-69]  | .92    |
| SARA score–to–DD ratio at last follow-up, a mean (SD) [range] | 1.5 (0.6) [0.5-3] | 1.9 (0.8) [1-3] | .33    |
| Cardiomyopathy, % of patients                  | 68              | 64              | .66    |
| GAA triplet repeat sequences, mean (SD) [range], No. | 634 (163) [367-1000] | 780 (256) [340-1070] | .10, b |
|                                              | 878 (220) [367-1667] c  |                  | .34 c |

Abbreviations: DD, disease duration; FA, Friedreich ataxia; FAexdel, FA due to an exonic deletion mutation; SARA, Scale for the Assessment and Rating of Ataxia.

This ratio calculation gives a good indication of the severity of the disease progression.

Short expansion vs 780 (256) [340-1070].

Large expansion vs 780 (256) [340-1070].

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patients with FAexdel in our series are all of French ancestry: 2 patients are from southern France, and the other 4 patients, with a deletion of exons 4 and 5, were originally from Brittany, which suggests that the multiple occurrence of this deletion is the consequence of a founder effect.

COMMENT

To our knowledge, we report the first series of 6 patients affected with FAexdel, which is characterized by a particularly young age at onset in comparison with the age at onset of typical FA. This comparison holds true because there is no difference between the 2 groups regarding the size of GAA expansion (Table 2). The disease is more severe in patients with FAexdel, mainly because of the early onset of the disease and the early age at which the patients become confined to a wheelchair but also because there appears to be a trend toward a faster progression of disability and a higher incidence of cardiomyopathy in FAexdel. For so far, only 1 patient with FAexdel (with a deletion surrounding exon 5a) had been reported with the disease at 9 years, sensory axonal neuropathy, foot deformity, loss of independent walking at 15 years, and cardiomyopathy at 21 years. Early-onset FA is a rare variant of FA, and patients whose age at onset is younger than 7 years represent approximately 12% of all patients. We found that FAexdel is exquisitely rare and accounts for approximately 1% of cases of FA diagnosed in France.

It appears that extensor plantar reflexes are an optional finding in FAexdel. Moreover, the incidence of extraneurological signs is particularly high in such patients. All but 1 of our 6 patients were affected with hypertrophic cardiomyopathy, albeit this is not significantly different from the 60% of patients with typical FA who have cardiomyopathy. However, the cardiomyopathy may be particularly disabling in FAexdel, as it lead to severe angina and early loss of ambulation in patient 2. The incidence of diabetes mellitus was higher in our patients with FAexdel (half of our patients were affected) than in previously reported series of patients with FA (about 30%). The incidence of scoliosis in our patients with FAexdel seems to be the same as in patients with typical FA, but the severity of scoliosis could be higher in FAexdel because 2 patients needed spine surgery only a few years after onset of the disease. Moreover, scoliosis may reveal FAexdel (patient 2). Interestingly, 3 patients had autonomic disturbances, especially repeated ileus, which is an unusual finding in FA. The severe phenotype associated with exonic deletion mutation is in agreement with the predicted absence of functional frataxin made from the deleted allele.

It appears that exonic deletion significantly contributes to the clinical picture of patients with FA, especially to the young age at onset of the disease, whereas the FA phenotype and the age at onset are usually conditioned by the size of the smaller GAA expansion (in case of homozygous expansions). For instance, patients with a short size of the smaller expansion may present with an FA variant such as late-onset FA defined by an age at onset of the disease of older than 25 and 40 years, respectively. Within our series, it seems that the size of the GAA expansion also contributes to the phenotype. Indeed, the 3 patients with the larger expansions (patients 1, 3, and 6, presenting with 1070, 970, and 830 GAA triplet repeat sequences, respectively) are those affected with both diabetes mellitus and autonomic disturbances, and patient 2 (800 GAA triplet repeat sequences) had had the first signs at 2 years of age, a severe cardiomyopathy at 4 years with angina at 9 years, and the shorter disease duration before becoming wheelchair bound. Moreover, patient 5, with the shortest expansion (340 GAA triplet repeat sequences) and deletion of exons 4 and 5, does not show symptoms of an extensor planter response, cardiomyopathy, or diabetes mellitus.

FXN should be investigated for exonic deletion in patients with early-onset FA in which only 1 GAA expansion without a point mutation is found. In clinical practice, patients with FAexdel have to be carefully observed using cardiological, orthopaedic, endocrinological, gastroenterological, and ophthalmological data. Moreover, FAexdel should be suspected in young patients presenting with severe scoliosis.

Accepted for Publication: August 10, 2011.
Published Online: March 12, 2012. doi:10.1001/archneurol.2011.834

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Author Contributions: Dr Anheim had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. Study concept and design: Anheim, Mariani, and Koenig. Acquisition of data: Anheim, Mariani, Calvas, Cheuret, Zagnoli, Odent, Seguela, Marelli, Fritsch, Delaunoy, Brice,
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**Financial Disclosure:** None reported.

**Previous Presentation:** Presented in part at the Journées de Neurologie de Langue Française; April 28, 2011; Paris, France (abstract and poster).

### REFERENCES