Real time PCR quantification of frataxin mRNA in the peripheral blood leucocytes of Friedreich ataxia patients and carriers

L Pianese, M Turano, M S Lo Casale, I De Biase, M Giacchetti, A Monticelli, C Criscuolo, A Filla, S Cocozza

The most common causative mutation of Friedreich ataxia (FRDA) is the unstable hyperexpansion of an intronic GAA triplet repeat that impairs frataxin transcription. Using real time quantitative PCR, we showed that FRDA patients had residual levels of frataxin mRNA ranging between 13% and 30% and that FRDA carriers had about 40% of that of controls. Asymptomatic carriers also showed reduced frataxin mRNA levels. We found an inverse correlation between the number of GAA repeats and frataxin mRNA levels. Real-time quantitative PCR may represent an alternative assay for FRDA molecular diagnosis.

Friedreich ataxia (FRDA) is an autosomal recessive progressive neurodegenerative disorder. The disease presents with gait and limb ataxia with onset usually in the first two decades of life, but later forms have also been described. Extra neurological features include skeletal deformity, left ventricular hypertrophy, and diabetes. GAA repeat expansion within the first intron of the X25 gene is the most common mutation in FRDA and it is present in 98% of the disease alleles. Few patients are compound heterozygotes for a GAA expansion and a point mutation and no patient homozygous for point mutation has been described. A correlation was shown between GAA expansion size and the severity of disease. The gene X25 encodes a mitochondrial protein named frataxin, which seems to be involved in mitochondrial iron homeostasis. Expanded GAA tracts lead, at the transcriptional level, to a reduction of X25 mRNA that is related to the size of the expansion.

Up to now, traditional assays such as semi-quantitative PCR, RNase protection, and Western blot were used to investigate X25 mRNA and frataxin levels in FRDA patients and FRDA carriers. In this study, real time quantitative polymerase chain reaction (PCR), a highly sensitive and specific method, was used for the first time to evaluate frataxin mRNA expression in peripheral blood leucocytes from FRDA patients and FRDA carriers.

**METHODS**

Peripheral blood samples were collected, following written informed consent, from 10 FRDA patients (two male and eight female), homozygous for GAA repeat expansion and three FRDA carriers heterozygous for GAA expansion.

GAA molecular analysis was performed as previously described. Total RNA was extracted from 2.5 ml peripheral blood leucocytes using the PAXgene Blood RNA Kit (Qiagen, Valencia, CA, USA).

Total RNA (1 μg) was reverse transcribed with 100 U of Superscript II RNase H- Reverse Transcriptase (Invitrogen, Gaithersburg, MD, USA) according to the manufacturer’s instructions. A 1 μl sample of cDNA was amplified by real time PCR using PCR primers and TaqMan MGB probes for Friedreich ataxia (FRDA) and hypoxanthine phosphoribosyltransferase 1 (HPRT1) genes as reference gene (Perkin-Elmer Applied Biosystems (PE-ABI), Foster City, CA, USA). Each sample was run in triplicate for both FRDA and HPRT1 in 20 μl reaction using TaqMan Universal PCR Master Mix according to the manufacturer’s instructions (PE-ABI). Reactions were performed in an ABIPrism 7000 sequence detector system (PE-ABI).

Quantitative real time PCR analysis was carried out using the 2(−Delta Delta C(T)) method. The new relative expression software tool (REST) was used to calculate the relative expression ratios on the basis of group means for target frataxin transcript versus reference HPRT1 transcript. REST also tests the group ratio results for significance by a randomisation test.

Correlations analyses were performed by using the SPSS statistical software package.

**RESULTS**

To investigate frataxin mRNA expression, 10 FRDA patients, three FRDA carriers, and three controls were analysed. Five patients had typical FRDA with onset within the first 20 years of life. FRDA carriers had onset at 20 or more years of age. The most common causative mutation of Friedreich ataxia (FRDA) is the unstable hyperexpansion of an intronic GAA triplet repeat that impairs frataxin transcription. Using real time quantitative PCR, we showed that FRDA patients had residual levels of frataxin mRNA ranging between 13% and 30% and that FRDA carriers had about 40% of that of controls. Asymptomatic carriers also showed reduced frataxin mRNA levels. We found an inverse correlation between the number of GAA repeats and frataxin mRNA levels. Real-time quantitative PCR may represent an alternative assay for FRDA molecular diagnosis.

**Abbreviations:** FRDA, Friedreich ataxia; HPRT1, hypoxanthine phosphoribosyltransferase 1; PCR, polymerase chain reaction
of life and five had onset after 20 years of age (late-onset FRDA, LOFA).\textsuperscript{2} The mean (SD) length of the expanded alleles was 741 (229) repeats with range of 348–1057. Mean (SD) age at onset was 12.6 (5.4) years in typical patients and 21.0 (0.7) years in LOFA patients.

Using quantitative real time PCR we found that frataxin mRNA was down-regulated in typical FRDA patients in comparison to the control group by a factor of 6.04 (p = 0.001). We also found that frataxin was significantly down-regulated in LOFA patients by a factor of 4.65 (p = 0.022) and in FRDA carriers by a factor of 2.84 (p = 0.016) in comparison to the control group (fig 1).

**DISCUSSION**

The present study is the first to use real time PCR to evaluate frataxin mRNA expression in FRDA patients and carriers. Real-time PCR is the most sensitive method for the detection and quantification of gene expression levels. Using real time PCR we found that all FRDA patients (typical and LOFA) showed frataxin mRNA levels ranging from 13% to 30% of that of healthy individuals. Furthermore, we were able to detect differences in frataxin mRNA levels between LOFA and typical patients. Previous studies used less sensitive methods such as semi-quantitative PCR, RNase protection, and Western blot to identify reductions in the amount of X25 mRNA and frataxin protein in FRDA patients.\textsuperscript{3 8 10 11} In these studies the residual amount of frataxin varied between 4% and 29% of the level in normal controls.\textsuperscript{11} Our data are in accordance with these previously reported data.

Here we report, for the first time, accurate quantification of frataxin mRNA in healthy FRDA carriers. These individuals showed about 40% of the normal frataxin mRNA levels. It is intriguing that these levels are not so different from that found in LOFA patients. It should be emphasised that, however, no overlap between the two groups was observed.

In addition, we found a significant inverse correlation between frataxin mRNA levels and GAA repeat number. The best correlation was found for GAA mean (r\textsuperscript{2} = 0.80) and for GAA1 (r\textsuperscript{2} = −0.74). A correlation was previously reported between the residual frataxin protein level and the size of GAA1.\textsuperscript{11} Considering the technical difficulties in Friedreich ataxia molecular diagnosis by PCR on genomic DNA due to the long GAA repeat, our data suggest that real time quantitative PCR could represent an alternative assay for Friedreich ataxia molecular diagnosis.

**REFERENCES**

Real time PCR quantification of frataxin mRNA


References

Historical note

Lewis Carroll’s Humpty Dumpty: an early report of prosopagnosia?

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Prosopagnosia is a rare form of visual agnosia characterised by impaired recognition of familiar faces (or equivalent stimuli). The term was coined by Bodamer in 1847, although the phenomenon had been described towards the end of the 19th century by Quaglini (1867), Hughlings Jackson (1872, 1876), and Charcot (1883). Brief accounts thought to be suggestive of prosopagnosia have been identified in writings from classical antiquity by Thucydides and Seneca. While there are dangers in this type of retrospective case identification, nonetheless I venture to suggest another early description of prosopagnosia.

The account is taken from Through the looking-glass and what Alice found there (1872) by Lewis Carroll (pseudonym of the Reverend Charles Lutwidge Dodgson). In chapter 6, Alice notices that the egg that she has just purchased had eyes and a nose and mouth; and when she had come close to it, she saw clearly that it was HUMPTY DUMPTY himself. “It can’t be anybody else!” she said to herself. “I’m as certain of it, as if his name were written all over his face.”

Discussion follows, in which Humpty Dumpty, sitting precariously balanced upon a wall, gives his famous definition of the meaning of a word (“just what I choose it to mean”) and coins the term “portmanteau word”. As Alice takes her leave of Humpty Dumpty, the subject of facial recognition recurs, in the following exchange:

“Good-bye, till we meet again!” she said as cheerfully as she could.
“I shouldn’t know you again if we did meet,” Humpty Dumpty replied in a discontented tone, giving her one of his fingers to shake: “you’re so exactly like other people.” “The face is what one goes by, generally,” Alice remarked in a thoughtful tone.
“That’s just what I complain of,” said Humpty Dumpty. “Your face is the same as everybody else has—the two eyes, so to speak (marking their places in the air with his thumb) ‘nose in the middle, mouth under. It’s always the same. Now if you had the two eyes on the same side of the nose, for instance—or the mouth at the top—that would be some help.’ “It wouldn’t look nice,” Alice objected.

Humpty Dumpty reports an inability to recognise a familiar face, yet is able to recognise eyes, nose, and mouth and their correct positions, as is also the case with prosopagnosics. In developmental or congenital prosopagnosia, where the neuropsychological deficit is perhaps most pure because acquired cases following pathological insults such as cerebrovascular disease may not respect functional boundaries and may be accompanied by additional neurological signs such as visual field defects, there are impairments in face identity matching tasks but the ability to identify sex, age, emotional facial expression, and eye gaze direction is preserved. As in these cases, Humpty Dumpty’s account seems to indicate preserved componential but impaired configural processing. There is also a suggestion that Humpty Dumpty might be able to use extraneous information to assist in facial recognition, his example being two eyes on one side of the nose or the mouth at the top of the face. Prosopagnosics may use extraneous visual cues such as spectacles, facial jewellery, and hair colour or style to aid facial recognition.

Whether Dodgson wrote this passage purely from imagination, or he based it upon observation of a prosopagnosic individual is not known. He did occasionally parody human idiosyncrasies, for example he himself appears as the Dodo because of his stammer (“Do-do-Dodgson”) in Alice’s Adventures in Wonderland (1865, chapters 2 and 3).

A J Larner
Cognitive Function Clinic, Walton Centre for Neurology and Neurosurgery, Lower Lane, Fazakerley, Liverpool, L9 7LU, UK; a.larner@thewalltoncentre.nhs.uk

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A J Larner

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