Automated measurement of unsaturated iron binding capacity is an effective screening strategy for C282Y homozygous haemochromatosis

P E Hickman, L F Hourigan, L W Powell, F Cordingley, G Dimeski, B Ormiston, J Shaw, W Ferguson, M Johnson, J Ascough, K McDonell, A Pink, D H G Crawford

Abstract

Background—C282Y hereditary haemochromatosis is an appropriate condition for population screening. Transferrin saturation, the best screening test to date, is relatively expensive, labour intensive, and cannot be automated. Unsaturated iron binding capacity is a surrogate marker of transferrin saturation and its measurement can be automated.

Aims—To evaluate a screening strategy for C282Y hereditary haemochromatosis in a tertiary hospital environment based on unsaturated iron binding capacity as the initial screening test.

Methods—Measurement of unsaturated iron binding capacity was adapted to the main laboratory analyser. An unsaturated iron binding capacity of less than 30 µmol/l was identified as an appropriate decision point and 5182 consecutive subjects were screened over 28 consecutive days.

Results—Of those screened, 697 had an unsaturated iron binding capacity less than 30 µmol/l. Of these, transferrin saturation was greater than 40% in 294. A total of 227 were able to be genotyped for the C282Y mutation. Nine subjects homozygous for C282Y were identified. Based on full cost recovery, affected persons were identified at a cost of Aus$2268 per case (approximately US$1496).

Conclusion—Automated measurement of unsaturated iron binding capacity enables a cost effective, large scale population screening programme for C282Y hereditary haemochromatosis to be developed.

Keywords: unsaturated iron binding capacity; haemochromatosis; screening

Hereditary haemochromatosis (HHC) is one of the most common autosomal recessive diseases in whites. The reported prevalence of the condition in this population is approximately 0.3%, and may be as high as 1.0% in populations of Celtic origin. It has been suggested that this high prevalence and the widespread availability of an acceptable and effective treatment make haemochromatosis an ideal condition for widespread population screening programmes. Indeed, population screening for HHC is an area of intense interest and a subject of some controversy as haemochromatosis fulfils almost all the criteria for screening programmes as defined by the World Health Organisation. However, the natural history of the disease is incompletely understood, and there is vigorous debate regarding optimal screening strategies for this condition. The best phenotypic marker for the condition seems to be transferrin saturation. However, the measurement of transferrin saturation is not automated, is labour intensive, and relatively expensive; these factors preclude long term screening programmes based on this test. Unsaturated iron binding capacity (UIBC) is a surrogate marker of transferrin saturation and has been reported to identify 100% of iron overloaded patients and 95% of patients with normal iron stores. It has been proposed as a suitable screening method for HHC.

In 1996, Feder et al described a novel gene (now known as HFE) located on chromosome 6. Eighty two per cent of their American patients with hepatic iron overload had a substitution of cysteine to tyrosine at position 282 of the gene product. Since this publication it has become apparent that in populations of a northern European extraction, most primary iron overload is associated with the C282Y mutation. An absolute requirement for establishing the diagnosis of expressing HHC is iron overload, which is identified universally by the presence of an increased transferrin saturation. However, not all C282Y homozygotes have biochemical markers of iron overload. Thus it seems logical to consider patients with C282Y homozygosity and increased transferrin saturation as expressing the disease and the population that should be targeted for detection in a screening programme.

In this study, we employed a strategy in a hospital based screening programme for expressed C282Y homozygous haemochromatosis whereby UIBC was used as the primary screening tool followed by C282Y genotyping in patients with a confirmed elevation of transferrin saturation. We report the clinical and economic efficiency of this strategy in a sample of 5182 consecutive patients at a major Australian tertiary hospital.

Materials and methods

SUBJECTS AND STUDY DESIGN

The study was performed over a four week period in November and December 1997, during which time any blood sample accompanied
by a request for analysis on the main laboratory analyser was included in the study. Over the four weeks of the study, 5182 separate persons had plasma samples analysed for UIBC measurement. A preliminary study to determine an appropriate cut off value for UIBC was performed with 73 samples that had a transferrin saturation of 40% or greater. Regression analysis indicated that a transferrin saturation of 40% corresponded to a UIBC of 25 µmol/l \( (r=-0.87). \) A UIBC cut off value of 30 µmol/l was selected to ensure maximum detection of affected subjects.

If UIBC was less than 30 µmol/l, the transferrin saturation was measured on the serum sample; if transferrin saturation was greater than 40%, serum ferritin concentration was measured and genotyping for the C282Y mutation was performed.

**LABORATORY PROCEDURES**

**Unsaturated iron binding capacity**

The following method was adapted to the Hitachi 747 analyser (Hitachi, Japan). A fixed amount of exogenous iron was added to a sample of plasma to fill all available iron binding sites on transferrin, with a small excess remaining free in plasma. After equilibration, the iron specific dye ferrozine was added, to react with the free iron. The colour that developed was proportional to the initial iron saturation of the transferrin molecule. At a UIBC of 17 µmol/l, the interassay coefficient of variation was 3.8%.

**Transferrin saturation and ferritin analysis**

Transferrin saturation and serum ferritin were measured by standard analytical procedures on a Hitachi 912 analyser (Hitachi, Japan).

**Computing**

To prevent repeated assay of the same person during the course of this study, the computer program controlling the laboratory information system was adjusted so that whenever a new sample was presented to the analyser for analysis, a check was performed to see whether previous measurement of UIBC had been made for that patient. If measurement had not been performed, then UIBC was added to the request. This result was not reported but was placed into a separate file which could be accessed at will.

**Genotyping**

DNA was extracted from peripheral blood lymphocytes, and the C282Y mutation was determined by SnaB1 restriction enzyme digestion, followed by polymerase chain reaction (PCR) and agarose gel electrophoresis.

**Calculation of costs**

All procedures within our laboratory have been costed as part of an Australia wide benchmarking process, and these costs were used to determine the real costs of all procedures used.

**ETHICAL APPROVAL**

The study was conducted with the approval of the Ethics Committee of the Princess Alexandra Hospital, Brisbane.

**Results**

Over the course of the study, UIBC was measured in 5182 patients. Figure 1 shows the distribution of UIBC in this hospital population. Figure 2 shows the relation between UIBC and the transferrin saturation. Some patients with a transferrin saturation greater than 40% had a UIBC greater than 30 µmol/l; however, essentially all patients with transferrin saturation greater than 50% were detected with this UIBC value, and all with a transferrin saturation greater than 60%.

A total of 697 of the 5182 subjects (13.5%) had a UIBC less than 30 µmol/l. The transferrin saturation was greater than 40% in 294 of these 697 subjects (42.2%); of these 294, genotyping for HFE mutations was able to be conducted in 227 (67 had either died or were lost to follow up). Initially we genotyped only those persons with no apparent cause for an increased transferrin saturation. Of 46 persons in this category we found two (4.34%) to be homozygous for the C282Y mutation. We decided to extend the genotyping to all persons with increased transferrin saturation regardless of cause. Of the 227 patients finally genotyped, nine (3.96%) were homozygous for the C282Y mutation, 44 were heterozygous, and 174 were homozygous normal.

Table 1 shows the iron related studies of the nine subjects with C282Y HHC. Two of these subjects had a previous diagnosis of expressed HHC as judged by histological and quantitative...
postnatal disease expression (raised transferrin saturation; its measurement cannot be readily automated and demand considerable resources). Therefore, the average cost per case detected (US$1347, using Aus$1.00 = US$0.66). Alternatively, the screening strategy employed in the current study may identify subjects with limited disease expression (raised transferrin saturation but normal serum ferritin concentration and homozygosity for the C282Y mutation; see case 9, table 1). However, our evidence indicates that subjects with these phenotypic characteristics are not common and occur in approximately 11% of the subjects detected by the screening strategy. Recently, an expert panel reached a consensus view stating that phenotypic measurement, rather than genetic testing, is recommended for population-based screening for haemochromatosis. Most authors agree that transferrin saturation is the best screening test for haemochromatosis. However, it is difficult to apply a large screening strategy for haemochromatosis based on transferrin saturation because its measurement cannot be readily automated and demand would overwhelm routine pathology laboratories. The data in the present study show that: UIBC is a satisfactory surrogate for transferrin saturation; its measurement can be automated; and application of a screening strategy based on UIBC can be successfully applied to a hospital-based population in a cost-effective manner.

The reference range for UIBC provided by the manufacturer was greater than 20 µmol/l. A preliminary analysis indicated that a UIBC of greater than 20 µmol/l corresponded to a transferrin saturation of approximately 55%. There have been several reports of the most appropriate “cut off” value of transferrin saturation for screening for haemochromatosis. Dadone and colleagues suggested a value of 62% whereas Leggett and colleagues suggested a value of 55%. There is now convincing evidence that some subjects with haemochromatosis have a transferrin saturation lower than these values. Recent mathematical modelling of data from a large Queensland population showed that a transferrin saturation of 45% would identify 98% of homozygotes of both sexes. Thus, to provide a reasonable safety margin and based on conclusions from a preliminary study, the UIBC decision point was extended to 30 µmol/l. Data presented in fig 2 indicate that even with this extended value, some persons with a transferrin saturation of between 40% and 50% may not be detected with a UIBC of greater than 30 µmol/l. However, as the transferrin saturations of these persons were close to the decision point of 30 µmol/l, they are still at risk of developing hepatic cirrhosis. The natural history of HHC is incompletely understood but the condition is characterised by a long latent period between onset of disease and the development of hepatic cirrhosis, during which time affected subjects are often asymptomatic. Venesection therapy is an acceptable treatment which prolongs life if instituted before the development of cirrhosis. These are important criteria endorsing the desirability of developing a widespread population screening programme for HHC in populations at risk. In this study, we showed the clinical and economic utility of unsaturated iron binding capacity as a screening tool for C282Y HHC in a hospital population. The results indicate that the actual cost of detecting each case was Aus$2268.77, which is approximately US$1496.

Table 1 Iron studies in homozygotes for the C282Y mutation

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<th>Sex</th>
<th>Age (y)</th>
<th>UIBC (µmol/l)</th>
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<th>Ferritin (µg/l)</th>
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*Haemochromatosis known. UIBC, unsaturated iron binding capacity.

Discussion

The natural history of HHC is incompletely understood but the condition is characterised by a long latent period between onset of disease and the development of hepatic cirrhosis, during which time affected subjects are often asymptomatic. Venesection therapy is an acceptable treatment which prolongs life if instituted before the development of cirrhosis. These are important criteria endorsing the desirability of developing a widespread population screening programme for HHC in populations at risk. In this study, we showed the clinical and economic utility of unsaturated iron binding capacity as a screening tool for C282Y HHC in a hospital population. The results indicate that the actual cost of detecting each case was Aus$2268.77, which is approximately US$1496.

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tion increased to more pathological values (above 50%), the proportion of subjects with an increased transferrin saturation who were not detected by a UIBC of less than 30 µmol/l fell substantially. Crawford and colleagues recently studied subjects homozygous for the C282Y mutation in the HFE and their data indicated that a transferrin saturation value of 50% would identify more than 90% of affected males and 80% of affected females. Clearly, screening programmes cannot identify 100% of all affected subjects as outlay costs to achieve such a result would be prohibitive.

Hereditary haemochromatosis is largely a disease of northern European populations with a reported prevalence of 1:300–1:400. The prevalence of haemochromatosis in the population studied in this hospital was 1:575. There are a number of reasons why the prevalence in the current study was less than expected. Fifteen per cent of subjects in the study were Asian or other ethnic grouping unassociated with C282Y. Furthermore, only 77.2% of the 294 subjects with a UIBC of less than 30 µmol/l were genotyped. This resulted either from patient death or loss to follow up in 67 subjects. The failure to perform genotype analysis in all subjects reflects the difficulties that will be encountered when this screening strategy is employed in a true clinical setting, as some subjects will choose not to pursue complete evaluation and others will not be contactable because of itinerant lifestyles. The rate of detecting C282Y homozygotes was remarkably constant as subjects were recalled for genotyping. We predicted that if all 294 subjects were genotyped, 11.65 C282Y homozygotes would be identified. If the subjects from ethnic groupings unassociated with C282Y are excluded from the analysis, this would provide a prevalence of approximately 1:380 which is similar to that found by Leggett and colleagues.

In addition to identifying affected haemochromatosis subjects, the implementation of a large scale screening programme using UIBC offers two other advantages. Firstly, a high UIBC occurs in iron deficiency and the widespread use of UIBC may result in earlier identification of subjects with colorectal cancer, peptic ulcer disease, and malabsorption. Secondly, the treatment of affected subjects by venesection and the use of this blood for transfusion purposes would help overcome the shortage of blood supplies which is a frequent occurrence in Australia and other countries. Blood transfusion policies in Australia are changing and blood from haemochromatosis subjects is now considered suitable for transfusion purposes.

Our experience of which persons to genotype is informative. In this large tertiary hospital, many patients have reasons other than HHC for an increased transferrin saturation, for example, transfusion dependency. Interestingly, the rate of detecting homozygotes was similar in those with and those without, clinical reasons for increased transferrin saturation. This observation suggests that completing the proposed screening strategy is important for subjects irrespective of whether there is an apparent underlying clinical reason for the increased transferrin saturation.

This study was designed to test the sensitivity or specificity of UIBC. Rather, the study was performed to test the economic efficiency of UIBC as the initial screening tool for C282Y haemochromatosis. However, it should be noted that almost 60% of those patients with a UIBC of less than 30 µmol/l did not have a transferrin saturation greater than 40%. In the preliminary regression analysis, a UIBC of 25 µmol/l corresponded to a transferrin saturation of 40%. As previously explained, a UIBC of less than 30 µmol/l was selected as the critical value to ensure maximum detection of affected subjects. A combination of error associated with regression analysis and the fact that 24% of subjects with a UIBC less than 30 µmol/l had a UIBC between 25 and 30 µmol/l explains, in part, the apparent poor relation of the UIBC and transferrin saturation near the cut off value of 30 µmol/l. Determining the precise cut off value to maximise efficiency of the UIBC screening test is the subject of the current study.

This study shows that the capacity now exists for large scale population screening for C282Y HHC that could be automated measurements of unsaturated iron binding capacity. Screening, even in a major hospital setting is clinically and cost effective. Furthermore, the cost per case detected is much less than other disease processes for which large scale population screening already exists. Our data suggest that automated measurement of UIBC is an appropriate strategy for large scale screening programmes for C282Y HHC in hospital settings.

We would like to acknowledge the assistance of Roche Diagnostics in the conduct of this study.


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*Gut* 2000 46: 405-409
doi: 10.1136/gut.46.3.405

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