Effect of genotype on changes in intelligence quotient after dietary relaxation in phenylketonuria and hyperphenylalaninaemia

Lindsay G Greeves, Christopher C Patterson, Dennis J Carson, Ruth Thom, Melanie C Wolfenden, Johannes Zschocke, Colin A Graham, Norman C Nevin, Elisabeth R Trimble

Abstract

Background—Associations between genotype and intellectual outcome in patients with phenylketonuria are complicated because intelligence is influenced by many variables, including environmental factors and other genetic determinants. Intellectual changes with age, both on and after relaxation of diet, vary within the patient population. This study aims to determine whether a significant association exists between genotype and change in intelligence after relaxation of diet.

Methods—125 patients with hyperphenylalaninaemia and phenylketonuria whose diet was relaxed after 8 years of age. Verbal, performance, and full scale intelligence quotients at 8, 14, and 18 years were expressed as standard deviation scores (IQ-SDS), and genotype as predicted residual enzyme activity (PRA) of phenylalanine hydroxylase.

Results—IQ-SDS at 8, 14, and 18 years were significantly below normal; no association was found between PRA and IQ-SDS. Significant reductions in verbal and full scale IQ-SDS occurred between 8 and 14 years and 8 and 18 years. There was a significant association between PRA and the reduction in verbal, performance, and full scale IQ between these years. Multiple regression analysis of 18 year results, using 8 year results as covariates, supported the association between PRA and IQ-SDS; after adjustment for phenylalanine control, both up to and after the age of 8 years, the full scale IQ-SDS at 14 and 18 years was 0.15 higher for each 10% increase in PRA.

Conclusions—Genotype might be useful in predicting the likelihood of intellectual change in patients with hyperphenylalaninaemia and phenylketonuria whose diet is relaxed after the age of 8 years.

Methods—We studied all patients with phenylketonuria and hyperphenylalaninaemia attending the regional centre who were born between 1969 and 1986 and identified by neonatal screening.
The following variables were considered: IQs, gender, father’s social class at diagnosis, age diagnostic venous sample was taken and/or diet commenced, number of days from birth until whole blood phenylalanine concentration was reduced to 500 µmol/litre or less, dietary content of phenylalanine prescribed at age 5 (mg/kg/day), and plasma/blood phenylalanine control. In addition, permission was requested from families for genotyping for mutational analysis of the phenylalanine hydroxylase gene. A few of these variables will be explained in more detail.

IQs AT 8, 14, AND 18 YEARS OF AGE

Intelligence testing was carried out on patients using prescribed tests at set ages (8, 14, 18 years). The “Wechsler intelligence scale for children” (WISC) or revised WISC (WISC-R) was used at 8 and 14 years, the “Wechsler adult intelligence scale” (WAIS) or the revised WAIS (WAIS-R) was used at 18 years of age. Verbal, performance, and full scale scores were recorded. The results were expressed as standard deviations from the population norms (IQ-SDS) and adjusted for long term secular trends using formulae described by Smith and Beasley.38 39 For full scale IQ, Population norms were estimated using results of a cross sectional analysis of population IQ trends of white Americans by Flynn.37 38 Flynn’s analysis has been shown to be applicable to children in the UK.39 40 In applying Flynn’s analysis to longitudinal data, it was assumed that individuals as well as populations show rises in IQ of 0.3 points each year and that the trends were the same for all tests used. In so doing, it is not possible to exclude a small, systematic error relating to change in absolute terms;41 however, the method is suitable for assessing change in intellectual outcome in relative terms within the patient group. Tests carried out within 9 months of 8th, 14th, or 18th birthdays were included.

SOCIAL CLASS

Social class was assigned according to the Registrar General’s classification 1980.41

GENOTYPE/PREDICTED RESIDUAL ENZYME ACTIVITY

Since 1994, automated sequencing has been used to detect the spectrum of mutations present in our phenylketonuria and hyperphenylalaninaemia population.42 43 Genotype was available on at least one sibling in a family; parents were also tested. The genotype of all affected siblings from the same family was then assumed to be the same. The phenylalanine hydroxylase residual enzyme activity of many mutations has been measured in vitro by various authors; however, this value is not known for all mutations. The PRA of each patient was calculated as the arithmetic mean of the residual enzyme activities (in vitro) of the two mutant alleles where these values were known.27 29 34 (table 1).

### Table 1: Phenylalanine hydroxylase mutations with corresponding in vitro predicted residual enzyme activities (PRA) used for subsequent analysis

<table>
<thead>
<tr>
<th>Mutation</th>
<th>Activity (% of normal)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R243X, R353W, R408W, F299C, IVS12nt1, IVS10nt546</td>
<td>0</td>
</tr>
<tr>
<td>E280K</td>
<td>3</td>
</tr>
<tr>
<td>R158Q</td>
<td>10</td>
</tr>
<tr>
<td>165TL 348VR261Q</td>
<td>30</td>
</tr>
<tr>
<td>Y414C</td>
<td>50</td>
</tr>
<tr>
<td>R408Q</td>
<td>55</td>
</tr>
</tbody>
</table>

**PLASMA/BLOOD PHENYLALANINE CONTROL**
Plasma/blood phenylalanine control was scored for 0–8, 8–14, and 8–18 years. In those under 8 years of age, all Guthrie results (carried out at least monthly) were considered once the blood phenylalanine concentration had fallen to 500 µmol/litre or less after commencement of dietary treatment. Phenylalanine concentrations 125–500 µmol/litre were considered good control. The proportion of results between 125 and 500 µmol/litre was calculated and a score given as follows: 1, 75–100%; 2, 50–74%; 3, 25–49%; and 4, 0–24%. An arithmetic mean phenylalanine concentration could not be calculated because some results fell outside the limits of sensitivity (low and high) of the Guthrie assay.

In this cohort of patients, diet was relaxed after the age of 8 years (1970s to early 1990s). Between the ages of 8 and 14 years and 8 and 18 years, phenylalanine control was measured for those on whom PRA could be calculated (see above). This was the mean of twice yearly plasma phenylalanine concentrations from amino acid analysis. Where a patient had more than two results in the year, the results used were those obtained nearest to the birthday visit and six months after that date.

**STATISTICS**
Associations between PRA, phenylalanine concentrations, and phenylalanine intake were initially assessed with Spearman’s rank correlation coefficients (r). The mean IQ-SDS at different ages were compared with normal values using the one sample t test. Changes in IQ-SDS were assessed using the paired samples t test. Comparisons of the mean IQ-SDS (and mean changes in IQ-SDS) between subgroups of children were obtained using either the independent samples t test or one way analysis of variance. Changes in IQ-SDS were initially examined in relation to PRA by simple regression analysis. Although the IQ results in this study were adjusted for age and sex to standard deviation scores (IQ-SDS), concerns remained about a possible lack of comparability between IQ assessments at different ages, particularly for the comparison of 8 and 18 year IQ results when different tests were used. For this reason analyses of the relation between change in IQ-SDS and PRA were repeated using multiple regression with IQ-SDS at age 14 or 18 years as the response variable and IQ-SDS at age 8 years as a covariate; in addition to PRA, levels of control both before and after dietary relaxation were included as predictor variables.
Table 2  Mutation frequency in 100 patients genotyped

<table>
<thead>
<tr>
<th>Mutation</th>
<th>Frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R408W</td>
<td>57 (28.5)</td>
</tr>
<tr>
<td>R408Q</td>
<td>42 (21.0)</td>
</tr>
<tr>
<td>F391L</td>
<td>17 (8.5)</td>
</tr>
<tr>
<td>Y414C</td>
<td>12 (6.0)</td>
</tr>
<tr>
<td>L348V, R408Q</td>
<td>10 (5.0)*</td>
</tr>
<tr>
<td>F299C</td>
<td>9 (4.5)</td>
</tr>
<tr>
<td>R241X, IVS121nt1</td>
<td>5 (2.5)*</td>
</tr>
<tr>
<td>G46S</td>
<td>4 (2.0)</td>
</tr>
<tr>
<td>E280K, S67P, T380M</td>
<td>3 (1.5)*</td>
</tr>
<tr>
<td>L249F, L333F, R241L, R252W, R243Q</td>
<td>2 (1.0)*</td>
</tr>
<tr>
<td>S273F, A309V, A300V, IVS0nt546, A308Q</td>
<td>1 (0.5)*</td>
</tr>
<tr>
<td>A309D, IVS10nt554, A300V, R261Q, S349P, unidentified</td>
<td>1 (0.5)*</td>
</tr>
</tbody>
</table>

*Frequency for each mutation in this group.

Results

POPULATION STUDIED

There were 125 patients, 64 boys and 61 girls (all white), who were born between 1969 and 1986. Sixtyseven per cent were from families where the head of the household was in a manual occupation at the time of diagnosis.

One hundred and eleven patients (93%) were diagnosed at ≤21 days of age. (There were six patients for whom this information was not available.) The remaining eight were diagnosed between 24 and 56 days; the reason for the delay was that they were mildly affected patients who were monitored by Guthrie test until phenylalanine concentration increased to a value requiring therapeutic intervention, at which stage a venous sample was taken. The time from birth until the whole blood (Guthrie) phenylalanine concentration fell to <500 µmol/litre ranged from nine to 63 (median, 21) days.

Permission was obtained for genotyping in 100 patients; the remainder failed to respond to the request. Forty six different genotypes were found in this group of patients, the most common being I65T/R408W (10%), R408W/R408W (9%), I65T/Y414C (7%), and I65T/I65T (6%). There were 27 different alleles of which R408W (28.5%), I65T (21%), F391L (8.5%), and Y414C (6%) were the most common (table 2).

PHENYLALANINE CONCENTRATIONS AT DIAGNOSIS, PHENYLALANINE INTAKE AT 5 YEARS, AND PHENYLALANINE CONTROL IN THE FIRST 8 YEARS WITH RESPECT TO PRA

PRA was calculated for 64 patients (64% of those genotyped); PRA values were not available for the mutations found in the remaining patients at the time of analysis.

In the patient group, plasma phenylalanine concentrations at diagnosis ranged from 4750 (median, 1945) µmol/litre (excluding siblings who were diagnosed early because of the known family history). There was a negative correlation between PRA and plasma phenylalanine concentrations at diagnosis, excluding siblings (n = 42; r = −0.53; p < 0.001).

Phenylalanine intake prescribed at 5 years ranged from 6.9 to 121.0 (median, 17.1) mg phenylalanine/kg body weight/day. There was a positive correlation between PRA and phenylalanine intake at 5 years (n = 63; r = 0.63; p < 0.001); there was also a negative correlation between plasma phenylalanine concentration at diagnosis and prescribed phenylalanine intake at 5 years (n = 77; r = −0.44; p < 0.001).

Phenylalanine control in the first 8 years was categorised into groups where 1 was best control and 4 was worst control (see Methods). There was a significant association between PRA and control in the first 8 years (n = 62; r = −0.38; p = 0.003).

INTELLECTUAL OUTCOME

IQ-SDS at 8, 14, and 18 years of age

IQ-SDS at 8, 14, and 18 years were each significantly below normal (p < 0.001). The mean full scale IQ-SDS was −0.79 (95% confidence interval (CI), −1.01 to −0.57). For 108 children examined at 8 years, −1.05 (95% CI, −1.32 to −0.78) for 76 children examined at 14 years, and −1.17 (95% CI, −1.42 to −0.93) for 51 children examined at 18 years. Similar deficits were seen for verbal and performance IQ-SDS (not shown). Sex had no effect on IQ-SDS but children of manual workers had consistently lower IQ scores than those of nonmanual workers; this was significant for verbal and full scale scores at 8 years of age and verbal scores at 14 years of age (p < 0.05) (results not shown). Significant change in verbal and full scale IQ-SDS occurred both between 8 and 14 years and 8 and 18 years (table 3).

Associations between IQ-SDS, phenylalanine control, and PRA

Of 112 patients who had blood phenylalanine concentrations available between 0 and 8 years, 18% of patients had results assigned to category 1 (best phenylalanine control), 47% to category 2, 23% to category 3, and 12% to category 4 (worst phenylalanine control). There was a significant association between phenylalanine control at 0–8 years and IQ-SDS at both 8 and 14 years; the association was no longer significant for IQ-SDS at 18 years (table 4).

Neither the change in IQ-SDS between 8 and 14 years nor the change between 8 and 18 years was significantly related to phenylalanine control in the first 8 years.

Dietary control was relaxed after 8 years. Mean plasma phenylalanine concentrations between 8 and 14 years ranged from 406 to 1623 (n = 43; mean, 912; SD, 293) µmol/litre, whereas those between 8 and 18 years ranged from 567 to 1627 (n = 31; mean, 1044; SD, 247) µmol/litre. There was a significant association between PRA and phenylalanine control between 8 and 14 years (n = 43; r = −0.37;
Genotype and intellectual change

Table 4  Association between dietary control between 0 and 8 years and verbal, performance, and full scale IQ standard deviation scores (IQ-SDS) at 8, 14, and 18 years

<table>
<thead>
<tr>
<th>Category</th>
<th>1 (best)</th>
<th>2</th>
<th>3</th>
<th>4 (worst)</th>
<th>p Value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>8 years IQ-SDS</td>
<td>(n = 15)</td>
<td>(n = 51)</td>
<td>(n = 25)</td>
<td>(n = 13)</td>
<td></td>
</tr>
<tr>
<td>Verbal</td>
<td>−0.22 (−0.69 to 0.24)</td>
<td>−0.26 (−0.58 to 0.06)</td>
<td>−1.05 (−1.58 to −0.52)</td>
<td>−1.11 (−1.69 to −0.53)</td>
<td>0.007</td>
</tr>
<tr>
<td>Performance</td>
<td>−0.01 (−0.95 to −0.28)</td>
<td>−0.71 (−1.02 to −0.39)</td>
<td>−1.43 (−1.88 to −0.98)</td>
<td>−1.66 (−2.04 to −1.29)</td>
<td>0.001</td>
</tr>
<tr>
<td>Full Scale</td>
<td>−0.39 (−0.77 to −0.01)</td>
<td>−0.47 (−0.80 to −0.15)</td>
<td>−1.32 (−1.82 to −0.82)</td>
<td>−1.47 (−1.92 to −1.01)</td>
<td>0.001</td>
</tr>
<tr>
<td>14 years IQ-SDS</td>
<td>(n = 6)</td>
<td>(n = 31)</td>
<td>(n = 20)</td>
<td>(n = 12)</td>
<td></td>
</tr>
<tr>
<td>Verbal</td>
<td>−1.03 (−2.36 to 0.31)</td>
<td>−0.46 (−0.91 to −0.01)</td>
<td>−1.33 (−1.82 to −0.84)</td>
<td>−1.70 (−2.15 to −1.25)</td>
<td>0.006</td>
</tr>
<tr>
<td>Performance</td>
<td>−0.61 (−2.03 to 0.81)</td>
<td>−0.67 (−1.13 to −0.21)</td>
<td>−1.36 (−1.79 to −0.92)</td>
<td>−1.72 (−2.32 to −1.11)</td>
<td>0.02</td>
</tr>
<tr>
<td>Full Scale</td>
<td>−0.85 (−2.29 to 0.59)</td>
<td>−0.52 (−0.99 to −0.05)</td>
<td>−1.41 (−1.85 to −0.97)</td>
<td>−1.82 (−2.34 to −1.30)</td>
<td>0.004</td>
</tr>
<tr>
<td>18 years IQ-SDS</td>
<td>(n = 2)</td>
<td>(n = 21)</td>
<td>(n = 14)</td>
<td>(n = 10)</td>
<td></td>
</tr>
<tr>
<td>Verbal</td>
<td>−1.57 (−3.53 to 0.40)</td>
<td>−0.91 (−1.32 to −0.51)</td>
<td>−1.06 (−1.60 to −0.52)</td>
<td>−1.49 (−2.08 to −0.90)</td>
<td>0.34</td>
</tr>
<tr>
<td>Performance</td>
<td>−1.57 (−9.89 to 6.76)</td>
<td>−0.91 (−1.30 to −0.52)</td>
<td>−0.96 (−1.61 to −0.31)</td>
<td>−1.83 (−2.28 to −1.37)</td>
<td>0.06</td>
</tr>
<tr>
<td>Full Scale</td>
<td>−1.64 (−6.27 to 3.00)</td>
<td>−0.94 (−1.33 to −0.55)</td>
<td>−1.09 (−1.65 to −0.53)</td>
<td>−1.72 (−2.23 to −1.20)</td>
<td>0.12</td>
</tr>
</tbody>
</table>

Results are expressed as mean (95% confidence interval).
Categories: 1, 75–100% phenylalanine concentrations between 125 and 500 µmol/litre; 2, 50–74% phenylalanine concentrations between 125 and 500 µmol/litre; 3, 25–49% phenylalanine concentrations between 125 and 500 µmol/litre; 4, 0–24% phenylalanine concentrations between 125 and 500 µmol/litre.

*One way analysis of variance.

Table 5  Regression coefficients for potential predictors of full scale IQ standard deviation scores (IQ-SDS) at 14 and 18 years

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Full scale IQ-SDS at 14 years (n = 42)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full scale IQ-SDS at 8 years</td>
<td>0.86 (±0.69 to 1.03)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>PRA (per 10% of normal)</td>
<td>0.15 (±0.00 to 0.29)</td>
<td>0.048</td>
</tr>
<tr>
<td>Control before 8 years (per scale step*)</td>
<td>−0.09 (±0.35 to 0.18)</td>
<td>0.52</td>
</tr>
<tr>
<td>Control after 8 years*</td>
<td>0.00 (±0.07 to 0.08)</td>
<td>0.94</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Full scale IQ-SDS at 18 years (n = 29)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full scale IQ-SDS at 8 years</td>
<td>0.86 (±0.69 to 1.03)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>PRA (per 10% of normal)</td>
<td>0.15 (±0.00 to 0.29)</td>
<td>0.048</td>
</tr>
<tr>
<td>Control before 8 years (per scale step*)</td>
<td>−0.09 (±0.35 to 0.18)</td>
<td>0.52</td>
</tr>
<tr>
<td>Control after 8 years*</td>
<td>0.00 (±0.07 to 0.08)</td>
<td>0.94</td>
</tr>
</tbody>
</table>

*pProportion of phenylalanine results between 125 and 500 µmol/litre scored as: 1, 75–100%; 2, 50–74%; 3, 25–49%; 4, 0–24%.
†For each 100 µmol/litre increase in phenylalanine.
CI, confidence interval.

and PRA, there was a significant association between the change in IQ-SDS and PRA, both between 8 and 14 years and 8 and 18 years (fig 1).

Table 5 summarises the multiple regression analysis of full scale IQ-SDS at 14 and 18 years in relation to PRA and phenylalanine control. Coefficients for PRA have been scaled to represent the difference in IQ-SDS associated with an increase in PRA of 10% of the normal value, whereas coefficients for control after the age of 8 years have been scaled to represent the difference in IQ-SDS associated with a worsening in phenylalanine concentrations of 100 µmol/litre. Both the 14 year and 18 year analyses suggest that PRA is an important predictor of IQ-SDS after adjustment for phenylalanine control up to and after the age of 8 years. The coefficients indicate that each 10% increase in PRA is associated with a 0.15 higher IQ-SDS at each age.

Discussion

The fact that 46 different genotypes were found in this patient group meant that it was impossible to consider directly the effect of a specific genotype on intellectual outcome; others who have attempted to do this in either treated or untreated patients have not shown a simple association.35 36 The method used to circumvent this problem was to express genotypes as the activities of the mutant enzymes, measured under in vitro conditions and termed predicted residual enzyme activity (PRA). It is recognised that this27 28 and other44 methodologies all have some limitation when extrapolated to the in vivo condition. It was assumed that affected
siblings from the same family have the same genotype, an assumption that was considered safe because parents were also genotyped; this avoided the possibility of three mutant alleles, one encoding mild hyperphenylalaninaemia, occurring within one family, as has been reported previously. All of the patients were diagnosed and treated early, and phenylalanine concentrations fell rapidly into the therapeutic range (median, 21 days after birth). Until the early 1990s, it was policy to relax the diet at the age of 8 years, and this is reflected in the plasma phenylalanine concentrations after this age. The associations between PRA and plasma phenylalanine concentration at diagnosis, and between PRA and prescribed phenylalanine intake at 5 years of age, which have been described previously, were confirmed in our study. The fact that mean IQ results were significantly below normal is also in keeping with previous studies.

Change in IQ-SDS was calculated as the difference between the 14 or 18 year results and the 8 year results and was found to be significant for verbal and full scale IQ-SDS. The limitations of this calculation have been detailed above; the multiple regression approach also takes into account some of the concerns about the validity of IQ changes. The results show that not all patients deteriorated intellectually when diet was relaxed after 8 years, and are in keeping with those of Waisbren and colleagues who, using absolute IQ results, found that 40% of those in their sample population gained IQ points or maintained their IQ value after discontinuation of diet at 5 years of age. In the present study, patients with a higher PRA were less likely to deteriorate intellectually and some even improved; this is most apparent in those with a PRA > 25% of normal values (genotypes R408Q/R408Q, L348V/R408Q, I65T/Y414C, I65T/L348V, and I65T/I65T). Simple regression analysis has confirmed an association between PRA and change in IQ-SDS. Multiple regression analysis of IQ-SDS at 14 and 18 years, using 8 year results as covariates, showed that PRA was an important predictor.

In a study of 64 patients still under strict dietary control, no correlation was found between PRA and IQ (WISC-R) at the age of 9 years; however, there was an association between low PRA and high fluctuations in phenylalanine concentrations. In another study of 55 older, untreated patients, a simple association was not found between PRA and IQ (assessed by the Binet test at undisclosed ages). However, both these studies measured IQ at a single point in time. We also found no significant association between PRA and IQ-SDS when assessed at a single time point. The lack of association could, in part, be the result of both differences in intrinsic intellectual endowment and other genetic interactions in the metabolism of phenylalanine, in addition to that exerted by phenylalanine hydroxylase. In patients with phenylketonuria and hyperphenylalaninaemia with different genotypes, longitudinal measurements evaluating change in IQ in the same individual render less significant the roles played by other genes and intrinsic intellectual endowment. This might explain why a strong association was found between PRA and change in IQ-SDS between 8 and 18 years.

It is clear that there is a complex relation between PRA, dietary management, and IQ. An association between PRA and control both before and after dietary relaxation was found in our study. However, Burgard et al found no relation between PRA and control. Their patients were all under good control, as measured by the mean of the first nine yearly phenylalanine medians. It is of interest that in our study the effects of control before 8 years of age were seen on IQ-SDS at both 8 and 14 years but no longer at 18 years of age.

This study has shown that patients with a PRA of 25% or greater are more likely to maintain or gain IQ points after relaxation of diet in the years 8–18 than those with a lower PRA. If further prospective and multicentre studies confirm these findings, the importance of genotype, PRA, and strict dietary control after the age of 8 years will need to be re-evaluated.

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