LETTERS TO THE EDITOR

The Gene Bomb

I would like to comment on Tom Wilkie's comments on 'The Gene Bomb' (J Med Genet 1997;34:438-9), since I feel he missed the whole point of the book and presented to the readers of this journal a distorted view of the issues I attempted to raise. They have implications for all clinicians.

First, the subtitle, which somewhat more accurately portrays the issues, Does Higher Education and Advanced Technology Accelerate the Selection of Genes for Learning Disorders, ADD and addictive and disruptive behaviors? was left out.

Second, no mention was made of part I of the book which reviews data showing the following.

(1) Epidemiological studies using structured diagnostic instruments have shown a highly significant increase in frequency, and decreased in the age of onset, in younger cohorts for all the major psychiatric disorders including, adult onset, manic depressive disorder, schizoaffective disorder, suicide, obsessive-compulsive disorders, panic disorder, phobias, alcoholism, and drug abuse.

(2) Using the Achenbach Child Behavior Checklist, Achenbach performed epidemiological studies in 1976 and 1989. Of 112 behavioural scores, 42 were worse in 1989 while none was worse in 1976. In addition, behaviour relating to specific diagnoses, such as ADHD and conduct disorder, increased from 10% in 1976 to 18.2% only 13 years later. In 1976, 8.2% of children had to repeat a grade while in 1989 17.8% had to repeat a grade.

(3) Studies of the same localised region in Sweden by Gillberg have shown an increase in the frequency of autism from 4/10 000 in 1980, to 7.5/10 000 in 1984, to 11.5/10 000 in 1988.

(4) US Department of Education studies have shown a progressive increase in the frequency of learning disabilities from 890 000 cases in 1976 to 2 714 000 cases in 1993. At the same time the population has only increased from 210 000 000 to 240 000 000.

To date there have been no adequate explanations for these trends and observations, very few professionals, including geneticists, have either been aware of or concerned about these trends, and no one has offered any satisfactory explanation for them.

I work with children with hereditary behavioural disorders (ADD, Tourette syndrome, learning disabilities, autism, Asperger syndrome, conduct disorder, and others) on a daily basis and I and their parents are concerned. I am particularly concerned about the trend in learning disabilities. These have increased from 2 000 000 in 1988 to 2 700 000 in 1993. At this rate the frequency of learning disorders is doubling every 15 years.

In attempting to understand these trends, I noticed the following. First, all of these disorders have a strong genetic component. Second, there is a strong inverse correlation between the frequency of these disorders, especially learning disorders, and years of education. Not surprisingly, those with learning disorders tend to drop out of school, while those without learning disorders are more likely to go to college and possibly graduate school. Third, there is an extraordinarily high inverse correlation between the number of years of education and the age of initiation of child birth. At the beginning of this century only 2% of the population attended college. Today, 38% attend college. All of these trends in behaviour disorders have occurred in the latter half of the century. At the present time the spread in years of age of onset of childbearing, for those who drop out of school before completing a high school education versus those who go to college, is approaching 20 years, and this will raise, anything that turns over more rapidly will more rapidly increase in frequency. This can provide a very strong selection for the genes associated with learning disability, totally independent of the number of children people have. Comparing 1920 to 1990 shows that the magnitude of the selective force (spread in years by percent attending college) has increased 40 fold.

How do we respond to these observations or concerns? There are several possibilities. One is to shoot the messenger, as Wilkie has chosen to do. A second is to provide alternative explanations for these trends. I would prefer that we all stop writing about the subject. The third is to accept it as a legitimate concern and either disprove the hypothesis or continue research into the genes involved in these disorders so that their frequencies in the population by age can be monitored.

In the book I suggested two approaches. The first were "motherhood and apple pie" approaches that no one should object to, such as special programmes to encourage children with ADHD and learning and related disorders to remain in school, and programmes for day care for children of college students so that if they want to they could start a family before completing their education. The second approach is to continue (or accelerate) the research towards providing a better understanding of the genes involved. Among other things this could involve couples participating in totally voluntary programmes by which they could make informed choices about reproductive decisions, as we now encourage people with a family history of cystic fibrosis or Huntington's disease to do.

Finally, I would like to point out that when the various behavioural disorders, most of which appear to be polygenic, are combined they occur in over 35% of the general population. This is 35 times the frequency of all the single gene disorders combined. I would suggest that those who propose that geneticists should not be involved in studies of the genetic etics of human behaviour and prefer that we do not engage in responsible talk about some of the social implications of these studies, because it may give geneticists a bad name, are guilty of sticking their head in the sand. If geneticists can't become involved and play some role in attempting to understand the cause of these trends, I suspect no one else. In his book, And the Band Played On, Randy Shilts provides us with the professional community for failing to mount an early response to the AIDS epidemic, because they preferred not to become involved or felt uncomfortable with the subject. Are we going to be similarly chastised in the next century because we also failed to care, or felt uncomfortable with the subject, and made no attempt to understand these trends?

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Gaucher disease plus

In a recent letter, Uyama et al discussed a Japanese sibship of three subjects with glucocerebrosidase deficiency. These patients had oculomotor abnormalities characteristic of neuroophthalmic Gaucher disease along with several more unusual clinical manifestations described in other populations, including valvular calcifications, communicating hydrocephalus, and corneal opacities. Yet, although the patients are reported to have glucocerebrosidase deficiency and point mutations on both glucocerebrosidase alleles, the authors suggest that the patients have a "Gaucher-like" disorder.

Gaucher disease is characterised by a vast phenotypic heterogeneity that is continually expanding. Recently, there has been a greater appreciation that diverse symptoms including congenital ichthyosis, hydrops fetalis, primary pulmonary hypertension, valvular calcifications, and even Parkinsonian symptoms can be seen in patients with Gaucher disease. Thus, we would suggest that the symptoms observed in the patients described by Uyama et al are just part of the broader Gaucher phenotype.

Although the authors mention other genes near to glucocerebrosidase, they discount the likelihood of a contiguous gene effect. Since the discovery of human metaxin, a gene located immediately downstream and transcriptionally coupled to the glucocerebroside pseudogene, we have characterised the locus more completely and have found that the region encompassing glucocerebrosidase is particularly gene rich. The physical relationship of metaxin and glucocerebrosidase on human chromosome 1q21 described by Uyama et al is incorrect, for actually the pseudogene for human metaxin and not the gene is immediately downstream to the glucocerebrosidase gene. We suggest that patients with these more unusual Gaucher phenotypes may indeed have "Gaucher disease plus", an interruption in a second contiguous gene, rather than a "Gaucher-like" illness. In Hunter syndrome, it has recently been shown that some patients with atypical phenotypes have deletions including segment of nearby genes. Further characterisations of all the genes around the glucocerebrosidase locus may provide a more complete explanation for the phenotypic variation observed in Gaucher disease.

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2 Uyama E, Takedashi K, Owada M, et al. Hydrocephalus, corneal opacities, deafness, valvular

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