could be made part of national curricula, and provide the basis for the continuing education of teachers in new developments in the biosciences.

The mountain of sceptics and anti-science people will not come to Mohammed

Although the EC has not yet developed a Europe-wide approach to the public communication of science, independent organizations have. The European Federation of Biotechnology, with remarkable prescience, set up a Task Group on Biotechnology 9 years ago, hence anticipating the GM debate. With a membership of 50,000, which more recently includes industrial interests, the EFB supports scientific conferences, public debates and press conferences. Another aspiring pan-European endeavour is the European Genetics Foundation (EGF) in Sestri Levante, Italy. After the success of the publicly open conference “Who owns the genome?” in 1999, the EGF is now coordinating the Genetics in Europe Open Days (GEOD) project in 2000. These publicly open conferences on the human genome and genetics take place in Milan, Barcelona, London and Heidelberg in November (www.geod.org). The open days will become an annual fixture.

However, science public relations is still a largely national rather than pan-European exercise. As in many other respects, European countries differ vastly in their efforts to break down the barriers between scientists and the public. Admittedly some have far greater economic concerns than others; public concern over scientific advances is in general positively correlated with GDP per capita. The UK has a particularly impressive record in the communication of science, which makes it all the more ironic that most of the recent public reaction against scientists has surfaced there. The three organizations that do most are the British Association for the Advancement of Science (BAAS), the Royal Society (RS) and the Royal Institution (RI), which together established COPUS. There are science buses—travelling exhibitions and laboratories—in several countries, and science on the buses in the UK, a campaign of brightly coloured, thought-provoking posters with very few words, which capture the interest of shoppers and commuters alike (http://www.uwe.ac.uk/fas/wavelength/wave18/burnet.html). The poster on cloning, for example, challenges its audience with the following words: ‘Identical twins are clones, and have the same genes…. …They may look like Hitler, but behave like Charlie Chaplin’.

The professions serving the public communication of science are starting to blossom in Europe. Universities are establishing courses in the communication of science, and journalists can now visit laboratories, look over a researcher’s shoulder for a few days and learn more of the method of science. Back pages of popular science magazines advertise lay courses and workshops given by active scientists. Predictably, the USA is a step ahead. America has had excellent science journalism schools and a highly respected association of science writers (NASW). Every provincial university and institute, no matter how small, has a PR department. National museums and visitor centres host web pages containing a variety of teaching and public information resources, e.g. Access Excellence hosted by the National Health Museum. On the national network scale, the North American Alliance for the Public Understanding of Science and Technology, a National Science Foundation project, concerns itself with the whole chain of science practice and communication, from basic research through teaching, and on to public understanding.

Once aware and engaged, the public expect to be involved in a dialogue concerning issues that could affect their lives

The activities briefly mentioned here, without doubt, make the public more aware of science, and give non-scientists the tools with which to discuss and think about science. But once aware and engaged, the public expect to be involved in a dialogue concerning issues that affect or could affect their lives. Scientists are starting to become more aware of, and engaged with, the public. Indeed, some approaches to teaching science are moving away from the ‘life in science’ paradigm of the past towards a ‘science in life’ model. Further up the career ladder, many now realize that the assessment of a researcher’s performance by national research councils should reward public communication of science as one of the researcher’s activities, rather than penalize it as time spent outside research. Contact with the public has been made. The next frontier is dialogue.

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Bear market slashes at human genome

The dropping guesses about the number of human genes challenges researchers to explain human complexity with so few genes

The media took fascinated note last May when genome researchers started gambling on the number of human genes. One trend was apparent immediately: the estimates of human gene number—made by the people who ought to know if anyone—were low compared with those made just a few years ago, when the number often tossed around was 100 000,
give or take. For some weeks, the median GeneSweep bet (http://www.ensembl.org/genesweep.html) stood at just under 54 000. In June this year, Nature Genetics acknowledged the same trend. It published three major papers estimating human gene number. Two of the three assessments were very low: ∼34 000 and 30 000 genes.

With nearly 300 wagers recorded as of early August, the GeneSweep estimates have trended upward again, which may have something to do with the announced completion of a working draft of the human genome sequence at the end of June. At that time, the National Human Genome Research Institute (NHGRI) confirmed the existence of 38 000 predicted genes. So it appears that anyone who bets below that is out of luck. Although there are contrary opinions, of course, the luminaries are at the head of the parade. NHGRI’s director Francis Collins has joined the accelerating trend to lowball guesses; his wager is 48 011. David Baltimore, perhaps taking his cue from the project leader, wrote in The New York Times that 50 000 seemed about right to him.

For the purposes of GeneSweep wagering, a gene is defined as a protein-coding sequence. Alternatively spliced transcripts are counted as one gene. Yet the numbers wagered are disconcerting, especially when compared with the gene count of ∼19 000 for the recently completed DNA sequence in Caenorhabditis elegans. Can it be that our illustrious species has only two or maybe three times as many genes as a transparent worm which dwells in the dirt beneath our feet? If this humiliating news is true, how did we get to be so majestic, so charming, so complicated? And what does it suggest about future directions for genome research?

Can it be that our illustrious species has only two or three times as many genes as a transparent worm which dwells in the dirt beneath our feet?

When Brent Ewing and Phil Green of the University of Washington in Seattle presented their low estimates of 35 000 genes in Nature Genetics, they speculated that the complexity of vertebrates is traceable to diversification of regulatory networks or alternative splicing, rather than to sheer gene number. If the low gene numbers for the human genome hold true, researchers will have to change their view of how genetic information is converted into diversity. It will also mean that in silico approaches to annotating the human genome may never reveal its entire complexity. These outcomes seem plausible enough, considering that an estimated 30% of human genes can be spliced to yield different forms of the protein product.

Paula Grabowski from the University of Pittsburgh has been pursuing alternative splicing, especially its regulation, for years. She and her colleagues investigate a number of genes in the nervous system—the place where most alternative splicing occurs—in order to uncover expression patterns and regulation strategies. One object of their study is a subunit of the receptor for the neurotransmitter GABA. The neuron-specific exons of the gene are spliced differently in different types of nerve cells during the same course of development.

In silico approaches to annotating the human genome may never reveal its entire complexity

‘The take-home message is that it’s extremely complex,’ she says. ‘The mode of regulation involves mechanisms that activate certain splicing events, and those mechanisms are intertwined with repressive modes of regulation. The sequences that are signals for this—to activate or to repress—are often interdigitated, and they bind many different factors.’ The regulatory machinery, she says, seems to be modular: ‘It is intricate and highly flexible. We see differences in regulation from cell type to cell type, from one brain region to other brain regions, during development and at other times.’

Grabowski has observed new interest in alternative splicing and its regulation especially among researchers studying neuromuscular diseases like amyotrophic lateral sclerosis and neurodegenerative diseases like dementia. ‘In these diseases there’s not a mistake in splicing per se, but the alternatively spliced isoforms are altered in their ratio. That strongly suggests that there’s misregulation’, she notes.

The question of balance between two forms of a protein appears to have wide implications. Most cases of Wilms’ tumor (a childhood cancer of the kidney), for instance, are caused by a mutation of WTI, a tumor suppressor gene essential for development of testis, ovary, heart and other organs. But there are Wilms’ tumor

The limits of clinical genomics

Veronica van Heyningen’s work with Pax-6 has made her an eloquent advocate for melding clinical investigations with laboratory research if we hope to lay bare our genetic complexity. ‘We can learn an awful lot about how the genome works just by observing variation in humans, including everything from moderate to severe disease. Some of it is not even disease, but just variation,’ she notes.

Genetics researchers have long advocated intensive study of single nucleotide polymorphisms (SNPs) to explain human variation. SNPs are, of course, important for the study of human disease, van Heyningen concedes. But her discovery that a control region 150 kb away from the Pax-6 gene can determine whether someone is born with an iris or not reminds us of the very elastic definition of a gene.

van Heyningen is convinced that only a multi-pronged research approach using several methodologies will do. ‘The idea that control can be so long-range is something that many people who are setting out to work with SNPs are not taking into account. It might be very difficult to say which change in a nucleotide is the functionally important one, because it could be a long way away from the gene whose function it’s affecting,’ she says. ‘Which is why we have to take many different directions to try to resolve what we mean by a gene and its whole region of influence—or the region that influences its expression.’

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patients whose only problem appears to be an alteration in the balance of two alternatively spliced WT1 transcripts, according to Veronica van Heyningen of the MRC Human Genetics Unit at Western General Hospital in Edinburgh. WT1 is also an example of how alternative splicing can explain the increasing complexity of organisms. According to van Heyningen’s colleague Nick Hastie, human WT1 codes for at least 24 different proteins. The zebrafish version of the gene makes only two, so evolution has been able to create complexity by figuring out ways to drive WT1 into diversifying.

**Evolution has been able to create complexity by figuring out ways to drive genes into diversifying**

It is not clear, however, that a gene must create a great many protein isoforms in order to do a great many things. van Heyningen works mostly on Pax-6, which, among other things, regulates eye development in species ranging from fruit flies to humans. When one copy of the gene is missing, the result is an eye disorder called aniridia—complete absence of the iris. So far, the Pax-6 gene is known to make only two protein isoforms, although there may be other forms too. Here, too, the ratio between isoforms may be significant. Alternative splicing adds or removes an extra 14 amino acids; van Heyningen and her colleagues are attempting to make mice with either one or two copies of the gene that include the extra 14 amino acids. They want to look separately at the effects of each form of the protein, but also at the effects of upsetting the balance of the two forms.

But there is more to Pax-6 than simple alternative splicing. Other levels of control regulate its temporal and spatial expression. Different enhancers seem to act in different tissues and probably bind a variety of regulatory proteins. In the eye, the gene is expressed in various cell types at different times during development, and then continues to be expressed in the adult retina, lens cells and cornea. The retina is brain tissue, so it is not surprising that Pax-6 is expressed in brain development as well: in the cortex, the thalamus and the cerebellum, among other places.

Researchers at The Salk Institute for Biological Studies in California and the Max-Planck-Institute for Biophysical Chemistry in Göttingen, Germany, reported in the 14 April issue of Science that Pax-6 helps to organize the mammalian neocortex into functionally specialized areas for sensory processing and motor control. To complicate the picture, Pax-6 confers this identity on cortical cells in co-operation with another gene, Emx-2, and the two appear to be regulated entirely separately.

Grabowski thinks that there is a lesson here for devotees of the currently fashionable *in silico* approach to genome investigation, and cautions that algorithms alone will not tell us everything we want to know about what genes do. ‘We’ll have to develop new methods and use some old ones also,’ she says. ‘The *in silico* approach by itself is limited in asking what are the natural gene functions and how are genes regulated.’ Grabowski is looking forward to the emerging field of proteomics: ‘People are really working hard to develop ways of taking whole batches of transcripts from cells or tissues and asking how one batch is different from another. Once you know that there are 3000 genes that are upregulated in a certain cancer relative to a non-cancerous tissue, then you can ask questions about the genes.’

The complexity of genes like WT1 and Pax-6—and the unknown number of their fellow genes whose actions are likely to be just as intricate—induces a bit of humility. No wonder that, in contrast to much of the media, Human Genome Project officials have been so cautious in forecasting how quickly medical applications will be forthcoming from knowledge of the human genome. At the press conference following the announcement of a working draft of the genome, someone asked Celera’s Craig Venter when we would know all about the human genome. Venter said he thought it would take most of this century. He may have been optimistic.

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