Complementary DNA Sequencing: Expressed Sequence Tags and Human Genome Project

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Automated partial DNA sequencing was conducted on more than 600 randomly selected human brain complementary DNA (cDNA) clones to generate expressed sequence tags (ESTs). ESTs have applications in the discovery of new human genes, mapping of the human genome, and identification of coding regions in genomic sequences. Of the sequences generated, 337 represent new genes, including 48 with significant similarity to genes from other organisms, such as a yeast RNA polymerase II subunit; Drosophila kinesin, Notch, and Enhancer of split; and a murine tyrosine kinase receptor. Forty-six ESTs were mapped to chromosomes after amplification by the polymerase chain reaction. This fast approach to cDNA characterization will facilitate the tagging of most human genes in a few years at a fraction of the cost of complete genomic sequencing, provide new genetic markers, and serve as a resource in diverse biological research fields.

The human genome is estimated to consist of 50,000 to 100,000 genes, up to 30,000 of which may be expressed in the brain (1). However, GenBank lists the sequence of only a few thousand human genes and <200 human brain messenger RNAs (mRNAs) (2). Once dedicated human chromosome sequencing begins in 5 years, it is expected that 12 to 15 years will be required to complete the sequence of the genome (3). It is therefore likely that the majority of human genes will remain unknown for at least the next decade. The merits of sequencing cDNA, reverse transcribed from mRNA, as a part of the human genome project have been vigorously debated since the idea of determining the complete nucleotide sequence of humans first surfaced. Proponents of cDNA sequencing have argued that because the coding sequences of genes represent the vast majority of the information content of the genome, but only 3% of the DNA, cDNA sequencing should take precedence over genomic sequencing (4). Proponents of genomic sequencing have argued the difficulty of finding every mRNA expressed in all tissues, cell types, and developmental stages and have pointed out that much valuable information from intronic and intergenic regions, including control and regulatory sequences, will be missed by cDNA sequencing (5). However, many genome enthusiasts have incorrectly stated that gene coding regions, and therefore mRNA sequences, are readily predictable from genomic sequences and have concluded that there is no need for large-scale cDNA sequencing. In fact, prediction of transcribed regions of human genomic sequence is currently feasible only for relatively large exons (6).

On the basis of our high output with automated DNA sequence analysis of 96 templates per day and consideration of the above issues, we initiated a pilot project to test the use of partial cDNA sequences (ESTs) in a comprehensive survey of expressed genes. Sequence-tagged sites (STSs) are becoming standard markers for the physical mapping of the human genome (7). These short sequences from physically mapped clones represent uniquely identified map positions. ESTs can serve the same purpose as the random genomic DNA STTs and provide the additional feature of pointing directly to an expressed gene. An EST is simply a segment of a sequence from a cDNA clone that corresponds to an mRNA. ESTs longer than 150 bp were found to be the most useful for similarity searches and mapping.
Libraries of Complementary DNAs

Of the estimated 30,000 genes expressed in the human brain, as many as 20,000 may encode low-abundance, brain-specific transcripts (1). The fact that up to one-fourth of all genetic diseases affect neurological functions is an indication of the diversity and importance of genes expressed in the brain (8).

An assumption in our choice of cDNA libraries was that randomly primed and partial cDNA clones would be more informative in identifying genes and constructing a useful EST database than sequencing from the ends of full-length cDNAs (which contain 5' and 3' untranslated sequences) would be. By obtaining coding sequences, we hoped to take advantage of more sensitive peptide comparisons, in addition to nucleotide sequence comparisons. To discover the inherent limitations to be overcome in a large-scale cDNA sequencing project, we wanted to examine the diversity of representative cDNA libraries, identify desirable and undesirable characteristics of the libraries, and determine the information content and accuracy of single-run sequencing from both coding and flanking regions. Single-run sequencing involves performing a single sequence reaction, rather than relying on multiple, redundant reactions from each strand. We chose three commercial human brain cDNA libraries made from mRNA isolated from the hippocampus and temporal cortex of a 2-year-old female and from a fetal brain (9).

Single-run DNA sequence data were obtained from 609 randomly chosen cDNA clones (Table 1). Double-stranded cDNA clones in the pBluescript vector (Stratagene) were sequenced by a cycle sequencing protocol (10) with dye-labeled primers and 373A DNA Sequencers (Applied Biosystems). The average length of usable sequence was 397 bases with a standard deviation of 99 bases.

Subtractive hybridization has been used by researchers to reduce the population of highly represented sequences in a cDNA library (11, 12) by selectively removing sequences shared by another library. We tested subtractive hybridization as a way of enhancing the number of brain-specific clones in the hippocampus library by hybridizing the hippocampus cDNA library with a WI38 human lung fibroblast cell line cDNA library and removing the common sequences (Table 1) (12, 13).

### EST Characterization

Initially, EST sequences were examined for similarities in the GenBank nucleic acid database (14). ESTs without exact GenBank matches to human genes. Matches of at least 97% were considered to indicate that the EST corresponds directly to the human gene. Number, GenBank accession number of the matched sequence. Map positions are from (8), except where indicated. LDL, low-density lipoprotein; EST names in GenBank are the three-digit number given here preceded by “EST00.”

<table>
<thead>
<tr>
<th>EST</th>
<th>Identification</th>
<th>Number</th>
<th>Map location</th>
</tr>
</thead>
<tbody>
<tr>
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<td>M10277</td>
<td></td>
</tr>
<tr>
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<td>β-Actin (cytoplasmic)</td>
<td>M10277</td>
<td></td>
</tr>
<tr>
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<td>M10277</td>
<td></td>
</tr>
<tr>
<td>264</td>
<td>T-Actin (nucleus)</td>
<td>M24241</td>
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<td>265</td>
<td>T-Actin (nucleus)</td>
<td>M24241</td>
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</tbody>
</table>

Table 1. Composition of cDNA libraries determined by random clone sequencing. Each a ZAP library (Stratagene) was converted en masse to pBluescript plasmids, transduced into Escherichia coli XL1-Blue (Stratagene) cells, and plated on plates with X-gal, isopropyl-β-D-thiogalactoside, and ampicillin. A total of 1652 clones were picked at random from three human brain cDNA libraries: fetal brain, 2-year-old hippocampus, and 2-year-old temporal cortex (9). Clones selected from the hippocampus library after subtraction with the fibroblast library are listed in the "Subtracted" column. Templates for DNA sequencing were PCR products or plasmids prepared by the alkaline lysis method. About half of the templates prepared by PCR failed to yield an amplified fragment suitable for sequencing. This was primarily due to use of PCR conditions that minimized the need for further purification of the product but selected against amplification of long inserts (5 I~I of each deoxynucleotide triphosphate, and 0.1 μM each primer for 35 cycles: 94°C, 40 s; 55°C, 40 s; 72°C, 90 s). A further percentage of the PCR-generated templates failed to sequence, largely because of primer-dimer or other amplification artifacts. Qiangen columns (Studio City, California) improved the percentage of plasmid templates that yielded usable sequences from about 60% with a standard alkaline lysis protocol to over 90%. Overall, 117 PCR-generated templates and 497 plasmid templates gave usable sequences. Deoxy chain-termination sequencing reactions were performed with fluorescent dye-labeled M13 universal or reverse primers (Applied Biosystems). After a cycle sequencing protocol (10), carried out in a Perkin-Elmer Thermal Cycler, sequencing reactions were run on a 373A automated DNA sequencer (Applied Biosystems). Some sequencing reactions were performed on an Applied Biosystems robotic workstation (28). For each column, numbers are indicated followed by percents in parentheses.
matches were translated in all six reading frames, and each translation was compared with the protein sequence database Protein Information Resource (PIR) and the ProSite protein motif database.

**Table 3.** EST similarities in the GenBank and PIR databases. All significant similarities (P < 0.01) with GenBank or PIR entries are listed. Matches indicate percent identical bases for nucleotide sequence similarity and percent identity (identical plus conservative substitutions) for peptides. Number indicates the accession number or locus name of the matched sequence. Abbreviations used are as follows: B, bovine; BM, Brevibacterium maltose; BMDV, bovine mucosal disease virus; C, chicken; CE, Caenorhabditis elegans; D, Drosophila melanogaster; E, E. coli; H, human; I, lamprey; M, mouse; N, Neospora crassa; P, pig; PP, Pseudomonas putida; PRV, Pseudorabies virus; R, rat; S, squid; T, Torpedo californica; TN, transposon Tr 4556; X, Xenopus laevis; Y, yeast; UT, untranslated; MARCS, myristoylated alanine-rich C kinase substrate; HPRT, hypoxanthine-guanine phosphoribosyltransferase; GTP, guanosine triphosphate; LAMP, lysosomal-associated membrane protein; RNA, transfer RNA; snRNP, small nuclear ribonucleoprotein; IGF, insulin growth factor; Mito, mitochondrial; DBP, albumin promoter D site-binding protein; and Pol, polymerase. EST names in GenBank are the three-digit number given here preceded by "EST00." (14). Comparisons with the ProSite motif database were done by means of the program MacPattern from the EMBL Data Library (14a). GenBank and PIR searches were conducted with our modifications of the "basic local alignment search tool" programs for nucleotide (BLASTN) and peptide (BLASTX) comparisons (15). These modifications permit many query sequences to be automatically searched in a sequential fashion. PIR searches were run on the National Center for Biotechnology Information Blast network service. The BLAST programs contain a rapid database-searching algorithm that searches for local areas of similarity between two sequences and then extends the alignments on the basis of defined match and mismatch criteria. The algorithm does not consider the potential of gaps to improve the alignment, thus sacrificing some sensitivity for 60- to 80-fold increase in speed over other database-searching programs such as FASTA (16).

Sequence similarities identified by the BLAST programs were considered statistically significant with a Poisson P-value < 0.01. The Poisson P-value is the probability of as high a score occurring by chance, given the number of residues in the query sequence and the database. After the BLAST search, 30 unmatched ESTs were compared against GenBank by FASTA to determine if significant matches were missed because of the use of BLASTN for the database search. No additional statistically significant matches were found. Statistical significance does not necessarily mean functional similarity; some of the matches reported here may indicate the presence of a conserved domain or motif or simply a common protein structure pattern. Statistically significant matches to GenBank and PIR are reported in Tables 2 and 3. The length and percent identity or similarity of each alignment is given in Table 3 to aid in evaluation of match quality.

On the basis of database searches, the 609 EST sequences were classified into eight groups as shown in Table 1. Four groups, with 197 of the sequences (32% of the total), consist of matches to human sequences: repetitive elements, mitochondrial genes, ribosomal RNA genes, and other nuclear genes. Forty-eight of the sequences (8%) matched nonhuman entries in GenBank or PIR, whereas 230 (38%) had no significant matches. The remaining 134 (22%) sequences contained no insert between the Eco R1 cloning sites or consisted entirely of polyadenylate.

**Table 4.** Matches to the ProSite motif database. Pattern matches from the ProSite database (except posttranslational modification sites) are shown. Abbreviations used are as follows: AA, amino acyl; HIGH, motif consensus in single-letter amino acid code (30); ILGF, insulin-like growth factor; DHRF, dihydrofolate reductase; EGFR, epidermal growth factor; Gal-P-UDP, galactose-1-phosphate-uridyl; C2H2, two Cys and two His residues.
Table 5. Accuracy of single-run double-stranded automated sequencing. ESTs listed in Table 2 and those matching mitochondrial and ribosomal genes were aligned with sequences from GenBank with the GCG program BESTFIT. The first 85 nucleotides were the polylinker sequence that was not aligned with the pBluescript SK reference sequence. Tabulation of errors began 15 bases into the BESTFIT alignment and thus is reported beginning with bases 101 to 200.

<table>
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<th>Bases from primer</th>
<th>Mismatches-ambiguities*</th>
<th>Gaps*</th>
<th>Accuracy %</th>
<th>Aligned bases</th>
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<td></td>
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<td>Deletions</td>
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</tr>
<tr>
<td>&gt;400</td>
<td>3.53</td>
<td>2.63</td>
<td>92.8</td>
<td>3197</td>
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</table>

*Error rates are reported as number of mismatches, insertions, or deletions per hundred aligned bases. "Mismatches" includes ambiguous base calls.

Thirty-six ESTs matched previously sequenced human nuclear genes with more than 97% identity (Table 2). Four of these ESTs were from genes encoding enzymes involved in maintaining metabolic energy, including ADP/ATP (adenosine diphosphate/adenosine triphosphate) translocase, aldolase C, hexokinase, and phosphoglycerate kinase. Human homologs of genes for the bovine mitochondrial ATP synthase F$_0$F$_1$ subunit and porcine aconitase were also found (Table 3). Brain-specific cDNAs included synapto-physin, glial fibrillary acidic protein (GFAP), and neurofilament light chain. At least six ESTs were from genes encoding proteins involved in signal transduction: 2',3'-cyclic nucleotide 3'-phosphodiesterase (GNPase) (two ESTs), calmodulin, c-erbA-α-2, G stimulating protein (G$_S$) α subunit, and Na$^+$,K$^+$-ATPase α subunit. Other ESTs were matches to genes for ubiquitous structural proteins—actins, tubulins, and fodrin (nonerythroid spectrin). Eight ESTs were from genes known to be associated with genetic disorders (8). More than half of the human-matched ESTs have been mapped to chromosomes, indicating the bias of GenBank entries toward well-studied genes and proteins.

ESTs without significant GenBank matches were also compared to the ProSite database of recognized protein motifs. Not counting posttranslational modification signatures, 54 sequences contained motifs from the database (Table 4). Some patterns are found in scores or even, as in the case of the leucine zipper, hundreds of proteins that do not share the functional property implied by the presence of the motif.

Similarities to sequences from other organisms were also detected in the BLAST searches of GenBank and PIR (Table 3). Several ESTs were similar to "housekeeping" genes, including the ribosomal proteins (RP) S10 and L30 (in the rat) and the above glycolytic enzymes. EST00257 showed strong nucleotide sequence similarity to the squid (67.4%) and Drosophila (70.4%) kinesin heavy chain. Kinesin was first described as a microtubule-associated motor protein involved in organelle transport in the squid giant axon (17). Six oncogene-related sequences were also among the cDNA clones sequenced. EST00299 and EST00283 showed similarity to several ras-related genes, and EST00248 matched the 3' untranslated region of the bovine substrate of borulin toxin ADP-ribosyltransferase. We also observed similarities with a Saccharomyces cerevisiae RNA polymerase subunit and Torpedo californica electromotor neuron-associated protein. Two ESTs may represent new members of known human gene families: EST00270 matched the three β-tubulin genes with 88 to 91% identity and EST00271 matched α-actinin with 85% identity at the nucleotide level.

Among the most interesting of the primary sequence relationships was the similarity of ESTs to the Drosophila genes Notch and Enhancer of split. Nucleotide and peptide alignments of EST00256 and EST00259 with the Drosophila genes are shown in Fig. 1. Both genes are part of a signal cascade encoded by the "neurogenic" genes that are involved in the differentiation of neuronal and epidermal cell lineages in the neuroectoderm of the developing Drosophila embryo (18). It has been proposed that the Enhancer of split protein interacts with a membrane protein that is the product of the Notch gene to convert a developmental signal into an altered pattern of gene expression (18). EST00256 matched near the 5’ end of the Enhancer of split coding sequence, away from the mammalian G protein β subunit and yeast cyd4-like elements (19). Part of the EST00259 match to Notch is in the cd10/SW16 region that is similar to three
cell-cycle control genes in yeast and is tightly conserved in the *Xenopus laevis* Notch homolog, *Notch*. In Drosophila, *Enhancer of split* is required for formation of epidermal tissue. *Notch* contains several epidermal growth factor–like repeats and appears to be involved in cell-cell communication during development (20).

Seven genes were represented by more than one EST. Comparisons of all the ESTs against one another revealed two overlaps of unknown ESTs: EST00223 and EST00234 matched in opposite orientations, and EST00235 and EST00236 matched in the same orientation beginning at the same nucleotide. Five human genes were represented by more than one EST: β-actin (three), γ-actin (two), α-tubulin (two), α-2-macroglobulin (two), and CNPase (two).

Mapping of ESTs to Human Chromosomes

We used the polymerase chain reaction (PCR) to screen a series of somatic cell hybrid cell lines containing defined sets of human chromosomes for the presence of a given EST (21). In this process, only the hybrids that contain the human gene corresponding to the EST will yield an amplified fragment. An EST is assigned to a chromosome by analysis of the segregation pattern of PCR products from hybrid DNA templates. The single human chromosome present in all hybrids that give rise to an amplified fragment is the location of the EST.

PCR mapping has been applied to 46 clones, as summarized in Table 2. The EST of the human gene for apolipoprotein J (also called SF-40, 40 complement-associated protein and sulfated glycoprotein 2) was localized to chromosome 8. Eleven other ESTs with GenBank or PIR similarities were mapped to chromosomes. Although PCR mapping of somatic cell hybrids is relatively rapid—up to three clones can be assigned per day with a single thermal cycler—it is relatively expensive, costing about ten times as much as EST sequencing. With the same oligonucleotide primers, sublocalization can be achieved with panels of fragments from specific chromosomes or pools of large genomic clones in an analogous manner. Other mapping strategies that have been proposed are multiplex in situ hybridization, prescreening with labeled flow-sorted chromosomes, and preselection by hybridization to construct chromosome-specific cDNA libraries. However, these methods are limited by the purity of the chromosome-specific material or the specific activity necessary for detection.

Automated DNA Sequencing Accuracy and GenBank Submission

ESTs that match human sequences in GenBank are excellent tools for the analysis of the accuracy of double-strand automated DNA sequencing. Ninety EST-GenBank matches were examined for the number of nucleotide mismatches and gaps required to achieve optimal alignment by the Genetics Computer Group (GCG) program BESTFIT (22). The number of mismatches, insertions, and deletions was counted for each hundred bases of the sequence (Table S). As expected, the sequence quality was best closest to the primer and decreased rapidly after about 400 bases. The number of deletions and insertions relative to the GenBank reference sequence increased five- to tenfold beyond 400 bases, whereas the number of mismatches doubled. The average accuracy rate for individual double-stranded sequencing runs was 97.7% for up to 400 bases. The minimum criteria for submission of ESTs to GenBank were that sequences be at least 150 bases in length and contain <3% ambiguous base calls. The overall accuracy of sequences submitted from each template group was at least 97%, based on matches to known human genes. Three hundred forty-eight ESTs met these criteria and were submitted to GenBank with accession numbers M61953 through M62300, inclusive. All ESTs except those matching mitochondrial or ribosomal RNA (rRNA) genes and simple repetitive elements were submitted to GenBank.

Conclusions and Prospects

Single-run DNA sequencing has proven to be an efficient method of obtaining preliminary data on cDNA clones. Our results demonstrate that sufficient information is contained in 150 to 400 bases of a nucleotide sequence from one sequencing run for preliminary identification of the cDNA and localization to a chromosome. In addition to the 35 EST's homologous to known human genes, 48 ESTs matched sequences in GenBank or PIR with moderate to striking similarity, including high-quality matches with genes from such evolutionarily distant organisms as yeast (EST00374) and Neurospora (EST00287) (Table 3).

Two hundred thirty ESTs did not match any current database entries and therefore represent new, previously uncharacterized genes. A multitude of approaches for classifying these genes exists, including complete sequencing and expression, chromosome mapping, tissue distribution, and immunological characterization. Currently unidentified cDNAs will also be classified by similarity to genes from other organisms as those sequences become available. Three ESTs reported here (EST00257, EST00259, and EST0374) were identified by similarity to sequences that have appeared since the last full release of GenBank.

The random selection approach used here revealed an unacceptably large number of highly represented clones in these cDNA libraries. Over 30% of the clones from the hippocampus cDNA library consisted of rRNA, mitochondrial cDNAs, or inserts consisting entirely of polyA. Sixty-eight ESTs matched 12 different mitochondrial genes, including 18 matches to cytochrome oxidase I. Although elimination of these uninformative clones is a priority for developing ideal cDNA libraries, techniques to reduce repeated sequencing of clones will become increasingly important as large numbers of cDNAs are sequenced. The use of library preprocessing techniques such as subtraction, which preferentially reduces the
population of certain sequences in the library (11, 12), and normalization, which results in all sequences being represented in approximately equal proportions in the library (23), should reduce repeated sequencing of high and intermediate abundance clones and maximize the chances of finding rare messages from specific cell populations. In our initial experiments with subtractive hybridization of the hippocampus library with a human fibroblast cDNA library, CNPase and GFAP clones were enriched greater than tenfold and twofold, respectively. Another characteristic of the ideal cDNA library would be directional cloning so that either a coding sequence or a 3' noncoding sequence could be selectively obtained.

The EST data, in conjunction with physical mapping, will provide a high resolution map of the location of genes along chromosomes, a map that would be more costly to construct by genomic sequencing and analysis. By performing a single DNA sequencing reaction on each cDNA clone, a key piece of information was obtained for the relatively low cost of about $0.12 to $0.15 per base. The EST approach will provide a new resource for the analysis of chromosome sequence and for human gene discovery.

The screening of cDNA clones to identify the protein complement of a tissue has been explored by others to a limited extent. In 1983, Putney and co-workers sequenced over 150 clones from a rabbit muscle cDNA library and identified clones for 13 of the 19 known muscle proteins, including one new isotype, but no unknown coding sequences (24). Over 400 adult head-specific cDNA clones from Drosophila have been identified by differential screening of cDNA libraries from different developmental stages (25). Improvements in DNA sequencing technologies have now made feasible essentially complete screening of the expressed gene complement of an organism.

In our own laboratory, the EST approach should result in the partial sequencing of most human brain cDNAs in a few years. Similar approaches begun elsewhere (26) could result in a database of most human expressed genes in less than 5 years. The presence of these minimally characterized sequences in GenBank will assist research efforts in several areas of biology. The EST database will provide identification and confirmation of coding regions in naive genomic sequences. Sublocalization of cDNAs that have been mapped to chromosomes will help define the genetic content of specific chromosomal regions and permit correlation with patterns of inheritance in genetic disease. In a related experiment, chromosome sublocalization was the key to establishing that the γ-aminobutyric acid–benzodiazepine receptor β3 subunit is deleted in individuals with Angelman–Prader–Willi syndrome (27). We anticipate that ESTs from human brain will further the identification of genes associated with other neurological diseases and will provide a more complete view of gene expression in the brain.

REFERENCES AND NOTES

8. V. A. McKusick, Online Mendelian Inheritance in Man, from the Welch Medical Library, Johns Hopkins University School of Medicine, Baltimore, MD.
9. Stratagene catalog numbers 936206, 936205, and 935205, respectively.
10. Cycle sequencing was performed in a Perkin-Elmer Thermal Cycler for 15 cycles of 95°C, 30 s; 60°C, 1 s; 70°C, 60 s; and 15 cycles of 95°C, 30 s; 70°C, 60 s with the Taq Dye Primer Cycle Sequencing Core Kit protocol (Applied Biosystems).
14. GenBank Release 65.0 and collective updates through 21 January 1990, PIR Release 26.0, and ProSite (EMBL protein motif database) Release 5.0 were used.
22. We designed oligonucleotide primer pairs from EST sequences to minimize the chance of amplifying through an intron. Introns are less likely to be present in 3' untranslated sequences, so ESTs were examined for the presence of stop codons in each reading frame and for consensus splice junctions (J. D. Hawkins, Nucleic Acids Res. 16, 9893 (1988); M. B. Shapiro and P. Senapathy, ibid. 15, 7159 (1987)). Also desirable for mapping are 3' untranslated sequences because they are less likely to be identical in other members of a gene family or in pseudogenes. We used the primers in a PCR to amplify from total human genomic DNA templates as described in the legend to Fig. 2. If the resulting product was of the expected size, then the PCR reaction was repeated with DNA templates from two panels of human-rodent somatic cell hybrids: Bios PCRable DNA (BIOS Corporation) and NIGMS Human-Rodent Somatic Cell Hybrid Mapping Panel Number 1 (NIGMS, Camden, NJ).
31. Abbreviations for the amino acid residues are as follows: A, Ala; C, Cys; D, Asp; E, Glu; F, Phe; G, Gly; H, His; I, Ile; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; T, Thr; V, Val; W, Trp; and Y, Tyr.
33. Dedicated to the memory of J. E. Venter, 18 August 1923 to 10 June 1982.