Hunting with Traps: Genome-Wide Strategies for Gene Discovery and Functional Analysis

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With sequence analysis of the human genome well underway, there is an increasingly urgent challenge to understand the fundamental function and interplay of genes that build and maintain an organism. Several approaches will be critical for interpreting gene function, including random cDNA sequencing, expression profiling in different tissues, genetic analysis of human or model organism phenotypes, and creation of transgenic or “knockout” animals. Traditional gene-trapping approaches, in which genes are randomly disrupted with DNA elements inserted throughout the genome, have been used to generate large numbers of mutant organisms for genetic analysis. Recent modifications of gene-trapping methods and their increased use in mammalian systems are likely to result in a wealth of new information on gene function. Various trapping strategies allow genes to be segregated based on criteria like the specific subcellular location of an encoded protein, the tissue expression profile, or responsiveness to specific stimuli. Genome-wide gene-trapping strategies, which integrate gene discovery and expression profiling, can be applied in a massively parallel format to produce living assays for drug discovery.

Classical Genetics, Genomics, and Analysis of Gene Function

The term “genomics” was proposed in 1986 by Thomas Roderick (McKusick 1997a) to describe the study of a genome by molecular means distinct from traditional genetic approaches. Genomics evolved from a much older word, genome (McKusick 1997a). A fusion of gene and chromosome, a genome is the complete collection of genes possessed by an organism. Living creatures result from the delicate interplay between a “functional” genome and environmental factors. With complete sequence of many genomes, including biological workhorses like Escherichia coli (Blattner et al. 1997), Saccharomyces cerevisiae (Mewes et al. 1997), and Caenorhabditis elegans (The C. elegans Sequencing Consortium 1998), and with the Human Genome Project “ahead of schedule and under budget” (Collins 1995; Collins et al. 1998), the scientific focus is shifting towards development of methods that effectively use this wealth of structural genomic information (Collins et al. 1998).

The ultimate goal of the Human Genome Project is to understand how the genome builds, maintains, and operates an organism. Multiple strategies at the DNA, RNA, protein, cellular, and organism level will be key in achieving this goal. Hybridization-based techniques, like oligonucleotide chips (Fodor et al. 1991) or cDNA arrays (Schena et al. 1995), can reveal the differential expression profiles of numerous genes in response to a stimulus (DeRisi et al. 1997; Wodicka et al. 1997). Gene expression profiles can also be elucidated through random sequencing of cDNA libraries derived from specific tissues or cell types (Okubo et al. 1992; Fannon 1996; Okubo and Matsubara 1997). Analysis of these expressed sequence tags (ESTs) can provide an “in-silico” gene expression landscape. Shifting towards the gene products, large two-dimensional polyacrylamide gels can be used to monitor the expression of at least a subset of proteins (Geisow 1998; Persidis 1998). In addition, various methods are available for elucidating physical interactions between proteins in live cells (Fields and Song 1989; Miyawaki et al. 1997).

Classical and molecular genetics provide a variety of powerful tools useful for understanding gene function and studying complex developmental events. Many structure–function relationships have been elucidated using elegant genetic approaches and by cloning disease associated genes. However, these strategies are too time consuming for efficient genome-wide investigation. This review will focus on the use of random gene-trapping techniques to gain insights into gene function. These methods are producing valuable raw material for current and future bioinformatics databases that will catalog many biological processes. Ultimately, this information should help researchers predict gene function in much the same way sequence databases are currently used in structural genomics efforts.

Gene Capturing and Mutagenesis

Conceptually, one straightforward way to discern the function of a gene is to observe the effects of an impaired or mutated gene on an organism over a desired number of generations. Many gene mutations have consequently been implicated in a variety of human
medical conditions. In addition, chromosomal markers identified through arduous structural genomics approaches have led to remarkable successes in positional cloning of disease related genes. The important functional insights gained through time-consuming map-based discovery of genes like the cystic fibrosis transmembrane conductance regulator (Riordan et al. 1989; Rommens et al. 1989) or BRCA-1 (Miki et al. 1994) have resulted in the mapping of many phenotypic variations to a specific gene. Much of the genetic basis of disease work has culminated in the Online Mendelian Inheritance in Man (OMIM) database provided on the National Center for Biotechnology Information (NCBI) Web site (http://www.ncbi.nlm.nih.gov/Omim/), which currently lists ~500 human genes that have disease-producing mutations (McKusick 1997b).

Nonhuman organisms have been exploited to gather additional information about gene function. Each organism offers its own unique advantages and drawbacks. Yeast, as the simplest eukaryote, is a logical place to start when searching for basic understanding of cell biology. Studies with C. elegans, Drosophila, zebrafish, and other model systems have revealed the functions of many genes during embryonic development or complex intercellular signaling. These organisms, however, are very distantly related to humans; so mammalian systems are required to expand the knowledge base and ultimately for pharmacological evaluations. The combined use of embryonic stem (ES) cells with homologous recombination in mice has created a very useful system for functional genomic study, allowing researchers to modify or eliminate any known gene and analyze its function (Capecchi 1989; Joyner 1993).

In contrast to the rational “one gene at a time” approaches, enhancer-trapping methods evolved to probe the entire genome simultaneously. Originally described in bacteria, enhancer trapping was first demonstrated using bacteriophage transposable elements to insert a reporter gene at scattered sites throughout the E. coli genome (Casadaban and Cohen 1979; Bellofatto et al. 1984). Chromosomal integration of a transposable element tagged the integration site and often mutated the gene into which it inserted. Mutants resulting from this approach could be selected using genetic screens. Because the insertion site was tagged, the mutated genes could be readily identified, which made this approach more efficient than traditional chemical mutagenesis (Meneely and Herman 1979; Rinchik 1991). The success and conceptual simplicity of the enhancer trap method was quickly adapted for use in other model systems including plants (Schell 1987), C. elegans (Hope 1991), and Drosophila (O’Kane and Gehring 1987). When the enhancer-trapping element consists of a cDNA encoding an easily monitored reporter gene like β-galactosidase, the expression pattern of the trapped gene can be visualized. In Drosophila, the P-transposon system has been used with great success to create transformed animals carrying enhancer detectors (Rubin and Spradling 1982; O’Kane and Gehring 1987). Large numbers of enhancer trap lines have been established and evaluated using both phenotypic and expression analysis (Bier et al. 1989; Spradling et al. 1995). Specific mutant lines can then be chosen for further study when there is a correlation between expression and phenotype.

Although enhancer traps are widely used, certain gene traps were developed as an alternative to enhancer traps to capture open reading frame information. The identification of target genes using enhancer trapping was sometimes problematic because the site of reporter insertion could be as much as 100 kb from the target gene. This would require extensive characterization of the genomic insertion site to identify candidate target genes. Gene trapping varies from enhancer trapping in that, instead of using a minimal promoter, gene trap vectors provide specific sequences that generate fusion RNA transcripts when inserted into a gene (see Fig. 1). This feature makes gene trapping (vs. enhancer trapping) especially advantageous in mammalian cells that have complex genomic organization, including large introns and small exons, because the trapped gene can be identified by mRNA sequence.

It is believed that the use of random “knockout” mutagenesis by gene trapping in mice will result in further enhancements for functional analysis of the genome. Typical vectors used for knockout gene trapping contain an acceptor site for RNA splicing followed by a reporter gene and then a transcript-terminating polyadenylation sequence (Brenner et al. 1989; Gossler et al. 1989; Friedrich and Soriano 1991; Skarnes et al. 1992). These gene-trapping vectors can be introduced into the genome of murine ES cells by electroporation or replication-deficient self-inactivating retroviruses. Insertion of this vector into an intron results in premature termination of the captured allele in which the splice donor at the 3’ end of an endogenous exon is “trapped” into splicing with the splice acceptor from the gene trap vector. The result is a fusion mRNA in which the reporter gene from the vector becomes transcriptionally regulated by the trapped promoter. Expression of the reporter gene can be used to monitor the spatial and developmental transcription profile of the trapped locus (see Fig. 2).

Sequence Acquisition: Identification of Trapped Genes
Recent reports have demonstrated the feasibility of using knockout gene-trapping approaches to create libraries of murine ES cells in which each individual clone has one gene disrupted and tagged. In gene-
trapping approaches, a fusion RNA is generated so the trapped endogenous gene from each isolated clone of interest can be identified using rapid amplification of cDNA ends (RACE), a method pioneered by Michael Frohman (Frohman et al. 1988). With this approach, PCR is used to amplify the unknown 5' or 3' end of the fusion mRNA. The design of the vector dictates whether 5' and/or 3' fusions with endogenous gene transcripts are produced.

Several reports have demonstrated that a gene trap method can effectively knock out genes and that a 5' RACE protocol can be used to obtain sequence information about the disrupted genes (Skarnes et al. 1992, 1995; Forrester et al. 1996; Chowdhury et al. 1997). In the largest study reported to date using 5' RACE, 115 sequences were successfully recovered from 153 cell lines (Townley et al. 1997). Sequence information from some of these murine ES cell clones is available on the Web (http://socrates.berkeley.edu/~skarnes/), and this set currently includes >250 characterized lines. In addition, details on a large number of other academic mouse ES cell tagging efforts have also recently been reported (Chowdhury et al. 1997; Hicks et al. 1997; Couldrey et al. 1998; Voss et al. 1998).

Researchers at Lexicon Genetics (Woodlands, TX) have reported sequence information of 2000 genes in murine ES cells (Zambrowicz et al. 1998). The vector design used in this study provides an internal constitutive promoter that drives expression of a selectable marker sequence with a 3' splice donor in place of a poly(A) signal. This vector should only confer resistance to ES cells where the vector traps an endogenous splice acceptor and polyadenylation signal. The higher rate of "novel" sequence recovery reported in this study, compared with that obtained previously (Skarnes et al. 1992; Townley et al. 1997), may suggest that these vectors tag genes that are under-represented in the EST databases or that tagging occurs in genes that are not normally transcribed (pseudogenes, line elements, interspersed repeats). Because these poly(A) trap vectors do not require integration into an actively transcribed gene for selection, they can be useful for identifying genes that are inactive in undifferentiated ES cells and become actively transcribed during differentiation (Salminen et al. 1998).

Although it may be possible to generate a complete set of murine gene trap ES cell lines, that would serve only as a starting point for functional analysis of the genome. The benefits of this approach include the potential to discover novel phenotypes and create useful in vivo model systems for the study of disease. The strategy is also well suited for studying embryonic development (Friedrich and Soriano 1991; Skarnes et al. 1995).

Figure 1  Schematic overview of enhancer and gene trapping. Normally a wild-type gene produces an unspliced RNA transcript that is processed by linking splice acceptor (SA) sites to splice donor (SD) sites. Transcription initiates at the promoter (Pro) and is regulated by enhancer sequences that can be located thousands of base pairs away from the promoter. (AAAAAAA) Poly-(A)-tail. (A) An enhancer-trapping element contains a minimal promoter and a translation start site. When the element integrates near an endogenous enhancer, that enhancer controls expression of the reporter gene. (B) In the basic gene-trapping element, a reporter gene sequence is flanked by splice acceptor (SA) and polyadenylation (pA) sequences. When the trapping element integrates within an intron, a fusion transcript is generated as a result of splicing of the endogenous splice donor with the vector-provided acceptor. As with enhancer trapping, the result is expression of the reporter gene that is faithfully controlled by the regulatory elements of the trapped gene locus. (C) The downstream sequences of the trapped gene can also be captured in a fusion transcript by providing a splice donor in the trapping element.
limitation to using the gene trap approach for generating knockout mice results from the fact that a gene trap vector may “tag” a gene, reliably tracking its expression with a reporter gene, without completely disrupting it. In some cases the full-length trapped gene may still be expressed as a result of differential or inefficient splicing, or an active fragment could be expressed when the trapping element inserts in the 3’ region of a gene.

**Genome-Wide Functional Analysis in Mammalian Cells**

Molecular expression profiling by hybridization and mutational analysis via traditional or gene trap methods are providing valuable information that will help index functions of many genes. The information gained may ultimately lead to better medicines and treatments. However, if one hopes to rapidly move from the vast amount of genomic information to therapeutic intervention, additional technologies are required.

Modern drug discovery efforts require bioassays to sort through the large number of potentially bioactive compounds generated by conventional and combinatorial chemistry. Large-scale mutagenesis in mice can provide functional information about genes and generate model systems for the study of some diseases, but mice are not well suited for high-throughput screening. The hybridization DNA array technologies can observe changes in gene expression on a genome-wide basis, but they are limited to only those genes that have already been cloned. To assess the gene expression effects of numerous test compounds using DNA arrays would require RNA isolation from pools of treated cells, probe generation from this material, and then hybridization to a chip for each assay. Such an approach is not currently cost effective for large-scale screening. The need for an integrated technology platform to discover genes, pathways, and corresponding lead drug candidates drove development of novel approaches, termed “gene-to-screen” genomics. Gene-“tagging” strategies...
have been described that may provide a link between trapping methods and drug discovery (Whitney et al. 1998).

The first step in this gene-to-screen approach is creation of a reporter gene-tagged library in a particular cell line. A library consists of millions of cells, each carrying the same tagging element integrated into a different site within the genome. The gene-tagging element typically contains a splice acceptor and reporter gene. As with knockout approaches in ES cells, the gene-tagging element can be introduced into the genome using physical methods such as transfection and electroporation (Skarnes et al. 1992; Friedrich and Soriano 1991; Gogos et al. 1997; Hicks et al. 1997; Zambrowicz et al. 1998). The result is reporter gene expression in those cells in which the tagging element has inserted into an actively transcribed gene.

A new and highly sensitive reporter system has recently been described and is well suited for gene tagging. A nontoxic fluorescent substrate of β-lactamase, CCF2-AM, enables real-time and sensitive monitoring of transcription in live cells (Zlokarnik et al. 1998). A β-lactamase tagged library can be used to clone genes with a variety of interesting expression characteristics (Whitney et al. 1998). Any β-lactamase tagged cell library can be subjected to fluorescent activated cell sorting to rapidly separate clones where the tagged gene is constitutively expressed. In addition, sequential rounds of sorting can be used to identify cell clones whose genes are induced or repressed by different agents including receptor ligands, drug candidates, and viruses (Rao 1998) (See Fig. 3).

The Future of the Trapping Trade
Gene-trapping methods are already proving to be indispensable for functional analysis of the genome. Versatility is one strength of this family of molecular tools, which is helpful when attempting to analyze complex genomes. With this goal in mind, gene-trapping schemes that can act as a filter and allow discovery of a functionally related set of genes are useful. One example of a fairly “broad-pass” filter would be a technique to trap only genes that encode secreted or membrane proteins (Skarnes et al. 1995). Such a “secretory-trap” approach can ultimately reveal the proteins important for communication between cells. Another strategy that has been used in mice is the “induction-trap” in which ES cells are selected based on responsiveness of the trapped gene to growth and differentiation stimuli like retinoic acid or nerve growth factor (Forrester et al. 1996; Bonaldo et al. 1998; Salminen et al. 1998). Techniques that can specifically trap and monitor genes expressed at low levels will be valuable for “genome closure” because these genes are currently under-represented in compiled EST databases. Reporter cell lines generated by some trapping methods will be useful tools for investigation of the signal transduction pathways that regulate expression of the trapped genes. This will ultimately provide another layer of functional genomic information.
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REFERENCES


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