DNA Mapping in Growth and Developmental Disorders

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Abstract. DNA mapping techniques are being increasingly applied to familial and acquired disorders affecting growth and development. To understand the potential applications of these techniques, one must first have a good understanding of the components and basic structure of DNA, the genetic code, the basic structure of genes and the roles that various components of gene structure play in regulating gene expression. Following review of this basic information, a variety of DNA mapping techniques including in situ hybridization, Southern blotting, polymerase chain reaction amplification, DNA sequencing and linkage analysis will be covered. Applications of these techniques can enable mapping of unknown genes by detection of loss of allelic heterozygosity or use of linkage analysis and genetic maps. With increasing ease, mutations such as deletions, expansions, rearrangements and point mutations can be detected in diseases such as congenital adrenal hyperplasia, cystic fibrosis, diabetes insipidus, growth hormone deficiency, fragile X syndrome, Laron dwarfism and Turner syndrome. From this discussion, a better understanding of methods of gene localization, uses of genetic maps and rapid, convenient methods to detect a variety of molecular derangements causing familial disorders affecting growth and development will be gained.

Key words: DNA analysis, gene mapping, linkage and PCR analysis

Introduction

This material will review DNA structure and a variety of techniques that are used in mapping genes that are important in growth and development. Examples of applications will include use of PCR amplification, Southern blotting, linkage analysis and DNA sequencing to map and detect derangements of the arginine vasopressin, growth hormone, growth hormone receptor, insulin, insulin receptor, Pit-1 and vitamin D receptor genes. From these examples the reader should better understand the molecular derangements causing some forms of familial diabetes insipidus, growth hormone deficiency, diabetes mellitus, Laron dwarfism, panhypopituitary dwarfism and vitamin D resistant rickets in humans.
Biochemistry of DNA

Components of DNA: DNA is composed of nucleotides which are made up of bases and deoxyribose sugar molecules. The purine and pyrimidine bases are shown in the table below.

<table>
<thead>
<tr>
<th>Purines</th>
<th>Adenine (A)</th>
<th>Guanine (G)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyrimidines</td>
<td>Cytosine (C)</td>
<td>Thymine (T)</td>
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The sugar component of DNA is a pentose (2-deoxyribose) that is connected at position 1 to one of the nitrogenous bases (A, C, G or T) (see Fig. 1). Also, note that the addition of a hydroxy1 group to carbon two of 2-deoxyribose converts it to ribose which is the sugar that is found in RNA. This sugar hydroxylation accounts for the alkali sensitivity of RNA as opposed to the resistance of DNA.

Structure of the DNA Molecule: The DNA molecule consists of nucleotides (made up of a base and a 2-deoxyribose sugar) that are polymerized in a series of 5'-3' sugar phosphate links (see Fig. 1). This sugar phosphate backbone is composed 5'-3' of 1) a phosphate molecule, 2) the 5' carbon of the deoxyribose molecule, 3) the 3' deoxyribose carbon which attaches to 4) the next phosphate molecule. Note that the bases are attached to carbon 1 of 2-deoxyribose and project away from the sugar-phosphate backbone.

Hydrogen bonding between A·T and G·C bases on complementary DNA strands stabilizes formation of the double helix. Note that A=T pairing forms two hydrogen bonds while G≡C pairing forms three hydrogen bonds. Also note that the hydrogen bonding occurs between the bases on complementary strands that run in anti-parallel directions i.e. 5'→3' on one strand and 3'→5' on the complementary strand. Finally, as the temperature is increased denaturation of the two DNA strands occurs first between A=T pairs then between G≡C pairs because of the higher stability of the latter. Both types of base pairs melt at higher temperatures than mispaired (A·C and G·T) sequences. These differences in melting temperature are exploited by a variety of methods in the detection of various types of mutations.

Methods of Mapping

Molecular tools used to map and examine endocrine disorders include restriction endonuclease analysis, the polymerase chain reaction and DNA sequence analysis.

A. Restriction Endonuclease Analysis

Restriction endonuclease analysis can be used to detect gene deletions, certain point mutations, and in linkage analysis to determine if a gene and a genetic endocrine disorder are transmitted together. Restriction endonucleases are bacterial enzymes that recognize and cleave double-stranded DNA at specific recognition sequences. This property...
enables them to cut the long strands of cellular DNA into fragments of reproducible size. If the restriction endonuclease is one like EcoRI, it will cut DNA leaving the ends of the two strands offset from each other (1). This overhang of one DNA strand over the other produces "sticky ends". Such "sticky ends" promote pairing with fragments also cut with EcoRI to form recombinant molecules. Other endonucleases, such as HaeIII, cut the ends off equally producing so called "blunt ends".

1. Chromosome Specific Probes

An example of the use of restriction endonuclease analysis to detect the presence or absence of an extra chromosome is Turner syndrome. Usually the karyotype shows a 45, X pattern but some subjects have additional chromosomal fragments. The fragments represent, in some cases, an X chromosome with a deletion or alternatively a portion of a Y chromosome. Distinction between these two is important because the presence of Y chromosomal material predisposes to gonadoblastomas.

Fig. 2 illustrates the detection of Y chromosomal fragments by restriction analysis. The solid circle represents a Turner syndrome subject. Her chromosomes included a normal X and a fragment (shown above). The fragment is smaller than either an X or a Y (compare to her father's Y at far right). A DNA probe containing a Y chromosome sequence was used to examine restriction fragments from a control female, the subject's mother, the subject, and the subject's father (2-3). DNA isolated from peripheral blood was digested with EcoRI. The resulting DNA fragments are separated by electrophoresis in agarose gels, the large DNA fragments were located at the top of the gel and the small fragments at the bottom. The DNA fragments are transferred or blotted from the agarose gel to a filter membrane as described by Southern and the filter-bound DNA is then hybridized to a solution containing a Y chromosome probe (32P-radiolabeled single-stranded DNA or RNA copy of sequences specific to the Y chromosome). The probe molecules anneal to their complementary sequences on the filter to form double-stranded DNA. One strand of these molecules is from the subject's DNA while the other strand is the radiolabeled probe. After hybridization, the filter is washed to remove excess probe and subjected to autoradiography. The DNA fragments that annealed to the probe appear as bands on the developed film. By comparing the location of these bands with that of markers of known molecular weight, the size of hybridizing DNA fragments is determined. In Fig. 2 obtained...
after EcoRI digestion, the probe shows no hybridization to 3.3 kb fragments from a normal female or the subject's mother. However, the subject and her father's DNA both show intense bands indicating they both have Y chromosomal DNA sequences. The decreased amount of hybridization in the subject's DNA compared to that of her father results from her being a 45, X/46, X+frag mosaic so that she did not carry the Y fragment in all of her cells.

2. Gene Deletions

Familial isolated growth hormone deficiency type 1A (IGHD 1A) is an endocrine disorder that is caused by gene deletions. This rare disorder has an autosomal recessive mode of inheritance and affected individuals have severe growth retardation due to complete deficiency of GH. Most cases respond only briefly to GH replacement therapy due to their tendency to develop high titers of anti-GH antibodies (4-5).

Restriction analysis of the structural gene for GH (GH1) is complicated by the fact that it is one of the five GH related genes (5′-GH1:CSHP1:CSH1:GH2:CSH2-3′) contained in the GH gene cluster. Although these various genes share extensive sequence homology, only the GH1 locus encodes GH. The CSHP1 gene is of unknown function and encodes a chorionic somatomammotropin hormone (CSH)-like peptide, whereas CSH1 and CSH2 encode CSH, and GH2 encodes a variant GH of unknown function whose gene contains an internal BamHI site. The GH1 gene is flanked by consistent BamHI sites that are 3.8 kb apart. While the CSHP1, CSH1, GH2, and CSH2 genes are sufficiently homologous to hybridize to the GH1 probe, they all are contained in BamHI derived fragments that differ in size from that of GH1. Autoradiograms of DNAs from IGHD 1A subjects lack the 3.8 kb fragments that normally contain the GH1 genes (Fig. 3). In addition, the intensity of the 3.8 kb bands in DNA from the heterozygous parents is intermediate between that of controls and their affected children. These results show that IGHD 1A subjects are homozygous and their parents are heterozygous for GH1 gene deletions. Since these deletions preclude production of any GH, affected individuals tend to be immunologically intolerant to exogenous GH.

3. Point Mutations

The detection of certain insulin gene mutations is an example of the detection of point mutations by restriction analysis. Several such mutant insulins have been reported. The clinical features seen in heterozygotes include hyperglycemia with high insulin levels, in the absence of insulin antagonists or receptor antibodies, and a normal response to exogenous insulin. The mutant insulin first described has an 88% decrease in biological potency (6-8). This mutant insulin was found to have a leucine substituted for phenylalanine at residue 25 in its B chain and is called (Leu<sup>25</sup> insulin) (9). DNA analysis of the Leu<sup>25</sup> insulin gene revealed a nucleotide substitution at position 402. Interestingly, the restriction enzyme MboII which cleaves the
normal insulin gene at this point is unable to cleave Leu$^{bss}$ insulin because of its nucleotide change. Thus, DNA from a control yield fragments of 336 and 555 bp when hybridized to an insulin gene probe. An individual heterozygous for Leu$^{bss}$ insulin will have a 891 bp fragment (not seen in controls) because MboII cannot cut at the site that separates the 336 and 555 bp fragments (10-11). A second mutant insulin (Ser$^{bss}$ insulin) has a serine for phenylalanine substitution. This mutation can also be detected because it prevents cleavage by MboII (12).

4. Linkage Analysis

Studies of familial diabetes insipidus are an example of mapping the gene responsible for an endocrine disorder by linkage analysis. Linkage analysis is usually done using restriction fragment length polymorphisms (RFLPs) (Fig. 4). RFLPs result from DNA polymorphisms or base differences that normally occur without any clinical consequences about 1 in every 200 to 500 nucleotides not encoding proteins (13). These differences are usually single nucleotide substitutions some of which can affect a restriction site. A second or insertion/deletion type of DNA polymorphism also called variable numbers of tandem repeats or dinucleotide repeats can alter the DNA fragment size without affecting the enzyme recognition site. Finally the presence or absence of a pseudogene (a nonfunctional gene whose sequence is homologous to a functional gene) changes the length of the DNA fragment that is bounded by two restriction sites (14).

In an autosomal dominant form of diabetes insipidus due to arginine vasopressin deficiency (ADNDI) we could not detect deletions, insertions or rearrangements of an arginine vasopressin (AVP) allele (15). Using an XbaI RFLP closely linked to AVP we found apparent cosegregation with ADNDI in two families (Fig. 5). The overall LOD score of 2.70 indicates that the genetic locus for ADNDI maps within or near the AVP locus and suggests that a defective AVP allele may cause this form of diabetes insipidus.

B. Polymerase Chain Reaction

The polymerase chain reaction (PCR) technology is an important new tool for DNA diagnosis (Fig. 6) (16-18). This procedure enables the gene selected for study to be enzymatically amplified relative to the remainder of the genomic DNA. This amplification can yield $10^7$-$10^8$ copies of the original segment
which can allow direct visualization of the fragments under study thereby precluding the need for Southern blots and radiolabeled probes. PCR amplification makes it possible to amplify specific sequences directly from complex genomic samples without cloning. This ability has led to the development of many new techniques of gene analysis and has revolutionized the way in which inherited disorders are studied. Mutant alleles can be PCR amplified and then rapidly characterized using a combination of segregation analysis, gene scanning, direct DNA sequencing, and RNA transcript analysis (16-18).

An example of the utility of PCR amplification is detection of Y chromosome fragments in Turner syndrome. In a case similar to that shown in Fig. 2 we have used the PCR and Y specific oligonucleotide primers to determine if the chromosomal fragment was derived from an X or Y. Note the affected subject who was mosaic for 45, X/46X + frag has easily detectible Y chromosomal material in DNA derived from peripheral blood (Fig. 7). Such studies can be done in 3 hours, require small alquots (1-2μL) of blood, and avoid the use of Southern blots or probes.

**C. DNA Sequencing**

DNA sequencing is usually done by the dideoxy method using DNA polymerase to synthesize a complementary radioactive copy (Fig. 8) (19). During the DNA polymerase reaction 2', 3' dideoxynucleotides of A, C, G, or T are added to four separate reactions.
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Fig. 8. Schematic depiction of DNA sequencing by the dideoxy method. Following the DNA polymerase reaction in the presence of dideoxy analogs, the four reaction products are subjected to gel electrophoresis and autoradiography enabling sequence deduction.

DNA molecules that is radioactive with a common 5' end, but of varying length because of the incorporation of a specific 3' end. Next, each of the four reaction products are subjected to electrophoresis and autoradiography and sequence deduction are performed. Comparison of the mutant and wild-type sequence allows deduction of the mutation. Sequencing not only identifies the mutation but can enable derivation of short oligonucleotide sequences that can be used as probes to detect the presence of that specific mutation in genomic DNA samples from other individuals.

Status of Molecular Endocrinology

For a variety of endocrine disorders including diabetes insipidus, GH deficiency, diabetes mellitus, and vitamin D resistant rickets there are varying amounts of data from DNA studies. I will briefly summarize selected aspects of these data and their conclusions.

A. Diabetes Insipidus (DI)

Molecular studies have clarified the basis of genetic forms of DI. These have clarified the structure of the AVP gene (Fig. 9) and assigned it to chromosome 20 (20). Recently, a number of mutations causing autosomal dominant DI have been identified (Fig. 10) (21-24). The AVP gene (AVP-NPII) contains AVP in exon 1 and its carrier protein (neurophysin or NPII) in exons 1-3. Cosegregation of the ADNDI phenotype and DNA polymorphisms closely linked to the AVP gene was found in two unrelated families, suggesting that muta-
Fig. 10. The location and types of mutations found in the human AVP gene associated with autosomal dominant diabetes insipidus.

mutations in AVP cause some cases of ADNDI (15). Two mutations in NP II (Gly→valine and Gly→ser at codons 17 and 57) causing ADNDI have been reported (21,22). Gly17 mutation is thought to perturb the conformation and self-aggregation of AVP-NP II precursors that are required for normal intracellular transportation and secretion (21). The Gly57 mutation is speculated to perturb NP II’s ability to transport or protect AVP from proteolytic degradation (22). Recently, a novel mutation that affects AVP rather than NP II, was reported that should affect cleavage of AVP from its precursor (prepro AVP) as an additional cause of ADNDI (23).

B. Severe GH Deficiency

The most severe form of IGHD called IGHD IA, has an autosomal recessive mode of inheritance. Affected individuals occasionally have short lengths at birth and hypoglycemia in infancy, but uniformly develop severe dwarfism by 6 months of age. In response to replacement therapy with exogenous GH, IGHD IA subjects have a strong initial anabolic and growth response that is frequently followed by the development of anti-GH antibodies in sufficient titer to block the response to GH replacement (25-27). These features led Illig et al to hypothesize that IGHD IA caused complete prenatal and postnatal deficiency of endogenous GH secretion which resulted in an immune response to exogenous GH (27).

Deletions. Initially, all individuals with IGHD IA were found to be homozygous for GH1 gene deletions and developed anti-GH antibodies with treatment (4,28). Subsequently, additional cases with GH1 gene deletions have been described who also have complete GH deficiency, but respond well to GH replacement (4,29). Thus, the clinical outcomes of subjects with the same molecular findings (homozygosity for GH1 gene deletions) vary making the presence of anti-GH antibodies an inconsistent finding in IGHD IA cases.

At a molecular level, Southern blot analysis showed deletions of ~6.7, 7.0 or 7.6kb with most ~75% being 6.7kb (4). DNA sequence analysis of the fusion fragments associated with GH1 gene deletions have shown that homologous recombination between sequences flanking the GH1 gene cause these deletions (30-32). This same mechanism has been demonstrated to cause deletions of the β-globin and LDL receptor genes in some subjects with β-thalassemia or familial hypercholesterolemia, respectively. Currently, GH1 gene deletions are detected using polymerase chain reaction (PCR) amplification of the homologous regions flanking the GH1 gene and the fusion fragments associated with GH1 gene deletions (Fig. 11) (31-32). Since the fusion fragments associated with 6.7 kb deletions differ in the size of fragments produced by certain restriction enzymes (see Sma I sites indicated by solid circle in Fig. 11) homozgyosity or heterozygosity for these deletions can be easily detected by enzyme digestion of PCR products (31-33). A variety of studies suggest that 13-15% of subjects with severe IGHD (>−4.5 SD in height) have GH1 gene deletions (4,30-33). Recently, frameshift and nonsense mutations have also been found in subjects with the IGHD IA phenotype so that this disorder may be best described as complete GH deficiency due to GH1 gene defects, rather than gene deletions alone (34-35).
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Fig. 11. PCR amplification of homologous sequences that flank the GH1 gene using a single set of oligonucleotide primers. Note that the 5' and 3' homologous sequences (labelled A and B) differ in the location of restriction sites (indicated by solid circles). The sites also differ from those on junction or fusion fragments (labelled C) shown below that arise during recombination events that produce GH1 gene deletions (18).

C. Laron Dwarfism

The first examples of GH receptor or GHR mutations reported were deletions of portions of the gene encoding the extracellular domain (Fig. 12) (36-37). Southern blotting showed altered restriction patterns of the GHP genes from patients with Laron syndrome who had no detectable GH binding protein (GHBP) and very low levels of IGF-1. While these and other studies were interpreted as showing deletions of exons 3, 5-6 and part of 4 from the GHR gene, the mechanism by which these two non-contiguous deletions arose remains unclear. In two of nine patients studied by Godowski et al. GHR deletions were found and JS Parks detected GHR deletions in five of nine of the Laron syndrome probands studied (36-37).

Multiple point mutations have been detected within the GHR gene (Fig. 13) (38-40). Amselem et al detected a T to C substitution that converts the 96th residue of the extracellular domain from phenylalanine to serine (39). Duquesnoy et al. demonstrated that cells transfected with this mutant cDNA lacked GH binding activity (40).

Two different stop codon mutations of GHR genes in Laron dwarf patients have been reported (38). In a patients of Northern European origin, a cysteine (TGC)→stop codon (TGA) mutation was detected at codon 38 in exon 4 (Fig. 13), and an arginine (CGA)→stop codon (TGA) mutation was found at codon 43 in exon 4 of two Mediterranean patients who were products of consanguineous marriages. Both stop codons truncate the GHR protein and delete most of its GHBP domain and all of its transmembrane and intracellular domains. These findings are consistent with the lack of GHBP in each of the patients with Laron syndrome. The mechanism of the CGA to TGA mutation is consistent with deamination.

Fig. 12. Schematic representation of the GH receptor (GHR) gene showing the relationship of various exons to the extracellular, transmembrane and cytoplasmic GHR domains. Locations of deletions and selected point mutations are shown below.
Fig. 13. Schematic representation of the GHR gene showing the locations of CpG dinucleotides as small circles. Segments of normal sequence and selected mutations are shown below.

of 5-methyl-cytosine that preferentially occurs in CpG dinucleotides. Such dinucleotides often represent “hot-spots” for CG to TG or CG to CA mutations, and 17 occur within the GHR gene (Fig. 13). Two of these, at nucleotides 181 and 703, occur in CGA codons that could yield stop codons (36,38-39).

Rosenbloom et al. identified 20 patients with Laron syndrome in an inbred population of Spanish extraction in Southern Ecuador (41). These patients were -6.7 to -10 SDS below the mean height and had limited elbow extension, blue sclera, short limbs, hip degeneration, acrohypoplasia and normal or superior intelligence. To determine the associated defect in the GHR gene, Berg et al. used denaturing gradient gel electrophoresis (DGGE) to analyze each exon of the GHR gene (42). Unusual fragments derived from exon 6 showed abnormal mobility and DNA sequencing showed an A→G substitution in the third position of codon 180, which is 24 nucleotides from the 3' end of exon 6. While this mutation does not cause an amino acid substitution, it produces a consensus 5' splice sequence within exon 6. The resulting near consensus donor splice site within exon 6 causes aberrant splicing and deletion of eight amino acids of the 3' end of exon 6. Deletion of these residues is thought to reduce the function of the GHBP molecule (42).

D. Panhypopituitary Dwarfism

Panhypopituitary dwarfism is characterized by deficiency of one or more of the other pituitary trophic hormones (ACTH, FSH, LH or TSH) in addition to GH deficiency. While the great majority of cases are sporadic, there are both autosomal recessive and X-linked forms (4). Recently, at least four different Pit-1 mutations have been found in humans in a subtype of panhypopituitary dwarfism associated with GH, PrL and TSH deficiency. These include, first, a C→T substitution in codon 172 which changes a CGA (arginine)→TGA (stop) (43). Second, a patient heterozygous for a G→T substitution in codon 271 was reported. The mechanism of the dominant effect of this GGG (arginine)→TGG (tryptophan) mutation is not completely understood (44). The third and fourth Pit-1 mutations were found in two Dutch families in which affected individuals had postnatal growth failure with complete deficiencies of GH and PrL, while their T4 levels were low prior to or following GH replacement (45-46). The family having normal T4 levels prior to GH replacement, were homozygous for a G→C substitution in codon 158 changing GCA (alanine)→CCA (proline). This mutation interferes with formation of Pit-1 homodimers and dramatically reduces the altered Pit-1's ability to activate transcription. In the family with low T4 levels, the affected children were genetic compounds with one deleted and one Pit-1 gene with the previous mutation. These cases emphasize the importance of determining PrL levels as well as TSH responses to TRH administration in evaluating panhypopituitarism. Since GH and TSH deficiency often occurs together, finding low PrL levels and failure of subjects to have TSH responses should raise the question of their having Pit-1 gene defects.

E. Diabetes Mellitus

Recent molecular genetic investigations have begun to unravel the suspected molecular heterogeneity and pathogenesis of diabetes
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mellitus. Using a variety of molecular techniques, specific nuclear and mitochondrial DNA abnormalities have been identified (see Table 1) (47-61). Interestingly, genes important in causing susceptibility to or protection from insulin dependent diabetes have been elucidated within the HLA-D and the insulin gene regions. One group of patients with maturity onset diabetes mellitus of the young or MODY have been found to have missense or nonsense mutations of the glucokinase gene, while a second group demonstrated linkage to the ADA (adenosine deaminase) locus on chromosome 20q and a third group showed no linkage to either locus. In the maternally inherited syndrome of diabetes mellitus and deafness a deletion and a point mutation of mitochondrial DNA (mtDNA) have been found. Finally patients with several insulin-resistance syndromes have been found to have mutations in the insulin receptor gene. These recent discoveries (Table 1) demonstrate the heterogeneity of diabetes mellitus and provide the potential to use molecular techniques in making early preclinical diagnosis as well as more accurate determinations of risk to develop diabetes mellitus possible in certain families (47-61).

F. Vitamin D Resistant Rickets

Vitamin D resistant rickets is an autosomal recessive disorder which is characterized by a target organ resistance to the action of 1,25-dihydroxyvitamin D$_3$. Interestingly, while the intracellular vitamin D receptor (VDR) from affected individuals in some families displays normal binding of 1, 25-dihydroxyvitamin D$_3$, it has decreased affinity for DNA (62). To determine the status of this biologically inactive VDR gene Hughes et al PCR amplified and sequenced exons 2 and 3. Sequence analysis showed single base substitutions that destabilize the ZN finger domain of the mutant VDR (62).

<p>| Table 1. |
| GENES AND REGIONS CONTRIBUTING TO DIABETES MELLITUS |</p>
<table>
<thead>
<tr>
<th>Type</th>
<th>Gene</th>
<th>Chromosome</th>
<th>Type of Abnormality</th>
<th>Ref No.</th>
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<tbody>
<tr>
<td>IDDM</td>
<td>HLA</td>
<td>6P</td>
<td>HLA-DRB3 + DR4 susceptibility</td>
<td>47-49</td>
</tr>
<tr>
<td></td>
<td>INS</td>
<td>11P</td>
<td>RFLP linkage to insulin gene</td>
<td>51-52</td>
</tr>
<tr>
<td>MODY</td>
<td>Unknown</td>
<td>20q</td>
<td>Linkage to ADA (adenosine deaminase) gene</td>
<td>53</td>
</tr>
<tr>
<td>Gluco-</td>
<td></td>
<td>7P</td>
<td>*Missense mutation (G to C substitution) in codon 299</td>
<td>54</td>
</tr>
<tr>
<td>kinase</td>
<td></td>
<td></td>
<td>*Nonsense mutation (G to T substitution) at codon 279</td>
<td>55</td>
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<tr>
<td>NIDDM</td>
<td>INS</td>
<td>11p</td>
<td>*TTCC→TTG substitution at residue 25 of insulin gene β chain (Insulin Chicago)</td>
<td>11</td>
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<tr>
<td></td>
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<td>*TTCC→TTC substitution at residue 24 of insulin gene β chain (Insulin Los Angeles)</td>
<td>12</td>
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<td></td>
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<td></td>
<td>*Leucine→Valine substitution at residue 3 of insulin a chain (Ittelson Wakeshima)</td>
<td>56-57</td>
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<tr>
<td>Secondary DM</td>
<td>INS</td>
<td>19P</td>
<td>*Decreased insulin receptor mRNA due to missense and nonsense mutations</td>
<td>58-60</td>
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<tr>
<td>MODY &amp; Deafness</td>
<td>Multiple</td>
<td>Mitochondrial DNA</td>
<td>*10.4 kb deletion (removes 20 of mRNA replication)</td>
<td>61</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>*A to G transition at nucleotide 3,243 [DNA→RNA gene]</td>
<td>62</td>
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<tr>
<td>INS = insulin</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td>INSR = insulin receptor</td>
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Summary

Several examples of molecular studies of hereditary endocrine disorders have been reviewed. These DNA studies have revealed the location and types of molecular derangements causing these diseases. The mutations discovered have in turn explained the alterations in the hormone products. These same methods should reveal the basis of many other endocrine disorders for which specific DNA probes are or will become available. I thank the Research Society for Growth Disturbance in Children for inviting me.

Acknowledgements

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