High-Resolution Physical and Genetic Mapping of the Critical Region for Meckel Syndrome and Mulibrey Nanism on Chromosome 17q22–q23

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Previously, we assigned the genes for two autosomal recessive disorders, Meckel syndrome (MKS; MIM 249000) and Mulibrey Nanism [MUL (muscle–liver–brain–eye Nanism); MIM 253250] that are enriched in the Finnish population, to overlapping genomic regions on chromosome 17q. Now, we report the construction of a bacterial clone contig over the critical region for both disorders. Several novel CA-repeat markers were isolated from these clones, which allowed refined mapping of the MKS and MUL loci using haplotype and linkage disequilibrium analysis. The localization of the MKS locus was narrowed to <1 cM between markers D17S1290 and 132-CA, within an ~800-kb region. The MUL locus was refined into an ~1400-kb interval between markers D17S1290 and 52-CA. The whole MKS region falls within the MUL region. In the common critical region, the conserved haplotypes were different in MKS and MUL patients. A transcript map was constructed by assigning expressed sequence tags (ESTs) and genes, derived from the human gene map, to the bacterial clone contig. Altogether, four genes and a total of 20 ESTs were precisely localized. These data provide the molecular tools for the final identification of the MKS and the MUL genes.

[The sequence data described in this paper have been submitted to the GenBank data library under accession nos. G42608–G42611, G42376–G42388, and G42200–G42250. The online supplement for primer sequences and PCR product sizes, as well as the STS-content table, are available at http://www.cshl.org/gr.]

Meckel syndrome (MKS; MIM 249000) and Mulibrey Nanism [MUL (muscle–liver–brain–eye Nanism); MIM 253250] are rare autosomal recessive disorders in which the clinical presentation involves multiple tissues. MKS, which leads to death soon after birth, is characterized by central nervous system (CNS) malformations, in most cases occipital encephalocele, polykyphic kidneys, cystic and fibrotic changes of the liver, and polydactyly (Salonen 1984). MKS has been suggested to be caused by a failure of mesodermal induction (Blankenberg et al. 1987), and in MUL, several tissues of mesodermal origin are affected. The typical clinical features for MUL are growth failure of prenatal onset, dysmorphic features, constrictive pericardium, with consequent hepatomegaly, yellowish dots in the ocular fundi, and J-shaped sella turcica (Perheentupa et al. 1973; Lipsanen-Nyman 1986; Lapunzina et al. 1995). The prognosis of MUL patients is determined mostly by the degree of the constrictive pericardium (Perheentupa et al. 1973).

Both MKS and MUL are typical diseases of the “Finnish disease heritage”; that is, a number of hereditary disorders that occur more frequently in Finland than elsewhere (Perheentupa et al. 1970). The birth prevalence of MKS in Finland is estimated to be 1:9000; however, even higher incidences are reported in some other isolated populations (Salonen and Norio 1984; Salonen 1984). Although the Finnish MKS patients represent both a clinically and genetically uniform syndrome, MKS, globally, seems to be heterogenous with a wide spectrum of phenotypes and locus heterogeneity (Paavola et al. 1997; Roume et al. 1997, 1998). Some 75 MUL patients have been described worldwide, most of them from Finland, where the incidence of MUL is estimated to be 1:40,000 (Lipsanen-Nyman 1986). The
reported non-Finnish MUL patients are mostly sporadic cases (Lapunzina et al. 1995).

The genes for both MKS and MUL have been assigned previously to chromosome 17q (Paavola et al. 1995; Avela et al. 1997). The MKS locus covered a 13-cM region between markers D17S1607 and D17S807, and no allelic association was initially observed with the markers available for the study. Obligate recombinations recently detected in two new Meckel families restricted the locus within ~5 cM between D17S1606 and D17S807 (P. Paavola, unpubl.). The MUL locus was assigned previously to a 7-cM region within the MKS locus interval between marker loci D17S1799 and D17S948 where initial linkage disequilibrium analysis suggested a localization near marker D17S1853 (Avela et al. 1997). The haplotype analysis further refined the localization between D17S1853 and D17S1604 (K. Avela, unpubl.).

Here, we report the physical and transcript map of the critical region for both MKS and MUL, which provides the material for the final identification of the disease genes. The construction of the physical map allowed us to isolate several novel polymorphic markers and to significantly refine the localization of the MKS and MUL loci by exploiting haplotype and linkage disequilibrium analysis. The high-resolution physical and genetic mapping also enabled the positional exclusion of a serine–arginine-rich splicing factor gene, SFRS1, and a T-box transcription factor gene, TBX2, a putative candidate based on the expression pattern of the homologous mouse gene during development.

RESULTS

Construction of the Physical and Transcript Map

The contig skeleton for our region of interest was derived from the Whitehead/MIT database (Hudson et al. 1995; http://www.genome.wi.mit.edu). A total of 19 previously mapped yeast artificial chromosomes (YACs) (Chumakov et al. 1995) were localized using metaphase fluorescence in situ hybridization (FISH) analyses. The following YACs on the MKS–MUL region were shown to be nonchimeric: 925H8, 791A5, 898E7, 905A4 proved to be chimeric. Only nonchimeric clones were used for sequence-tagged site (STS)-content mapping to confirm the order of the clones and STSs.

To construct a more detailed map and to generate additional microsatellite markers for further narrowing of the critical disease gene regions, we screened PAC and BAC libraries (Shizuya et al. 1992; Ioannou et al. 1994) with the available STSs and expressed sequence tags (ESTs). The ends of the isolated PAC and BAC clones were sequenced to develop new STSs (http://www.cshl.org/gr, Suppl. 1) for further screening of clones and for securing the overlaps between the clones. In the critical region for MKS and MUL, between the markers D17S1290 and S2-CA, we were able to construct a bacterial clone contig with at least two times clone coverage consisting of both PAC and BAC clones (see Fig. 1). Three bacterial clones, PAC hRPC.1171110, BAC hRPK.506h21, and PAC hRPC.1096f1, that have been sequenced at the Whitehead Institute/MIT (http://www.seq.wi.mit.edu), were found by using BLAST comparisons with the STS sequences (Fig. 1). The sequence data from the clone hRPC.1171110 was used to fill the remaining region of 1 × clone coverage in our contig. The contig extended outside our region of interest both on the centromeric and telomeric side where it partially reached only 1 × clone coverage.

Six novel dinucleotide repeat markers were isolated from three PAC (95i19, 52i20, and 58i18) and two BAC (272g3 and 132i10) clones of the region (Fig. 1). The primer sequences and the allele sizes and frequencies are shown in Table 1. We also assigned a CHLC marker, D17S1290, into the genomic clones within this region (925H8, 791A5, and S3n20, 194012, 17b7, 313g19, and hRPC.1171110). In addition, the detailed physical map enabled us to precisely localize a total of 20 ESTs of both known and unknown homology, which were previously only roughly assigned either by radiation hybrid mapping or by STS-content mapping to the YAC clones in the region (see Fig. 1; http://www.cshl.org/gr, Suppl. 2). These ESTs were derived from the gene map of the human genome (Schuler et al. 1996). One EST (WI-7318 or SGC30655) showed considerable homology to a mouse gene encoding for clathrin heavy-chain protein, and two other ESTs (WI-18519 and WI-18389) represented the recently cloned human RAD51c gene (Dosanjh et al. 1998). Three other genes, the B-cell leukemia/lymphoma-5 gene, BCL5 (GenBank accession no. M31826), the myeloperoxidase gene, MPO (GenBank accession no. M19507; the corresponding EST is WI-6053), and the serine–arginine-rich splicing factor gene, SFRS1 (GenBank accession no. M69040), were also precisely localized (see Fig. 1; http://www.cshl.org/gr, Suppl. 2).

Simultaneously with the STS-content mapping of the isolated PAC and BAC clones, we positioned clones in relation to each other by the fiber-FISH method (Fig. 2; Heiskanen et al. 1994, 1996). Before the fiber-FISH analysis, the localization of the clones on 17q22-q23 was confirmed by metaphase-FISH. By hybridizing two to four PAC or BAC clones simultaneously on free DNA fibers, the clone order and orientation in relation to each other as well as the clone overlaps and gaps between different clones were determined. This enabled the construction of a fine-resolution visual clone contig across the MKS–MUL region.
The insert sizes of the PAC and BAC clones constituting the minimum tiling path of the contig were determined by using restriction digestion followed by pulsed-field gel electrophoresis (PFGE; Table 2). Fiber-FISH analyses were used to verify and confirm these size estimations. In fiber-FISH, one or two bacterial clones, whose sizes were determined by PFGE, were used as a standard for calibration in each hybridization image to estimate the sizes of other clones. According to both the PFGE data and fiber-FISH, the size of the contig from D17S1290 to 52-CA, that is also the critical region for MUL, was estimated to be ~1.4 Mb. Furthermore, the total physical length of the critical region for MKS was determined to be ~800 kb (between markers D17S1290 and 132-CA; Figs. 1 and 2).

**High-Resolution Mapping of MKS and MUL by Haplotype and Linkage Disequilibrium Analysis**

**MKS**

Before the establishment of the physical map and isolation of novel polymorphic markers, the critical chromosomal region for MKS was indicated by a core haplotype (cen-3-3-6-tel for markers D17S957, D17S1606, and D17S1604) that was observed in 10 out of 40 MKS chromosomes but in none of the control chromosomes. Later, significant allelic association was detected between MKS and the novel marker loci D17S1290, 95-CA, 272a-CA, 272b-CA, 132-CA, and 58-CA. Using the DISLAMB program (Terwilliger 1995), the strongest associations between MKS and an allele were detected at the marker loci 95-CA and 132-CA ($\lambda$-value > 0.80 with $P$-value < 10$^{-5}$; see Table 3). At 132-CA the disease associated allele (n:o 5, uncommon allele in the controls) was observed also in four chromosomes (Table 3) that represent other, rare disease haplotypes (see below). Therefore, the $\lambda$-value for 132-CA may be misleadingly high (for predicting the distance of the gene from the marker) as compared with the other marker loci.

Three telomeric and two centromeric historical recombinations in MKS chromosomes defined a minimum genetic region for the MKS locus between the markers D17S1290 and 132-CA (Table 3) that represent other, rare disease haplotypes (see below). Therefore, the $\lambda$-value for 132-CA may be misleadingly high (for predicting the distance of the gene from the marker) as compared with the other marker loci.

Three telomeric and two centromeric historical recombinations in MKS chromosomes defined a minimum genetic region for MKS chromosomes located between the markers D17S1290 and 132-CA (Table 3). Five patients were compound heterozygotes for the common MKS haplotype and another haplotype, and in two patients, both chromosomes differed from the common MKS haplotype. In these seven patients, four different rare MKS haplotypes were observed on the minimum genetic region, possibly suggesting “minor” MKS mutations (in Table 3 the haplotypes are defined by a solid line).
The analysis of the novel markers from the region between D17S1853 and D17S1604 revealed significant allelic association between MUL and marker loci D17S1290, 95-CA, 272a-CA, 272b-CA, 132-CA, 58-CA, and 52-CA. Using the DISLAMB program (Terwilliger 1995) the strongest associations between MUL and an allele were detected at 95-CA, 272a-CA, 272b-CA, 132-CA, and 58-CA (l-value > 0.95 with P-value < 10^{-5}; see Table 3).

Analysis of historical recombinations in haplotypes of the MUL chromosomes (Table 3) refined the critical MUL region between markers D17S1290 and 52-CA. Altogether, five centromeric and four telomeric historical recombinations were detected in MUL chromosomes. One patient was a compound heterozygote for the ancestral haplotype and a different haplotype that most likely represents a “minor” MUL mutation in the Finnish population.

**Exclusion of TBX2 and SFRS1 as Candidate Genes for MKS and MUL**

TBX2, a gene encoding a T-box transcription factor that in mouse is expressed in many stages and tissues during the embryonal development (Chapman et al. 1996), was recently cloned and assigned to 17q23 (Cammell et al. 1995). Thus, it presented a potential candidate gene for both MKS and MUL. Using PCR with primers designed from the 3’ UTR of the TBX2 cDNA, we located TBX2 to YAC clones (731E9 and 935C4) on 17q23 that did not overlap with the MKS-MUL clone contig (Fig. 1). A PAC clone, 210a19, isolated for TBX2 was also assigned by metaphase FISH on chromosome 17q23. The hybridization of 210a19 on
free DNA fibers showed that TBX2 is located at least 2 Mb telomeric to the marker D17S1604 (data not shown). Thus, TBX2 could be positionally excluded as the causative gene for MKS and MUL. The serine–arginine-rich splicing factor gene, SFRS1 (GenBank accession no. M69040), was localized to the MKS–MUL clone contig, but it resided centromeric to D17S1290 that defined the centromeric border of the critical regions for both MKS and MUL. Thus, SFRS1 was also positionally excluded.

DISCUSSION

We have created a defined physical and EST-based transcript map for the region on chromosome 17q harboring the genes for MKS and MUL, two distinct disorders displaying symptoms from multiple organs, for the eventual cloning of the corresponding genes.

The bacterial clone contig has been confirmed by STS-content mapping using previously only roughly assigned STSs and ESTs as well as several novel PAC/BAC-end sequence-derived STSs (see Fig. 1; http://www.cshl.org/gr, Suppl. 2). The contig assembly was further confirmed visually by the fiber-FISH technique. Furthermore, the length of the contig in the critical MKS–MUL region (from D17S1290 to 52-CA) was estimated to be ~1.4 kb. This estimate was primarily based on the clone size determinations by PFGE, but it was also concordant with the measurements made by the fiber-FISH analyses. The fiber-FISH method has earlier been shown reliable in estimating physical distances of various disease linked genomic regions (Heiskanen et al. 1995; Klockars et al. 1996; Aaltonen et al. 1997). The entire contig spanning the MKS–MUL critical region has at least 2 × clone coverage. The contig has thus enough coverage to enable long-range sequencing as a tool to identify novel genes, provided that the disease locus regions as determined by linkage disequilibrium mapping are small enough to justify such an effort.

The detailed physical map that we constructed allowed us to isolate and precisely map several polymorphic microsatellite markers, which enabled the significant refinement of the MKS and MUL loci. The MKS locus was narrowed from 13 cM to a <1-cM region between markers D17S1290 and 132-CA, whereas the MUL locus was refined between the markers D17S1290 and 52-CA, to a region that overlaps the MKS critical region (Fig. 1). Taking advantage of the genomic clones isolated from this region and their size estimates, the MKS locus interval was estimated to cover ~800 kb and the MUL locus interval approximately 1400 kb of DNA.

Table 2. The Estimated PAC/BAC Insert Sizes of the Minimum Tiling Pathway of the Contig

| Clone name | Estimated size (kb) |
|------------|---------------------|
| 6i20       | 130                 |
| 40h14      | 120                 |
| 53n20      | 240                 |
| 58p18      | 80                  |
| 95i19      | 170                 |
| 104c2      | 80                  |
| 158s5      | 110                 |
| 180k5      | 170                 |
| 194o12     | 80                  |
| 229h9      | 200                 |
| 261p2      | 50                  |
| 272g3      | 180                 |
| 303g1      | 140                 |
| 459c10     | 240                 |
| 471n15     | 180                 |
| 520n16     | 130                 |
| 523e2      | 140                 |
### Table 3. Linkage Disequilibrium and Haplotype Data in MKS and MUL

| MKS Markers | λ | Confidence intervals for λ | P-value | MKS-associated haplotypes from unrelated patients |
|-------------|---|---------------------------|---------|--------------------------------------------------|
| D17S1606    | 0.32 | 0.32–0.85 | N.S. | 3 3 × 3 3 3 3 5 4 5 8 7 10 7 N.D. 3 N.D. 5 13 7 2 |
| D17S1290    | 0.64 | 0.43–0.85 | N.S. | 5 6a 5 5 5 5 5 5 5 5b 5 2 3 8 8 6 2 1 |
| 95-CA       | 0.82 | 0.62–0.95 | 4.3 × 10^-6 | 3 3 3 3 3 3 3 3 3 3 3 3 3 3 2a 3 3 5 5 5 4 4 4 2 2 1 |
| 272a-CA     | 0.78 | 0.61–0.90 | 1.1 × 10^-8 | 5 5 5 5 5 5 5 5 5 5 5 5 3 4 4 4 5 5 4 6 4 5 5b 3 |
| 272b-CA     | 0.68 | 0.45–0.89 | N.D. | 7 7 7 7 7 7 7 7 7 6 6 3 N.D. 2 6 6 N.D. 5 6 5 6 5 |
| 132-CA      | 0.84 | 0.67–0.94 | 1.3 × 10^-12 | 5 5 5 5 5 5 5 5 5 5 5 5 4 4 4 2 2 1 |
| 58-CA       | 0.71 | 0.53–0.85 | N.S. | 3 3 3 3 3 6 5 6 5 6b 2 3 3 6 5 6 N.D. 6 6 6 2 6 2 |
| 52-CA       | 0.32 | N.S. | 6 6 6 × 5 9 9 5 8 6b 4 9 14 N.D. 2 N.D. 4 1 6 2 |
| D17S1604    | 0.12 | N.S. | | Number of chromosomes detected: 13 1 3 4 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
| | | | | Total number of unrelated MKS chromosomes 48 |

| MUL Markers | λ | Confidence intervals for λ | P-value | MUL-associated haplotypes from unrelated patients |
|-------------|---|---------------------------|---------|--------------------------------------------------|
| D17S1290    | 0.80 | 0.61–0.92 | 8.3 × 10^-10 | 7 7 7 5 5 7 6 6 4 4 4 4 4 4 3 |
| 95-CA       | 0.97 | 0.84–1.00 | 6.5 × 10^-13 | 4 4 4 4 4 4 4 1 4 4 4 4 4 4 5 |
| 272a-CA     | 0.96 | 0.80–1.00 | 5.7 × 10^-10 | 7 7 7 7 7 7 7 4 7 7 7 7 4 4 4 5 |
| 272b-CA     | 0.97 | 0.87–1.00 | <10^-15 | 3 3 3 3 3 3 3 4 |
| 132-CA      | 0.96 | 0.81–1.00 | <10^-8 | 1 1 1 1 1 1 1 1 |
| 58-CA       | 0.97 | 0.86–1.00 | <10^-8 | 6 6 2 6 6 6 6 6 |
| 52-CA       | 0.83 | 0.61–0.95 | 1.1 × 10^-7 | 4 4 4 4 4 4 4 15 |
| D17S1604    | 0.57 | 0.36–0.75 | 1.7 × 10^-6 | |
| | | | | Number of chromosomes detected: 19 11 4 4 1 2 1 |
| | | | | Total number of unrelated MUL chromosomes 42 |

The haplotypes consist of nine (MKS) and eight (MUL) microsatellite repeat markers covering the region for the disease loci. The λ values, confidence intervals for λ and P-values are shown for each marker.

The conserved haplotype.

Haplotypes most likely representing other mutations on the minimum genetic region.

(N.D.) Not determined.
(N.S.) Not significant.
*Possible allele mutation.
Uninformative meiosis.
(x) Other than the ancient allele.
Previously, the MUL locus was reported to map to an ~500-kb region around marker locus D17S1853 (Avela et al. 1997). This estimate was based on the assumption of 1 cm corresponding to 1 Mb in the region. The construction of the contig has enabled a more accurate estimate of the physical distance across the region. It is evident that in the MKS–MUL region the ratio of physical to genetic distance is well above the average of 1.0, maybe even as high as 2.0. Considering this and the confidence limits of the linkage disequilibrium analysis, the original MUL region estimate falls within the present, more accurate estimate.

In MKS, linkage disequilibrium was detectable only within an ~1000-kb region (D17S1290–S8-CA), whereas in MUL, linkage disequilibrium was detectable across ~2000 kb from D17S1606 to D17S1604. These data would suggest an earlier introduction of the MKS mutation into the Finnish population. The later appearance of MUL into the Finnish population is also supported by the genealogy of the disorders. In MUL, the disease is clustered in the restricted Savo and North Carelia regions of Eastern Finland (Lipsanen-Nyman 1986), representing partly early and partly late-settled areas (de la Chapelle 1993), whereas in MKS, the ancestors are spread all over Finland also on the coastal, early settlement regions (Salonen and Norio 1984). Furthermore, the haplotype data on MKS and MUL patients refers to multiple mutations in MKS versus one major ancestral mutation in MUL. In MUL, one major haplotype was found, and only a single patient was a compound heterozygote for this major haplotype and another very different haplotype. In MKS, one major haplotype was associated with the disease, but also four different haplotypes (on the minimum genetic region) were observed in seven heterozygous patients (Table 3).

The detailed physical map that we have constructed has now created valuable tools for isolating candidate genes for MKS and MUL. We have shown that the TBX2 and SFRS1 genes are located outside the critical regions of MKS and MUL thus excluding these genes as candidates. Even though the MPO gene maps to the MKS critical region, it does not appear as a relevant candidate because mutations in MPO have already been shown to cause defective cellular immunity (Kizaki et al. 1994). BCLS may be considered a relatively good candidate because BCL2-deficient mice have a polycystic kidney disease-like change of renal tubules (Nakayama et al. 1994). However, the nervous system was reported to be normal in mice despite the fact that this organ shows high levels of endogenous expression. The recently cloned RAD51c gene belongs to a group of strand transfer proteins that are thought to be involved in both recombinational repair of DNA damage and meiotic recombination (Dosanjh et al. 1998). No specific phenotypes are described to be caused by the mutations in the RAD51c gene or the other members of this gene family; thus, the probability of this positional candidate being the MKS or the MUL gene is difficult to evaluate. Additionally, the contig of genomic clones that we have constructed has enabled the localization of several ESTs and EST contigs of unknown homology on the critical MKS–MUL region. We are currently testing these positional candidates for mutations in the MKS and MUL patients to identify the genes causing these two disorders.

**METHODS**

**Isolation of Genomic Clones**

As a first step, we identified YAC clones for the MKS–MUL region. Clones positive for the previously mapped STSs and ESTs (Hudson et al. 1995; Dib et al. 1996; http://www.genome.wi.mit.edu) between markers D17S957 and D17S1604 were provided by the Centre d’Etude du Polymorphisme Humain (CEPH) in Paris. Individual clones were cultured, and total yeast DNA was extracted and stored in agarose beads or extracted by the alkaline lysis method. The chimera of YAC clones was analyzed by the metaphase FISH technique (Bray-Ward et al. 1996), and clones mapping only to 17q22–q23 were used for further mapping. The presence of STSs and ESTs (Hudson et al. 1995; Schuler et al. 1996) in the YACs was tested by PCR amplification.

The STSs and ESTs present in the YACs were used for screening of PAC and BAC libraries by PCR amplification (http://www.cshl.org/gr, Suppls. 1 and 2). Primer sequences for the genes used for bacterial clone screenings are as follows: BCLS (GenBank accession no. M31826), I: 5’-gagaacctttggccatgtg-3’, II: 5’-tggctgattcgctgtcgtc-3’, fragment size 180 bp; SFRS1 (GenBank accession no. M69040), I: 5’-gctgtattcatctttgaagttc-3’, II: 5’-gcctgtgattcatctttgaagttc-3’, fragment size 196 bp; TBX2 (GenBank accession no. M19507), I: 5’-ctgccccagcgctcctgcc-3’, II: 5’-ccgctgacccacactttc-3’, fragment size 120 bp. The PAC library consisting of 321 multiwell microtiter plates was kindly provided by Professor Pieter J. De Jong (Roswell Park Institute, Buffalo, NY). The commercially available P1 library (Genome Systems, Inc., St. Louis, MO) and BAC library DNA pools (release III, Research Genetics, Inc., Huntsville, AL) were used for the P1 (SFRS1) and BAC screenings. PCR-positive clones were picked and cultured in 100 ml of Terrific Broth media or 400 ml of Luria–Bertani media, supplemented with 25 mg/ml of kanamycin (PACs) (Ioannou et al. 1994) or 12.5 mg/ml of chloramphenicol (BACs) (Shizuya et al. 1992). DNA was extracted using a Plasmid Maxiprep (tip 500 kit) (Qiagen).

**Generation of New STSs**

The end fragments of the PAC and BAC clones were
sequenced directly from the clone DNA with SP6 and T7 vector primers at the DNA Sequencing Laboratory of the Institute of Biotechnology, University of Helsinki. New STSs were designed from the nonrepetitive end sequences using Primer3 software (Whitehead Institute). PCRs were done in a 50-µl reaction mix with 50 pmoles of each primer using the annealing temperatures 3–5°C degrees less than \(T_m\). Primer sequences and PCR product sizes for the novel STSs are presented in Supplement 1 at http://www.cshl.org/gr.

**Isolation of Novel CA-Repeats**

Novel CA-repeats were isolated from the PAC and BAC clones as described earlier (Klockars et al. 1996; Ranta et al. 1997). Shortly, PAC DNA was digested with Sau3A1 and ligated to pGEM7, transformed into DH5α cells and plated onto selective agar plates. BAC DNA was digested with Rsal and ligated to Smal-cut pUC18 vector (Pharmacia, Biotech), transformed into Epicurian Coli SURE competent cells (Stratagene), and plated onto selective agar plates. The colonies were transferred onto nylon filters (Hybond-N, Amersham) and hybridized with a radioactively labeled \((GT)_{16}\) oligonucleotide at 65°C overnight. After autoradiography, positive colonies were selected, and DNA was isolated using a Plasmid Miniprep kit (Qiagen) or Wizard Plus Miniprep kit (Promega). The inserts were sequenced with SP6 and T7 vector primers. Oligonucleotide primers flanking the CA-repeats were selected with Primer3 software (Whitehead Institute).

**FISH**

**Targets for FISH**

The metaphase chromosome targets were achieved by culturing human peripheral blood lymphocytes according to standard protocols (Lemieux et al. 1992; Tenhunen et al. 1995). Agarose-embedded lymphocytes were used as a source of free DNA fibers according to the method described earlier (Heiskanen et al. 1994, 1996).

**Probes for FISH**

All clones used for the FISH experiments (PACs and BACs) were labeled with biotin 11-dUTP (Sigma) and digoxigenin 11-dUTP (Boehringer Mannheim) by nick translation according to standard protocols.

**FISH Procedure**

The FISH procedure was performed as described earlier (Pinkel et al. 1986; Lichter et al. 1988; Aaltonen et al. 1997). Briefly, the hybridization mixture contained 50% formamide and 10% dextran sulfate in 2× SSC. Repetitive sequences were suppressed with 10-fold excess of Cot-1 DNA (BRL, Gaithersburg, MD). After overnight incubation at 37°C the slides were washed at 44°C three times in 50% formamide and 2× SSC, twice in 2× SSC, and once in 0.5× SSC. The slides were counterstained with DAPI (Sigma).

**Digital Imaging**

A multicolor image analysis was used for acquisition, display and quantitative analysis of fiber-FISH as described elsewhere (Heiskanen et al. 1996; Aaltonen et al. 1997). In fiber-FISH, the clone sizes determined by PFGE were used as a standard for calibration in each individual fiber-FISH image. The distance measurements were done by applying the distance measurement option of the IPLab software (Signal Analytics Corp., Vienna, VA). In each fiber-FISH image, at least one clone was used as a calibration standard to measure the sizes of other clones, gaps separating different clones, and overlaps between these clones. The known clone sizes enabled compatible and therefore reliable measurements to be made. If two known clone sizes were in the same hybridization, the results of the distance measurements were verified with each other (data not shown).

**Estimation of Clone Insert Sizes by PFGE**

Nol-digested BAC/PAC DNA was separated in 1% agarose using a workshop made pulsed-field gel electrophoresis apparatus designed according to the principles described by Chu et al. (1986). The electrophoretic condition was 120 V, 60-sec pulse time for 48 hr at 16°C. The fragments were visualized by staining the gel with ethidium bromide, and their sizes were estimated by comparing them with the Lambda Ladder (New England BioLabs) size marker.

**Family Material**

Previously, 17 Finnish Meckel families including 25 patients have been described (Paavola et al. 1995). Eight new families with 11 patients were included in this study. The diagnostic criteria for inclusion were as described (Salonen 1984). In MUL, 19 patients from 13 previously described families (Avela et al. 1997) and 8 new families with one affected individual in each were studied. The diagnostic criteria of MUL were as described previously (Lipsanen-Nyman 1986; Lapunzina et al. 1995). DNA was extracted from venous blood or in the case of a deceased patient, from paraffin-embedded autopsy samples using standard procedures (Isola et al. 1994). All the blood samples were drawn in accordance with the Helsinki declaration.

**Genotyping**

Markers D17S957, D17S1853, D17S1606, and D17S1604 originated from the Génethon marker collection (Dib et al. 1996). Marker D17S1290 originated from the CHLC marker collection (http://www.chlc.org). Marker 95-CA was detected in the T7
end sequence of PAC 95i19. Markers 272a-CA, 272b-CA, 132-CA, 58-CA, and 52-CA were discovered by a CA-repeat screening protocol described above. The PCR amplification of the polymorphic microsatellite markers using radiolabeled primers and the size analysis of the PCR products were performed as described earlier (Aaltonen et al. 1993; Sulisalo et al. 1993).

Linkage Disequilibrium Analysis

The significance of linkage disequilibrium between the disease and marker loci was analyzed using the DIS-LAMB program (Terwilliger 1995). This program applies a likelihood-based test for linkage disequilibrium and has only one degree of freedom, irrespective of the number of alleles at any given marker. The calculations are based on the parameter $\lambda$, which expresses the proportion of the increase of a certain allele in the disease chromosomes, relative to its population frequency. Confidence intervals for $\lambda$ were calculated for 1-lod units. The most likely haplotypes were constructed manually, assuming a minimum number of recombinations in each family.

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