Father-to-Daughter Transmission of Focal Dermal Hypoplasia Associated With Nonrandom X-Inactivation: Support for X-Linked Inheritance and Paternal X Chromosome Mosaicism

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Focal dermal hypoplasia (FDH) is a rare syndrome of severe developmental anomalies of the tissues and organs derived from ectoderm and mesoderm. Though data have suggested that FDH is an X-linked dominant trait associated with male hemizygote lethality, a hypothesis supported by the observation of three unrelated infants with FDH manifestations and de novo chromosome rearrangements involving Xp22, observations of father-to-daughter transmission have suggested possible genetic heterogeneity and autosomal dominant inheritance with sex limitation. We hypothesize that, if FDH is an X-linked disorder, cells expressing an active disease locus might experience a selective disadvantage resulting in a nonrandom pattern of X-inactivation in patient tissue. To test this hypothesis, we studied one of the two previously described families demonstrating father-to-daughter inheritance of FDH. To determine if the affected daughter had a skewed pattern of X-inactivation consistent with X-linked inheritance of FDH, somatic cell hybrids were constructed by fusing hypoxanthine phosphoribosyl transferase (HPRT)-deficient rodent fibroblasts with either patient dermal fibroblasts or peripheral white blood cells (WBCs); hybrid clones retaining an active X chromosome were analyzed to determine the parental origin of the active X chromosome. Analyses of resulting hybrid clones showed that while hybrids constructed from skin fibroblasts contained an active X chromosome inherited from either of the patient’s parents, hybrids constructed from WBCs showed a skewed pattern of X-inactivation; 11 of 11 hybrids contained an active maternal X chromosome ($\chi^2 = 12.2, P = .001$). These findings indicated that, in this family, FDH was associated with a nonrandom pattern of X-inactivation consistent with X-linked inheritance, suggesting that the patient’s father was mosaic for a mutant FDH allele.

KEY WORDS: ectodermal dysplasia, somatic cell hybrid, developmental anomalies

INTRODUCTION

Several putatively X-linked diseases have been identified which primarily affect women [Wettkke-Schafer and Kantner, 1983]. The reason for the observed excess of affected women has not been determined in most of these disorders. In some X-linked dominant disorders, the loss of hemizygous males has been well documented; for example, most males with severe ornithine transcarbamylase deficiency die shortly after birth. In other disorders, it has been assumed that the death of hemizygous males occurs during gestation; the observed excess of male spontaneous miscarriages in some kindreds segregating a putative X-linked dominant trait has supported this presumption [Wettkke-Schafer and Kantner, 1983]. However, the observations of similarly affected 46,XY males and kindreds demonstrating father-to-daughter inheritance has suggested that some disorders may be phenotypically or genetically heterogeneous [Goltz et al., 1970].

One disorder that is predominantly limited to women is focal dermal hypoplasia (FDH) or Goltz-Gorlin syndrome (MIM 30560). This rare severe developmental disorder is characterized by focal areas of partial underdevelopment or thinning of the dermis with consequent herniation of subcutaneous tissue [Goltz et al., 1962]. Affected individuals have multiple congenital anomalies of tissues and organs derived from embryonic ecto-

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dermatologic abnormalities including linear hypo- or derm and mesoderm including skeletal defects, eye anomalies such as colobomas and microphthalmia, and dermatologic abnormalities including linear hypo- or hyperpigmented macules, telangiectasias, and papil- lomas [Freire-Maia and Pinheiro, 1984]. Though 95% of FDH cases have been estimated to be nonfamilial, pre- sumably representing new mutations [Fryns et al., 1978], two alternative modes of inheritance have been postulated. It has been proposed that FDH is an X-linked dominant trait with male hemizygote lethality because most affected individuals are women, affected women experience an increased incidence of sponta- neous midgestational miscarriages of male fetuses [Goltz et al., 1962], and four pedigrees have demon- strated mother-to-daughter transmission [Freeman, 1955; Wodiansky, 1957; Ginsburg et al., 1970; Ruiz- Maldonado et al., 1974]. The observation of three unre- lated females with manifestations suggestive of FDH and de novo cytologic abnormalities involving region Xp22 [Al-Gazali et al., 1990; Temple et al., 1990a] pro- vides additional support that FDH is X-linked. Alter- natively, an autosomal dominant sex-limited mode of inheritance was proposed [Goltz et al., 1970] based on the observations of multiple affected males and the ob- servation of two pedigrees with father-to-daughter in- heritance [Larregue et al., 1971; Burgdorf et al., 1981]; however, these observations did not exclude the possi- bility that the affected males were mosaic for FDH [Tem- ple et al., 1990b].

Several X-linked disorders have been identified in which carrier women demonstrate a nonrandom pattern of X chromosome inactivation, presumably on the basis of a selective disadvantage to a subpopulation of cells with an expressed disease allele [Hall, 1988]. We hypothesize that, if FDH is an X-linked disorder, cells expressing an active disease locus might experience a selective disadvantage resulting in a nonrandom pat- tern of X-inactivation in heterozygote tissue. To test this hypothesis, we studied one of the two previously de- scribed families with demonstrated father-to-daughter inheritance of FDH [Burgdorf et al., 1981]. We now report our results which demonstrate that, in this family, FDH is associated with a nonrandom pattern of X-inactivation consistent with X-linked dominant in- heritance and paternal X chromosome mosaicism for FDH.

MATERIALS AND METHODS

Cell Lines

Dermal explants were dissected from sterile skin bi- opsy samples and cultured in standard conditions. Per- ipheral white blood cells (WBCs) were biopsied by veni- puncture in heparinized tubes and separated from other cellular constituents by ficoll gradient centrifugation (LSM, Organon Teknika, Durham NC). Hypoxanthine phosphoribosyltransferase (HPRT)-deficient Chinese hamster fibroblasts, cell line RJK-88 [Fuscoe et al., 1983], were fused with human peripheral WBCs, and HPRT-deficient murine fibroblasts, cell line M613 [Kabara et al., 1974], were fused with human skin fibroblasts using polyethylene glycol as previously de- scribed [Gorski et al., 1989]. Independent somatic cell hybrid clones, one per plate, were isolated and main- tained in DMEM media supplemented with 15% fetal bovine serum, penicillin-streptomycin (100 μg/ml), 4 mM glutamine, HAT (10^{-4} M hypoxanthine, 10^{-5} M amethopterin, 10^{-2} M thymidine), and 1 μM ouabain.

DNA Isolation, Digestion, Blotting, and Hybridization

DNA was isolated from cells by phenol/chloroform extraction of sodium dodecylsulfate (SDS) and proteinase K-treated cell lysates followed by ammonium acetate/ethanol precipitation [Hermann and Frischau, 1987]. Restriction endonuclease digestions were performed by digesting 10 μg aliquots of DNA to completion with a restriction enzyme per manufacturer's suggestions (Bethesda Research Laboratories, Bethesda, MD), as judged by minigel agarose electrophoresis. Digested DNA was fractionated by agarose electrophoresis, and depurinated DNA was transferred to Zeta-probe nylon membranes (BioRad, Richmond, CA) by alkaline trans- fer [Reed and Mann, 1985]. Radiolabeled probes [Feinberg and Vogelstein, 1983] were hybridized to the immo- bilized DNA at 65°C in 1.5 × SSPE [Maniatis et al., 1982], 0.5% nonfat dry milk, 1% SDS, and 100 μg/ml denatured salmon sperm DNA. Filters were washed at a final stringency of 0.1 × SSC [Maniatis et al., 1982], 1% SDS at 65°C, and exposed to Kodak XAR-5 film with an intensifying screen for 3–72 hours.

CLINICAL REPORTS

Patient 1

The proposita had multiple skeletal and dermatologic anomalies at birth. She was the 2,800 g product of a normal uneventful 40 week gestation to a 33-year-old woman who had one previous pregnancy resulting in a normal male infant. There was no known exposure to recognized teratogens. Congenital anomalies included a coloboma of the right iris and retina, a hypoplastic right nasal ala, perioral papillomas, right hemiatrophy, mul- tiple telangiectasias, linear hyperpigmented macules, and linear streaks of dermal thinning with focal hernia- tions of subcutaneous tissue. Her right hand had a single mobile digit with a normal nail and a second small flaccid digit located radially. Both feet had deep clefts. Medial to the cleft, both feet had normal first digits; lateral to the clefts, the right foot had two nailless sym- dactylyous digits and the left had two digits curved plan- tarward. Roentgenographic examinations showed a hypo- plastic right clavicle and osteopathia striata of the long bones; histologic examination of an atrophic macule showed normal stratified squamous epithelia with under- lying adipose tissue and no discernible dermis [Burg- dorf et al., 1981].

Her major motor development was normal, and at 12 years, she was in an age-appropriate class. Her major cutaneous defects have required numerous skin-graft- ing procedures to repair superficial ulcerations over atrophic regions (Fig. 1). She had a surgical revision of her right lacrimal duct and recurrent bilateral otitis media resulting in bilateral conductive hearing loss and necessitating bilateral tympanostomies and surgical os- sicular chain repair. She had hypodontia; teeth were
Fig. 1. Patient 1 at age 20 months (A) and the dorsal surface of her legs at age 14 years (B). Focal hypotrichosis, right inferiornasal coloboma, labial papillomas, right upper limb ectrodactyly, linear dermal atrophy in associated herniations of subcutaneous tissue, and linear hyperpigmented macules are shown.

dysplastic with hypoplastic enamel and were delayed in eruption. She had an atraumatic fracture of the left tibia and a surgically reduced dislocated of the right radial head.

Examination at 12 years showed all growth parameters below the 3rd centile with an occipitofrontal circumference (OFC) of 49 cm, height of 129.5 cm, and weight of 23.6 kg. Additional findings included fine sparse hair with focal alopecia, right facial hemiatrophy, a highly arched palate, thoracolumbar scoliosis, and a narrow anterior-posterior thoracic diameter. Prometaphase chromosomal analysis showed a 46,XX chromosomal constitution.

Patient 2

The father of Patient 1, a 54-year-old man of normal proportions, had linear telanectasias, atrophic dermal depressions, and streaked hyperpigmented macules of his right chest, abdomen, and leg recognized during infancy. The herniation of subcutaneous tissue beneath the atrophic lesions of his right axilla and inguinal region developed later in life (Fig. 2). When he was 14, he had a traumatic fractured left femoral epiphysis and right humerus. Roentgenologic examination failed to show osteopathia striata of his long bones; histologic examination of an atrophic macule from the forearm showed an absence of papillary dermis with fatty replacement [Burgdorf et al., 1981]. Physical examinations failed to show other stigmata of FDH; hair and teeth were normal, and his nails showed mild linear streaking which, though consistent with FDH, may have been acquired. He had a normal prometaphase karyotype showing a 46,XY chromosomal constitution.

RESULTS

To distinguish the proposita’s paternally and maternally derived X chromosomes, DNAs isolated from peripheral WBCs of the propositus (Patient 1), her mother, and her father (Patient 2) were analyzed using oli-
goradiolabeled DNA probes detecting X chromosome restriction fragment length polymorphisms. Analyses showed that both the propositus and her mother were heterozygous for the DXS14 MspI polymorphism detected with probe p58-1. DNA samples from the propositus and her mother both contained the 3.8 and 2.4 kilobase (kb) alleles; the DNA of the propositus's father contained only the 3.8 kb allele, suggesting that the propositus inherited the 2.4 kb allele from her mother and the 3.8 kb from her father (Fig. 3).

In order to determine the parental origin of the active X chromosome in cells isolated from the propositus (skin fibroblasts isolated from an atrophic lesion and peripheral WBCs), somatic cell hybrids were constructed; biopsied cells were fused with HGPRT-deficient rodent fibroblast cell lines using polyethylene glycol. Somatic and the 3.8 kb from her father (Fig. 3). DNA samples from the proposita's dermal fibroblasts containing both the paternal 2.4 kb allele (hybrids SA3 and SA4) and the maternal 3.8 kb allele (hybrid SC5); these results were consistent with a random pattern of X-inactivation within the proposita's fibroblasts. Most of the isolated hybrid clones were stable upon passage; DNA was isolated from 14 different hybrids derived from venous WBCs.

To determine the parental origin of the active X chromosome retained in each of the resultant hybrid clones, hybrid DNA was digested with MspI and hybridized with probe p58-1 to identify the DXS14 parental allele present (Fig. 3). Somatic cell hybrids derived from the proposita's dermal fibroblasts contained both the maternal 2.4 kb allele (hybrids SA3 and SA4) and the paternal 3.8 kb allele (hybrid SC5); these results were consistent with a random pattern of X-inactivation within the proposita's fibroblasts. Most of the isolated hybrid clones were stable upon passage; DNA was isolated from 14 different hybrids derived from venous WBCs.

DISCUSSION

These analyses sought to answer two interrelated questions regarding the genetics and biology of FDH: 1) Is FDH X-linked? and if so, 2) Are cells expressing a mutant FDH allele at a selective disadvantage? To address these questions, we used recombinant-DNA probes to study a family demonstrating father-to-daughter inheritance of FDH and analyzed somatic cell hybrids retaining an active X chromosome derived from the affected daughter. Though an apparently random pattern of X inactivation in hybrids was observed in hybrids constructed from dermal fibroblasts, preferential X inactivation of the paternal X chromosome was observed in hybrids derived from the proposita's WBCs. Within this family, these findings are consistent with the hypothesis that FDH was inherited as an X-linked dominant trait and that the proposita's father was mosaic for the mutant FDH gene. These findings suggest that the expression of a mutant FDH allele was associated with a selective disadvantage in hematopoietic cells. The mild
clinical manifestations of the affected father are consistent with mosaicism and are similar to the mild changes observed in males mosaic for other X-linked, hemizygote lethal disorders such as ornithine transcarbamylase deficiency [Maddalena et al., 1988; Legius et al., 1990] and X-linked agammaglobulinemia [Hendriks et al., 1989].

Significant evidence indicates that, within the cells of an inner cell mass, X chromosome inactivation is initially random, resulting in a mosaic population of cells; however, the proportion of the two mosaic populations may be subsequently skewed as a result of selection for alleles at loci influencing cell proliferation [Chapman, 1986]. In heterozygous women, the absence and presumed loss of cells expressing a mutant X-chromosomal allele is well documented: B cells expressing a mutant allele are absent from carriers of X-linked agammaglobulinemia [Conley et al., 1987; Fearon et al., 1987], abnormal platelets are apparently lost from Wiskott-Aldrich carriers [Gealy et al., 1980; Prchal et al., 1980], dermal fibroblasts expressing a mutant incontinentia pigmenti allele are underrepresented [Wieacker et al., 1985; Migeon et al., 1989], and HPRT-deficient hematopoietic cells are absent in carriers of Lesch-Nyhan syndrome [Albertini and DeMars, 1974].

In addition to providing a means to detect heterozygous women, these observations have assisted in providing information regarding the nature of the mutant gene product, the tissue in which the mutant gene is expressed, and the developmental pattern of gene expression and regulation. However, the demonstration of a nonrandom pattern of X-inactivation has not necessarily provided a plausible explanation for a disease phenotype. For example, among heterozygotes, while the demonstrated deficiency of B cells expressing a mutant agammaglobulinemia allele corresponds to the disease phenotype of the affected hemizygote [Conley et al., 1987; Fearon et al., 1987], the absence of hematopoietic cells expressing a mutant HPRT in Lesch-Nyhan carriers does not correspond to the observed phenotype or play an obvious causative role in disease pathogenesis [Albertini and DeMars, 1974].

Our analyses of a family demonstrating father-to-daughter transmission of FDH showed that, in somatic cell hybrids derived from the proposita's WBCs, the maternally derived X chromosome was preferentially retained. These findings suggest that mutations at the FDH locus may be detrimental to the proliferation of hematopoietic cells expressing the mutation, resulting in an underrepresentation of WBCs containing an active paternally derived X chromosome. Presumably, the loss of WBCs or precursors containing an active paternally derived X chromosome resulted from cell selection following random X-inactivation. Because X chromosome inactivation patterns in females follow a binomial distribution curve [Fialkow, 1975], occasional normal females are expected to have apparently nonrandom patterns of X-inactivation by chance alone. However, finding such a significantly skewed pattern of X-inactivation by chance, with selection against the paternally derived X chromosome in particular, is unlikely and provides new evidence that FDH is an X-linked disorder.

The observation that WBCs, but not dermal fibroblasts, demonstrate a nonrandom pattern of X-inactivation suggests that in FDH, like HPRT deficiency [Albertini and DeMars, 1974], different cell types exhibit different tolerances toward a mutant disease locus. The apparent absence of nonrandom X-inactivation in dermal fibroblasts suggests that, unlike incontinentia pigmenti in which observed dermatologic anomalies have been proposed to be secondary to the loss of cells expressing a mutant allele [Wieacker et al., 1985; Migeon et al., 1989], the dermatologic anomalies observed in FDH may be associated with cells expressing a mutant FDH allele. Resolution of this issue will require the analysis of additional FDH patients and multiple tissues from affected and nonaffected areas.

We conclude that the nonrandom pattern of X-inactivation observed in the hematopoietic cells of the propositus of this family segregating FDH suggests, but does not prove, the existence of an X-linked locus for this disease. These analyses cannot rule out potential genetic heterogeneity, such as that recently demonstrated for Aicardi syndrome [Neidich et al., 1990]; further analyses with additional patients will be required to verify these findings. The potential application of such analyses to carrier detection and prenatal diagnosis will be dependent upon verification.

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