
**Cutaneous mosaicism: right before our eyes**

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Autosomal recessive cutaneous disorders, including various types of epidermolysis bullosa (EB), usually manifest shortly after birth. The clinical course of these diseases is often characterized by severe complications, limited therapeutic options, and a poor prognosis. A study by Pasmooj et al. reported in this issue of the *JCI* unravels the molecular mechanisms by which germline mutations in non-Herlitz junctional EB can be corrected in vivo by multiple spontaneously occurring somatic mutational events, a phenomenon known as revertant mosaicism (see the related article beginning on page 1240). These insights open new avenues of thinking for the design of future gene therapy strategies for skin diseases.

“I see a bright future for genetics in dermatology and for dermatology in genetics . . . ” — this was the vision of Victor A. McKusick, widely regarded as one of the founding fathers of medical genetics, some 35 years ago (1). Today, it is justified to say that he was absolutely right. During the past 20 years, the genetic basis of more than 350 monogenic skin diseases has been elucidated (2), and the number of disorders awaiting discovery of the underlying gene defect is steadily increasing. However, research in the field of inherited cutaneous disorders is by no means limited to finding novel genes and mutations. Because the skin is easily accessible and, for the trained observer, everything lies “right before your eyes,” genetic skin diseases offer unique opportunities to study complex clinical expression patterns and uncommon inheritance modes and the ability to explore molecular therapeutic regimens such as enzyme replacement strategies and gene therapy (2, 3). This notion holds particularly true for cutaneous disorders that clinically present with mosaicism.

**Mosaicism**

The term mosaicism refers to the occurrence in an individual of two or more cell populations that are karyotypically or genotypically different and yet are derived from a single zygote (4). Mosaicism can result from a mutation during development that is propagated in only a limited number of the adult cells. In general, any type of cell may be affected by such a mutational event, including gametes (egg and sperm cells), blood cells, and skin cells. The best-known example of mosaicism is mammalian females (normal karyotype 46,XX), which are functional mosaics because one of their X chromosomes is randomly inactivated during embryogenesis (5).

In autosomal recessive mendelian disorders, the underlying mutations often cause embryonic or early postnatal death. Notwithstanding, studying the effects of such mutations at later developmental stages may be possible in individuals carrying the mutation in a mosaic form because gain or loss of genetic functions might be limited to specific cells and tissues or to selected stages of development, as recently shown in various organisms, including, for example, *Caenorhabditis elegans*, *Drosophila*, mice, and zebrafish (6–9). Therefore, mosaic organisms are perfectly suited for the investigation of the molecular processes orchestrating very early developmental phases. Further, they may be of use to study a specific cell type or tissue in which a given gene is required to assure proper functioning of signaling pathways and metabolic processes. With respect to organ systems, mosaic organisms can serve to determine whether a particular gene is cell autonomous, i.e., whether the gene is exerting its action exclusively within the cell in which it is expressed or if it also affects neighboring cells that do not manifest a phenotype themselves when carrying a mutation in that gene (8). In cutaneous mosaicism, the underlying mutations often cause embryonic or early postnatal death. Notwithstanding, studying the effects of such mutations at later developmental stages may be possible in individuals carrying the mutation in a mosaic form because gain or loss of genetic functions might be limited to specific cells and tissues or to selected stages of development, as recently shown in various organisms, including, for example, *Caenorhabditis elegans*, *Drosophila*, mice, and zebrafish (6–9). Therefore, mosaic organisms are perfectly suited for the investigation of the molecular processes orchestrating very early developmental phases. Further, they may be of use to study a specific cell type or tissue in which a given gene is required to assure proper functioning of signaling pathways and metabolic processes. With respect to organ systems, mosaic organisms can serve to determine whether a particular gene is cell autonomous, i.e., whether the gene is exerting its action exclusively within the cell in which it is expressed or if it also affects neighboring cells that do not manifest a phenotype themselves when carrying a mutation in that gene (8). In cutaneous mosaicism, the underlying mutations often cause embryonic or early postnatal death. Notwithstanding, studying the effects of such mutations at later developmental stages may be possible in individuals carrying the mutation in a mosaic form because gain or loss of genetic functions might be limited to specific cells and tissues or to selected stages of development, as recently shown in various organisms, including, for example, *Caenorhabditis elegans*, *Drosophila*, mice, and zebrafish (6–9). Therefore, mosaic organisms are perfectly suited for the investigation of the molecular processes orchestrating very early developmental phases. Further, they may be of use to study a specific cell type or tissue in which a given gene is required to assure proper functioning of signaling pathways and metabolic processes. With respect to organ systems, mosaic organisms can serve to determine whether a particular gene is cell autonomous, i.e., whether the gene is exerting its action exclusively within the cell in which it is expressed or if it also affects neighboring cells that do not manifest a phenotype themselves when carrying a mutation in that gene (8).

**Mosaicism in monogenic skin disorders**

Mosaic skin diseases may show different patterns of clinical involvement such as lines of Blaschko, a checkerboard pattern, a phylloid pattern, a patchy pattern without midline separation, and a lateralization pattern.

**Nonstandard abbreviations used:** EB, epidermolysis bullosa; LAMBS, laminin β3; LOH, loss of heterozygosity.

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Mosaicism has been demonstrated at the cellular level in more than 15 monogenic skin disorders to date (10). Recently, the specific molecular mechanisms governing three particular forms of cutaneous mosaicism have been unraveled and reported in the JCI.

In 2004, we showed at the cellular and molecular level that the type 2 segmental manifestation of Hailey-Hailey disease, being superimposed on the ordinary nonsegmental lesions of this autosomal dominant trait, results from postzygotic loss of heterozygosity (LOH) (11). These data provided the first molecular evidence supporting this genetic concept, which was postulated ten years ago (12). If LOH results from postzygotic crossing-over, this would give rise to two different daughter cells, one of them being homozygous for the underlying mutation. Conversely, the other cell would be homozygous for the wild-type allele, resulting in a band or patch of completely healthy tissue (13). This would reflect “natural gene therapy” in an autosomal dominant skin disease in a way analogous to that proposed by Pasmooij et al. in this issue of the JCI (14) for autosomal recessive cutaneous traits. After ex vivo expansion of cells from such healthy skin areas, cell sheets could be generated and used for autologous transplantation.

More recently, the molecular basis of linear epidermal nevi of the common, nonorganoid and nonepidermolytic type was also reported in the JCI (15). These benign keratinocytic skin lesions are present at birth but become more conspicuous during early childhood. Using a SNaPshot Multiplex assay, Hafner et al. detected activating somatic mutations in the FGF receptor 3 (FGFR3) gene in about one-third of the nevi studied (15). These findings could provide a basis for the future development of noninvasive therapeutic strategies for skin tumors, such as, for example, siRNA specific to the mutant allele (16). A significant therapeutic benefit of RNA interference in vivo has recently been demonstrated for amyotrophic lateral sclerosis (17).

**Figure 1**

Schematic representation of postzygotic mechanisms that may give rise to revertant mosaicism in an individual compound heterozygous for an autosomal recessive phenotype. Two corresponding chromosomes carry either the paternal (blue) or maternal (pink) mutation. Various mutational events may result in cells showing loss of compound heterozygosity (bottom row). As a consequence, these cells will produce either a functional protein (black boxes), a semifunctional protein (gray box), or no protein at all (white boxes). Reproduced with permission from the American Journal of Medical Genetics (19).
governing this kind of naturally occurring phenotypic rescue have been proposed and demonstrated, including true reverse point mutations, deletions, nondisjunction, crossing-over events, and gene conversion (Figure 1) (19). Moreover, revertant mosaicism can be induced by transposable elements known as retrotransposons (20) as documented in Duchenne muscular dystrophy (21, 22).

In 1997, Jonkman et al. were the first to demonstrate at the molecular level the occurrence of revertant mosaicism in human skin in a patient suffering from generalized atrophic benign epidermolysis bullosa (EB), who showed some healthy patches of skin in which blistering could not be evoked (23). The authors proved that within these areas, phenotypic reversion was caused by mitotic gene conversion in one of the two mutated collagen type XVII α1 (COL17A1) alleles. Based on these data, they suggested that revertant mosaicism in autosomal recessive skin diseases is as an example of natural gene therapy.

Revertant mosaicism in EB: guiding the path toward gene therapy

In this issue of the JCI, Pasmooij et al. (14) convincingly demonstrate that several distinct revertant mutational events can occur in one individual at different time points in life, all contributing to amelioration of the disease phenotype by different mechanisms (14). This notion is supported by recent observations in other hereditary diseases such as tyrosinemia type I (24) and Omenn syndrome (25).

The two patients presented here had autosomal recessive non-Herlitz junctional EB caused by compound heterozygous or homozygous germline mutations in the laminin β3 (LAMB3) gene, respectively (14). Molecular analysis of biopsy specimens derived from areas of healthy-appearing skin led to the identification of five different correcting somatic mutations that predominantly affected mRNA splicing. Interestingly, one of the patients studied by these authors revealed expansion of the healthy skin regions harboring a spontaneously occurring correcting second-site mutation. It is tempting to speculate that some of the keratinocytes in which revertant mutational events occurred must have been epidermal stem cells. Over time, clonal expansion of these stem cells led to amelioration of the phenotype within a defined area of skin due to a selection advantage of revertant stem cells compared with their deficient counterparts (14).

These observations have far-reaching consequences for possible future strategies of gene therapy for autosomal recessive cutaneous disorders. In mosaic individuals, autologous skin grafts derived from areas with normal-appearing skin could be transplanted to affected dermepidermized skin regions on the same patient, thereby following the example of natural gene therapy and avoiding immunologic reactions that might cause graft rejection. Pasmooij et al. (14) conclude that in LAMB3 revertant mosaicism, one might expand in vitro the patient’s own revertant cells and use such naturally corrected cells for grafting.

Gene therapy in EB: where are we now?

When designing gene therapy approaches as a future method for causal treatment of severe inherited cutaneous diseases some 10–15 years ago, investigators convinced of the benefits of this therapeutic regimen were often confronted with skepticism and concerns. Today, gene therapy for EB is no longer wishful thinking but is at the verge of being introduced into clinical practice.

Very recently, Mavilio et al. (26) reported the first successful ex vivo gene therapy approach in a patient suffering from LAM5-β3–deficient junctional EB, a disorder that is severe and often lethal due to dysfunctional skin adhesion, manifesting with extensive blistering at birth and serious complications such as recurrent infections. In a phase I/II clinical trial, epidermal stem cells from this patient were transduced with a retroviral vector expressing LAMB3 cDNA (encoding LAM5-β3) and used to create genetically corrected epidermal grafts. After surgical preparation, these grafts were transplanted onto the patient’s legs, resulting in development of fully functional and adherent epidermis. One year later, the grafts were stable and did not show any blistering, indicating that this therapeutic approach was curative (26).

This encouraging report (26) is just the beginning of a new era in which laboratory researchers and clinicians will intensify their efforts to develop and improve strategies of gene therapy for potentially fatal skin diseases. The concept of revertant mosaicism as presented in this issue by Pasmooij et al. (14) will be an important contribution toward this goal. In particular, the discovery that postzygotic mutations giving rise to revertant mosaicism occur far more frequently than so far assumed may serve as a starting point to develop new strategies of gene therapy that we are sorely missing in our clinical practice today.

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Diabetic foot ulcers (DFUs), a leading cause of amputations, affect 15% of people with diabetes. A series of multiple mechanisms, including decreased cell and growth factor response, lead to diminished peripheral blood flow and decreased local angiogenesis, all of which can contribute to lack of healing in persons with DFUs. In this issue of the JCI, Gallagher and colleagues demonstrate that in diabetic mice, hyperoxia enhances the mobilization of circulating endothelial progenitor cells (EPCs) from the bone marrow to the peripheral circulation (see the related article beginning on page 1249). Local injection of the chemokine stromal cell–derived factor–1α then recruits these EPCs to the cutaneous wound site, resulting in accelerated wound healing. Thus, Gallagher et al. have identified novel potential targets for therapeutic intervention in diabetic wound healing.

Despite the existence of protocols to standardize care, the physiological impairments that can result in a DFU complicate the healing process. Currently, the only FDA-approved growth factor and cell therapies for DFUs are not routinely used during treatment, preventing professionals from implementing evidence-based protocols (5).

Molecular pathogenesis of diabetic wound healing

The moment a person with diabetes suffers a break in the skin of their foot, they become at danger for amputation. Most commonly, patients have neuropathy, which could be causative. When coupled with an impaired ability to fight infection, these patients become largely unable to mount an adequate inflammatory response. Thus, the DFU that may look like a healing wound becomes a portal for infection that can lead to sepsis and require limb amputation.

Over 100 known physiologic factors contribute to wound healing deficiencies in individuals with diabetes (Figure 1). These include decreased or impaired growth factor production (6–8), angiogenic response (8, 9), macrophage function (10), collagen accumulation, epidermal barrier function, quantity of granulation tissue (8), keratinocyte and fibroblast migration and proliferation, number of epidermal nerves (11), bone healing, and balance between the accumulation of ECM components and their remodeling by MMPs (12). Wound healing occurs as a cellular response to injury and involves activation of keratinocytes, fibroblasts, endothelial cells, macrophages, and platelets. Many growth factors and cytokines released by these cell types are needed to coordinate and maintain healing.

Molecular analyses of biopsies from the epidermis of patients have identified pathogenic markers that correlate with delayed wound healing. These include overexpression of c-myc and nuclear localization of β-catenin (13). Coupled with a reduction in and abnormal localization of EGFR and activation of the glucocorticoid pathway, keratinocyte migration is inhibited (13, 14). At the nonhealing edge (callus) of DFUs, keratinocytes show an absence of migration, hyperproliferation, and incomplete differentiation (13, 14). Fibroblasts demonstrate a phenotypic change as well as decreased migration and proliferation.

Cellular and molecular basis of wound healing in diabetes

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Diabetes affects approximately 170 million people worldwide, including 20.8 million in the USA (1), and by 2030 these numbers are projected to double (2). The foot ulcer is a leading cause of hospital admissions for people with diabetes in the developed world (3) and is a major morbidity associated with diabetes, often leading to pain, suffering, and a poor quality of life for patients. Diabetic foot ulcers (DFUs) are estimated to occur in 15% of all patients with diabetes (3) and precede 84% of all diabetes-related lower-leg amputations (4).

Nonstandard abbreviations used: DFU, diabetic foot ulcer; EPC, endothelial progenitor cell; HBO, hyperbaric oxygen therapy; SDF–1α, stromal cell–derived factor–1α.

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1219