Abnormal Epidermal Keratinization in the Repeated Epilation Mutant Mouse

KAREN A. HOLBROOK, BEVERLY A. DALE, and KENNETH S. BROWN
Departments of Biological Structure, Medicine (Dermatology), and Periodontics, University of Washington, Seattle, Washington 98195, and the Laboratory of Developmental Biology and Anomalies, National Institute of Dental Research, National Institutes of Health, Bethesda, Maryland 20205

ABSTRACT Repeated epilation (Er) is a radiation-induced, autosomal, incomplete dominant mutation in mice which is expressed in heterozygotes but is lethal in the homozygous condition. Many effects of the mutation occur in skin: the epidermis in Er/Er mice is adhesive (oral and nasal orifices fuse, limbs adhere to the body wall), hyperplastic, and fails to undergo terminal differentiation. Skin from fetal +/+ , Er/+ and Er/Er mice at ages pre- and postkeratinization examined by light, scanning, and transmission electron microscopy showed marked abnormalities in tissue architecture, differentiation, and cell structure; light and dark basal epidermal cells were separated by wide intercellular spaces, joined by few desmosomes, and contained phagolysosomes. The numbers of spinous, granular, and superficial layers were highly variable within any given region and among various regions of the body. In some areas, 2–8 layers of granular cells, containing large or diminutive keratohyalin granules, extended to the epidermal surface; in others, the granular layers were covered by several layers of partially keratinized or nonkeratinized cells. In rare instances, a single or small group of cornified cells was present among the granular layers but was not associated with the epidermal surface. Both the granular and nonkeratinized/partially keratinized upper epidermal layers in Er/Er skin gave positive immunofluorescence with antisera to the histidine-rich, basic protein, filagrin.

Proteins in epidermal extracts from +/+ , Er/+ and Er/Er mice were separated and identified by radio- and immunolabeling techniques. The Er/Er extract was missing a 26.5-kdalton protein and had an altered ratio of bands in the keratin region. The 26.5-kdalton band was histidine-rich and cross-reacted with the antisera to rat filagrin. Several high molecular weight bands present in both Er/Er and +/+ extracts also reacted with the antisera. These are presumed to be the precursors of filagrin and to account for the immunofluorescence in Er/Er epidermis even though the product protein is absent. The morphologic and biochemical data indicated that the genetic defect has a general and profound influence on epidermal differentiation, including alteration of two proteins (filagrin and keratin) important in normal terminal differentiation, tissue architecture, and cytology. Identification of epidermal abnormalities at early stages of development (prekeratinization) and defective structure of other tissues and gross anatomy suggest that the mutation is responsible for a defect in some regulatory step important in many processes of differentiation and development.

The repeated epilation mutation is thought to have been induced by γ-irradiation (19) of the Er gene on chromosome 4 (12, 16). The condition is inherited as an autosomal, incomplete dominant and is named for the pattern of hair loss and regrowth that occurs in viable, fertile Er/+ adults (19). The mutant Er/Er animals die in utero or moments after birth. The oral cavity is fused, giving the impression of no mouth (17) and, although they make respiratory movements at birth, the external nares are sealed or are limited to pinhole-sized openings. There are marked skeletal abnormalities of the limbs and in the facial region; the paws and tail are stunted and all appendages adhere to the body. The palate is cleft and fused with the tongue. The
Er/Er mice have an unusual, tight, thickened skin that was originally described as lacking granular and cornified layers (17).

In contrast with the observations of Guenet et al. (17), our preliminary histologic examination of Er/Er epidermis revealed hyperplastic (instead of absent) granular layers and prominent keratohyalin granules in granular cells, yet an almost complete absence of keratinization in more superficial cells, cells which normally form the keratinized stratum corneum. The purpose of the present investigation was to extend these observations and to define the morphologic and biochemical properties of the epidermis in fetal and newborn +/+ , Er/+ and Er/Er mice, specifically, to determine (a) what effect the mutation has on epidermal architecture, cytology, and pattern of differentiation, (b) whether the defect is expressed before keratinization or simultaneous with the onset of keratinization, and (c) whether, and how, either or both of the two major proteins (filaggrin and keratin) associated with keratinization are affected (3, 7, 15, 33).

MATERIALS AND METHODS

Animals

Er/+ breeding stock was obtained from Jackson Laboratories, Bar Harbor, Maine. The background genotype was C57Bl/6 ByEi. Animals were maintained in an air-conditioned, windowless room with 12-12 light cycle and fed NIH-007 diet and water ad libitum. At age 10 wk, the Er/+ animals were distinguished from normal by sparse hair and regions of hairless skin. Matings of two Er/+ animals were confirmed by observation of a vaginal plug on the morning of day zero of gestation. The mothers were killed on the designated day of gestation and the uterus was quickly excised. The embryos or fetuses were dissected into saline under a microscope and classified as to type. At day 13, only two phenotypes were separable, Er/Er and others combining Er/+ and +/- genotypes. By day 18 or 19, three phenotypes were clearly recognizable; Er/+ animals had edema of the feet and a hemorrhagic tail tip; the latter persists one day after birth. Most Er/Er mice were alive at birth and active in their nest boxes. The maternal littermates were usually killed at birth and dissected. The uterus was quickly excised. The embryos or fetuses were dissected into saline and blotted on filter paper. Sections were stained with saturated uranyl acetate and lead hydroxide, dried flat on slides, and viewed under a microscope and classified as to type. At day 13, only two phenotypes were separable, Er/Er and others combining Er/+ and +/- genotypes. By day 18 or 19, three phenotypes were clearly recognizable; Er/+ animals had edema of the feet and a hemorrhagic tail tip; the latter persists one day after birth. Most Er/Er mice were alive at birth and active in their nest boxes. The maternal littermates were usually killed at birth and dissected. The uterus was quickly excised. The embryos or fetuses were dissected into saline and blotted on filter paper.

Histology and Ultrastructure

Animals removed from the uteri of 13-, 18-, and 19-d pregnant females were immediately immersed in half-strength Karnovsky's fixative (22) buffered in 0.1 M cacodylate buffer and fixed for several hours. Small segments of the head and dorsal and lateral body wall from one-half of several animals of each genotype were dissected for histologic and transmission electron microscopy (TEM) studies. The remaining half of each animal was photographed from internal and external aspects, then processed for scanning electron microscopy (SEM). All samples were washed several times in buffer, postfixed in 2% OsO4 in distilled water, and dehydrated through a graded series of alcohol into 100% ethanol and (EiOH). Samples for light microscopy (LM) and TEM were embedded in Epon by conventional procedures (27) and cut into 1-μm (LM) and 300-nm (TEM) sections. The 1-μm sections were examined by conventional and periodic acid-Schiff (PAS) procedures. Thin sections were stained with saturated uranyl acetate and lead citrate and viewed in a Phillips 201 transmission electron microscope. Specimens for TEM were dehydrated further through a graded EiOH-Geneveis series into 100% Geneveis, then critical-point-dried in the pressure chamber of a Bomar Critical Processor. Thin sections were stained with saturated uranyl acetate and lead hydroxide and viewed in a Hitachi H-800 electron microscope operated at 80 kV in the secondary electron mode.

Immunofluorescence

Samples from 19-d fetal mice of the three genotypes were covered with medium for cryostat sectioning and frozen in isopentane in liquid nitrogen. 8-μm sections from the frozen blocks were cut on a cryostat, fixed for 10 min in acetone at -20°C, washed with phosphate-buffered saline (PBS), preincubated with normal goat serum for 30 min, washed in PBS, then reacted with the IgG fraction of rabbit antiserum against rat epidermal stratum corneum-derived histidine-rich protein, filaggrin, for 30 min at 4°C. Sections were washed in PBS and incubated with fluorescein-conjugated goat anti-rabbit IgG for 30 min at 4°C, washed again, and mounted in 90% glycerol at pH 9. They were studied with a Zeiss WL research photomicroscope with fluorescent excitation of 440-490 nm and barrier filter of 520 nm.

Extraction and Separation of Epidermal Proteins

The skin from newborn mice was incubated in 5 mM EDTA in PBS for 3 min at 52°C to separate dermis and epidermis (modified from Harris et al. [18]). The epidermis was homogenized in 8 M urea containing 0.1 M Tris-HCl, pH 7.5, 0.1 M 2-Mercaptoethanol, 1 mM diithiothreitol, and 20 μg/ml phenylmethylsulfonyl fluoride (PMSF). Samples were stored for 2-4 h and then centrifuged at 20,000 g to remove the insoluble material.

Extracted proteins were separated by SDS PAGE using the method of Laemmli (24). Protein samples were incubated at 100°C for 2-5 min in buffer containing 1% SDS and 1% 2-Mercaptoethanol. Gels were stained with Coomassie Brilliant Blue, destained, and photographed. Protein concentration was determined by the Bio-Rad technique (Bio-Rad Laboratories, Richmond, Calif.) using chymotrypsinogen as a standard.

Radiolabeling Studies In Vivo

25 μCi of [3H]histidine (339 mCi/mmol) was injected intraperitoneally into newborn or 2-d-old +/- and Er/+ mice. Some of the animals were killed at 2 h and others at 24 h and the skins were removed. The epidermal proteins were extracted, separated by electrophoresis, and identified by fluorography. Gels were processed for fluorography using En3Hance® (New England Nuclear, Boston, Mass.); X-OmatAR film was exposed according to Laskey and Mills (25).

Immunologic Detection of Antigen

Detection of antigens (histidine-rich proteins: filaggrin) on SDS gels was done by the method of Renart et al. (31). Samples of epidermal extracts were electrophoresed on gels containing N,N'-diallyltartardiamide (DATT) cross-linker. The gel was then sandwiched between two sheets of diazobenzoylomethyl (DBM) paper (prepared according to manufacturer's instructions [Schleicher & Schuell, Inc., Keene, NH]) for transfer of proteins. The paper containing the transferred proteins was incubated with antiserum to rat epidermal filaggrin or preimmune control serum, each diluted 1:50 in buffer 1 (31), washed with two changes of buffer 1, incubated with 125I-protein A (~0.25 μCi/gel slot) for 2 h, washed, dried, and fluorographed. Fluorographs were scanned in white light on a densitometer (Helena Laboratories, Beaumont, Tex.)

RESULTS

Morphology

SURFACE MORPHOLOGY—SEM: The skin from the normal mouse was loose, with puckered wrinkles in a regular pattern and cells that were similar in size and beaded in surface appearance (Fig. 1 a). The Er/+ mouse skin showed dorsoventrally oriented wrinkles (Fig. 1 b) while the skin of the Er/Er mouse appeared smooth, taut, and ragged in appearance (Fig. 1 c). Individual cells of both Er/+ and Er/Er mice were variable in size, irregular in shape, and had a smooth membrane surface. These variations from normal were more marked in Er/Er epidermis than in Er/+ tissue.

HISTOLOGY OF THE EPIDERMIS: The normal 19-d fetal mouse trunk epidermis was a 50-μm-thick epithelium with one basal, two spinous, three to four granular and six to eight cornified layers (Fig. 2 a). Hair follicles extended from the basal layer at regular intervals into the dermis. The epidermis of Er/+ animals was indistinguishable from the +/- epidermis. Small amounts of glycogen were detected in spinous cells and in developing follicles of +/- and Er/+ epidermis. Marked abnormalities in epidermal thickness and organization were seen in skin from the Er/Er littermates (Figs. 2 b and 3 a-c). Overall, the epidermis was hyperplastic, yet highly inconsistent in thickness in any given region. Basal cells often stained more darkly compared with those of the +/- and Er/+...
epidermis and with cells from other layers of the Er/Er epidermis (Figs. 2b and 3a). In the thickened areas, increases in one or more of the spinous, granular, or superficial regions were seen. A true cornified zone was not seen, although isolated areas of cornified cells (Fig. 3b) were present within the epidermis, surrounded by cells which were morphologically less differentiated (Fig. 3b). Cells of the granular and superficial layers, in particular, were irregular in shape and oriented at random (Fig. 2b and 3a and b). Large amounts of glycogen were seen as PAS-positive deposits in spinous and granular cells (data not shown).

**ULTRASTRUCTURE OF THE EPIDERMIS:** By TEM, marked abnormalities were seen in cells of all layers of the Er/Er mouse epidermis in comparison with the +/+ (Fig. 4) or Er/+ animals. Normal mouse basal cells were joined tightly together and with cells of the spinous layer. In Er/Er mice, the number of desmosomes among basal cells was sparse. The basal lamina was intact and bound to basal cells by typical hemidesmosomes. Basal cells contained phagolysosomes (Fig. 5a) and mitochondria with large, electron-opaque, membrane-bounded granules (Fig. 5a, inset). All cell layers above the basal layer in the Er/Er mouse were variable in number and morphology (Figs. 2b, 3, and 5). In some regions a single spinous layer separated basal and granular layers (Fig. 5a); in others, even adjacent regions, there were several layers of spinous cells (Fig. 5b). The only notable difference in the fine structure of spinous cells compared with those of +/+ or Er/+ animals was in the mitochondria which, like those in the basal cells, contained large, electron-opaque spheres.

Granular layers in the Er/Er epidermis varied in number (2–8), in content of keratohyalin granules, and in position relative to the epidermal surface (Figs. 2b, 3, and 5). In the +/+ mouse (Figs. 2a and 4), keratohyalin granules were present in cells of each of three to four granular layers and increased in size progressively toward the skin surface. They were typically associated with keratin filaments. By contrast, granular cells in the Er/Er epidermis were variable in size, shape, and association with filaments regardless of the position of the cell within the epidermis (Figs. 2b, 3, and 5a). In a few regions, kerato-
hyalin granules were not evident histologically (Fig. 3 c) but at the ultrastructural level were observed as small, round bodies (Fig. 5 b). Granular cells were generally located subjacent to one or more layers of nongranular, surface-related cells (Fig. 3 a and b), but in a few instances they were totally superficial or covered by only a single cell layer (Figs. 3 b and c, 5 a, and 6 a). Thus, in contrast with an earlier report (17), the numbers of granular cell layers appeared exaggerated in the mutant rather than lacking.

Fully cornified cells were identified only as isolated cells or multiple layers of keratinized cells within deeper epidermal strata (Figs. 2 b, 3, and 6 a), some of these "embedded" squames were nearly normal in morphology (Fig. 6 a); the cell contents were electron-dense, nuclear and cytoplasmic remnants were not evident, and a cornified envelope (32) formed the cell boundary. Contents of lamellar granules were abundant in the extracellular space surrounding such cells (Fig. 6, inset).

Cells with a transitional morphology ("T" cells) were occasionally identified among upper strata cells. These cells showed partial nuclear disintegration, loss of organelles, condensation of filaments with keratohyalin, and a cornified cell envelope. Other surface cells had only some of the cornified features of "T" cells (Fig. 6 b), while still others were completely noncornified, having an unmodified plasma membrane and a full complement of cytoplasmic organelles (Fig. 5).

Samples of skin from the head of all animals were compared with body epidermis to determine whether there was regional

![Figure 3](image_url)

**Figure 3** Epidermis from the trunk of Er/Er mice. Different regions of the epidermis show: (a) thickened zone of irregularly shaped granular cells (G) beneath noncornified superficial cells (nC). (b) A spiderlike zone of cornified cells (C) within the epidermis, granular cells at the epidermal surface (arrows) with a "cap" of noncornified superficial cells and (c) a variable position and abundance of granular cells (arrows) appear among the epidermal strata. Widened intercellular spaces in basal and spinous regions are evident in all micrographs. X 325.

![Figure 4](image_url)

**Figure 4** Full-thickness epidermis from the trunk of a +/+ mouse showing the ultrastructure and normal arrangement of basal, spinous, granular, and cornified cells. X 2,900.
variation in epidermal structure or pattern of differentiation. Only in the +/+ animal was the epidermis of the head identical with that of the trunk. Head epidermis from Er/+ animals was thickened as a consequence of increases in all suprabasal cell layers, whereas in the Er/Er animals thickening was due primarily to a greater number of cell layers in granular and cornified regions (Fig. 7). Granular cells of the Er/Er epidermis were flattened, electron-dense (Fig. 7a), and demonstrated a gradual morphologic transition into superficial cells that retained a nucleus, cellular organelles, vacuoles, and bundles of electron-dense filaments, embedded in an electron-opaque matrix (Fig. 7b). Such cells lacked a cornified cell envelope. A broader spectrum of superficial cell types occurred in Er/+ animals (data not shown) but, like the Er/Er superficial cells, they contained a variety of organelles and often a nucleus.

Observations of the head by SEM revealed alterations in the surface properties and in the development of epidermal appendages of Er/Er and Er/+ animals when compared with the +/+ mice. Vibrissae were fully exposed in the +/+ mice, reduced in Er/+ mice, and absent from Er/Er animals (data not shown). It is characteristic of the adult Er/+ mice to have fewer vibrissae than normal.

Fetal mice from 13-d-gestation litters were examined to determine whether the epidermis of Er/Er animals is normal before keratinization, in which case, the genetic defect would be a primary keratinization defect and would be expressed only with this event, or whether the mutation affects epidermal development more generally before keratinization is initiated. 13-d Er/Er mice were identified by their blunt limbs and stumpy tail. The Er/+ mice, however, could not be distinguished from the genetically normal animals. As in the older fetus, epidermal thickness was irregular (two to seven layers) in the Er/Er mutant compared with the normal (Figs. 8a and b). Cells of the lowermost layers were widely separated (Fig. 8b) and attached by few desmosomes, as compared ultrastructurally with tissue from the normal 13-d fetal three-layered epidermis (Figs. 9a and b). In addition, a diffuse matrixlike material was observed in the intracellular spaces among Er/Er

![Figure 5](https://example.com/figure5.png)

**FIGURE 5** Two regions of full-thickness epidermis from the trunk of a Er/Er mouse. (a) Basal (B) and lower spinous (S) cells are widely separated and contain phagolysosomes (PhLy). One layer of spinous cells separates basal and granular (G) cells. Granular cells extend all the way to the epidermal surface as indicated by the keratohyalin granules (KHG and arrows) in the cytoplasm. X 2,500. Inset: spherical inclusions within mitochondria found in cells of basal, spinous, and granular layers. X 21,000 (b) Granular cells have minute KHG that would not be distinguishable at the LM level. The intercellular spaces are wide among basal and lower spinous cells. X 2,500. Nonkeratinized cells form the epidermal surface in both samples.
epidermal cells (Fig. 9b); morphologically similar material was associated with the basal lamina in greater amounts in Er/Er animals than in the normal (Fig. 9b). Cell shapes were highly irregular in the Er/Er epidermis but the cytologic characteristics of basal, intermediate, and peridermal cells were equivalent in Er/Er and +/+ animals. The irregularity in cell size, shape, and organization was also recognized at the epidermal surface by the aberrant shape, arrangement, and distribution of microvilli of peridermal cells (Fig. 10).

**Biochemistry**

The proteins in epidermal extracts from newborn +/+, Er/+
+ and Er/Er mice were compared by SDS PAGE. The +/+ and Er/+ extracts had almost identical protein profiles (Fig. 11, lanes a and b). The keratin region contained prominent bands of 65, 62.5, 59, and 56 kdaltons, minor bands of 62, 57, and 53 kdaltons, and several additional bands between 53 and 44 kdaltons. A prominent band was seen at 26.5 kdaltons but was reduced in intensity in the heterozygote. The extract of the Er/Er epidermis (Fig. 11, lane c) had multiple protein differences from the normal (Fig. 11, lanes a and b). The 26.5-kdalton protein was absent from the Er/Er animal. In the keratin region, the band at 62.5 kdaltons was missing; several bands (53, 49, 46, and 44 kdaltons) were much stronger in the Er/Er extract than in the +/+ and occasionally appeared as

![Figure 6](image-url)
doublets (Fig. 12, lane c; compare high molecular weight bands). There were additional alterations in proteins of a wide range of molecular weights.

Incorporation of radiolabeled histidine was performed in vivo in the +/+ and Er/+ animals to determine whether any of these proteins could be characterized as histidine-rich. In extracts from mice which had been labeled for 2 h, multiple high molecular weight bands were labeled (Fig. 12, lanes a and a'). In contrast, labeling for 24 h (Fig. 12, lanes b and b') resulted in the incorporation of histidine into the 26.5-kdalton band as well as the high molecular weight bands. Since the 26.5-kdalton band was strongly labeled after 24 h (Fig. 12, lanes b and b'), it was considered a histidine-rich protein. This band corresponded to the band which was absent from the Er/Er extract (Fig. 11, lane c).

Detection of Antigens

Epidermal proteins which are immunologically related to rat epidermal filaggrin were identified in the gels by reaction with antiserum, followed by 125I-protein A. A densitometric scan of the fluorograph is shown in Fig. 13. The 26.5-kdalton band previously identified as a histidine-containing protein in the +/+ and Er/+ samples (peak f) was cross-reactive. This band was missing from the Er/Er extract. Additional cross-reactive bands in all three extracts were found at the origin (peak a) and at approximately 145 (peak b), 116 (peak c), 90 (peak d), and 59 kdaltons (peak e). The high molecular weight cross-reactive bands were most intense in the Er/Er and were broader (compare peaks d and e) when compared with the heterozygote and normal extracts.

Immunofluorescence

An indirect immunofluorescent reaction using antibody to rat epidermal filaggrin is shown in Fig. 14. Skin from both the +/+ and the Er/Er mice gave a strong positive reaction. In the normal, the positive reaction was found in the stratum corneum, and was associated with granules in the granular layers (Fig. 14a). In the Er/Er, a both diffuse and granular immunofluorescent reaction was seen throughout the cytoplasm of granular and superficial cells (Fig. 14b).

DISCUSSION

The present study has shown abnormalities in epidermal mor-

![Figure 7](image_url)
The epidermis is hyperplastic, has a variable and aberrant pattern of organization, fails to synthesize the histidine-rich product, filaggrin, produces abnormal keratins, and does not carry out the normal events of terminal differentiation.

Many of the morphologic findings of the present study differ from those of Guenet et al. (17) in their original report of the Er/Er mouse. They described an epidermis with only basal and spinous layers and rare development of hair follicles. In contrast, we have observed an epidermis which is hyperplastic and has exaggerated numbers of granular layers, superficial noncornified cells, isolated foci of partially cornified cells, and hair follicles spaced at irregular intervals. The discrepancies between our findings and those of Guenet et al. (17) may be due to differences in sampling and/or in breeding stock. Our mice were bred on a uniform genetic background, whereas Guenet et al. used two genetic types of matings to produce the homozygous embryos.

Epidermal hyperplasia in the Er mutant mouse is associated with tissue edema, hence widened intercellular spaces, increased accumulations of glycogen, large intramitochondrial spheres, phagolysosomes in keratinocytes, the presence of dark keratinocytes, and increased numbers of keratohyalin granules in granular cells (2, 4, 13, 20, 30). Such features are also characteristics of epidermal hyperplasia associated with pathologic conditions (e.g., benign keratoses and neoplasias), can be induced by mechanical, chemical, thermal, or radiation assault (2), and are believed to be associated with a general state of...
FIGURE 10 Surface view of peridermal cells from 13-d-gestation normal (a) and Er/Er (b) mice. (a) Cells from the normal mouse have a regular polygonal shape and an even distribution and density of microvilli. (b) Er/Er peridermal cells have irregular shapes and variable densities of microvilli, and appear loosely joined in certain regions. ×1,300.

FIGURE 11 Electrophoresis of epidermal extracts on 8.75% SDS polyacrylamide gels. Lane a, +/+; lane b, Er/+; lane c, Er/Er. Molecular weights (× 10^{-3}) were calculated by comparison with protein standards. Note the relative increase in intensity of bands of 59, 53, 49 kdaltons, and the absence of the 26.5-kdalton band in the Er/Er extract.

FIGURE 12 Electrophoresis on SDS polyacrylamide gel on Er/+ epidermal extracts labeled in vivo with [^{14}C]histidine. Lanes a and a' were labeled for 2 h; lanes b and b' were labeled for 24 h. Lanes a and b are stained with Coomassie Brilliant Blue for protein. Lanes a' and b' are the fluorograph showing the radiolabeled bands in the same gel. Note the strong labeling of the 26.5-kdalton (arrow) band in sample b' labeled for 24 h.

cellular injury and altered metabolism. The intramitochondrial granules, for example, have been observed not only in hyperplastic mouse epidermis (14) but also in fetal mouse epidermal cells (35), cultured mouse epidermal cells (our unpublished observation), and in normal, in metabolically deficient and in transformed epithelial cells from a wide variety of organs (bladder [23], stomach, colon, uterus [reviewed in reference 21] and in healing wounds [34]). In many of these instances there is an undifferentiated or dedifferentiated state of the tissue as in the Er/Er mouse.

The histologic and cytologic similarities between pathologic, induced, and Er/Er hyperplastic epidermis are probably the consequence of very different underlying defects. While some of the induced hyperplastic conditions are combined with a subterminally differentiated or dedifferentiated state of the tissue (e.g., chemically induced neoplasia [30]), there seem to be no examples where terminal differentiation fails to occur as it does, almost uniformly, in the Er/Er mouse. It may not be possible to find such a comparison because the full effects of the mutation in the Er/Er mouse are fatal.

In normal epidermis, differentiation progresses uniformly throughout the tissue in an orderly sequence of basal-spinous-granular-cornified cells. A “pattern” of epidermal differentiation in Er/Er tissue cannot be described because of marked variability in the presence and numbers of granular layers, and presence and position of cornified, partially cornified, and noncornified cells among cell strata. The variability of the products of terminal differentiation raises the question of whether there is differing ability of cells derived from different epidermal stem cells (different epidermal proliferating units [1, 28, 29]) to carry out the events of differentiation or whether some local, even microenvironmental, factors influence differentiation.

Assuming that the granular cells differentiate into the overlying (more superficial) noncornified cells, then the fate of the keratohyalin granules during the transition must be explained. During normal keratinization, a high molecular weight, phos-
phorylated, histidine-rich precursor protein (10, 26) in kerato-
hyalin granules is converted via a multienzyme-regulated se-
quence to a lower molecular weight product, filaggrin. Filag-
 grin is believed to be the electron-dense matrix protein (8) 
which embeds electron-lucent keratin filaments in the cornified 
cell to form a filament-matrix assembly described ultrastruc-
turally as the keratin pattern (5). Although filaggrin was absent 
from Er/Er epidermal extracts (Figs. 12 and 13) and present in 
diminished amounts in Er/+ extracts (Figs. 11 and 13), the 
immunofluorescence studies (Fig. 14b) showed that some form 
of a histidine-rich protein was present in the noncornified 
surface cells. For the following reasons, we suspect that this 
protein is the precursor of filaggrin and that the mutation 
blocks its normal conversion to the product. First, a series of 
high molecular weight, histidine-rich proteins was identified 
on gels of epidermal extracts from all three genotypes; secondly, 
the precursor protein in keratohyalin granules cross-reacts with 
anti-protein A-labeled blot of an SDS polyacrylamide gel acted 
with antibody to the rat epidermal filaggrin. +/+ extract, top; Er/+ 
extract, center; Er/Er extract, bottom. Peaks a-f are discussed in the 

Thus we conclude that the epidermis of the Er/Er mutant 
mouse contains precursor histidine-rich matrix protein which 
cannot undergo the posttranslational modifications necessary 
to generate the 26.5-kdalton functional protein found in normal 
stratum corneum. This could be due to a mutation in the primary 
structure of the histidine-rich protein gene product or to an alteration in an enzyme responsible for some posttrans-

We favor the latter possibility for the reason that, although 
the alterations in both the histidine-rich and keratin proteins 
might explain the abnormal keratinization, they are insufficient 
to explain the organizational defects in the tissue, the structural 
changes in the prekeratinized embryonic epidermis, and the 
various other phenotypic changes in face and appendages of 
the mutant animal. The alterations in filaggrin and keratin are 

The authors gratefully acknowledge the technical expertise of Mary 
Hoff, Carolyn Foster, Julie Scofield, and Alexis Lynley. We thank L.C. 
Harne for careful observation, breeding, and care of the animals. We 
thank Drs. George Odland, Peter Byers, John Lonsdale-Eccles, John 
Stanley, David Woodley, and Mark Mamrack for helpful suggestions 
and review of this work. The manuscript was skillfully prepared by 
Doris Ringer and Marion Brown.

The study was supported by U. S. Public Health Service grants AM-
21557 and DE-04660 from the National Institutes of Health.

Received for publication 9 February 1981, and in revised form 18 August 
1981.

REFERENCES

5. Brody, I. 1959. The keratinization of epidermal cells of normal guinea pig skin as revealed 
fluorocarbons ("freon") as intermediate and transitional fluids. J. Microsc. (Oxf.). 13:31-
342.
7. Dale, B. A. 1977. Purification and characterization of a basic protein from the stratum 