Advances in understanding the genetic basis of inherited single gene skin barrier disorders: new clues to key genes that may be involved in the pathogenesis of atopic dermatitis

Avanços no entendimento da base genética de doenças hereditárias monogênicas da barreira epidérmica: novas pistas para os principais genes que podem estar envolvidos na patogênese da dermatite atópica.

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Abstract: Increasing knowledge of genomic DNA sequences and genetic databases has led to the characterization of the molecular basis of several inherited skin disorders. In this review we summarize some of the major recent discoveries that have been made in defining the pathogenic mutations that cause inherited disorders of the skin barrier leading to skin scaling or increased transepidermal water loss in either rare disorders (Netherton’s syndrome or harlequin ichthyosis) or more common genodermatoses (ichthyosis vulgaris). These molecular breakthroughs have led to more accurate diagnoses, better genetic counselling and, where appropriate, the feasibility of DNA-based prenatal diagnosis, as well as the possibility of developing newer forms of treatment, including gene or protein therapy. Identifying the molecular basis of these conditions, especially ichthyosis vulgaris, has also provided dramatic new insight into the genetic abnormalities in the common disorder, atopic dermatitis. Thus research on the relatively rare single gene inherited skin disorders not only has benefits for patients and their families with these uncommon conditions but also has the potential to yield fresh and significant new information about very common skin diseases.

Key words: Eczema; Epidermis; Ichthyosis; Mutation

Resumo: O maior conhecimento sobre as de sequências genómicas de DNA e as bases de dados genéticas levou à caracterização da base molecular de várias doenças hereditárias de pele. Nesta revisão resumimos algumas das descobertas recentes mais importantes quanto à definição das mutações patogênicas que causam as doenças hereditárias da barreira cutânea, levando à descamação ou aumento da perda bidrônica transepidérmica, seja em doenças raras, (síndrome de Netherton ou ictiose em Arlequim) ou genodermatoses mais comuns (ictiose vulgar). Estas descobertas moleculares têm conduzido a diagnósticos mais acurados, melhor aconselhamento genético e, quando apropriado, à possibilidade de diagnóstico pré-natal baseado em DNA, assim como desenvolvimento de novas formas de tratamento, incluindo terapia gênica ou proteica. A identificação da base molecular destas doenças, especialmente ictiose vulgar, também propiciou importante entendimento das anormalidades genéticas de doença comum, a dermatite atópica. Assim, as pesquisas em doenças hereditárias monogênicas de pele relativamente raras não apenas trazem benefícios para os pacientes e familiares, mas também têm o potencial de trazer informações novas e significativas sobre doenças de pele muito comuns.

Palavras chave: Eczema; Epiderme; Ictiose; Mutação
INTRODUCTION

During the last two decades the molecular basis of over 2000 inherited single gene disorders has been determined. More than 25% of these have a skin phenotype, making the inherited skin disorders one of the best characterized groups of genetic diseases. For clinicians, the accumulating molecular data have several benefits for patients. First, access to DNA sequencing laboratories that can identify pathogenic mutations in a particular gene allows more accurate diagnoses to be made. Moreover, the nature of the mutation(s) may provide additional information about inheritance patterns as well as the likely clinical course and prognosis, collectively improving genetic counselling. Secondly, knowledge of family-specific gene mutations makes it possible to undertake DNA-based prenatal diagnosis in subsequent at risk pregnancies. Examination of fetal DNA during the first trimester following chorionic villus biopsy is now feasible for a broad range of severe genodermatoses. Furthermore, preimplantation genetic diagnosis has become a viable option in certain countries. Finally, a key objective of molecular research on the genodermatoses is to develop new treatments that benefit patients. Gene therapy is a science that is very much in its infancy: there are several technical, practical and safety issues that still need to be developed and optimized, and therefore the next few years are likely to see pilot trials of human gene therapy rather than widespread clinical application of such treatments. In essence, the big recent advances in the genodermatoses have been in the plethora of new laboratories that can identify pathogenic mutations in the whole skin leading to impaired thermoregulation, fluid balance and protection against infection. The thick keratin layer also leads to contraction of the skin around the eyes causing ectropion, and the lips resulting in eclabion, and the chest leading to respiratory difficulties. Infants born with HI often die within the first few days of life but some survive with the appropriate clinical care and the early use of retinoids. In HI there is clearly a grossly abnormal epidermal bar-
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rrier but it is plausible that less disruptive abnormalities in the gene responsible for HI might also provide some clues to epidermal barrier dysfunction in AD. In 2005, the genetic basis of HI was resolved by two independent groups using single nucleotide polymorphism chip technology with homozygosity mapping or a candidate gene approach.\textsuperscript{12,13} The gene was mapped to chromosome 2q35 and is known as \textit{ABCA12}, which encodes a transmembranous regulatory protein involved in lipid transport. Deletion, truncation and some missense mutations result in a dysfunctional protein and lead to HI whilst replacing the wild type allele in cultured null keratinocytes leads to recovery of lamellar granule-mediated lipid secretion.\textsuperscript{12,14} In contrast to the highly disruptive mutations in \textit{ABCA12} that underlie HI, it is possible that less severe functional polymorphisms might be present in some individuals with AD. Such sequence alterations could compromise epidermal barrier integrity, increase transepidermal water loss and, perhaps in combination with other factors, lead to AD. Thus far, however, no specific \textit{ABCA12} genotypes have been linked to AD.

\textbf{Ichthyosis Vulgaris}

In contrast to the very rare HI, ichthyosis vulgaris (IV) (Figure 1C) is the most common disorder of keratinization, estimated to affect about 1 in 250 people.\textsuperscript{15} The clinical features are variable and include keratosis pilaris, fine scale on arms and legs and palmar hyperlinearity, although in some cases the scaling can be extensive and severe. Moreover, the co-existence of IV and AD is well known, with up to 50% of subjects with IV having an atopic tendency,\textsuperscript{16} and therefore elucidating the molecular basis of IV might be expected to provide new insight into the genetics of AD. Linkage of IV to 1q21 and the epidermal differentiation complex has been previously demonstrated\textsuperscript{17} but it was not until 2006 that pathogenic mutations were identified in the gene encoding filaggrin (\textit{FLG}). Notably, homozygous and compound heterozygous mutations including the recurrent mutations R501X and 2282del4 were found in several kindreds with IV. Indeed, these two recurrent mutations were found in about 9% of the European population thereby showing just how common deleterious mutations in \textit{FLG} actually are. In addition, dominant and semi-dominant mutations in \textit{FLG} were identified to explain both the differing phenotypic severities of IV and the pattern of inheritance. Notably, a number of mutations were identified that led to ablation of processed filaggrin products.\textsuperscript{18} Filaggrin is an essential component of the stratum corneum and it is cleaved from profilaggrin causing aggregation of the keratin cytoskeleton during terminal differentiation of the granular layer of the epidermis. Filaggrin is then cross-linked by transglutaminases to contribute to the formation of the cornified cell envelope. In the absence of filaggrin, the epidermal barrier fails to function normally, increasing transepidermal water loss and giving rise to the ichthyosis.

\textbf{Atopic Dermatitis}

Atopic dermatitis (Figure 1D) is a common skin disorder, the aetiology of which is complex and interplay between genetic and environmental factors is believed to be important. The finding of common mutations in \textit{FLG} in a substantial proportion of the European population led investigators to assess the prevalence of filaggrin pathology in AD. For the mutations R501X and 2282del4, the combined allele frequency in AD is 0.330 compared to 0.042 in the non-atopic population, thus demonstrating just how common these mutations actually are.\textsuperscript{16} The reason why these discoveries were not realized until 2006 was largely due to technical difficulties experienced by many researchers in amplifying the \textit{FLG} gene (lots of homologous repeat units with little sequence dissimilarity). Once DNA sequencing became possible, however, the common finding of \textit{FLG} mutations in AD was a truly remarkable discovery. Of course, loss of filaggrin expression does not provide a full explanation for the genetic basis of AD in all subjects. It is likely that AD is in fact a heterogeneous disorder with some individuals having a primary epidermal barrier defect due to \textit{FLG} gene mutations, whereas others will have primary immunological abnormalities or alternative primary susceptibilities to inflammation or infection or, most probably, a combination of several genetic factors influencing these skin responses. Interestingly, \textit{FLG} mutations are also a predisposing factor for asthma, but only when associated with AD rather than clinical asthma as a separate entity.\textsuperscript{16} 

common disorders of epidermal dysfunction such as AD. In the future, systematic analysis of the roles of the various proteins involved such as LEKTI, glucosylceramide and, in particular, filaggrin in the maintenance of epidermal integrity is expected to contribute to the development of novel therapeutic strategies and thus represent a direct benefit of translational research on rare genodermatoses to the management of very common skin problems encountered by dermatologists in daily clinical practice.

CONCLUSIONS

The molecular characterization of inherited single gene skin disorders such as HI, NS and IV is important not only for patients and their families but also for our understanding of epidermal physiology and pathology. This has direct implications for more common disorders of epidermal dysfunction such as AD. In the future, systematic analysis of the roles of the various proteins involved such as LEKTI, glucosylceramide and, in particular, filaggrin in the maintenance of epidermal integrity is expected to contribute to the development of novel therapeutic strategies and thus represent a direct benefit of translational research on rare genodermatoses to the management of very common skin problems encountered by dermatologists in daily clinical practice.

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