Dystrophic Epidermolysis Bullosa: Secondary Disease Mechanisms and Disease Modifiers

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The phenotypic presentation of monogenetic diseases is determined not only by the nature of the causative mutations but also is influenced by manifold cellular, microenvironmental, and external factors. Here, heritable extracellular matrix diseases, including dystrophic epidermolysis bullosa (DEB), are no exceptions. Dystrophic epidermolysis bullosa is caused by mutations in the COL7A1 gene encoding collagen VII. Deficiency of collagen VII leads to skin and mucosal fragility, which progresses from skin blistering to severe fibrosis and cancer. Clinical and pre-clinical studies suggest that targeting of secondary disease mechanisms or employment of natural disease modifiers can alleviate DEB severity and progression. However, since many of these mechanisms are needed for tissue homeostasis, informed, selective targeting is essential for safe and efficacious treatment. Here, we discuss a selection of key disease modifiers and modifying processes active in DEB, summarize the still scattered knowledge of them, and reflect on ways forward toward their utilization for symptom-relief or enhancement of curative therapies.

Keywords: collagen VII, fibrosis, inflammation, transforming growth factor β, skin

INTRODUCTION

Epidermolysis bullosa (EB) comprises a group of genetic disorders manifested by mechanically induced blistering and fragility of the skin and other stratified epithelia. Over 20 genes have been described to be causative of EB and related skin fragility disorders (Has et al., 2020). These span genes encoding intracellular transporters, via cytoskeletal proteins and integrin receptors, to extracellular matrix (ECM) proteins. Depending on the level of separation in the skin, EB is divided into four main types: Epidermolysis bullosa simplex with blistering occurring in the epidermal basal keratinocyte layer, junctional EB with blistering within the lamina lucida of the epidermal basement membrane, dystrophic EB (DEB) with blistering below the epidermal basement membrane, and Kindler EB with blistering occurring in all layers (Bardhan et al., 2020).

This mini-review will focus on DEB, which can be inherited in a dominant or a recessive (RDEB) manner (Bardhan et al., 2020). Dystrophic epidermolysis bullosa is primarily caused by genetic loss of function or abundance of collagen VII, encoded by the COL7A1 gene (Has et al., 2018), but a number of secondary, molecular and cellular, events modify the disease phenotype.
COLLAGEN VII

Collagen VII is a large, in parts atypical collagen, which attaches the epidermal basement membrane to the dermal ECM. It is synthesized as pro-collagen VII consisting of three pro-α1 chains that fold into one pro-collagen VII molecule. The molecule has sizable N-terminal non-collagenous 1 (NC1) domains, followed by an extended collagenous domain, and ends with minor NC2 domains at the C-terminus (Has et al., 2018). Although minor, the NC2 domain is essential for intermolecular assembly (Bruckner-Tuderman et al., 1995; Chen et al., 2001). Extracellularly in the skin, pro-collagen VII molecules align as antiparallel dimers, which laterally aggregate to form anchoring fibrils (Chung and Uitto, 2010). The antiparallel dimer formation occurs over the NC2 domain, and in a stretched configuration, anchoring fibrils consequently contain NC1 domains at both ends and the NC2 domains in the middle. Proper removal of a large part of the NC2 domain is essential for stable anchoring fibril formation (Bruckner-Tuderman et al., 1995; Chen et al., 2001). Astacin-like proteases BMP-1/mTLD and meprins have been shown to cut pro-collagen VII to collagen VII (Rattenholl et al., 2002; Moali et al., 2005; Krupa et al., 2021).

In the skin, epidermal keratinocytes and papillary dermal fibroblasts produce collagen VII and contribute to its deposition at the dermal-epidermal junction (DEJ; Twaroski et al., 2019). The NC1 domains are generally positioned in the epidermal basement membrane and the collagenous domains loop down into the superficial papillary ECM (Has et al., 2018). Anchoring fibrils provide skin stability by attaching the epidermal basement membrane, via high-affinity interactions of their collagen VII NC1 domains with laminin-332 and collagen IV, and simultaneous binding to collagen fibrils in the papillary dermis (Chen et al., 1997, 1999; Brittingham et al., 2006; Villone et al., 2008). It appears that the sixth or seventh fibronectin type III-like domain in the NC1 domain harbors the major binding sites for laminin-332, collagen IV, and even the weaker interaction partner thrombospondin-1 (Aho and Uitto, 1998; Chen et al., 1999; Brittingham et al., 2006; Atanasova et al., 2019).

For the correct function, deposition and stability of collagen VII and anchoring fibrils posttranslational modifications are needed. Currently, there is an insufficient understanding on the exact role of these modifications. Collagen VII appears to be a substrate of the cross-linking enzyme transglutaminase 2 (TGM2), and TGM2-mediated crosslink formation may stabilize anchoring fibrils (Raghunath et al., 1996; Küttnner et al., 2014). In addition, collagen VII has been suggested to be modified by the multi-functional enzyme lysyl hydroxylase 3 (LH3) also known as pro-collagen-lysine, 2-oxoglutarate 5-dioxygenase 3 (Watt et al., 2015; Vahidnezhad et al., 2019). Lysyl hydroxylase 3 both hydroxylates lysyl residues and then further O-glycosylates these (Salo et al., 2006; Risteli et al., 2009). Lysyl hydroxylase 3 deficiency is linked to altered deposition and reduced functionality of multiple tissue-stabilizing collagens, including collagen VII. Interestingly, skin blistering in LH3-deficient skin shows similarities to that of collagen VII-deficient RDEB skin (Salo et al., 2008; Vahidnezhad et al., 2019). In addition to posttranslational modifications, coordinated production and deposition of collagen VII by both keratinocytes and fibroblasts have been indicated to facilitate anchoring fibril assembly (Supp et al., 2019). Collectively, the complex synthesis and modification of collagen VII, the contribution of multiple cellular sources to anchoring fibril formation and the incomplete understanding of these processes, pose challenges for RNA, gene, protein, or cellular therapies aiming to restore collagen VII and anchoring fibrils in RDEB.

GENOTYPE-PHENOTYPE CORRELATIONS

More than 1,000 distinct mutations have been reported to cause DEB. The most severe phenotypes are associated with COL7A1 mutations causing complete loss of translated collagen VII. However, the genotype–phenotype correlations are not completely clear and there is a vast phenotypic variability (Figure 1). There are cases with sole nail dystrophy or mild localized disease. These are mostly associated with glycine substitutions in the collagenous domain (Dang and Murrell, 2008). Patients might suffer from an inverse phenotype, with skin fragility mostly in the flexural skin areas and the mucosa. This seems to be caused by recessive arginine and glycine substitutions in the collagenous domain (van den Akker et al., 2011), and the hypothesis was proposed that the higher temperature in the body flexures impairs stability of the glycine-substituted collagen VII, leading to skin lesions in these specific areas (van den Akker et al., 2011). The rare subtype, DEB pruriginosa, clinically characterized by intensively itchy,
hypertrophic, prurigo-like papules, and nodules, is in more than 50% of the cases associated with glycine substitutions, followed by in-frame skipping mutations in around 30% (Kim et al., 2015). These data highlight that the type of mutation but also its position within collagen VII are phenotypic determinants (Dang and Murrell, 2008) with a broad range of cutaneous manifestations. Further, the nature of amino acid change for a given position is important, as shown by the different phenotypes caused by mutations in the same position (Almaani et al., 2011). Nonetheless, even in patients with the same COL7A1 mutations the phenotypes might differ, as shown by studies in siblings (Hovnanian et al., 1997; Bodemer et al., 2003; Titeux et al., 2008; Odorisio et al., 2014), disclosing that in addition to the causative mutations, other genetic, epigenetic, microenvironmental, and environmental factors contribute to the phenotype. As discussed below, these factors remain limitedly known. A better understanding of them is important for improved prognostication of the disease severity and therapeutic exploitation for disease-modifying therapies. Pathological COL7A1 mutations could also be used to provide insights on the collagen VII intertome; however, it should in this context be mentioned that missense mutations in known protein–protein interacting domains, such as the NC1 domain, are exceedingly rare [COL7A1(gene) - ClinVar – NCBI, 2020].

**CUTANEOUS MANIFESTATIONS IN RDEB**

The disease hallmark of RDEB is skin fragility, which manifests with the mechanically induced development of blisters and erosions, especially in trauma-prone skin areas, such as hands, feet, and knees (Figure 1). Acute or chronic wounds occur at different body sites with itch and pain as a consequence (Solis et al., 2021). Also the mucosa and adnexal structures of the skin might be affected, with features of intraoral soft-tissue involvement and dental abnormalities (Krämer et al., 2020), nail dystrophy, and secondary alopecia. Ocular involvement encompasses corneal erosions and subsequent scarring, as well as eye lid erosions, followed by ectropion or symblephara (Figueira et al., 2010). These may result in severe impairment of visual acuity.

The cutaneous and mucosal blisters heal with scarring. At sites, meeting frequent mechanical and frictional challenges, typically the extremities, fibrosis driven by a chronic state of injury and inflammation leads to joint contractures and webbing of fingers and toes and the formation pseudosyndactylies – so called mitten deformities – allowing only limited function and mobility (Figure 1B).

The intermittent blistering, impaired wound healing with inflammation, high bacterial load, wound infections, and subsequent fibrosis of the skin are considered to be main events creating a microenvironment that promotes mutagenesis of keratinocytes through, e.g., activation of cell-intrinsic DNA editing enzymes, such as the APOBECs (Hoste et al., 2015; Cho et al., 2018). These events also establish a stiffened and hyper-vascularized environment, facilitating the growth, progression, and metastasis of established squamous cell carcinomas (SCCs; Martins et al., 2016; Mittapalli et al., 2016; Föll et al., 2017; Condorelli et al., 2019). These arise already in early adulthood and are the main cause of death in patients with severe generalized RDEB (Montaudié et al., 2016). Notably, in that specific patient, population more than 90% will have developed an SCC by the age of 55 years and have a 78% cumulative risk of death from metastatic SCCs (Fine et al., 2009).

**EXTRACUTANEOUS MANIFESTATIONS IN RDEB**

Besides the disease manifestations in skin and mucosa, complications arise in other organs. Such extracutaneous manifestations are more prominent in patients with the severe subtypes of RDEB. The perpetual cycles of trauma-induced blistering of the esophagus result in stenoses, requiring dilatations, and/or gastrostoma to ensure proper food intake. Consequences are anemia, partially due to iron deficiency, deficiencies in vitamins and minerals, and severe failure to thrive (Reimer et al., 2020). With progressive disease, connected to the systemic impact of cutaneous wounding and inflammation, internal organs can become affected. Renal parenchymal disease might arise, linked to amyloidosis or the autoimmune disease IgA nephritis (Bardhan et al., 2020). A subset of individuals with RDEB develops life-threatening dilated cardiomyopathy at an early age (Fine et al., 2008). A less considered change is external auditory canal stenosis, which may result in hearing loss (Brown et al., 2017).

**Immune Anomalies in RDEB**

People with RDEB have elevated bacterial colonization of wounds compared to non-RDEB individuals with large, chronic wounds (Levin et al., 2021), with *Staphylococcus aureus* and *Streptococcus pyogenes* being the most common colonizers (Fuentes et al., 2018; Levin et al., 2021). In addition, RDEB wounds and skin show a reduced diversity of their bacterial microbiome (Bar et al., 2021). Our studies of RDEB patients and RDEB mouse models indicate that the increased susceptibility to bacterial infections is not only associated with wounding, but also to a large extent due to loss of collagen VII from secondary lymphoid organs (Nyström et al., 2018). One mechanism making people with RDEB unable to respond appropriately to bacterial challenges is through loss of interactions between collagen VII and the ECM protein cochlin in the conduits of secondary lymphoid organs. Upon bacterial challenges, systemic danger signals, through, e.g., TNF, increase expression of aggrecanase 1 and 2 in the spleen and other secondary lymphoid organs. Aggrecanases in turn release the N-terminal LCCL domain of cochlin into the circulation. The LCCL domain promotes activation and boosting of antibacterial immunity exerted by innate immune cells (Py et al., 2013). Collagen VII binds cochlin with high affinity and appears to serve as its major anchor in secondary lymphoid organs. Loss of collagen VII evokes a dramatic loss of cochlin from these organs and, thus, an inability to mount a cochlin LCCL...
domain-mediated innate immune response toward bacteria (Nyström et al., 2018).

Intriguingly, besides the abnormalities in the innate immune system, also anomalies in adaptive immunity including presence of autoantibodies against several proteins of the DEJ have been reported (see below; Tampoia et al., 2013; Annicchiarico et al., 2015; Esposito et al., 2016). If these changes are bystanders of chronic injury or if they represent active participants in disease pathogenesis needs to be elucidated; however, observation of alleviation of disease manifestations in RDEB patients receiving various forms of immunosuppressive or immunomodulatory treatment – mild to harsh (Wagner et al., 2010; Petrof et al., 2015; Ebens et al., 2019) – could suggest active participation to disease.

### DISEASE MODIFIERS

There is a general tendency that higher levels of collagen VII expression positively correlate with milder disease in RDEB (van den Akker et al., 2009). Upon closer inspection, this tendency is not constant and the mechanisms regulating disease severity are not well understood. Unexpectedly, the same mutation combinations even in the same family can result in widely disparate disease severity (Hovnanian et al., 1997), indicating the presence of strong environmental, genetic, and epigenetic modifiers of disease in RDEB. The detailed nature of such modifiers remains for most elusive or not definitely proven, reflecting the challenges in generating robust data and statistically assessing these in a small subset of a rare disease (Kern et al., 2009). Below, we discuss molecular and cellular players that are thought to act as disease modifiers in DEB.

### Proteolytic Activities

Dystrophic epidermolysis bullosa has been associated with increased proteolytic activity in epidermal and dermal microenvironments (Eisen, 1969; Lazarus, 1972). Human and murine RDEB skin and cultured RDEB skin-derived keratinocytes and fibroblasts display alterations in the expression and the activity of several proteases and inhibitors, including matrix metalloproteinase (MMP)-1, MMP-2, MMP-3, MMP-9, MMP-7, MMP-13, MMP-14, TIMP-1, TIMP-3, cathepsin B and Z, meprins, and fetuin B (Valle and Bauer, 1980; Akasaka et al., 2021), which suggests long-lasting intrinsic changes attributed to genetics, epigenetics, cellular responsiveness, or cellular memory. However, this is not a general phenomenon observed in all studies and whether or not these activities are heighted compared to donor cells appear to depend on donors and culture conditions (Winberg et al., 1989; Küttner et al., 2013; Akasaka et al., 2021).

The variability in collagenase and gelatinase activity between individuals with DEB, together with the fact that they may degrade collagen VII, resulted in the hypothesis that MMP activity could be a disease modifier for DEB (Bodemer et al., 2003). In three brothers with RDEB with dramatically discordant severity, the abundance of MMP-1, −2, −3, and −9 was increased in both blisters and non-blistered skin, as compared to healthy control skin. Skin from the injuredly affected brother showed the highest MMP-2 and -9 activity. However, the abundance of MMP-1 was greatest in the most severely affected brother and this correlated with MMP-1 activity being uniquely detected in his blistered skin. Building on these studies, a subsequent investigation identified a SNP in the MMP-1 promoter associated with higher MMP-1 expression in the more severely affected brothers and further linked this SNP to more severe RDEB in a larger cohort (Titeux et al., 2008). Higher frequency of the specific MMP-1 SNP was also detected in RDEB compared to healthy controls in another study (Almaani et al., 2009). Analysis of a larger DEB cohort confirmed higher frequency of the MMP-1 SNP in RDEB, however, could not make a correlation between the MMP-1 SNP and more severe RDEB (Kern et al., 2009).

Early clinical trials on cancer treatment with MMP inhibitors were unsuccessful, which was due to multiple factors including low selectivity of the inhibitors, challenges with bioavailability and insufficient understanding of MMP function, and their interconnectivity in the protease web (Winer et al., 2018; Fields, 2019). Progress of optimized inhibitor synthesis and overall enhanced knowledge of protease biology has created a renewed interest in MMP targeting in multiple diseases from autoimmune diseases to cancers (Winer et al., 2018; Fields, 2019). Research for the application of MMP targeting for DEB could ride this wave. However, to avoid repeating past failures related to uninformed targeting, it is important to first gain a comprehensive understanding of the protease web in DEB and its dysregulation, as proteases are active on both sides of the disease spectrum – they are degenerative and may promote inflammation but also...
FIGURE 2 | Secondary disease mechanisms and potential disease modifiers in dystrophic epidermolysis bullosa (DEB). Disease progression in DEB is jointly driven by external and tissue-intrinsic factors. Frictional and mechanical challenges induce blistering of a pre-destabilized tissue, this together with high bacterial load trigger an exaggerated inflammatory and wound healing response. Bacteria may promote direct destruction of tissue through release of bacterial proteases. Epidermal keratinocytes, activated by impaired adhesion, injury, and inflammation, increase production of proteases, including matrix metalloproteinases (MMPs), which further evoke tissue degradation and inflammation. In addition, the keratinocytes secrete pro-inflammatory factors and trigger activation of dermal fibroblasts. Inflammatory cells – inflammatory macrophages and neutrophils – promote tissue degradation through secretion of MMPs and elastase, and activation of fibroblasts through release of, e.g., TGFβ and IL-6. Activated fibroblasts respond by increasing production of provisional and interstitial extracellular matrix (ECM) and altering the ECM architecture to a stiffer structure. Vascular leakage enhanced by the hyper-vascularized tissue may cause a transient increase in stiffness triggering stiffness response in fibroblasts. Increasing stiffness and TGFβ, which is released by multiple cell types – keratinocytes, fibroblasts, inflammatory cells, and platelets – and activated by several mechanisms – integrins, proteases, thrombospondin-1, and ROS – promote conversion of fibroblasts to contractile myofibroblasts. With time, adaptive immune activation occurs which may further enhance chronic inflammation and injury response. Collectively, these multifaceted mechanisms promote the establishment of a fragile, fibrotic, and contractile tissue that supports the growth of squamous cell carcinomas.

essential for tissue regeneration and inflammatory homeostasis. Apart from their processing and degrading the ECM, they are essential regulators of transforming growth factor (TGFβ) activity, inflammation, and angiogenesis – three other potential phenotype modifiers in RDEB.

TGFβ Activity

Increased TGFβ activity has since long been associated with fibrotic diseases (Kim et al., 2018). In a chronically injured tissue setting as in RDEB, it is contributed by both activated tissue-resident cells as well as inflammatory cells and platelets (Kiritsi and Nyström, 2018; Figure 2). TGFβ is a pleiotropic cytokine, part of the larger TGFβ superfamily. Humans have three isoforms TGFβ1-3, which signal through TGFβ receptors type I and II, and in the canonical pathway phosphorylate SMAD-2/3, which in a complex together with SMAD4 translocate to nucleus and promote gene expression. Non-canonical TGFβ signaling also occurs, activating multiple other pathways including JNK, MAPK, and AKT. TGFβ has a variety of essential roles in organ and tissue homeostasis. It exerts immunoregulatory actions, stimulates epithelial cell mobility, epithelial-to-mesenchymal transition, activation of fibroblasts, and production of ECM proteins (Ramirez et al., 2014; Hinz, 2015; Sanjabi et al., 2017; Figure 2).

TGFβ is secreted in a latent form that needs to be activated to allow for signaling through its receptors (Kim et al., 2018). Multiple mechanisms can activate TGFβ; the reliance on these mechanisms differs depending on the context and the TGFβ isoforms. These mechanisms include activation through application of mechanical forces to the latency-associated pro-peptide via RGD-binding integrins, breaking of chemical bonds via reactive oxygen species, induction of sterical shifts via binding to the ECM protein thrombospondin-1, proteolytic release via proteases including MMPs, and on lymphocytes via binding to GARP (Wang et al., 2012, 2017; Kim et al., 2018).

Recent data suggest an intricate relationship between collagen VII and TGFβ. Collagen VII can downregulate TGFβ activity in vitro and in vivo reducing scarring and it also seems to limit conversion of latent to active TGFβ (Wang et al., 2013; Nyström et al., 2013b; Akasaka et al., 2021). In a viable mouse model of severe RDEB – the collagen VII hypomorphic mouse – increased abundance of TGFβ ligands was seen in fibrotic forepaws (Fritsch et al., 2008). Healing skin wounds in the same model showed elevated canonical TGFβ signaling (Nyström et al., 2013b). Interestingly, increased TGFβ ligand expression was not disclosed in heavily scarred deformities of RDEB patients (Breitenbach et al., 2015a), whereas we observed elevated
levels of circulating TGFβ in RDEB, as well as tissue-bound TGFβ and increased canonical TGFβ signaling in human RDEB wounds (Nyström et al., 2015). Collectively, these results may reflect differences in the dependence of TGFβ during the onset and after establishment of fibrosis.

Cultured RDEB keratinocytes and fibroblasts display signs of increased TGFβ ligand expression and activity (Knaup et al., 2011; Küttner et al., 2013; Odorisio et al., 2014; Nyström et al., 2015; Atanasova et al., 2019). Keratinocytes contribute to fibrosis by facilitating TGFβ activation (Thriene et al., 2018; Akasaka et al., 2021). Furthermore, TGFβ activity appears to be strongly dependent on culture conditions and time (Akasaka et al., 2021). In RDEB fibroblasts and keratinocytes, RGD-binding integrins, ROS, thrombospondin-1, and MMP-2 and 9 have been suggested to participate in latent TGFβ activation (Atanasova et al., 2019; Chacón-Solano et al., 2019; Akasaka et al., 2021) and the reliance of these changes with culture time (Akasaka et al., 2021). Consequently, these processes and interactions can themselves be disease modulating.

A groundbreaking paper from Odorisio and colleagues linked TGFβ activity to disparate disease presentation in identical twins with RDEB (Odorisio et al., 2014). The twins synthesized and secreted similarly reduced levels of collagen VII, but the TGFβ activity in the skin differed. Low TGFβ activity was associated with milder disease and high TGFβ activity with a severe phenotype. Mechanistically, the TGFβ activity correlated inversely with the production of the small leucine-rich proteoglycan decorin (Odorisio et al., 2014). Decorin is a multi-functional protein with a large interactome and many biological functions assigned to its name (Neill et al., 2012). The focus of decorin in the context of the discordant RDEB phenotype was on its ability to sequester TGFβ and hide it from presentation to its cognate receptors (Yamaguchi et al., 1990; Odorisio et al., 2014). Subsequently, experimental pre-clinical support for the abilities of decorin to modify RDEB severity was obtained from systemic overexpression of decorin in collagen VII hypomorphic mice (Cianfarani et al., 2019). Expression of human wild-type decorin in newborn collagen VII hypomorphic mice significantly improved survival and reduced formation of mitten deformities of forepaws (Cianfarani et al., 2019). In addition, systemic injections of recombinant decorin fused to a wound target-seeking peptide (Järvinen and Ruoslahti, 2010) extended the life span of completely collagen VII-deficient mouse pups, reduced TGFβ ligand gene expression and markers associated with fibrosis (Pemmari et al., 2020).

TGFβ activity in RDEB is a double-edged sword – on the one hand, it increases fibrosis by activating fibroblasts and promoting ECM production, and on the other hand, it is a strong inductor of collagen VII expression. In the setting of tumors, it inhibits keratinocyte proliferation but also creates a stiffened and hypervascular microenvironment supporting tumor progression, the stiffness combined with immunosuppression may reduce anti-tumor immune surveillance (Ryynänen et al., 1991; Salonen et al., 2012; Martins et al., 2016; Mittapalli et al., 2016; Sanjabi et al., 2017). This makes direct TGFβ targeting for treatment of RDEB challenging.

In an attempt to limit TGFβ activity with a therapeutically relevant pharmacological approach, we treated collagen VII hypomorphic mice with the angiotensin II type 1 receptor antagonist losartan (Nyström et al., 2015). Losartan had in previous studies been shown to reduce TGFβ activity in fibrotic conditions, including other genetic ECM diseases (Habashi et al., 2006). RDEB fibroblasts responded to losartan by reducing pro-fibrotic activation status. Prolonged losartan treatment in adult collagen VII hypomorphic mice protected against progression of dermal fibrosis (Nyström et al., 2015). The outcome of this pre-clinical study prompted the initiation of a clinical trial testing the safety and efficacy of losartan for treatment of children with RDEB (EudraCT Number: 2015–003670-32, interim results presented in First World Congress on Epidermolysis Bullosa, January 19–23, 2020, London, United Kingdom (2020).

While clear effects on reduction of TGFβ activity were seen in the losartan-treated collagen VII hypomorphic mice, the most prominent was on inflammation (Nyström et al., 2015). Thus, a major benefit of losartan appeared to be from reduction of inflammation.

### Inflammation

Peripheral and systemic alterations in immune cell number and activity have been observed in RDEB (Lazarus, 1972; Tyring et al., 1989; Chopra et al., 1990, 1992). Because RDEB is associated with chronic tissue damage, inflammation could be a consequence of the damage and not involved in modifying disease manifestations. However, different lines of evidence from human and animal models suggest that DEB can be considered a systemic inflammatory disease (Annichiarico et al., 2015; Nyström et al., 2015; Figure 2).

In collagen VII hypomorphic mice, natural disease severity positively correlated with myeloid cell numbers and immunoglobulin content in fibrotic skin (Nyström et al., 2015). Transcriptomic investigation of biopsies from scarred DEB skin revealed signs of heightened inflammation of DEB skin and posited mitten deformities to be in part inflammation-driven (Breitenbach et al., 2015a, 2015b). There is an element of autoimmune in DEB. Analyses have revealed presence of antibodies against DEJ proteins including collagen VII in people with RDEB (Tampoia et al., 2013; Woodley et al., 2014; Annichiarico et al., 2015). The level of autoantibodies and level of interleukin (IL)-6 and –12 showed positive correlation with disease severity in EB and specifically in DEB (Tampoia et al., 2013; Annichiarico et al., 2015). A subsequent study indicated that the IL-6 to IL-10 ratio in serum could be used as a prognostic marker of disease severity in RDEB (Tampoia et al., 2017).

Direct evidence of inflammation as an active modifier of disease in RDEB is still missing. As suggested by proteomics, the major effect from losartan treatment of RDEB model mice was reduction of inflammation; lower tissue inflammation and lower levels of circulating IL-6 and TGFβ correlated with milder disease. Mesenchymal stromal cells (MSCs) have been of interest for treatment of RDEB, since they, on the one hand, have the ability to produce collagen VII, on the other hand are
immunomodulatory (Tolarova and Tolar, 2015). Using the same animal model, we disclosed that the major benefit of MSCs in promoting healing and reducing fibrotic aspects of RDEB wounds was from reduction in tissue inflammation (Kühl et al., 2015). Clinical trials provide a similar picture. It appears that immunosuppression and -modulation after allogeneic bone marrow transplantation, which has been evaluated as a potentially curative treatment for RDEB, confer symptom-relief effects (Wagner et al., 2010; Ebens et al., 2019). Similarly, systemic administration of allogeneic MSCs alone has improved the wellbeing of people with RDEB by modulating inflammation, without increasing collagen VII deposition (Petrof et al., 2015; Maseda et al., 2020). Since RDEB is associated with a heavily heightened risk of developing aggressive SCC, it is important to not apply general immunosuppression but to keep anti-tumor immunity intact. Therefore, more studies are needed to better understand which aspects of inflammation and immunity in RDEB correlate with and potentially drive disease.

**Angiogenesis**

While not yet functionally assessed in studies, another potential disease modifier in RDEB could be increased angiogenesis (Arbiser et al., 1998; Martins et al., 2016). As shown in the context of cancer development and growth, angiogenesis is increased in RDEB skin prior to the occurrence of tumors (Martins et al., 2016). Mechanistically, a higher vascularization of the tissue may lead to increased vascular leakage after damage, which could increase the stiffness of the tissue and evoke pro-fibrotic stiffness responses of fibroblasts that, in turn, support tumorigenesis (Figure 2).

**Therapeutic Perspective**

Like in many monogenic diseases, several factors outside the mutational status determine the disease severity and phenotypic presentation of RDEB. With disease progression, the reliance on such factors is likely to change. Because of the rarity of the disease and the large mutational spectra of RDEB, it is challenging to identify factors in detail and on a mechanistic level in the human system. Here, the use of small-animal models greatly facilitates the discovery and functional assessment of disease-modulating mechanisms, as spontaneous and genetically engineered small-animal models with identical with COL7A1 mutations display diversity in phenotypic presentations (Nyström et al., 2013a, 2015; Smith et al., 2021).

A feared complication of RDEB is the early occurrence of highly aggressive squamous cell carcinoma (Guerra et al., 2017; Cho et al., 2018). Evidence points to that the aggressiveness is in large caused by changes in the microenvironment (Ng et al., 2012; Mittapalli et al., 2016; Guerra et al., 2017). It should be emphasized that all the discussed disease modifiers have been implicated in supporting the progression of SCC in the context of RDEB-associated SCCs and other SCCs (Ng et al., 2012; Martins et al., 2016; Mittapalli et al., 2016; Föll et al., 2017; Rahmati Nezhad et al., 2021). Thus, therapies aiming at modulating the activity of these disease modifiers could create a microenvironment less supportive of malignant conversion of keratinocytes and high-risk tumor behavior.

An improved knowledge of disease-modulating mechanisms in RDEB will facilitate the development of new disease-stage specific, symptom-relief therapies. These could also have potential to improve the efficacy of curative therapies. For example, reducing inflammation and tissue-damaging processes in the dermal microenvironment should improve the take and healing of gene-corrected keratinocyte grafts that have been tested for treatment of EB (Nyström and Bruckner-Tuderman, 2016; Marinkovich and Tang, 2019). Since many of the already known disease-modulating or disease-associated mechanisms in RDEB are naturally active and important for wound healing and tissue regeneration, it is important to carefully test treatment combinations. Furthermore, as RDEB is a life-long and systemic disease, it is essential to consider the nature and actions of potential drug candidates on a system level over an extended time. For example, targeting certain aspects of inflammation, such as strong suppression of adaptive immunity, may reduce fibrosis and itch and improve wound healing but could also increase occurrence of cancers.

The aim of future research in the disease-modulating space of RDEB should be to obtain careful mechanistic knowledge and translate this into evidence-based therapies that will relieve symptoms and slow disease progression.

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