T-cell activation and bacterial infection in skin wounds of recessive dystrophic epidermolysis bullosa patients

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Funding information
U.S. Department of Defense, Grant/Award Number: W81XWH1810628

Abstract
Recessive dystrophic epidermolysis bullosa (RDEB) patients develop poorly healing skin wounds that are frequently colonized with microbiota. Because T cells play an important role in clearing such pathogens, we aimed to define the status of adaptive T cell-mediated immunity in RDEB wounds. Using a non-invasive approach for sampling of wound-associated constituents, we evaluated microbial contaminants in cellular fraction and exudates obtained from RDEB wounds. Infectivity and intracellular trafficking of inactivated *Staphylococcus aureus* was accessed in RDEB keratinocytes. *S. aureus* and microbial antigen-specific activation of RDEB wound-derived T cells were investigated by fluorescence-activated cell sorting-based immune-phenotyping and T-cell functional assays. We found that RDEB wounds and epithelial cells are most frequently infected with *Staphylococcus* sp. and *Pseudomonas* sp. and that *S. aureus* essentially infects more RDEB keratinocytes and RDEB-derived squamous cell carcinoma cells than keratinocytes from healthy donors. The RDEB wound-associated T cells contain populations of CD4+ and CD8+ peripheral memory T cells that respond to soluble microbial antigens by proliferating and secreting interferon gamma (IFNγ). Moreover, CD8+ cytotoxic T lymphocytes recognize *S. aureus*-infected RDEB keratinocytes and respond by producing interleukin-2 (IL-2) and IFNγ and degranulating and cytotoxicly killing infected cells. Prolonged exposure of RDEB-derived T cells to microbial antigens in vitro does not trigger PD-1-mediated T-cell exhaustion but induces differentiation of the CD4<sup>hi</sup> population into CD4<sup>hi</sup>CD25<sup>+</sup>FoxP3<sup>+</sup> regulatory T cells. Our data demonstrated that adaptive T cell-mediated immunity could clear infected cells from wound sites, but these effects might be inhibited by PD-1/Treg-mediated immuno-suppression in RDEB.

KEYWORDS
infection, RDEB, T-cell immunity, wound healing
1 | INTRODUCTION

Recessive dystrophic epidermolysis bullosa (RDEB) is a hereditary mechanobullous disease caused by mutations in COL7A1 gene that lead to a lack or dysfunction of type VII collagen (Col7), weak adhesion of skin layers and extreme fragility of the skin. RDEB patients commonly develop skin lesions following minor mechanical stress to the skin and blisters that progress to poorly healing wounds, having a detrimental effect on quality of life. Human skin is home to a wide array of commensal microorganisms. Most common commensal species include Staphylococcus and Micrococcus bacteria, and Malassezia fungi. These and many other commensals establish symbiotic relations with host immune system and prevent potentially harmful foreign bacteria from colonizing the skin. When skin integrity is compromised, wounds are frequently invaded by microorganisms that acquire access to abundant nutrients and become pathogenic, allowing for uncontrolled growth, proliferation and colonization of the lesions. Prior clinical investigations have shown that the pathogens that mostly commonly colonize RDEB skin lesions are Staphylococcus, Pseudomonas, Streptococcus, diphtheroids and Candida species.

Uncontrolled growth of pathogenic microorganisms at a wound site activates the immune system. Innate immunity, particularly neutrophils, plays a leading role in harnessing bacterial infection, however, activation of adaptive T cell-mediated immunity is essential for effective anti-microbial response. Bacterial peptides activate CD4+ T helper (Th) cells presented by antigen-presenting cells (APC) in the context of major histocompatibility complex (MHC) class II, enabling effective B cell-mediated immunity and enhancing clearance of the pathogens. Cytotoxic CD8+ T lymphocytes (CTL) also help to clear some microbial infection. Unlike Th cells, CTL recognize antigens (Ag) presented in the context of MHC class I (MHC I) molecules. Although CTL are commonly activated after viral infections, several studies showed that bacterial Ag could undergo MHC I restricted CD4+ and CD8+ T cells priming and activation if bacteria could escape intracellular lysis and survive inside the host. Yet, only professional antigen presenting cells (APC), such as epidermal Langerhans cells (LC) and dermal dendritic cells (DC), could activate MHC class I-restricted adaptive T cell-mediated immunity.

Little is known about the state of the adaptive immunity in RDEB skin wounds. We recently established an experimental protocol that enabled isolation of wound bed-associated cellular and molecular constituents, and we demonstrated that RDEB wound progression is associated with the accumulation of CD80+CD86+ mature APC, antigen-experienced CD45RO+ peripheral memory T cells and their transition to CD45RA+ effector cells. In this study, we investigated the status of T cell-mediated immunity in RDEB lesions and its contribution to harnessing microbial pathogens.

2 | MATERIALS AND METHODS

More detailed experimental procedures are described in Supporting Information and Methods.

2.1 | Subjects and samples

Regulatory approval for the collection of patient material has been obtained from the local ethics committee and is in accordance with local laws and regulations for the protection of human subjects. All patients received counselling pertaining to the study, and written informed consents were obtained from all patients or their legal guardians. Information on wounds and patients from which dressings were collected is provided in Supporting Information Table S3.

2.2 | Isolation of wound associated cells from wound dressing

RDEB wound-associated cells were isolated from wound-covering bandages as described previously. FD3+ pan T cells were isolated using positive selection following expansion using Immunocult media supplemented with IL-2 and T-cell activator (Stem Cell Technologies, Cambridge, MA).

2.3 | Immuno-phenotyping and fluorescence-activated cell sorting analyses

RDEB wound-derived and control T cells isolated from peripheral blood of healthy donors (about 5 × 10⁶ cells/sample) were incubated with fluorophore-labelled leukocyte marker-specific antibodies (BioLegend, San Diego, CA) for 30 min at +4°C. Fluorescence was assessed by fluorescence-activated cell sorting (FACS) on the Guava EasyCyte system and analysed using GuavaSoft 2.7 software (Millipore, Billerica, MA).

2.4 | Preparation of microbial antigens

Samples of the wound-populating microbiota were collected from transport media during isolation of cellular constituents. All collected media was filtered through 0.22 μm filters, and filters were intensively scraped in PBS. To obtain pooled soluble antigens, the resultant bulk microbiota samples or Staphylococcus aureus (Invitrogen, Carlsbad, CA) were heat inactivated at 95°C for 1 h. Concentration of the resultant antigenic mixture was assessed calorimetrically using 660 reagent (Thermo/Fisher, Waltham, MA).
2.5 | Intracellular cytokine staining

Analysis of IL-2 and IFNγ induction in T cells was performed on antigen-exposed and control T cells (1 × 10^5 cells/reaction) using ICS detection kit with brefeldin A (Golgiplug; BD Bioscience, San Jose, CA) according to the manufacturer protocol. Cytokines, CD4, CD8 and CD107a were detected using protein-specific antibodies (BioLegend). Stained cells were evaluated and analysed on Guava EasyCyte FACS System (Millipore).

2.6 | ELISA-based quantitation of cytokine release

Analysis of cytokine secretion from T cells was done using Quantikine cytokine-specific ELISA (IFNγ, IL-2, IL-17), according to manufacturer protocol (R&D Systems, Minneapolis, MN).

2.7 | T-cell stimulation

RDEB-derived and control T cells were incubated with Ag (50 μg/ml) in Immnocult-XL media (Stem Cell Technologies). At variable time points, media and cells were collected for the analyses of cytokine secretion and T-cell activation status by FACS. Micrographs of the proliferating T cells were taken during observational period.

2.8 | T-cell proliferation assay

T-cell proliferation was evaluated by the standard CFSE dilution assay. Cells were labelled with CFSE according to manufacturer protocol (Invitrogen). Labelled T cells were exposed to microbial Ag for various times, and samples were collected to define CFSE dilution (changes in mean fluorescence intensity) using Guava EasyCyte FACS System (Millipore).

2.9 | Analysis of the microbial contaminants

Microbial contaminants were identified by PCR using species-specific primers (Table S1). To evaluate microbial contaminants in exudates and media from cultured wound-derived cells, liquids were filtered through 0.22 μm nylon filters that were then crashed and boiled in H_2O for 15 min to recover DNA. To define intracellular bacteria, wound-derived cells were cultured for 48 hr in HighBind 48-well plates. Then, cells were washed twice in acid buffer (pH 3.0) to remove unbound bacteria (S. aureus) and twice in PBS. Cell pellets were resuspended in 150 μl H_2O and boiled for 15 min to recover DNA. Recovered total DNA was used for PCR (100 ng per reaction).

2.10 | Indirect immunofluorescence and confocal microscopy

Indirect immunofluorescence (IFC) was done using standard protocol. A list of antibodies, dilutions, source and catalogue number are provided in Table S2. Images were acquired on an inverted fluorescent microscope and analysed using NIS elements software (Nikon TS100; Melville, NY). To evaluate bacterial infectivity and intracellular S. aureus routing, cells were incubated in serum-free EpiLife media without calcium and magnesium (Thermo/Fisher) for 2 h. Then, inactivated, fluorescently-labelled S. aureus (Invitrogen) was added to cultures (1 × 10^6 particles/ml) for 4 h or overnight. After extensive washes, cells were fixed in 4% PFA and subjected for IFC detection of the intracellular structures using protein-specific antibodies. Slides were imaged on confocal Zeiss LSM780 NLO confocal microscope and analysed using Zeiss LSM imaging software (White Plains, NY).

2.11 | S. aureus uptake by keratinocytes

Control and RDEB-derived keratinocytes (Supporting Information Table S4) at 90% confluence were infected with AlexaFluor594 labelled inactivated S. aureus (Wood strain; Invitrogen/Molecular Probes; 6 × 10^6 bacterial particles per well). Cells were exposed to bacteria for designated periods. After infection, cells were extensively rinsed with citric acid buffer (pH 4.4) and PBS. Wells were covered with FluorSafe reagent (Calbiochem/Millipore).

2.12 | Multiplex cytokine and chemokine analysis

Secretion of pro-inflammatory chemokines (CCL2, CCL3, CCL4, CCL5, CCL11, CCL17, CCL20, CXCL1, CXCL5, CXCL9, CXCL10, CXCL11) and cytokines (TNF-α, MCP-1, IL-6, IL-8, IL-10, IL-12, IL-17, IL-18, IL-23, IL-33) in culture media from antigen-exposed and control T cells was done using LegendPlex MultiPlex ELISA (Biolegend). Control and RDEB-derived T cells were cultured in the presence or absence of soluble S. aureus antigens for 72 and 120 h. Then, culture media was clarified by subsequent centrifugation at 300 and 10 000 g to remove cells and debris, respectively. Twenty-five microlitres of clarified media were used for MultiPlex ELISA according to manufacturer protocols.

2.13 | Statistical analysis

Comparison of the data was performed using Student 2-tailed t-test; p < 0.05 was considered statistically significant.
3 | RESULTS

3.1 | Wound-associated *S. aureus* could serve as a common target for T cell-mediated immunity in RDEB skin

To define microbial species whose antigens could be recognized by the adaptive T cell-mediated immunity, we evaluated extracellular and intracellular microbial contaminants in cellular fraction and exudates obtained from RDED wounds with no documented active infection, as described previously. The samples were subjected to PCR-based analysis of the most common microbial contaminants identified in RDEB wounds with species-specific primers (Table S1). This evaluation showed that *S. aureus* is detected in more than 50% of the exudates (n = 57) with high, medium and low levels of contamination as defined by semi-quantitative PCR, whereas *Staphylococcus epidermidis* and *Pseudomonas aeruginosa* were detected in a substantially lower percent of samples with mostly high and medium levels of infection (Figure 1A,B). Other examined bacterial and fungal contaminants (Table S1) were not detected. When bandage-derived cells were placed in culture conditions, one-third of samples showed the presence of *S. aureus* either in conditioned media or intracellularly (Figure 1C,D). These findings demonstrated that *S. aureus* can infect and survive in keratinocytes at RDEB wounds.

Unlike viruses which most often utilize cell surface receptors to infect the host, bacteria are engulfed by the host cells. Macrophages detected on cell surface (Figure RDEB cells. In control keratinocytes, bacterial particles were mostly an overwhelming intracellular presence of bacterial particles in the- tially lower percent of samples with mostly high and medium lev- alization as defined by semi-quantitative PCR, whereas *Staphylococcus epidermidis* and *Pseudomonas aeruginosa* were detected in a substantially lower percent of samples with mostly high and medium levels of infection (Figure 1A,B). Other examined bacterial and fungal contaminants (Table S1) were not detected. When bandage-derived cells were placed in culture conditions, one-third of samples showed the presence of *S. aureus* either in conditioned media or intracellularly (Figure 1C,D). These findings demonstrated that *S. aureus* can infect and survive in keratinocytes at RDEB wounds.

Unlike viruses which most often utilize cell surface receptors to infect the host, bacteria are engulfed by the host cells. Macrophages and neutrophils are well-equipped for bacterial phagocytosis. However, studies have shown that other cell types, particularly keratinocytes, could engulf the bacteria. Using AlexaFluor 594-labelled *S. aureus*, we evaluated its ability to infect control and RDEB-derived keratinocytes. Our assessment showed that overnight exposure of the cells to bacteria leads to a more robust infection of RDEB- (6-times) and RDEB-associated squamous cell carcinoma (RDEB SCC)- derived (10-times) keratinocytes with *S. aureus* when compared with keratinocytes from healthy donors (Figure 1D). Using confocal microscopy to scan infected cells at 5 μm Z-position, we observed an overwhelming intracellular presence of bacterial particles in the RDEB cells. In control keratinocytes, bacterial particles were mostly detected on cell surface (Figure 2A). *S. aureus* co-localization with Rab7+ endosomes was very prominent in RDEB SCC (Figure 2B). Further analysis of the intracellular bacterial trafficking showed that *S. aureus* could be detected in association with LC3+ autophagosomes (Figure 2C) and routed toward LAMP2+ phagolysosomes (Figure 2D).

3.2 | Microbial antigen-specific CD4+ and CD8+ T cells are present in RDEB skin wounds

To evaluate whether microbial antigen-specific CD4+ and CD8+ T cells are present in RDEB skin wounds, we isolated a pool of CD3+ T cells by positive selection from bandage-derived leukocytes and propagated them in vitro. FACS-based immuno-phenotyping showed that CD4+ and CD8+ T cells are present in the CD3+ population at an average ratio of 6 to 1 (Figure 3A). Exposure of RDEB-derived T cells to wound-associated pooled microbial Ag led to clonal expansion that was most prominent after 48 h of exposure (Figure 3B). CFSE dilution assay confirmed these observations by showing that 2-day exposure to microbial Ag induced vigorous proliferation in about 30% of RDEB wound-derived T cells and did not substantially affect proliferation of control T cells in which anti-CD3/CD28 stimulation triggered robust proliferation (Figure 3C). Intracellular staining for IFN-γ combined with detection of CD107a degranulation marker confirmed microbial Ag-specific T-cell activation in RDEB T cells (n = 5). Two days of exposure induced IFN-γ expression in about 50% of RDEB T cells, whereas induction of IL-2 was detected in only 10% of cells (Figure 3D,F). Quantitative ELISA showed a significant increase in IFN-γ secretion from RDEB T cells detected at 100 pg/ml after 1 day of exposure and 275 pg/ml after 7 days of exposure (Figure 3E). However, analysis of IL-2 and IL-17 in culture media did not display any appreciable secretion of these cytokines. Population profiles showed the presence of a small CD107a+IFN-γ+ population of T cells that was identified as CD8+ T cells (Figure 3E). Collectively, these data demonstrated that RDEB wounds contain a pool of primed T cells that react to microbial antigens by proliferation, degranulation and secretion of IFN-γ.

3.3 | Wound-derived T cells can recognize *S. aureus* antigens and target infected keratinocytes in HLA-dependent manner

Considering our findings that *S. aureus* infects RDEB cells and that the presence of *S. aureus* super antigens could activate adaptive immunity, we tested the capacity of RDEB T cells to recognize and target *S. aureus*-infected keratinocytes. To set-up an IFN-γ ELISpot assay, we screened five primary RDEB-derived keratinocyte cell lines and patient-derived T cells (n = 5) for HLA-A02.1 and HLA-A03.1 expression for HLA- matching (data not shown). *S. aureus*-infected and uninfected keratinocytes were used as targets and control and RDEB T cells were used as effectors. Exposure of control or patient-derived T cells to soluble *S. aureus* Ag slightly increased a number IFN-γ spot-forming cells (SFC) (Figure 4A). Minor activation of control and RDEB T cells in response to uninfected keratinocytes was attributed to allogeneic response. Yet, a significantly greater number of SFC were detected when RDEB T cells were exposed to *S. aureus*-infected RDEB keratinocytes as compared with control (Figure 4A). T cells isolated from established RDEB lesions contained significantly greater number of IFN-γ-producing T cells than T cells from early and chronic lesions. IL-2 and IFN-γ intracellular staining combined with detection of the CD107a degranulation marker confirmed that a significantly higher percentage of cytokine-expressing RDEB T cells responded to *S. aureus* (Figure 4B). CTL capable of target- ing and killing infected keratinocytes were also identified among RDEB T cells (Figure 4C). Moreover, CTL response to infected cells appeared to be HLA-restricted. Four independent patient-derived
HLA-A02+ T cells showed higher CTL activity towards HLA-A02+ S. aureus infected targets. One of these T-cell pools, while having specific CTL activity against HLA-A02+ targets, showed unspecific allogeneic response towards HLA-A3+ targets. Conversely, HLA-A03+ patient-derived T cells were more specific against HLA-A03+ than towards HLA-A2+ infected targets (Figure 4C). Evaluation of pro-inflammatory chemokines and cytokines showed that 72 h exposure of RDEB-derived T cells to S. aureus antigens induces secretion of CCL5, CCL3 and CCL11, which coincided with down-regulation of IL-12 and IL-10. Additional 48 h of exposure did not significantly alter secretion of the chemokines, yet, a significantly higher levels of IL-10 were detected (Figure 4D). Comparison between T cell isolated from early, established and chronic wounds did not reveal substantial differences in secretion of cytokines or chemokines, although somewhat higher IL-10 levels were produced by T cells isolated from chronic wounds. Conversely, lower secretion of CCL5 and CCL3 and higher secretion of CCL4 and IL-12 were detected in antigen-exposed control T cells (Figure 4D). No changes in other examined chemokines and cytokines were detected. Collectively, these data demonstrated that S. aureus-specific memory/effector T cells capable of producing IL-2 and IFN-γ and killing infected keratinocytes are present in RDEB wounds. These data also point out that exposure of RDEB wound-derived T cells leads to the secretion of chemokines and cytokines that influence on T-cell responses.

3.4 | PD-1 and Treg-dependent mechanisms may contribute to inhibition of bacterial antigen-specific T cells in chronic RDEB wounds

In chronic infections, T cells exposure to persistent antigenic load could result in down-modulation of robust T cell effector function via activation of programmed cell death protein 1 (PD-1) and T cell exhaustion or through induction of regulatory T cells (Treg). 11–13
One-day exposure of RDEB T cells to the pooled microbial Ag led to induction of PD-1 expression on about 20% of the cells with no substantial differences in the CD8^+ T-cell population (Figure 5A). By day 2, the CD69 T-cell activation marker was induced along with PD-1 on about 30% of CD4^+ T cells. By day 3, about 60% of CD4^+ T cells were identified as CD69^+PD-1^+ cells. However by day 6, CD69 expression was diminished, yet, about 60% of CD4^+ T cells continue to express PD-1. During the observed period, no induction of CD57, a marker of PD-1-mediated T-cell exhaustion, was detected in control (not shown) and RDEB T cells (Figure 5B). Exposure to pooled S. aureus-derived Ag for 2 days also triggered prominent PD-1 expression in about 40% of RDEB T cells, a level of expression that was significantly higher than in control T cells (Figure 5C). However, 6 days of exposure of both T cell types led to PD-1 expression in more than 60% of T cells. Unlike exposure to pooled microbial Ag, exposure to S. aureus Ag led to a separation resulting in two distinct populations with lower and higher levels of CD4. The entire CD4^high population was PD-1 positive (Figure 5C). CD57 was expressed on about 2% of PD-1^+ T cells (Figure 5B). A separation of CD4^+ T cells into 2 distinct populations was detected. Suggesting...
that PD-1 could be activated by PD-1 ligand (PD-L1) expressed by infected keratinocytes, we evaluated its expression in infected and non-infected RDEB SCC. Indirect immuno-fluorescence and Western blot analyses showed no significant induction of PD-L1 in the cells after infection. However, in uninfected RDEB cells, PD-L1 was detected mostly intracellularly, whereas in infected cells, it was localized to the membrane as defined by light and confocal microscopy (Figure 5E).

Since immune-inhibitory activity of regulatory T cells (Treg) may also play a role in diminishing T cell-mediated response, we evaluated the presence of Treg in control and RDEB T cells in similar settings. FACS-based analysis showed that both control (not shown) and RDEB T cells contain a small (~2%) population of CD4+CD25+FOXP3+ Treg among CD4-gated T cells and that this percentage was increased after 6 days of exposure to microbial antigens and S. aureus Ag up to 10% in the control and 20% in the RDEB T cells (Figure 5F). The majority of CD25+FoxP3+ Treg were identified in the CD4high population (Figure 5F). When T cells were exposed to S. aureus-infected keratinocytes for 6 days, much lower (i.e., up to 7%) induction of Treg was detected in the RDEB T cells (Figure 5F). No substantial Treg induction was detected in control T cells (not shown). All detected differences between control and RDEB T cells to microbial, S. aureus and S. aureus-infected keratinocytes were significant (Figure 5G).
It is well known that skin wounds of RDEB patients are frequently colonized by bacteria and fungi that impaire wound healing. Critical colonization leads to increased wound size, pain, erythema and development of chronic wounds.\(^3,6\) Yet, the role of innate and adaptive immunity in controlling microbiota in RDEB-associated wounds remains incompletely understood. Recently, animal studies demonstrated that poor control of infection in RDEB wounds could be associated with altered macrophage activity due to lower levels of cochl in the circulation.\(^14\) However, it is plausible that inaptitude of the immune system to control wound infection could be caused by other post-infectious immunopathology in RDEB skin.

In support of this notion, we previously demonstrated that blister formation coincides with increased levels of CXCL2, CCL2, CCL4 and CCL5 and with elevated recruitment of granulocytes and T cells.\(^15\) Furthermore, we showed that progression of RDEB wounds to an established and chronic state is accompanied by reduction of macrophages, continuous recruitment of mature neutrophils, accumulation of antigen-experience APC and differentiation of peripheral memory T cells into effector T cells.\(^5\) Thus far, functional activity of adaptive T cell-mediated immunity remains uncharacterized in RDEB skin lesions. Based on our current knowledge, we proposed that RDEB wound-associated APC play a role in the acquisition of microbial antigens, T-cell priming and activation as it was shown for naïve T cells that differentiate into effector and memory T-cell pools\(^16\) and support B cell-mediated antibody response and innate immunity to clear bacterial contamination.\(^17\) Consistent with this notion, data presented here showed that RDEB wounds contain antigen-experienced CD4\(^+\) T cells that proliferate and secrete IFN\(\gamma\), a potent activator of polymorphonuclear neutrophils (PMN)\(^18\) in response to microbial antigens. Higher number of IFN\(\gamma\)-producing T cells in established RDEB wounds (Figure 4A) correlates extremely well with our prior data demonstrating accumulation of differentiated PMN to these advanced RDEB lesions.\(^5\) T cell-derived IFN\(\gamma\) is also heightens macrophage response to microbial products and is a potent inducer of macrophage M1 polarization.\(^17\) However, paucity of the macrophages in RDEB advanced skin lesions detected in our prior studies\(^5\) and impaired macrophage response in infected RDEB wounds\(^19\) suggest that T cell-derived IFN\(\gamma\) primarily activates PMN in RDEB lesions. Reduction of IFN\(\gamma\)-producing T cells in chronic RDEB wounds (Figure 4A) suggests inadequate IFN\(\gamma\)-mediated PMN stimulation and their functional impairment in chronic lesions.

Besides detecting CD4\(^+\) Th cells, we have identified a small population of CD8\(^+\) CTL among RDEB wound-associated T cells. Although the role of CTL in harnessing bacterial infection in the skin, and particularly in RDEB wounds, remains poorly defined, we suggest that bacterial Ag-specific CTL could target host cells...
in which bacteria evade intracellular destruction. Such a situation has most frequently been described for professional phagocytes, such as macrophages. Bacteria have diverse mechanisms for avoiding autophagy, including escaping to cytoplasm where they can survive and replicate. Such escape leads to the introduction of bacterial antigens into the cytoplasmic compartment and MHC class
I-mediated presentation, making bacterial Ag "visible" to the CTL. Although this mechanism was described for some bacterial and protozoan infections more than 30 years ago, recent studies have shown that keratinocytes could engulf bacterial particles and that bacteria (e.g., *S. aureus*) could escape into the cytoplasm by inhibiting autophagosome maturation.

Among others, *S. aureus*, as one of the most frequent skin colonizers, was associated with progression of keratinocyte skin tumors. Clinical studies have shown that this bacteria frequently infects pre-malignant actinic keratosis and cutaneous SCC. Our data corroborate these findings in showing significantly higher susceptibility of the RDEB SCC to *S. aureus* infection and suggesting that malignant transformation of the cells could modulate bacterial engulfment. Although the exact mechanism of *S. aureus* engulfment by SCC remains to be further elucidated, it is plausible that some sphingolipids that are associated with advanced stage of SCC play an important role in bacterial engulfment via "lipid zipping," a mechanism described for *P. aeruginosa* infection. Moreover, identified association of the *S. aureus* particles with both LAMP2+ phagolysosomes and LC3+ autophagosomes suggests that, in RDEB keratinocytes, bacteria could be routed to degradation or escape to the cytoplasm.

In the latter scenario, adaptive immunity could recognize bacterial antigens presented in the context of MHC I by specific T-cell receptors developed against MHC-antigenic peptide complex. This concept is directly supported by our data showing typical T-cell responses towards *S. aureus*-infected keratinocytes (Figure 4). Of particular note are elevated CTL activity towards infected cells and significant induction of both IFNγ and IL-2 after exposure of T cells to infected keratinocytes. This is contrary to the lack of IL-2 response to soluble microbial antigens, where only IFNγ induction and secretion were significant (Figure 3). Considering that *S. aureus*-derived super-antigens could activate T cells in orders of magnitude above conventional process, it is plausible that a full array of T-cell activities could be engaged only against MHC I-mediated presentation of bacterial Ag and that exposure of peripheral memory T cells to soluble microbial Ag is limited to IFNγ production to support PMN activity.

In chronically infected wounds, T cells are often exposed to persistent antigenic load and inflammatory signals, resulting in over-stimulation and T cell exhaust and/or Treg-mediated immunosuppression. Our experiments showing transient induction of CD69, persistent expression of PD-1 in T cells exposed to microbial Ag and infected keratinocytes and a lack of CD57 induction within 6 days of exposure suggest that PD-1-mediated exhaustion may not be the primary mechanism of the T-cell inhibition. Nevertheless, presented data showing that the CD4high T cells differentiate into CD25+FoxP3+ Treg cells upon exposure to pooled microbial and *S. aureus* Ag but do not differentiate in experiments involving infected keratinocytes (Figure 5F) suggest that soluble microbial Ag at wounded sites could induce Treg and provide a primary immunoinhibitory effect. Because the CD4high population almost uniformly expresses PD-1 (Figure SC), it is plausible that PD-1 expression synergizes with Treg-mediated inhibition and enhances suppression of the CD8+ T-cell responses, as it was shown in chronic viral infection and neoplasms. Our findings showing that RDEB wound-derived T cells could present their anti-microbial activity ex vivo within 7 days of exposure and that CD1+ Treg cell differentiation during the same period suggest that synergistic Treg-PD-1-mediated inhibition of T-cell immunity could be a conservative mechanism that extends to cutaneous bacterial infection. These findings also indicate that re-vitalization of exhausted T cells or inhibition of Treg differentiation could re-invigorate anti-bacterial T cell-mediated immunity in RDEB wounds and, potentially, prevent bacterial survival in the host and RDEB-associated SCC progression via immune-mediated elimination of infected malignant cells. Activation of T cells by microbial antigens and inhibition of this response after prolong exposure is also supported by secretion of pro-inflammatory chemokines and cytokines by *S. aureus* antigen-exposed RDEB T cells. This is illustrated by reduced secretion of the immune-inhibitory cytokine IL-10 during first 2.5 days of antigen exposure and induction of this cytokine during consecutive 2 days. This process coincides with secretion of T cell-derived CCL11. It is known that CCL11 is expressed by skin-residing T cells and that it is chemotactic to eosinophils that participate in tissue repair and remodelling. Very recent studies have shown that CCL11 increases the proportion of CD4+CD25+FoxP3+ Treg cells. This data are in agreement with our findings showing differentiation of RDEB-derived T cells towards FoxP3+ Treg upon prolong exposure to microbial / *S. aureus* antigens (Figure 5).

Although acknowledging that ex vivo and in vitro experimental settings do not recapitulate the complexity of molecular and cellular interactions at wound sites, these studies demonstrated that microbial/*S. aureus*-specific T helper and cytotoxic T cells are present in RDEB skin wounds and that CTL could target infected keratinocytes. Our data indicate that prolong exposure of wound-associated T cells to soluble microbial antigens and other molecules (e.g., CCL11) could trigger differentiation of Treg cells that could inhibit T cell-mediated immunity at RDEB wound sites.

AUTHOR CONTRIBUTIONS

Taylor Phillips, Leonie Huitema, Julio Cesar Salas-Alanis, Jouni Uttto, Vitali Alexeev, and Olga Igoucheva in conceptualization. Taylor Phillips, Leonie Huitema, Julio Cesar Salas-Alanis, Vitali Alexeev, and Olga Igoucheva involved in data curation. Taylor Phillips, Leonie Huitema, Julio Cesar Salas-Alanis, Vitali Alexeev and Olga Igoucheva involved in formal analysis. Taylor Phillips, Leonie Huitema, Vitali Alexeev, and Olga Igoucheva involved in funding acquisition and project administration. TP, Leonie Huitema, RC, Diego de los Cobos, Regina Isabella Matus Perez, Mauricio Salas-Garza, Oscar R. Fajardo-Ramirez, Franziska Ringpfeil, Julio Cesar Salas-Alanis, Vitali Alexeev, and Olga Igoucheva involved in investigation and resources. Taylor Phillips, Leonie Huitema, Julio Cesar Salas-Alanis, Vitali Alexeev, and Olga Igoucheva involved in methodology. Taylor Phillips, Leonie Huitema, Vitali Alexeev, and Olga Igoucheva involved in software. Taylor Phillips, Leonie Huitema, Vitali Alexeev, and Olga Igoucheva involved in supervision, visualization, validation...
and writing—review and editing. Leonie Huitema, Taylor Phillips, Jouni Uitto, Vitali Alexeev, and Olga Igoucheva involved in writing—original draft preparation.

ACKNOWLEDGEMENTS
We would like to thank the Dystrophic Epidermolysis Bullosa Research Association (DEBRA) of Mexico for its efforts to collect patients’ samples. Research reported in this publication was supported by the grant from Department of Defence under Award Number W81XWH1810628 to Ol.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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SUPPORTING INFORMATION

Additional supporting information may be found in the online version of the article at the publisher’s website.

FIGURE S1 Analysis of RDEB keratinocyte infectivity with Staphylococcus aureus (red) in tricolour (RGB) image (panels to the left) as shown in Figure 1d and red channel only (panels to the right)

FIGURE S2 Analyses of the infectivity and intracellular trafficking of the Staphylococcus aureus in control and RDEB keratinocytes in tricolour (RGB) image as shown in Figure 2 and in split, red, green and blue channels as indicated

FIGURE S3 Indirect immunofluorescence analysis of PD-L1 (green) expression in Staphylococcus aureus (red)-infected and non-infected RDEB keratinocytes in tricolour (RGB) image, as shown in Figure 5e and in split red and green channels as indicated

Supplementary Material S1
TABLE S1 PCR primers for detection of microbial infection
TABLE S2 List of antibodies used in the study
TABLE S3 Wound dressing information
TABLE S4 Cells used in the study

How to cite this article: Alexeev V, Huitema L, Phillips T, et al. T-cell activation and bacterial infection in skin wounds of recessive dystrophic epidermolysis bullosa patients. Exp Dermatol. 2022;31:1431-1442. doi: 10.1111/exd.14615