**Ex Vivo Culture Models of Hidradenitis Suppurativa for Defining Molecular Pathogenesis and Treatment Efficacy of Novel Drugs**

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**Abstract**—— Hidradenitis suppurativa (HS) is a complex and debilitating inflammatory skin disease for which no effective treatment is available currently. This is partly because of the lack of adequate human or animal models for defining the pathobiology of the disease. Here, we describe the development of air–liquid (A-L) interface, liquid-submersion (L-S), and bioreactor (Bio) *ex vivo* skin culture models. All three *ex vivo* platforms were effective for culturing skin samples for up to 14 days. Tissue architecture and integrity remained intact for at least 3 days for healthy skin and 14 days for HS skin. Up to day 3, no significant differences were observed in % early apoptotic cells among all three platforms. However, late apoptotic/necrotic cell death was increased in HS skin at day 3 in A-L and Bio culture. These cultures efficiently support the growth of various cells populations, including keratinocytes and immune cells. Profiling inflammatory gene signatures in HS skin from these *ex vivo* cultures showed dynamic changes in expression at day 3 and day 14. All three culture platforms were necessary to represent the inflammatory gene status of HS skin at day 0, suggesting that not all gene clusters were identically altered in each culture method. Similarly, cytokine/chemokine profiling of the supernatants from vehicle- and drug-treated *ex vivo* HS cultures again showed a better prediction of drug efficacy against HS. Overall, development of these three culture systems collectively provides a powerful tool to uncover the pathobiology of HS progression and screen various drugs against HS.

**KEY WORDS:** Hidradenitis suppurativa; *Ex vivo* models; Cytokines/chemokines; Drug screening.

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INTRODUCTION

Hidradenitis suppurativa (HS) is a chronic inflammatory skin condition with painful lesions, including nodules, abscesses, draining tracts, and fibrotic scars. This debilitating disease affects 1–4% of the Western population [1, 2]. So far, no effective treatments have been identified particularly for severe disease. The pathologic process of HS is thought to develop from occlusion and rupture of defective hair follicles and the subsequent release of contents including bacteria into the dermis [3, 4]. A complex inflammatory response by surrounding neutrophils and lymphocytes is known to accelerate the formation of abscess and causes the subsequent destruction of the pilosebaceous unit [5, 6]. Although the presence of abscesses and malodorous drainage has implicated a role of microorganisms in HS pathogenesis, only certain antimicrobial therapies have been found temporarily and/or partially effective. In addition, the risk of drug resistance is high, if administered on a long-term basis [7]. Keratinocyte hyperproliferation is also a hallmark of HS [8]. Unlike other inflammatory skin diseases, the immunobiology of HS is poorly understood, which has also hindered the development of effective therapies. Studies defining the elevation of inflammatory cytokines, such as tumor necrosis factor alpha (TNF-α), interleukins (IL)-1β and IL17A, and various others in HS patients, have provided possible therapeutic targets for emerging treatments [9–12]. This is also based on the known success of biologics against these targets for therapeutic intervention of various other inflammatory diseases [13]. The TNF-α antagonist adalimumab is currently the only biological approved by the United States Food and Drug Administration [13, 14] and by European Medicines Agency for treatment of HS [15]. The lack of optimal pre-clinical models of HS has hampered the pace of development of novel treatments to effectively target immune dysregulation and the associated pathobiology of HS disease progression [16].

In this study, we report the establishment of three different ex vivo HS skin culture models that enable efficient characterization of immune infiltrates and hyperproliferative keratinocytes in HS tissues alone and following treatment with various drugs. These models include air–liquid interface (A-L) culture, liquid submersion (L-S) culture, and culture using a three-dimensional perfusion bioreactor (Bio) platform with well-defined culture condition. L-S culture is very similar to traditional cell culture, in which the entire three-dimensional tissue is submerged in culture media. A-L culture better mimics the native tissue environment, with the apical tissue surface exposed to air and the basal surface covered by media. Both the A-L culture and the L-S culture are static culture platforms, whereas the Bio platform allows for continuous perfusion of oxygen/nutrients to mimic the human circulatory system. Thus, each of these culture methods has some benefits over others as well as some disadvantages. We demonstrate that the tissue architecture and viability of both healthy and HS skin tissues are maintained ex vivo in these models, over a period of 14 days. CD4+ and CD8+ lymphocyte populations, including the regulatory T cells and memory T cells, as well as keratinocytes and monocytes/macrophages could be characterized and enumerated by FACS analysis in these models. Importantly, we assess the feasibility of these models to test the effects of drugs on the inflammatory cytokines/chemokine signatures and signaling pathways associated with immune dysfunction in HS. Utilization of multiple ex vivo HS skin culture platforms will facilitate the development of novel therapeutics and unravel unique mechanisms underlying the disease pathogenesis.

METHODS

Human Tissue

The Institutional Review Boards of the University of Alabama at Birmingham approved the protocol (IRB-300005214) for obtaining discarded surgically removed skin tissues from Healthy and HS (Hurley stage 2 or stage 3) subjects.

Sample Processing and Ex Vivo Culture

Clinical specimen were divided into ~ 5 mm × 5 mm tissue pieces. Tissue pieces were then either (1) placed on to a 0.4-µm transwell filter (Merck Millipore) with a thin layer of extracellular matrix (ECM, 90% collagen type 1 (Advanced Biomatrix, USA) + 10% growth factor reduced Matrigel (Corning, USA)) for stability or (2) placed into the central chamber of a polydimethylsiloxane (PDMS, Krayden, USA) bioreactor containing a mixture of ECM for structural support as previously described [17]. The tissue/ECM volume within the bioreactor was penetrated with five 400-µm Teflon-coated stainless steel wires to generate through-channels for tissue perfusion. Following ECM polymerization, the wires were removed and the through-channels were filled with tissue culture media.
(1:1 mixture of X-Vivo15 and bronchial epithelial growth media (Lonza, USA) with antibiotics (MP Biomedicals, USA)). The bioreactor was then connected to a perfusion system that contained a media reservoir, peroxide-cured silicon tubing (Cole Parmer, USA), a collection reservoir, and peristaltic pump (ESI, USA), and tissue culture media was perfused through the tissue volume for 3 to 14 days (37 °C, 5% CO₂), with media changed every 3 days. Following ECM polymerization, tissue culture media was added to the transwell filters and to the bottom of the well to generate L-S (tissue fully submerged in media) and A-L (the media volume did not cover the top tissue layer) cultures. When applicable, treatment with lenalidomide (1.5 mg/mL, Selleckchem), CPI-0610 (1.5 mg/mL, Selleckchem), or vehicle control (dimethyl sulfoxide (DMSO)) occurred daily for 3 days of culture. For the end of each experiment, a portion of each tissue was fixed for histologic processing, digested with collagenase B (Roche, Switzerland) for flow cytometry analysis, or stored in RNAlater (Sigma Aldrich, USA) for RNA extraction. The supernatant culture media was collected for assaying cytokine/chemokines levels.

Histologic Processing and Analysis

Following ex vivo culture, a portion of each cultured tissue was fixed with neutral buffered formalin, processed to paraffin, and histological sections were prepared, as previously described [17]. Five-micron sections were stained with hematoxylin and eosin (H&E) to evaluate tissue morphology.

Analysis of Cell Viability via Flow Cytometry

The PE Annexin V Apoptosis Detection Kit I (BD, Germany) was used to analyze cell viability following manufacturer’s instructions. Analyses were performed on FACSsymphony A3 Cell Analyzer with FACSdiva software version 8.0.1 (BD, Germany). Data were analyzed with FlowJo 10.7.1 (Treestar, USA).

Multiparametric Flow Cytometry

The following antibodies were used for multiparametric flow cytometry for T cell analysis: anti-CD3-Alexa Fluor 700 (Clone: UCHT1); anti-CD4-PE-Cy7 (Clone: SK3); anti-FoxP3-PerCP-Cy5.5 (Clone: PCH101); anti-CD8-FITC (Clone: SK1); anti-CD-127-APC (Clone: eBioRDR5) from ebioscience (Thermo Fisher, Germany). Anti-Ki-67-Dylight350 (Clone: 1297A) from Novus (USA). Anti-CD45-PE-Cy7 (Clone: 2D1); anti-CCR7-Pacific Blue (Clone: G043H7); anti-CD45RA-BV510 (Clone: HI100); anti-CD45RO-PerCy-Cy5.5 (Clone: UCHL1); anti-CD62L-BV650 (Clone: DREG-56); anti-CD103-PE (Clone: Ber-Act8); and anti-CD154-PE/Dazzle (Clone: 24–31) from Biolegend (USA). The following antibodies were used for multiparametric flow cytometry for analysis of resident immune and structural cells: anti-CD64-PerCp-eFluor710 (Clone: 10.1); anti-CD11b-APC-Cy7 (Clone: ICRF44); anti-HLA-DR-FITC (Clone: LN3); anti-E-Cadherin-PerCP-eFluor 710 (Clone: DECMA-1); and anti-CD16-PE (Clone: eBioCB16) from ebioscience (Thermo Fisher, Germany). Anti-CD45-Pacific Blue (Clone: H130); anti-CD15-Alexa Fluor 700 (Clone: WCD3) from Biolegend (USA). Anti-PanCytokeratin-APC (Clone: C-11) and anti-Ki-67-Dylight350 (Clone: 1297A) from Novus Biologicals (USA). The Foxp3/transcription factor staining buffer set (Thermo Fisher, Germany) and the cytotoxix/cytoperm fixation/permeabilization kit (BD, Germany) were used according to the manufacturer’s protocol to stain for intracellular molecules (intracellular and cytoplasmic molecules, respectively). Analyses were performed on FACSsymphony A3 Cell Analyzer with FACSdiva software version 8.0.1 (BD, Germany). Data were analyzed with FlowJo 10.7.1 (Treestar, USA). For analysis of cell populations, initial gating on the forward scatter (FSC) vs. side scatter (SSC) plot was completed (as shown in Sup. Figs. 1 and 2) to remove dead cells and cell debris from analysis.

Expression Profiling of Inflammatory Markers

To assess the expression of inflammation-related genes, we employed the TaqMan-based human inflammation open array panel (Cat # 4475389, Thermo Fischer Scientific, USA). This panel contained 607 targets, 586 inflammatory genes, and 21 endogenous controls. For this, total RNA from tissues from various ex vivo cultures of HS skins was isolated using TRIzol reagent
A total of 2 µg of RNA was reverse transcribed into cDNA using the SuperScript® VILO™ cDNA Synthesis Kit (Cat#11754250, Life Technologies, USA). Following the reverse transcription reaction, the mixture was denatured by incubating it at 85 °C. Pre-amplification of cDNA using custom TaqMan preamp primer pools (Cat#4441856, Thermo Fischer Scientific, USA) was performed in 25 µl total volume containing 12.5 µl of TaqMan PreAmp master mixture (Cat#4391128), 2.5-µl PreAmp primer pools, 2.5-µl reverse transcription product, and the remaining volume was adjusted by nuclease-free water. Pre-amplification thermal cycling conditions include incubation of products at 95 °C for 10 min followed by 12 × cycles at 95 °C for 15 s and 4 min at 60 °C. The pre-amplification products obtained after the cycling were incubated for 10 min at 99.9 °C. A 20-fold diluted pre-amplification product was used in the final amplification reaction as per the manufacturer’s instructions. OpenArray chip containing the pre-coated primers for 607 targets was read on a 12-K Flex RT-PCR machine (Thermo Fischer Scientific, Life Technologies Corporation, Grand Island, New York). Data analysis was performed through the online available ExpressionSuite v 1.3 software (Thermo Fisher Scientific, Life Technologies Corporation, Grand Island, NY) using global gene normalization method. Bioinformatics analysis was carried out using Ingenuity Pathway analysis (IPA, Qiagen) application.

**Chemokine Profiling**

To identify the drug response on HS skin, multiplex cytokine/chemokine analysis was performed using the conditioned medium procured from various *ex vivo* cultures. Cytokine/chemokine/growth factor 45-Plex Human ProcartaPlex™ Panel 1 (Cat# EPX450-12171–901, Thermo Fisher, USA) was used. Briefly, the conditioned medium was centrifuged at 10,000 rpm for 10 min and stored at −80 °C until assayed. A total of forty-five target proteins containing (i) Th1/Th2 markers: GM-CSF, IFN gamma, IL-1 beta, IL-2, IL-4, IL-5, IL-6, IL-8, IL-12p70, IL-13, IL-18, TNF alpha, and LIF; (ii) Th9/Th17/Th22/Treg markers: IL-9, IL-10, IL-17A (CTLA-8), IL-21, IL-22, IL-23, and IL-27; (iii) Inflammatory cytokines: IFNα, IL-1α, IL-1RA, IL-7, IL-15, IL-31, and TNFβ; (iv) chemokines: eotaxin (CCL11), GR0α (CXCL1), IP-10 (CXCL10), MCP-1 (CCL2), MIP-1α (CCL3), MIP-1β (CCL4), RANTES (CCL5), SDF-1 alpha; and (v) growth factors: BDNF, EGF, FGF-2, HGF, NGF beta, PDGF-BB, PIGF-1, SCF, VEGF-A, and VEGF-D were assessed. 50 µl of condition medium was used for multiplexing on the Luminex 200 instrument (Luminex Corporation, USA) as described earlier [18].

**Fig. 1** *Ex Vivo* Culture Platforms for healthy and HS skin. Schematics of three *ex vivo* culture platforms utilized to culture human skin (top) with images of each platform (bottom). A Air–liquid interface (A-L) culture utilized 0.4-µm transwell filters in which an extracellular matrix (ECM) scaffold stabilized tissue specimen for static culture. Media was then added to the top and bottom chambers but did not fully cover the tissue so that the apical surface had exposure to air. B Liquid submersion culture (L-S) was set up in a similar method to A-L, with tissues stabilized within ECM on transwell filters; however, media was added to cover the entire tissue. C For perfusion bioreactor culture (Bio), a portion of tissue was placed into the central chamber of the polydimethylsiloxane (PDMS) bioreactor which contained an ECM scaffold for stabilization. Through-channels were generated in the tissue/ECM volume using five 400-µm Teflon-coated stainless steel wires during ECM polymerization for continuous tissue perfusion. Wires were removed following ECM polymerization, and the bioreactor was then connected to a perfusion system, in which a serum-free defined tissue culture media was perfused from a media reservoir through the tissue volume and collected in a collection reservoir.
Statistical analysis was performed using GraphPad Prism software (La Jolla, CA, USA). Data are presented as mean ± standard error of the mean unless indicated otherwise in the figure legend. A $p$-value less than 0.05 was considered statistically significant. A two-tailed unpaired Student’s $t$-test was used to evaluate statistical differences between two groups. One-way ANOVA with Tukey’s multiple comparison testing was utilized to evaluate statistical difference for data with more than two groups.

RESULTS

Tissue Integrity

Skin tissue specimen were collected from healthy individuals (without evidence of any skin disease) and patients with HS. Three culture platforms were established and tested for ex vivo maintenance of tissues. These culture platforms include air–liquid interface culture (A-L), liquid submersion culture (L-S), and perfusion bioreactor culture (Bio), which we have previously utilized for human lung tissue (Fig. 1A–C) [19]. Histologic analysis
of tissues demonstrates that all three platforms are effective for culturing these skin samples, with maintenance of tissue integrity for at least 3 days (Fig. 2A, left). When the culture period was further extended to 14 days, the healthy skin tissues showed some signs of loss of tissue architecture, whereas HS skin specimen maintained most of the tissue structure and tissue integrity when H&E stained cross-sections were analyzed (Fig. 2A, right).

**Apoptosis Analysis**

Flow cytometry data did not show significant differences in early apoptosis among the three culture platforms up to day 3. However, the percentage of early apoptotic cells was higher in HS skin culture (L-S: 32.8% ± 4.26%, Bio: 26.4% ± 2.14%) relative to healthy skin (L-S: 12.2% ± 2.9%, Bio: 9.47% ± 2.1%; p = 0.017 Bio culture) (Fig. 2B). No significant differences were observed in the percentage of late apoptotic/necrotic cells among the three platforms of healthy skin cultures, but late apoptotic/necrotic cells were significantly higher in Bio culture (3.83% ± 0.69%) of HS skin relative to A-L culture (2.29% ± 0.4%; p = 0.026) at 3 days (Fig. 2C). Interestingly, at day 14, a moderate increase in apoptotic cells was observed in the healthy skin cultures relative to HS skin cultures. When healthy and HS skin cultures were compared, a significant increase in apoptotic cells was observed in A-L culture (36.37% ± 9.27%) and Bio culture (40.8% ± 15.7%) from healthy skin when compared to A-L culture (8.9% ± 2.26%) and Bio culture (12.18% ± 3.2%) of HS skin (A-L culture: p = 0.0074, Bio culture: p = 0.0069) (Fig. 2D). Similarly, an increase in the late apoptotic population stained with Annexin V+ 7-amino-actinomycin D+ (7-AAD) was observed at day 14 relative to day 3 of healthy skin cultures. L-S culture of healthy skin contained the highest proportion of late apoptotic cells (14.5% ± 1.83%) when compared to A-L culture (7.17% ± 2.38%; p = 0.0001) and Bio culture (4.68% ± 0.15%; p < 0.0001). Additionally, when healthy and HS skin culture were compared, a significant increase in the late apoptotic population was observed in healthy skin samples when using all culture platforms (Fig. 2E).

**Characterization of Various Cell Populations**

T cell populations were evaluated following 3 days of culture. Although the CD4+ population was very high in HS skin (11.27 ± 3.86) relative to healthy skin (2.95 ± 0.168; p = 0.037) at day 0, no significant difference was observed in the proportion of CD4+ T cells between healthy and HS skin (Fig. 3A and B). Upon further analysis, Bio culture was shown to best represent the CD4+ T cell populations observed in the starting tissue of healthy skin, while a significant decrease in this population was found with A-L (2.92 ± 1.08; p = 0.011) and L-S (2.65 ± 0.93; p = 0.011) culture at day 3 relative to HS skin at day 0 (11.27 ± 3.86). Interestingly, CD4+ tissue-resident memory (TRM) cells, regulatory T cells (T_reg), and central memory T cells were expanded in all culture platforms in HS specimen when compared to their levels in the starting tissue (Fig. 3C), whereas TRM, effector, and central memory T cells were decreased in healthy skin. When CD8+ T cell populations were compared in healthy skin, the A-L and L-S culture platforms were found to best maintain similar proportions to those seen in the starting tissue. However, T_reg increased and TRM and central memory T cells decreased when using all culture platforms (Fig. 3D). In HS skin, CD8+ T cell populations remained similar to the starting tissue across culture platforms; however, slight increases in T_reg and central memory T cells, as well as slight decreases in effector memory cells, were seen with culture. When activated T cell populations were evaluated, a significant increase in activated CD4+ T cells (A-L culture: 16.83 ± 4.37, p = 0.0001; L-S culture: 12.03 ± 2.07, p = 0.035; Bio culture: 12.07 ± 3.19, p = 0.034) was observed following culture at day 3 relative to healthy skin tissues at day 0 (2.94 ± 0.62) (Fig. 3E). Similarly, activated CD8+ T cells were elevated with culture, but the number did not reach significance in either tissue type or across culture platforms (Fig. 3F). Proliferating CD4+ T cells were increased with Bio culture in HS tissue, but no other changes were observed and no changes in proliferating CD8+ T cells were found (Fig. 3G and H).

When the proportions of keratinocytes and proliferating keratinocytes were evaluated, a significant increase in keratinocytes was found in A-L culture (11.74 ± 2.17) relative to the starting tissue (4.6 ± 0.48) in HS skin (p = 0.035); however, no other significant changes were observed between tissue type or across culture platforms (Fig. 3I and Sup. Fig. 3A). Similarly, no changes were observed in the proportions of Langerhans cells (Fig. 3J and Sup. Fig. 3B). Although the proportion of macrophages and proliferating macrophages were small within both healthy and HS skin specimen, these cell populations were found to increase with Bio culture in
healthy skin (Fig. 3K and Sup. Fig. 3C). No significant changes were observed in M2 macrophages and proliferating M2 macrophages (Fig. 3L and Sup. Fig. 3D). Similarly, the proportions of neutrophils (Fig. 3M) and natural killer (NK) cells (Fig. 3N) were found to be small within both healthy and HS skin. While no significant differences were observed in the NK cell population, a significant decrease in neutrophils was observed with all culture platforms in healthy skin tissue when compared to the starting tissue (Fig. 3M).

Status of Inflammatory Markers Genes in Ex Vivo HS Models

Next, we carried out mRNA expression profiling of 586 inflammatory genes using the TaqMan-based human inflammation open array panel to assess the dynamic regulation of the expression of inflammatory genes in these culture models. Data collected from inflammatory arrays demonstrate that HS skin tissue maintain the dynamics of the inflammatory signatures in ex vivo skin culture platforms (Fig. 4A), although these changes were not identical across the three culture platforms. The expression of many inflammatory signatures was progressively elevated at day 14, as shown by the heat map (Fig. 4A). Pathway analysis using Ingenuity Pathway Analysis (IPA) showed that the osteoarthritis pathway, IL-6 signaling, pathways associated with the role of IL-17A in psoriasis, acute phase response signaling, STAT3, NFκB, and TLR signaling remained elevated in all ex vivo culture platforms up to day 14 (Fig. 4B). On the other hand, downregulated gene signatures affected pathways such as...
as LXR/RXR activation, PPARα/RXRα activation, antioxidant action of vitamin C, CREB signaling in neurons, PI3K/AKT signaling in the pathogenesis of influenza, PTEN signaling, phagosome formation, eicosanoids signaling, and crosstalk between dendritic and NK cells (Fig. 4C). Many pathways were unchanged in culture at both time-points, including the G1/S checkpoint regulation pathway, Gxc signaling, apoptosis signaling, coagulation system, endocannabinoid cancer inhibition pathway, and the ferroptosis-signaling pathway (Fig. 4D).

Inflammatory pathways, including hepatic fibrosis signaling, cardiac hypertrophy signaling, production of NO and ROS in macrophages, dendritic cell maturation, TEC kinase signaling, HER-2 signaling in breast cancer, ILK signaling, type 1 diabetes mellitus signaling, iNOS signaling, HMGB1 signaling, and IL-9 signaling (Fig. 5A), were upregulated at day 3 in various cultures, but downregulated and/or returned to basal levels by day 14 of culture. The signaling pathways, which remained highly activated at day 14 of various *ex vivo* HS skin
cultures were the osteoarthritis pathway, the pathway associated with the role of IL-17A in psoriasis, and LPS/IL-1-mediated inhibition of RXR function. Interestingly, Fcγ receptor-mediated phagocytosis in macrophage and monocytes was only upregulated in various ex vivo cultures at day 14 (Fig. 6A). Overall, the data from the inflammatory array demonstrate that these ex vivo HS skin cultures are suitable to defining the molecular pathogenesis of inflammation in HS skin.

Verification of Suitability of Ex Vivo HS Skin Culture Models for Drug Screening

To verify the suitability of these ex vivo skin cultures to predict treatment efficacy, we employed an immunomodulatory drug, lenalidomide, and a bromodomain and extraterminal (BET) inhibitor, CPI-0610, in all three ex vivo culture platforms for 3 days of culture. Both of these drugs are expected to alter the inflammatory profile by different mechanisms [20, 21]. Histopathologic evaluation using H&E staining did not reflect major differences in tissue architecture due to drug treatment or associated toxicity in healthy skin when using any of the three culture methods (Fig. 6A and Sup. Fig. 4). Moreover, no significant change in early apoptotic (Annexin V+ 7AAD−) or late apoptotic/necrotic (Annexin V+ 7AAD+) cell populations was observed in comparisons between drug-treated- and vehicle-treated groups using any of all these culture platforms at day-3 (Fig. 6B). However, an overall increase in late apoptosis/necrosis in HS tissues was observed. Nonetheless, we did not establish a dose–response relationship for any of these drugs at this stage (Fig. 6C).

Employing Luminex analysis to evaluate cytokines/chemokines and growth factors in the supernatants procured from drug- and vehicle-treated cultures, we observed that both drugs effectively blocked the secretion of various cytokines/chemokines and growth factors in the supernatants (Fig. 7). Both drugs were able to reduce the levels of Th1/Th2/Th9/Treg associated cytokines such as IL2, IL4, IL5, IL9, and IL21, while inflammatory cytokines/chemokines such as IL-8, IFN-α, MIP-1α, SDF-1α, MCP-1, GROα/KC, etc., and various growth factors, namely, PDGF-1, VEGF, and SCF, were also diminished in the supernatant procured from A-L ex vivo HS skin cultures at day 3 relative to vehicle-treated controls (Fig. 7A). Secretion of some inflammatory cytokines/chemokines such as MIP-1α, IL-1RA, IL-6, and IFN-α were also blocked in the supernatant of ex vivo HS skin from Bio culture at day 3 of culture (Fig. 7B). However, only CPI-0610 was effective in blocking many cytokines/chemokines and growth factors, as assessed in the supernatant of ex vivo HS skin tissue of L-L culture at day 3 with exception to IL-2, IL-4, MCP-1, IL-9, and VEGF (Fig. 7C). Interestingly, these were not inhibited in the supernatant of A-L ex vivo HS skin cultures at day 3. Multiple studies have also evaluated cytokines and other inflammatory proteins in lesional, peri-lesional skin wound exudate, and serum of HS patients to find out the importance of these proteins in pathogenesis and treatment [22–25].

Reconstructing a heat map from these model cultures was achieved by dividing the heatmap of HS day 0 skin detailing the expression profile of inflammatory genes in eight clusters, as shown in (Fig. 4A (iii)). Thereafter, we searched the expression profiles of all three cultures at day 3 and day 14 and identified the segments where we did not find any alteration in the expression of the clusters compared to day 0 profile. These unaltered segments were sutured to make an artificial heatmap, as shown in Fig. 4A (iii). This heatmap could be utilized to predict the impact of drug treatment on the profile of inflammatory genes in HS skin (day 0). In sum, this reconstruction of the heatmap was necessary for drug evaluation, as our culture methods provide a dynamic environment for the gene expression and many of these genes are continuously altered.

Overall, various ex vivo cultures together provide a powerful system to assess the drug effects as exemplified by demonstrating the effect of two distinct classes of mechanistically anti-inflammatory drugs.

DISCUSSION

In this study, we established three different ex vivo culture model systems for healthy and HS skin and assessed suitability of these ex vivo models to screen drugs which can affect inflammatory signaling and progression of inflammation in HS skin. Previously, Vossen and colleagues used Transwell ex vivo skin culture from healthy and HS skin [26]. Using these cultures authors showed that this methodology could be used for the ex vivo evaluation of drugs against HS. In this Transwell culture, HS skin tissue was exposed to various anti-inflammatory drugs, including prednisolone or biologics targeting TNF-α (adalimumab and infliximab), interleukin (IL)-17A (secukinumab), IL-12/23p40 (ustekinumab), or CD20 (rituximab). Of these, TNF-α inhibitors and prednisolone
were found to be the most powerful inhibitors of proinflammatory cytokines. In comparison to the previous efforts to establish skin tissue cultures, our three different types of culture platforms, namely, air–liquid interface, liquid submersion, and a more controlled perfusion bioreactor, together provide a more powerful and rigorous tool to establish *ex vivo* cultures of healthy and HS skin. In an attempt to demonstrate the suitability of these cultures to aid in understanding the molecular pathogenesis of disease progression, particularly with respect to the dynamic regulation of inflammatory cascade, we characterized various cell populations including keratinocytes and immune cells in these cultures. Our initial findings demonstrated that these *ex vivo* cultures were able to maintain a significant level of the tissue architecture and integrity at least up to day 3 of culture. Furthermore, when cultures were extended to day 14, the tissue architecture and integrity were maintained in HS skin culture, but not in healthy skin culture. To avoid any complexity in group comparisons, the longer duration culture conditions were not modified for healthy skin.

These results are primarily based on H&E staining of the cultured skin. However, a further detailed analysis is needed to assess each cell type and their ability to maintain the molecular signatures, for example, using single-cell analyses. The observed differences in healthy and HS skin cells may be due to multiple factors, particularly due to differences in metabolic signatures. In this regard, vitamin D metabolism, steroid metabolism, biological processes associated with collagen and extracellular matrix remodeling, growth and tissue factors such as tissue factor 3 (T3), growth and differentiation factor

![Image]
2 (GDF2), midkine, and clusterin which are known to play important roles in vital cellular and physiological processes and are dysregulated in HS skin [27–30]. The compromised metabolic environment of HS skin is considered to be associated with HS disease progression [4, 31].

Next, evaluation of various cell populations in these ex vivo cultures showed that Bio culture at day 3 was able to best represent the CD4+ T cell populations seen in the starting HS tissue (day 0). However, with further evaluation of CD4+ T cells, TRM, Treg, and central memory T cells were found to expand in all culture platforms in HS specimen when compared to the starting (day 0) tissue. CD8+ T cell populations remained unaltered when compared to the starting tissue across all culture platforms; however, small increases in Treg and central memory T cells, as well as decreases in effector memory cells were seen. When activated T cell populations were evaluated, an increase in activated CD4+ T and CD8+ T cells was observed in all culture platforms, but statistical significance could only be achieved for CD4+ T cells in ex vivo HS skin models from A-L (A), Bio (B), L-S (C) platforms.

Molecular profiling of inflammatory genes in these cultures showed that there are dynamic expression profiles and that a single culture platform cannot precisely represent the inflammatory status of HS skin at day 0. However, using all three ex vivo culture platforms at day 3 and day 14 could recapitulate most of the inflammatory pattern seen in HS skin. Molecular expression data from these cultures also indicate that these cultures could serve as useful models to screen anti-inflammatory drugs. In this regard, our data employing the immunomodulatory drug, lenalidomide, and the anti-inflammatory/epigenetic modulatory drug, CPI-0610 (a BET inhibitor) in these ex vivo HS models showed that many inflammatory cytokines/chemokines were downregulated by these treatments. Although, as described earlier, a single culture method was not as powerful as the results obtained from the three culture methods together.

Thus, our combined ex vivo culture models provide a unique opportunity to employ these systems to...
understand the pathobiology of HS as well as screening and development of various therapeutic agents, particularly those which are already FDA approved for other inflammatory skin conditions and could be repurposed for HS treatment.

**CONCLUSION**

We believe that these three *ex vivo* culture systems collectively provide a powerful tool for defining the pathobiology of HS and assessing drugs for suppressing inflammatory responses.

**SUPPLEMENTARY INFORMATION**

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**AUTHOR CONTRIBUTION**

Conceived and designed the experiments: J.S.D., C.R., M.S.M., M.A. Performed the experiments: K.F.G., M.P.K., J.K., R.S., Z.W., A.O., L.J., V.A., M.B.L. Analyzed the data: K.F.G., M.P.K., J.K., R.S., Z.W., A.O., L.J., V.A., M.B.L., C.A.E., M.S.M., C.R., J.S.D., M.A. Contributed reagents/materials/analysis tools: C.A.E., M.S.M., C.R., J.S.D., M.A. Wrote the paper: K.F.G., M.P.K., J.S.D., and M.A.

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**AVAILABILITY OF DATA AND MATERIALS**

Data generated/analyzed during this study are included in this published article (and its supplementary information files).

**DECLARATIONS**

**Ethical Approval and Consent to Participate** The institutional Review Board of the University of Alabama at Birmingham approved the protocol (IRB-300005214) for obtaining surgically removed discarded skin tissues from breast/tummy reduction from otherwise healthy individuals and HS patients.

**Consent of Publication** All authors agree to submit this manuscript.

**Conflict of Interest** The authors declare no competing interests.

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