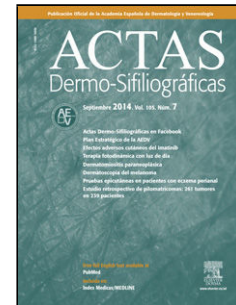


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Dermatología Práctica

Alteraciones mucocutáneas tras trasplante de progenitores hematopoyéticos: revisión y actualización de la literatura

[[Translated article]]Mucocutaneous Alterations After Hematopoietic Stem Cell Transplantation: Literature Update and Review

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RESUMEN

En los últimos años, el trasplante de progenitores hematopoyéticos (TPH) ha revolucionado el tratamiento de diversas enfermedades hematológicas y no hematológicas. Su realización no está exenta de riesgos y conlleva una tasa significativa de complicaciones, entre ellas mucocutáneas. Presentamos una revisión narrativa de las alteraciones mucocutáneas observadas tras TPH. Entre ellas, destacan la enfermedad de injerto contra receptor (EICR) aguda y crónica, cuyo diagnóstico y tratamiento puede ser notoriamente complejos. Otras patologías frecuentes son las toxicodermias y las infecciones con afectación mucocutánea. Además, diversos estudios muestran que estos individuos pueden tener una mayor tasa de neoplasias mucocutáneas. La identificación y manejo temprano de estas complicaciones, junto con un enfoque multidisciplinar, son esenciales para mejorar la calidad de vida y los resultados a largo plazo de estos pacientes. Asimismo, es recomendable el despistaje de cáncer cutáneo en estos individuos, especialmente si presentan otros factores de riesgo.

Palabras clave:

Trasplante de progenitores hematopoyéticos; Trasplante alogénico; Trasplante autólogo; Enfermedad injerto contra huésped; Cáncer cutáneo.

ABSTRACT

In recent years, hematopoietic stem cell transplantation (HSCT) has revolutionized the treatment of various hematological and non-hematological diseases. Its implementation is not stranger to risks and involves a significant rate of complications, including mucocutaneous adverse events. We present a narrative review of the mucocutaneous alterations observed after HSCT. Among these, acute and chronic graft-versus-host disease (GVHD) stand out, whose diagnosis and treatment can be challenging. Other common conditions include cutaneous adverse reactions and

infections with mucocutaneous involvement. Additionally, various studies indicate that these individuals may have a higher rate of mucocutaneous neoplasms. Early identification and management of these complications, along with a multidisciplinary approach, are essential to improving these patients' quality of life and long-term outcomes. Furthermore, it is advisable to screen for skin cancer in these individuals, especially if they have other associated risk factors.

Keywords:

Hematopoietic stem cell transplantation; Allogeneic transplant; Autologous transplant; Graft-versus-host disease; Skin cancer.

Introduction

Hematopoietic stem cell transplantation (HSCT), which includes bone marrow, peripheral blood, and umbilical cord blood transplants, involves administering healthy hematopoietic stem cells to patients with dysfunctional bone marrow due to malignant hematological diseases, bone marrow failure syndromes, or severe immunodeficiencies. HSCT can be autologous or allogeneic, depending on whether the hematopoietic cells come from the patient or a donor, respectively. Currently, HSCT is established as the treatment of choice for various severe malignant and non-malignant hematological conditions¹. According to data from the Spanish National Transplant Organization (ONT), more than 3,500 HSCTs were performed in 2022—double the number conducted in 2002².

HSCT includes a conditioning phase, in which chemotherapy and/or radiotherapy is administered to prepare the recipient's bone marrow and eliminate neoplastic cells (Fig. 1). One or two days later, the infusion phase is conducted. The patient then enters the aplasia period, during which immunity is significantly reduced due to lack of blood cell production by the bone marrow. This is a critical stage, during which the patient may experience severe anemia, bleeding, and infections. Finally, the engraftment phase occurs, beginning when the transplanted stem cells start producing new cells in the marrow. The time to engraftment varies depending on the type of transplant and patient-specific conditions, ranging from 11 to 40 days³.

After HSCT, patients may experience numerous mucocutaneous complications resulting from the transplant itself, the immunosuppressive therapy, or the graft-versus-host effect. A recent study reported that up to 45% of HSCT recipients developed some type of skin eruption within year 1 after receiving the transplant, with rates rising to 60–70% in the long term. These

complications can significantly impact patients' quality of life and may even be life-threatening⁴. This article reviews the mucocutaneous changes observed in HSCT recipients, focusing on acute and chronic graft-versus-host disease (GVHD), drug eruptions, infections, and mucocutaneous neoplasms.

Graft-Versus-Host Disease after Hematopoietic Stem Cell Transplantation

In GVHD, the immune cells from the graft (transplant) recognize the recipient (patient) as foreign and attack their tissues. GVHD is categorized into acute (aGVHD) and chronic (cGVHD) forms. Classically, they were differentiated by timing (before or after day 100 post-HSCT), but current classification is based on different pathophysiologic mechanisms and clinical presentations (Table 1)⁵⁻⁷. GVHD can affect any organ, although skin and mucous membranes are most widely involved (20–70%), and this involvement often aids diagnosis. It is an intrinsic complication of allogeneic HSCT (allo-HSCT), where donor cells differ from the recipient's, which significantly contributes to morbidity and mortality, being the most common cause of death after hematologic malignancy relapse⁶.

Acute Graft-Versus-Host Disease

aGVHD frequently first affects the skin and mucous membranes. The liver and intestines are typically affected next. The classic aGVHD triad includes skin rash, hyperbilirubinemia, and diarrhea. Traditionally, it appears and resolves within the first 100 days post-HSCT (often developing between days 30–40). However, it may also occur later (late onset), persist beyond 100 days, or recur after resolution⁷. It is graded into 4 stages: grade 1 (+): < 25% of total body surface area (TBSA) involved; grade 2 (++): 25–50% TBSA; grade 3 (+++): 50–75% TBSA; grade 4 (++++): >75% TBSA⁷. Cutaneous presentation (Fig. 2) typically begins with dysesthesias, pruritus, erythema, or edema, progressing into a morbilliform rash, often folliculotropic and trunk-predominant, then becoming confluent and spreading centrifugally. Palmar, plantar, and retroauricular involvement is a typical finding as well. Severe cases may develop epidermal detachment and blistering. The oral, genital, nasal, and ocular mucosa may also be involved, presenting as mucositis⁸.

Initial suspicion of aGVHD is based on clinical findings: the triad of rash, diarrhea, and hyperbilirubinemia—although not all signs may be present or other organs may be affected. Skin biopsy is not pathognomonic. Histologically, aGVHD can be categorized into grade I: focal vacuolar changes at the basal membrane, with sparse lymphocytic infiltrate; grade II: keratinocyte necrosis with more evident vacuolar damage; grade III: keratinocyte apoptosis, dermoepidermal junction obliteration, lichenoid dermal infiltrate; grade IV: total epidermal necrosis with dermoepidermal separation⁹.

Differential diagnosis is complex (Table 2, Fig. 2). Rashes due to drug eruptions or viral infections may mimic aGVHD. In this context, concurrent diarrhea and hyperbilirubinemia support the diagnosis of aGVHD; a new drug exposure supports drug eruption; and respiratory symptoms or PCR/serologic positivity support infection^{5, 8, 10}. Histologically, sparse eosinophils and absence of spongiosis in aGVHD may help distinguish it from drug eruptions^{11, 12}. Similarly, immunohistochemical markers such as elafin¹³ and more recently, microRNA expression¹⁴, have been proposed for aGVHD diagnosis, but validation is ongoing and diagnosis may remain uncertain despite thorough work-up.

The treatment of aGVHD depends on the grade and location of the disease. For localized grade I cutaneous aGVHD, topical corticosteroids may be used. In grade II aGVHD, systemic treatment with corticosteroids such as prednisone (1–2 mg/kg/day, although lower doses may be sufficient) is required. For corticosteroid-refractory cutaneous aGVHD, a therapeutic option is extracorporeal photopheresis, which achieves complete response rates > 80%, improves survival, and reduces mortality—especially when initiated within the first 35 days. UVA-1 and UVB phototherapy can also be beneficial in localized skin signs. Another alternative in refractory cases is the use of antithymocyte globulin (ATG). Among pharmacologic treatments, tacrolimus, mycophenolate mofetil, sirolimus, and Janus kinase inhibitors (JAK inhibitors), particularly ruxolitinib^{15, 16}, are notable. The latter has recently been approved for steroid-refractory aGVHD¹⁷.

Chronic Graft-Versus-Host Disease

cGVHD is a multisystem disease potentially affecting any organ, with skin and oral mucosa being the most commonly involved sites—up to 80% of cases. Other affected organs include liver, eyes (dry eye syndrome), gut, and lungs. Musculoskeletal and psychological involvement is also common due to the chronicity of the disease^{5, 10, 18, 19}.

Mucocutaneous cGVHD is polymorphic, affecting skin, oral, and genital mucosa (Table 3, Fig. 3). In 2014, the NIH Consensus Project proposed an organ-specific classification of cGVHD^{5, 7}.

Diagnosis is clinical and may be supported by skin biopsy. Histologically, 2 major patterns are recognized: lichenoid pattern: acanthosis, orthokeratotic and parakeratotic hyperkeratosis, band-like lymphocytic infiltrate, basal vacuolization, apoptotic keratinocytes, similar to lichen planus but with satellite cell necrosis Sclerodermiform pattern: dermal sclerosis and periadnexal fat loss, resembling morphea or lichen sclerosis Less common variants include fascial and psoriasiform patterns^{9, 20}. These findings are nonspecific and must be interpreted clinically. Biomarkers for cGVHD are under investigation²¹. Differential diagnoses include aGVHD, drug eruptions, viral infections, lichen planus, psoriasis, morphea, and systemic sclerosis^{5, 7}.

Regarding the treatment of mucocutaneous cGVHD, standardized therapeutic clinical practice guidelines are lacking, as most clinical trials exclude dermatologic outcomes. Proper skin care is essential, including general measures and emollients. First-line therapy for mild forms include topical corticosteroids. Topical calcineurin inhibitors may be used as corticosteroid-sparing agents. In more severe cases, phototherapy (UVB or UVA1), extracorporeal photopheresis, rituximab, imatinib (especially in sclerodermiform forms), and more recently, ibrutinib and ruxolitinib^{10,15,16} are widely used. For oral and genital involvement, treatment is similar, with particular benefit noted from tacrolimus mouth rinses for oral lichenoid lesions²². A multidisciplinary follow-up approach (hematology, dermatology, rheumatology, gynecology, among others) is essential given the chronicity of this disease and its potential complications.

Skin Cancer after Hematopoietic Stem Cell Transplantation

Chronic immunosuppression is clearly associated with skin cancer in solid organ transplant (SOT) recipients^{23–27}. However, the relationship between HSCT and skin cancer is less well documented. HSCT recipients have an increased risk of secondary malignancies vs the general population^{28–30}. Solid tumors develop in up to 15% of patients 15 years after HSCT with myeloablative conditioning and account for 5–10% of late deaths³⁰. Regarding skin cancer (Table 4)^{29,32,50,52–61,65–67}, published studies reveal an approximate incidence of 1% to 2% at the 5-year follow-up, 1% to 7% at the 10-year follow-up, and 6% to 10% at the 20-year follow-up³³. Several risk factors have been identified, including male sex, age at the time of HSCT, prior history of skin cancer, conditioning regimen, total body irradiation (TBI), use of voriconazole for antifungal prophylaxis, and presence of cGVHD^{29–33}. A recent systematic review and meta-analysis reported a standardized incidence ratio (SIR) for post-HSCT skin cancer of 7.21 (95%CI, 3.98–13.08), with an SIR of 2.25 (95%CI, 1.7–3.68) for autologous HSCT and 10.18 (95%CI, 5.07–20.43) for allogeneic HSCT. Risk factors for skin cancer included cGVHD—specifically for basal cell carcinoma and cutaneous squamous cell carcinoma (cSCC)—as well as male sex and voriconazole exposure for cSCC³⁴. GVHD, particularly cGVHD with mucocutaneous involvement, may be associated with increased skin cancer risk for several reasons: chronic inflammation of the skin and mucosa in cGVHD patients, and the greater need for immunosuppression in its treatment. Chronic inflammation has already been demonstrated to be an independent risk factor for skin cancer, particularly cSCC⁴⁹. Furthermore, voriconazole is a well-known phototoxic and carcinogenic drug, linked to the production of reactive oxygen species during its metabolism. Its use as antifungal prophylaxis in HSCT patients has been associated with skin cancer, particularly within the actinic keratosis–cSCC spectrum^{50–52}.

Due to the increased risk of skin cancer, several authors have recommended selective screening and ongoing dermatologic surveillance in these patients^{34–36}.

Other Mucocutaneous Alterations after HSCT (Table 5)^{37,41–44,45,47,50,70}

Patients who undergo HSCT may present with numerous mucocutaneous alterations^{44,45,47,50,70}. In addition to the already mentioned GVHD, viral infections, drug eruptions, and secondary mucocutaneous neoplasms, attention should be paid to other infections and less frequent entities. The most common mucocutaneous infections after HSCT are those caused by the varicella-zoster virus, tunnel or catheter exit-site infections, and

cutaneous signs of disseminated bacterial or fungal infections. Focal areas of bacterial cellulitis are common in the lower extremities, particularly in the context of edema due to heart failure, lymphedema, or impaired venous return^{37,38}. Molluscum contagiosum and cytomegalovirus infections are also relatively frequent. Of note, these infections may present atypically and more aggressively, given the immunosuppressed state and the polypharmacy in HSCT patients³⁹, and the possibility of post-transplant lymphoproliferative disorder associated with Epstein-Barr virus, though isolated skin lesions are rare⁴⁰.

Other dermatoses include de novo development of psoriasis⁴¹, vitiligo in patients without other signs of chronic GVHD⁴², alopecia in patients without other signs of cGVHD⁴³, or the appearance of melanocytic nevi in children⁴⁴. Recently, several cases of dermatomyositis have been reported in patients previously subjected to HSCT, some of them with severe pulmonary involvement^{45,46}, and 1 case of post-HSCT bullous pemphigoid⁴⁷. A recent study described late cutaneous alterations in children who had undergone HSCT, highlighting a high incidence of vitiligo, psoriasis/sebopsoriasis, alopecia, and nail changes—particularly in children with cGVHD, age < 10 years at the time of HSCT, and with primary immunodeficiency as the underlying condition at transplant⁴⁸.

Discussion

This review presents the main mucocutaneous changes in HSCT recipients. GVHD is the most prominent complication due to its frequency and severity. The acute form poses a complex differential diagnosis—especially vs drug eruptions and viral infections. Accurate diagnosis depends on a thorough clinical history^{5,8,10}. Treatment includes topical/systemic corticosteroids and, more recently, ruxolitinib, approved as second-line therapy¹⁵⁻¹⁷. Chronic GVHD is strikingly polymorphic, with more than 10 possible skin signs, including lichenoid and sclerodermiform types, and can also affect oral, genital, hair, or nail sites^{5,7}. This highlights the need for dermatologic evaluation and multidisciplinary care.

Conclusions

HSCT has revolutionized the treatment of various hematologic and non-hematologic diseases but carries a significant risk of mucocutaneous complications. Among them, GVHD—acute and chronic—remains a major diagnostic and therapeutic challenge. Other frequent issues include skin infections and drug eruptions. Early identification and a multidisciplinary approach are essential for improving quality of life and long-term outcomes. Skin cancer screening and photoprotection advice are also recommended, especially in at-risk individuals.

Conflicts of interest

None declared.

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Figure 1. Indications and process of HSCT. Chemo: chemotherapy; RT: radiotherapy; HSCT: hematopoietic stem cell transplantation.

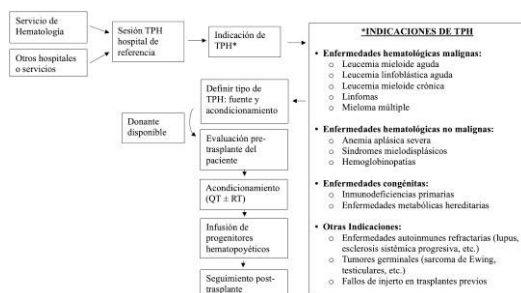


Figure 2. Acute cutaneous GVHD. A. Maculopapular rash affecting the trunk, with follicular predominance. B. Retroauricular involvement, characteristic of the disease. C. Epidermal detachment in a patient with grade 4 aGVHD. D. Maculopapular rash affecting the trunk, with follicular predominance. This patient was ultimately diagnosed with a piperacillin-tazobactam-induced drug eruption. Note the diagnostic challenge with GVHD, as this case is very similar to patient A.



Figure 3. Chronic mucocutaneous GVHD. A–C. Lichen planus–like pattern affecting the back (A), labial and buccal mucosa (B), and tongue (C). D. Nail involvement with onycholysis and pterygium formation. E. Scleroderma-like pattern with secondary hyperpigmentation. F. Fasciitis-like pattern. G. Genital mucosa involvement with sclerodermiform, lichen sclerosus–like changes. H. Poikiloderma-like pattern. I. Lichen sclerosus–like pattern with extragenital involvement on the back. J. Keratosis pilaris–like pattern. K. Psoriasiform pattern.



Table 1. GVHD Classification

Type		Time Since HSCT	Acute GVHD Signs/Symptoms ^(a)	Chronic GVHD Signs/Symptoms ^(b)
Acute GVHD	Classic	≤ 100 days	+	–
	Persistent, recurrent, or late-onset	> 100 days	+	–
Chronic GVHD	Classic lichenoid	No time limit, typically earlier	–	+
	Classic sclerodermiform	No time limit, typically later	–	+
	Other patterns of chronic GVHD	Variable	–	+
Overlap syndrome	—	—	+	+

GVHD: Graft-versus-host disease; HSCT: Hematopoietic stem cell transplantation

Source: Adapted from Jagasia et al.⁵ and Ballester-Sánchez et al.⁷.

^(a) Acute GVHD signs/symptoms: maculopapular rash, diarrhea, cholestatic hepatitis.

^(b) Chronic GVHD signs/symptoms: skin: sclerodermiform, lichenoid, or other patterns; mouth: dry syndrome, lichenoid, erosive, etc.; genital: dry syndrome, lichenoid, erosive, etc.; gastrointestinal: chronic diarrhea, abdominal pain, hepatic dysfunction; pulmonary: bronchiolitis obliterans; muscular, joint, neurological: peripheral or central neuropathy, etc.

Table 2. Differential Diagnosis of Mucocutaneous Signs of Acute GVHD^a

Differential Diagnosis	Key Clinical Clues	Additional Tests
Drug eruptions	Triggering drug exposure; absence of other suggestive GVHD symptoms. ^b	Histology: absence of adnexal involvement, presence of spongiosis and dermal eosinophils favors drug eruptions, though not specific.
	Atypical targets or epidermal detachment (SJS/TEN). Lymphadenopathy and facial edema (DRESS).	
	Pustules and fever (AGEP).	Lab test results: elevated liver enzymes, renal or cardiac dysfunction suggest DRESS; neutrophilia suggests AGEP; cholestatic liver pattern suggests GVHD.
	Retroauricular, folliculotropic trunk, or palmoplantar involvement favors GVHD.	
Viral exanthem	More common in children; associated cough, conjunctivitis, rhinorrhea, reactive lymphadenopathy; typically non-pruritic; absence of other GVHD signs.	Viral serologies, viral PCR.
Engraftment syndrome	Occurs within the first 2 weeks post-HSCT (including autologous transplants). Common features include fever, pulmonary edema, weight gain, absence of diarrhea.	Lab test results: absence of transaminitis supports engraftment syndrome.
Connective tissue autoimmune diseases (e.g., lupus, dermatomyositis, morphea, systemic sclerosis)	Signs/symptoms of lupus (mucocutaneous, joint, muscle, lung, neuropsychiatric, etc.), dermatomyositis (cutaneous, muscle), morphea (indurated plaques, usually with prior inflammatory violet halo, without HSCT history), systemic sclerosis (scleroderma, Raynaud, digital ulcers, telangiectasias, calcinosis, musculoskeletal, dysphagia, lung involvement). Absence of other GVHD signs.	Lab test results: autoantibodies may be present. Muscle enzyme elevation in dermatomyositis.

Differential Diagnosis	Key Clinical Clues	Additional Tests
Contact dermatitis	History of exposure to irritant/allergen, prior sensitization, sharply demarcated lesions, pruritus, absence of other GVHD signs.	Patch testing in allergic contact dermatitis.
Psoriasis	Well-defined erythematous-squamous plaques, Auspitz sign, predominance on extensor surfaces, scalp involvement, joint manifestations, and absence of other signs of GVHD.	Characteristic histology in psoriasis.
Lichen planus	Violaceous, pruritic, polygonal papules, often on wrists and ankles; Wickham striae; absence of GVHD signs.	Characteristic histology in lichen planus.
Zinc deficiency	Acral, periorificial dermatitis and alopecia; history of poor diet, alcoholism, or GI disease; improves with zinc supplementation; absence of GVHD signs.	Lab tests: serum zinc and alkaline phosphatase levels.

DRESS: Drug Reaction with Eosinophilia and Systemic Symptoms; GVHD: Graft-Versus-Host Disease; AGEP: Acute Generalized Exanthematous Pustulosis; SJS/TEN: Stevens-Johnson Syndrome / Toxic Epidermal Necrolysis; HSCT: Hematopoietic Stem Cell Transplantation

Source: Adapted from the authors' original work.

^a Retroauricular, palmoplantar, and folliculotropic trunk involvement are highly characteristic in suspected acute GVHD. Diarrhea or hyperbilirubinemia on lab testing may aid diagnosis.

^b Some drug eruptions such as DRESS may also cause liver involvement and fever with multiple organ features—patient context must always be evaluated holistically.

Table 3. Clinical Signs of Mucocutaneous Chronic GVHD

Skin	Hair	Nails	Oral Mucosa	Genital Mucosa
Lichen planus-like				
Lichen sclerosus-like				
Morphea-like				
Fasciitis-like				
Poikiloderma			Keratotic plaques, lichen planus-like lesions	Lichen planus-like lesions
Psoriasiform	New-onset alopecia (especially patchy > diffuse).	Roughness		Vulvovaginal stenosis or scarring
Eczematous/dyshidrotic		Thinning	Microstomia due to sclerosis Gingivitis	Fissures
SCLE-like	Can be scarring or non-scarring	Breakage		
Pityriasis rosea-like		Fragility	Mucositis, Pseudomembranes	Erosions
Ichthyosiform/keratosis pilaris-like		Onycholysis		Ulcers
Hypopigmentation	Premature gray hair discoloration	Dorsal pterygium	Ulcers	Balanitis
Hyperpigmentation		Anonychia	Xerostomia	Phimosis
Vitiligo			Mucosal atrophy Mucocele	
Angiomatoid nodules				
Calcinosis cutis				

GVHD: Graft-versus-host disease; SCLE: Subacute Cutaneous Lupus Erythematosus

Source: Own elaboration.

The “diagnostic” criteria from the NIH Consensus Project are highlighted in bold. These criteria refer to cutaneous signs that are sufficient for making a clinical diagnosis of chronic GVHD. The remaining criteria are referred to as “distinctive” for chronic GVHD, as they require the exclusion of other possible etiologies.

Table 4. Main Studies Evaluating the Risk of Actinic Keratoses and Skin Cancer After HSCT

Source	Primary Skin Cancers (n) or Patients with Primary Skin Cancers (n)	Total No. of Patients Included	Age at HSCT (years), Median (Range)	Primary Diseases	Time to Diagnosis (years), Median (Range)	Cumulative Incidence of Each Specific Skin Cancer	Identified Risk Factors
<i>Cutaneous Squamous Cell Carcinoma (cSCC)</i>							
Curtis et al. ⁵³ , 2005	19 cSCC	24,011	26.5 (3.5-61.3) (all cSCC cases)	ALL (6), AML (15), CML (14), lymphomas/MM (1), AA (17), FA (4), HGB (1) (all cSCC cases)	7 (0.9-22.9)	1.1% at 20 years	Combination of azathioprine + cyclosporine + steroids (all cSCC cases); azathioprine-containing therapies; long-term immunosuppression; chronic GVHD
Hasegawa et al. ⁵⁴ , 2005	4 cSCC	557	33.6	CML (2), NHL (1), AA (1)	4.37	ND	ND
Leisenring et al. ⁵⁵ , 2006	53 cSCC (includes mucosal)	211	41.6 (6.8-71.4) (skin and mucosal SCC)	Hematological/marrow failure (10), malignant hematological disease (84), other malignant neoplasms (1) (skin and mucosal SCC)	6.3 (0.3-24.8)	3.5% at 20 years	Acute GVHD, chronic GVHD, younger age at transplant (< 10 years)
Gallagher and Forrest ⁵⁵ , 2007	4 cSCC	926	49	CML (1), AML (1), MDS (1), NHL (1)	2.1	ND	ND
Rizzo et al. ²⁹ , 2009	19 cSCC	28,874	ND	ND	ND	ND	Chronic GVHD, male sex
Yokota et al. ³² , 2012	1 cSCC	2,062	46	CML	1.6	ND	ND
Wojenski et al. ⁵⁷ , 2015	27 cSCC	381	55 (39-71)	Most common were AML (7), CLL (9), and MDS (6)	ND	19% at 5 years	Male gender, underlying primary malignancy of CLL, transplant age, pre-HSCT skin cancer, extracorporeal photopheresis, UV therapy
Lupo-Stanghellini et al. ⁵² , 2016	6 cSCC	302	ND	ND	3.5 (0.9-20) (both cSCC and BCC)	3.2% at 3 years and 6.2% at 5 years (both cSCC and BCC)	Voriconazole
Omland et al. ⁵⁸ , 2016	4 cSCC (2 allogeneic HSCT and 2	3,302	ND	ND	ND	ND	ND

	autologous HSCT)						
Kuklinski et al. ⁵⁰ , 2017	78 cSCC	2,638	ND	ND	ND	ND	Chronic GVHD, male sex, voriconazole
Tanaka et al. ⁵⁹ , 2017	4 (all oral squamous cell carcinomas)	1,060	ND	ND	ND	24.8% at 2 years (all oral cases)	ND
Wu et al. ⁶⁰ , 2019	79 cSCC	1,974	58.1	AML, ALL, CML, CLL, lymphomas, others	ND	IRR, 9.8; 95%CI, 7.7-12.3	Age, CLL, chronic GVHD
Scott et al. ⁶¹ , 2020	62 cSCC	872	ND	AML, MPD, ALL, CLL, plasma cell disorders, CML, lymphomas, other non-malignant disorders	ND	12.3% at 5 years; 95% CI, 8.5-16.3	Chronic GVHD, Fitzpatrick skin type I
Cutaneous Squamous Cell Carcinoma (cSCC)							
Gruber et al. ⁶² , 2024	17 cSCC (includes 3 oral and 1 genital)	266	ND	AML	ND	4.2% [95% CI (2.2, 7.2)] and 8.1% [95% CI (4.6, 12.8)] at 10 and 15 years, respectively	ND
Squamous Cell Carcinoma In Situ (SCCis)							
Gruber et al. ⁶² , 2024	8 (includes 1 genital)	266	ND	AML	ND	ND	ND
Basal Cell Carcinoma (BCC)							
Hasegawa et al. ⁵⁴ , 2005	5 BCC	557	39.8	ALL (2), CML (2), NHL (1)	7.3	ND	ND
Leisenring et al. ⁵⁵ , 2006	201 BCC	211	38.1 (2.9-71.3)	Hematological/marrow failure (7), hematological cancer (150), other malignant neoplasms (1)	7.9 (0.5-30.2)	6.5% at 20 years	TBI, fair skin color, chronic GVHD, younger age at transplant (< 10 years), leukemia/lymphomas/blood or malignant bone marrow disease as primary diagnosis
Basal Cell Carcinoma (BCC)							
Gallagher and Forrest ⁵⁶ , 2007	8 BCC	926	41	CML (3), ALL (1), AML (1), MDS (1), MM (1), NHL (1)	7.6	ND	ND
Borgmann et al. ⁶³ , 2008	1 BCC	490	7.8	ALL	20.3	ND	ND
Schwartz et al. ⁶⁴ , 2009	282 BCC	6,306	ND	ND	ND	ND	TBI with higher risk for younger ages (< 10 years) at transplant and no excess risk for ages > 40 years at transplant, fair skin color for patients who had not

								undergone TBI, chronic GVHD in patients without TBI
Yokota et al.32, 2012	3 BCC	2,062	40 (17-50)	AML (2), ALL (1)	8.4 (7.1-17.6)	ND	ND	
Lupo-Stanghellini et al.52, 2016	19 BCC	302	ND	ND	3.5 (0.9-20) (both cSCC and BCC)	3.2% at 3 years and 6.2% at 5 years (both cSCC and BCC)	ND	
Omland et al.58, 2016	24 BCC (7 allogeneic HSCT and 17 autologous HSCT)	3,302	ND	ND	ND	ND	ND	
Kuklinski et al.50, 2017	35 BCC	2,638	ND	ND	ND	ND	Chronic GVHD, male sex, voriconazole	
Wu et al.60, 2019	54 BCC	1,974	54.5	AML, ALL, CML, CLL, lymphomas, others	ND	IRR, 2.5; 95%CI, 1.9-3.2	CLL, reduced-intensity conditioning, acute GVHD, chronic GVHD	
<i>Basal Cell Carcinoma (BCC)</i>								
Scott et al.61, 2020	62 BCC	872	ND	AML, MPD, ALL, CLL, plasma cell disorders, CML, lymphomas, other non-malignant disorders	ND	9.1% at 5 years; 95% CI, 5.4-12.8	Age, phototherapy exposure prior to allogeneic HSCT	
<i>Melanoma</i>								
Baker et al.65, 2003	8 melanomas	3,372	ND	ND	ND	O:E, 8:0.96, SIR, 8.3 (95% CI, 3.6-15.1), SIR, 6.7	ND	
Curtis et al.53, 2005	22 melanomas	24,011	ND	ND	ND	ND	ND	
Brown et al.66, 2005	5 melanomas	605	ND	ND	North (O:3, 5:0.85)	ND	ND	
Rizzo et al.29, 2009	18 melanomas	28,874	ND	ND	1-to-4 year group (< 1 to > 10 years)	O:E ratio, 3.47, SIR, 1.5	T-cell depletion, TBI, short latency period (< 1 year), female sex	
Yokota et al.32, 2012	1 melanoma	2,062	51	NHL	2.1	ND	ND	
Mahindra et al.67, 2015	19 melanomas	4,161	ND	ND	SIR 3.58 (CI, 1.82-6.29)	ND	ND	
Omland et al.58, 2016	6 melanomas (2 allogeneic HSCT and 4	3,302	ND	ND	ND	ND	ND	

		autologous HSCT)						
Tanaka et al. ⁵⁹ , 2017	1 melanoma	1,060	ND	ND	ND	ND	ND	ND
Wu et al. ⁶⁰ , 2019	11 melanomas	1,974	48.6	AML, ALL, CML, CLL, lymphomas, others	ND	IRR, 3.3; 95%CI, 1.7-5.9	None	

AA: Aplastic Anemia; FA: Fanconi Anemia; ALL: Acute Lymphoblastic Leukemia; AML: Acute Myeloid Leukemia; BCC: Basal Cell Carcinoma; CI: Confidence Interval; CLL: Chronic Lymphatic Leukemia; CML: Chronic Myeloid Leukemia; cSCC: Cutaneous Squamous Cell Carcinoma; GVHD: Graft-versus-Host Disease; HGB: Hemoglobinopathies; HSCT: Hematopoietic Stem Cell Transplantation; IRR: Incidence Rate Ratio; MDS: Myelodysplastic Syndrome; MM: Multiple Myeloma; MPD: myeloproliferative disorder; ND: Not Described; NHL: Non-Hodgkin Lymphomas; SIR: Standardized Incidence Ratio; TBI: Total Body Irradiation

Table 5. Main Studies Evaluating Mucocutaneous Alterations (Other than GVHD) After HSCT

Author/Year	Sample Size	Study Type and Conditions Evaluated	Results
Canninga-van Dijk et al. ³⁷ , 2003	ND	Narrative Review - All types of indications	<p>Patients undergoing HSCT, in addition to viral rashes, are at higher risk of bacterial infections (especially <i>Staphylococcus aureus</i> pyoderma), herpes simplex and varicella-zoster virus infections, cytomegalovirus reactivation, molluscum contagiosum, or fungal infections.</p> <p>- Their presentation can be atypical and more aggressive given the context.</p> <p>- Of note, the possibility of EBV-associated post-transplant lymphoma, although presentation with isolated cutaneous lesions is rare.</p>
Mabuchi et al. ⁴¹ , 2012	1	Isolated Case Report - Single-center - NHL	<p>Psoriasis can develop de novo after HSCT from a non-psoriatic donor.^a In this case, 10 years after allogeneic HSCT in a patient with NHL.</p>
Khalil et al. ⁶⁸ , 2014	92	- Retrospective Cohort Study - Single-center - Non-malignant diseases	<p>Six patients (6.5%) were diagnosed with vitiligo unrelated to chronic GVHD, 6 (6.5%) with autoimmune hemolytic anemia, 6 (6.5%) with idiopathic thrombocytopenia, 3 (3.3%) with mild leukopenia, 2 (2.2%) with aplastic anemia, and one (1.1%) with autoimmune thyroid disease.</p> <p>- Autoimmune complications were more frequent in patients who underwent HSCT for metabolic disorders.</p>
Kato et al. ⁴⁷ , 2015	1	Isolated Case Report - Single-center - T-cell Lymphoblastic Leukemia	<p>- HSCT could be a risk factor for bullous pemphigoid in patients undergoing HSCT. Therefore, in cases of suspicious clinical presentation, it should be ruled out with clinical examination, biopsy, and serology for anti-epidermal basement membrane antibodies.</p>
Bae et al. ⁴² , 2016	2,457 HSCT recipients vs 8,241 controls	- Retrospective Cohort Study - Multi-center - All diagnoses	<p>- An association with vitiligo was found in HSCT patients, independently of the presence of GVHD and higher than in the control group.</p> <p>- The risk factors most associated with the development of vitiligo were allogeneic HSCT and bone marrow as the source.</p>

Song et al. ⁴⁴ , 2017	85 HSCT recipients vs 85 controls	Prospective Cohort Study	- Children who underwent HSCT had significantly more nevi than controls (median [range]: 44 (0-150) vs. 11 (0-94), p = 0.0001).
		- Single-center	- Children with HSCT had significantly more nevi > 5 mm in diameter and more atypical nevi than controls.
Bresters et al. ⁴³ , 2017	263	- All diagnoses	- Factors associated with a higher number of nevi included malignant indication for HSCT, pre-transplant chemo, TBI exposure, and myeloablative conditioning.
		Retrospective Cohort Study	- The percentage of permanent alopecia was 15.6% (41/263 patients).
Huang et al. ⁴⁸ , 2018	85	- Single-center	- A conditioning regimen with busulfan and busulfan plus fludarabine (OR, 5.7 [CI, 2.5-12.7] and OR; 7.4 [CI, 3.3-16.2], respectively) was the main risk factor and was associated with alopecia regardless on the presence of acute/chronic GVHD.
		- All diagnoses	- 14% (n = 12) of patients developed vitiligo; 16% (n = 14) developed psoriasis/sebopsoriasis; 25% (n = 21) developed alopecia; and 6% (n = 5) developed nail alterations.
Miyagi et al. ⁴⁵ , 2023	2	Retrospective Cohort Study	- Factors significantly associated with vitiligo, independent of GVHD, included an indication of primary immunodeficiency and younger age at transplant (< 10 years).
		- Single-center	- Factors significantly associated with alopecia, independent of GVHD, were busulfan conditioning and a family history of early-onset androgenic alopecia.
		- All diagnoses	- The only risk factor identified for nail alterations was a history of chronic GVHD.
		- Two Case Reports and Literature Review	- Dermatomyositis can be a late complication of HSCT, and it's essential to rule it out in patients undergoing HSCT who present with compatible cutaneous and muscular symptoms.
		- Single-center	

GVHD: Graft-versus-Host Disease; CI: Confidence Interval; NHL: Non-Hodgkin Lymphoma; ND: Not Described; OR: Odds Ratio; Chemo: Chemotherapy; TBI: Total Body Irradiation; HSCT: Hematopoietic Stem Cell Transplantation

^a Previously, 2 cases of psoriasis after HSCT had been described in patients who received a transplant from a psoriatic donor.

TRADUCCIÓN DE LA FIGURA 1

(negro: español · azul: inglés)

Servicio de hematología

Otros hospitales o servicios

Sesión TPH hospital de referencia

Indicación de TPH

Donante disponible

Definit tipo de TPH: fuente y acondicionamiento

Evaluación pre-transplatent del paciente

Acondicionamiento (QT \pm RT)

Infusión de progenitores hematopoyéticos

Seguimiento post-transplante

Indicaciones de TPH

Leucemia mieloide aguda

Leucemia linfoblástica aguda

Leucemia mieloide crónica

Linfomas

Mieloma múltiple

Enfermedades hematológicas no malignas

Anemia aplásica severa

Síndromes mielodisplásicos

Hemoglobinopatías

Enfermedades congénitas

Inmunodeficiencias primarias

Enfermedades metabólicas hereditarias

Otras indicaciones

Enfermedades autoinmunes refractarias (lupus, esclerosis sistémica progresiva, etc)

Tumores germinales (sacroma de Ewing, testiculares, etc.)

Fallos de injerto en trasplantes previos

Hematology Department

Other Hospitals or Services

HSCT Session at Reference Hospital

HSCT Indication

Donor Availability

Define HSCT Type: Source and Conditioning

Patient Pre-transplant Evaluation

Conditioning (Chemo \pm RT)

Hematopoietic Progenitor Infusion

Post-transplant Follow-up

HSCT Indications

Acute Myeloid Leukemia

Acute Lymphoblastic Leukemia

Chronic Myeloid Leukemia

Lymphomas

Multiple Myeloma

Non-Malignant Hematological Diseases

Severe Aplastic Anemia

Myelodysplastic Syndromes

Hemoglobinopathies

Congenital Diseases

Primary Immunodeficiencies

Hereditary Metabolic Diseases

Other Indications

Refractory Autoimmune Diseases (lupus, progressive systemic sclerosis, etc.)

Germ Cell Tumors (Ewing's sarcoma, testicular, etc.)

Graft Failure in Previous Transplants