Severe cutaneous adverse reactions to drugs

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During the past decade, major advances have been made in the accurate diagnosis of severe cutaneous adverse reactions (SCARs) to drugs, management of their manifestations, and identification of their pathogenetic mechanisms and at-risk populations. Early recognition and diagnosis of SCARs are key in the identification of culprit drugs. SCARs are potentially life threatening, and associated with various clinical patterns and morbidity during the acute stage of Stevens-Johnson syndrome and toxic epidermal necrolysis, drug reactions with eosinophilia and systemic symptoms, and acute generalised exanthematous pustulosis. Early drug withdrawal is mandatory in all SCARs. Physicians’ knowledge is essential to the improvement of diagnosis and management, and in the limitation and prevention of long-term sequelae. This Seminar provides the tools to help physicians in their clinical approach and investigations of SCARs.

Introduction
Severe cutaneous adverse reactions (SCARs) to drugs are associated with morbidity, mortality, health-care costs, and drug development challenges. SCARs to drugs cover a broad spectrum of entities mainly consisting of Stevens-Johnson syndrome and toxic epidermal necrolysis, and drug reaction with eosinophilia and systemic symptoms (DRESS) syndrome.1 Because of the extensive eruption or the possibility of systemic symptoms, physicians also consider acute generalised exanthematous pustulosis (AGEP) a SCAR. This Seminar focuses on these three main entities.

Despite their low annual incidence, SCARs, especially Stevens-Johnson syndrome, toxic epidermal necrolysis and DRESS syndrome, can be life threatening and responsible for severe, potentially chronic sequelae. The incidence of Stevens-Johnson syndrome and toxic epidermal necrolysis is estimated to be two per 1 million people, whereas the incidence of DRESS syndrome in new users of antiepileptic drugs (eg, carbamazepine or phenytoin) is estimated to be one per 1000 to one per 10 000.2

Although they are rare, physicians need to be able to recognise SCARs to enable early drug withdrawal and appropriate management.

Classification and diagnosis
SCAR classification tools and adequate identification of SCARs have been widely emphasised as being key to identification and assessment of the potential culprit drug (table 1).14 During the past decade, several retrospective validation scores have been developed by a European network of SCAR experts, such as EuroSCAR and RegiSCAR (appendix pp 1–2).12

Stevens-Johnson syndrome and toxic epidermal necrolysis
Stevens-Johnson syndrome and toxic epidermal necrolysis are considered variants of epidermal necrolysis. They occur 4–28 days after drug exposure. Clinical classification is defined by the extent of body surface area with skin detachment—ie, Stevens-Johnson syndrome in cases of less than 10% skin detachment, toxic epidermal necrolysis in cases of 30% or greater, and Stevens-Johnson syndrome—
toxic epidermal necrolysis (SJS–TEN) for anything in between.1 In about 30% of cases of Stevens-Johnson syndrome and toxic epidermal necrolysis, no causative drug is identified,1,15 and in 15%, drug responsibility is deemed unlikely.16 Mycoplasma pneumoniae has been associated with Stevens-Johnson syndrome and toxic epidermal necrolysis, mainly in children.17 General physical deterioration, fever, influenza-like illness, ocular symptoms, ear, nose, and throat (ENT) events, and skin pain frequently precede dermatological manifestations, and are key in early diagnosis.18 Initially, the eruption is distributed on the face, upper trunk (appendix p 7), and proximal extremities,1 whereas distal portions of upper and lower limbs are relatively spared.1 Initial lesions are characterised as erythematous, irregularly shaped, dusky-red macules. Atypical target lesions with dark centres can often be observed without the typical three concentric
Table 1: Main clinical and histological characteristics of SCARs

<table>
<thead>
<tr>
<th>Drug-to-SCAR interval</th>
<th>General symptoms*</th>
<th>Skin features</th>
<th>Laboratory values</th>
<th>Main organs involved</th>
<th>Severity score</th>
<th>Score system for classification</th>
<th>Histological features</th>
</tr>
</thead>
<tbody>
<tr>
<td>SJS and TEN†‡ 4–28 days</td>
<td>Fever ≥38°C, influenza-like syndrome, respiratory tract symptoms</td>
<td>Blisters, large skin detachment, confluent erythema, atypical target lesions, purpura, Nikolsky’s sign, skin detachment</td>
<td>Lymphopenia, transitory neutropenia, mild cytolytic, renal impairment</td>
<td>Ear, nose, and throat, lung, intestinal tract, liver, kidney</td>
<td>SCORTEN!</td>
<td>No‡</td>
<td>Full-thickness epidermal necrosis, focal adnexal necrosis, necrotic keratinocytes, mild mononuclear cell dermal infiltrate, negative direct immunofluorescence test</td>
</tr>
<tr>
<td>DRESS syndrome† 2–6 weeks</td>
<td>Fever ≥38°C, influenza-like syndrome</td>
<td>Maculopapular rash, erythema, facial or extremity oedema, purpura, pustules, focal mononuclear mucocutaneous membrane involvement</td>
<td>Eosinophilia &gt;700 cells per µL, atypical lymphocytes, elevated transaminase concentration, impaired renal function, herpesvirus family reactivation (HHV6, HHV7, EBV, CMV), parvovirus B19 reactivation</td>
<td>Liver, kidney, lung, muscle, heart, pancreas, medulla, lymph nodes at two or more sites</td>
<td>None</td>
<td>Yes</td>
<td>Lichenoid infiltrate or eczematous pattern (spongiosis, oedema), focal necrotic keratinocytes, mononuclear infiltrate, focal eosinophil and neutrophil infiltrates, mild vasculitis</td>
</tr>
<tr>
<td>AGEP (2–13) 1–11 days</td>
<td>Fever ≥38°C</td>
<td>Intertriginous erythema, oedema, widespread non-follicular sterile pustules, post-pustular pinpoint desquamation, Nikolsky’s sign, rare oral mucocutaneous membrane involvement</td>
<td>Hyperleukocytosis, neutrophils ≥7000 cells per µL, mild eosinophilia</td>
<td>Rare: liver, lung</td>
<td>None</td>
<td>Yes</td>
<td>Subcorneal or intraepidermal spongiform or non-spongiform pustules with or without papillary oedema, focal necrotic keratinocytes, neutrophilic sometimes with eosinophils, mild vasculitis</td>
</tr>
</tbody>
</table>

SCAR—severe cutaneous adverse reaction. SJS—Stevens-Johnson syndrome. TEN—toxic epidermal necrolysis. SCORTEN—Scoring for Cutaneous Toxic Epidermal Necrolysis. DRESS—drug reaction with eosinophilia and systemic symptoms. HHV—human herpesvirus. EBV—Epstein-Barr virus. CMV—cytomegalovirus. AGEP—acute generalised exanthematous pustulosis. *General symptoms can precede or occur at the same time as skin manifestations. †See appendix. ‡Not published.

Rings of erythema multiforme major (appendix pp 7–8). Necrotic lesion confluence leads to extensive erythema, flaccid blisters, and large epidermal sheets, revealing areas of red dermis (appendix p 9). Nikolsky’s sign—when the epidermis sloughs off under lateral pressure—is positive on erythematous areas (appendix p 10). Two or more mucous membranes are involved in 80% of cases, often preceding skin lesions. Erythema, blisters, or erosions involve the nasopharynx, oropharynx, eyes, genitalia, or anus mucous membranes, and occur during the early stage associated with pain and dysfunction (appendix pp 11–12). The lips can develop a vermillion border, and greyish-white pseudomembranes coat oral-cavity haemorrhagic erosions, with crusts being the main lesions (appendix p 13). Conjunctival lesions, including hyperaemia, erosions, chemosis, photophobia, and tearing comprise eye involvement. Severe forms lead to corneal ulceration, anterior uveitis, or purulent conjunctivitis (table 1). Disease progression is time limited (7–10 days).

Visceral involvement associated with Stevens-Johnson syndrome and toxic epidermal necrolysis consists of transient liver or renal enzyme increases or bronchial and digestive tract epithelial necrosis. Although rare, specific acute visceral failures in Stevens-Johnson syndrome and toxic epidermal necrolysis should be suspected and documented after eliminating bacterial or viral super-infection. No specific score or diagnostic test is available for the diagnosis of Stevens-Johnson syndrome and toxic epidermal necrolysis. The diagnosis mainly relies on identification of a broad range of clinical signs and symptoms and histological tests (tables 1, 2). Full-thickness epidermal necrosis (appendix p 14) and a negative direct immunofluorescence test are mandatory for diagnosis. Differential diagnoses include erythema multiforme major, linear IgA bullous dermatosis (appendix p 15; spontaneous or drug-related), generalised fixed drug eruption (appendix p 16), superficial burns, cytotoxic drug (eg, methotrexate) toxicity, and acute graft-versus-host disease (table 2).

### DRESS syndrome

Since its description in 1996, the R of DRESS syndrome has changed from rash to reaction. It is also known as hypersensitivity syndrome or drug-induced hypersensitivity syndrome. It usually begins 2–6 weeks after drug exposure. The difficulty in diagnosing DRESS syndrome is mainly due to its complex natural course and heterogeneous clinical presentation, involving visceral symptoms with or without dermatological involvement and biological abnormalities. The prodromal stage,
including fever, lymphadenopathy, influenza-like symptoms, burning pain, or pruritus, can precede the skin eruption by up to 2 weeks. Clinical dermatological symptoms consist of facial oedema, erythroderma, distal oedema, purpura, pustules, and sometimes focal mucosal involvement (table 1; appendix pp 17–18). To reduce misdiagnosis, several investigations are recommended (table 2), such as those to identify blood abnormalities indicating visceral involvement, virus reactivation, hypereosinophilia, and rare respiratory failure. Heart involvement—inie, myocarditis and pericarditis—with electrocardiogram, CT scan, or cardiac enzyme abnormalities, can be fatal. Poor prognoses are also associated with rare visceral effects that can be neurological, muscular, haemophagocytic, or pancreatic. To rule out misdiagnosis, several investigations are recommended (table 2), such as those to identify blood abnormalities indicating visceral involvement, virus reactivation, hypereosinophilia, and rare respiratory failure. Heart involvement—inie, myocarditis and pericarditis—with electrocardiogram, CT scan, or cardiac enzyme abnormalities, can be fatal. Poor prognoses are also associated with rare visceral effects that can be neurological, muscular, haemophagocytic, or pancreatic.

### Table 2: Severe cutaneous adverse reactions: suggested confirmation tests and main differential diagnoses

<table>
<thead>
<tr>
<th>Suggested confirmation tests</th>
<th>Differential diagnosis</th>
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<tbody>
<tr>
<td><strong>Clinical diagnosis</strong></td>
<td><strong>Tests</strong></td>
</tr>
<tr>
<td><strong>SJS and TEN</strong></td>
<td><strong>Histology</strong></td>
</tr>
<tr>
<td>Erythema multiforme major, <em>Mycoplasma pneumoniae</em> infection*, coxsackievirus infection, linear IgA bullous dermatosis, generalised bullous fixed drug eruption, methotretate toxicity, graft-vs-host disease, staphylococcal skin scalded syndrome, systemic lupus erythematosus, pemphigus</td>
<td><em>Mycoplasma pneumoniae</em> serology and PCR, coxsackievirus serology and PCR, direct and indirect immunofluorescence, anti-epidermal basement membrane zone and intercellular antibodies, methotretate plasma concentration, antinuclear antibodies, anti-Ro-SSA antibodies</td>
</tr>
<tr>
<td><strong>DRESS syndrome</strong></td>
<td>Blood cell counts, liver enzyme tests, serum urea, creatatinemia, proteinuria, arterial oxygen saturation, blood gas, chest radiograph, CT scan, heart assessments (troponin I, electrocardiogram, echocardiography)</td>
</tr>
<tr>
<td>T-cell lymphoma, pseudolymphoma, viral rash</td>
<td>Sézary cells (ie, atypical T cells found in Sézary disease), cutaneous clonal T-cell rearrangements, viral serology</td>
</tr>
<tr>
<td><strong>AGEP</strong></td>
<td>Histology</td>
</tr>
<tr>
<td>Pustular pustasis, cutaneous localisation of fungal or bacterial septicemia, neutrophilic dermatosis, postular vasculitis</td>
<td>Bacterial or fungal pustule analysis, blood cultures</td>
</tr>
</tbody>
</table>

SJS=S-Johnson syndrome. TEN=toxic epidermal necrolysis. SSA=Sjogren’s-syndrome-related antigen A. DRESS=drug reaction with eosinophilia and systemic symptoms. AGEP=acute generalised exanthematous pustulosis. *Mycoplasma pneumoniae* can trigger SJS and TEN, erythema multiforme, and a specific skin rash with mucosal erosions.
keratinocytes, and larger mixed dermal and interstitial infiltrates than in pustular psoriasis, and the absence of diluted blood vessels.\(^\text{[15,28]}\) For physicians, relapse of pustular eruption without drug re-challenge is the most reliable sign to reject a diagnosis of AGEP, even though authentic drug-induced and non-drug-induced pustulosis have been observed in specific patients with AGEP.\(^\text{[11,31]}\)

In the absence of specific tests, SCAR diagnosis mainly relies on the analysis of clinical and histological patterns (table 1). Despite some specific patterns, misclassification can still occur—eg, AGEP instead of toxic epidermal necrolysis, AGEP as an initial feature of DRESS syndrome—and even authentic SCARs can have clinical or histological features that overlap with each other—ie, DRESS syndrome with Stevens-Johnson syndrome and toxic epidermal necrolysis, AGEP with DRESS syndrome, and AGEP with Stevens-Johnson syndrome and toxic epidermal necrolysis.\(^\text{[12]}\)

**Pathogenesis**

The clinical heterogeneity of SCARs might be explained by the activation of different effector or regulatory cells secreting specific cytokines.\(^\text{[15-22]}\) SCARs are considered to be non-immediate hypersensitivity reactions with four proposed subgroups: IVa, mediated by type 1 T helper (Th1) T cells; IVb, mediated by Th2 T cells and interleukins 5, 4, and 13, and eotaxin cytokines (as occurs in DRESS syndrome); IVc, mediated by cytotoxic T cells (as occurs in Stevens-Johnson syndrome and toxic epidermal necrolysis); and IVd, mediated by T cells and neutrophils via chemokine (C-X-C motif) ligand 8 (CXCL-8) and granulocyte-macrophage colony-stimulating factor cytokines (as occurs in AGEP).\(^\text{[14]}\)

Several mechanistic models have been proposed to explain the recognition by T cells of small compounds (ie, drugs) and the ability of T cells to promote an immune response (figure I).

In the hapten model, covalent bonds are established between drug molecules and autologous proteins or peptides, leading to a drug-specific humoral or cellular immune response. Haptens are chemically reactive small molecules that are able to bind covalently with larger proteins or peptides, initiating an immune reaction. By contrast, pro-haptens are not chemically reactive; they become chemically active compounds after being metabolised. An example of a hapten model is the penicillin hypersensitivity model, in which penicillin derivatives bind to serum albumin that then undergoes intracellular processing to generate chemically modified peptides that elicit an immune reaction.\(^\text{[14]}\)

Another mechanism involves the pharmacological interaction of drugs with immune receptors: the so-called p-i concept. The drug, in its native form or as a metabolite, binds directly and non-covalently to immune receptors such as T-cell receptors, or to specific HLA molecules (ie, MHC proteins), without a specific peptide ligand.\(^\text{[14]}\) For example, in the carbamazepine-induced Stevens-Johnson syndrome and toxic epidermal necrolysis model, carbamazepine directly binds to the protein encoded by the HLA-B*15:02 allele via a non-peptide processing pathway; this was demonstrated by showing that fixation of antigen-presenting cells could still elicit an immune reaction.\(^\text{[25]}\) In the p-i model, the antigenic peptide-processing pathway in antigen-presenting cells is not necessary.

Lastly, a new physiopathological hypothesis has emerged, known as the altered peptide repertoire model. In this model, the drug binds non-covalently within the binding pocket of MHC, leading to alteration of both the chemistry of the binding cleft and the self-peptide repertoire. This new self-peptide presentation can lead to cytotoxic T-cell activation.\(^\text{[26,27]}\) For example, in the abacavir hypersensitivity model, abacavir alters the repertoire of self-peptides by triggering conformational changes in endogenous peptides presented by the protein encoded by the HLA-B*57:01 allele, resulting in the generation of a polyclonal T-cell response and induction of hypersensitivity reactions.\(^\text{[28]}\) In this model, the offending drug does not directly interact with the HLA repertoire; rather, the peptides that change the binding cleft of the HLA repertoire, induced by the offending drug, are treated as foreign antigens by antigen-presenting cells and therefore elicit T-cell activation.

**Genetic factors**

Several genetic factors that cause a predisposition to SCARs have been previously reported—eg, metabolic enzyme mutations, or specific HLA-A, B, or C alleles (appendix p 3).\(^\text{[28-30]}\)

In AGEP, a mutation of the interleukin-36 receptor antagonist gene (IL36RN) was proposed to be a genetic factor in rare cases.\(^\text{[29]}\) A strong (100%) association has been established between the HLA-B*15:02 allele and carbamazepine-triggered Stevens-Johnson syndrome and toxic epidermal necrolysis, and the HLA-B*58:01 allele and allopurinol-induced Stevens-Johnson syndrome and toxic epidermal necrolysis or DRESS syndrome.\(^\text{[31]}\) These associations were first identified in countries with a high prevalence of one specific allele and few ethnic groups (eg, Taiwan [Han Chinese]). European studies did not find such an association.\(^\text{[32]}\) In patients with carbamazepine-induced SCARs, the HLA-A*31:01 allele was reported in patients of northern European ancestry and Japanese ancestry, but not in Taiwanese patients.\(^\text{[32]}\) These results emphasise that various HLA alleles could be associated with a drug-specific clinical pattern, maybe owing to a similar distribution of key aminoacids at the binding sites (eg, as in carbamazepine-induced SCARs).\(^\text{[33]}\) In carbamazepine-induced SCARs, a restricted T-cell receptor clonotype role has been suggested, since a T-cell receptor clonotype role has been associated with individuals with Stevens-Johnson syndrome or toxic epidermal necrolysis who are HLA-B*15:02 positive, whereas it is absent in all carbamazepine-tolerant HLA-B*15:02 carriers.\(^\text{[34]}\)
In allopurinol-induced SCARs, studies have identified a specific T-cell receptor clonotype reacting to its metabolite oxypurinol in addition to the HLA-B*5801 allele.52

HLA studies have identified various SCAR phenotypes with the same drug–HLA association (eg, allopurinol-induced Stevens-Johnson syndrome and toxic epidermal necrolysis or DRESS syndrome and HLA-B*58:01, and carbamazepine-induced Stevens-Johnson syndrome and toxic epidermal necrolysis or DRESS syndrome and HLA-B*15:02), and exclusive drug–HLA associations with one phenotype—eg, dapsone hypersensitivity with the HLA-B*13:01 allele, or phenytoin-induced Stevens-Johnson syndrome and toxic epidermal necrolysis with the HLA-B*15:02 allele.40

Adding to the complexity of the mechanism of SCARs, genome-wide association studies have identified a variant of the cytochrome P450 2C9 enzyme, known to reduce drug clearance, as being an important genetic factor in phenytoin-related SCARs,53 and other studies54 have identified an ABC transporter and proteasome pathway.

Figure 1: Immune mechanisms of SCARs
Schematic diagram showing the immune stimulation, cytotoxic T-cell activation, and key actors in the development of SCARs, adapted from Pichler and colleagues.34 In the hapten (A) and pro-hapten (B) models, the drug or its metabolite, respectively, binds covalently to a peptide carrier (peptide A) and is then presented by MHC proteins to the T-cell receptor (TCR). In the p-i concept (C), the drug or its metabolite interacts directly and non-covalently with the TCR or a peptide-loaded MHC protein. In the altered peptide repertoire model (D), the drug binds directly to the MHC binding pocket and alters its specificity, resulting in presentation of novel ligands (peptide B), leading to cytotoxic T-cell activation. Pre-existing virus-specific T cells might become reactivated during the drug-induced immune response (eg, HHV6, HHV-7, EBV, parvovirus B19). The key actors involved in drug-induced immune reactions leading to SJS or TEN, AGEP, and DRESS syndrome are shown at the bottom of the figure. AGEP=acute generalised exanthematous pustulosis. CXCL8=chemokine (C–X–C motif) ligand 8. DRESS=drug reaction with eosinophilia and systemic symptoms. EBV=Epstein-Barr virus. GM-CSF=granulocyte-macrophage colony-stimulating factor. NK=natural killer. p-i=pharmacological interaction. SCAR=severe cutaneous adverse reaction. SJS=Stevens-Johnson syndrome. TEN=toxic epidermal necrolysis. TRAIL=TNF-related apoptosis-inducing ligand. T-reg=regulatory T cell. TWEAK=TNF-related weak inducer of apoptosis.
mutation in non-drug-specific Stevens-Johnson syndrome and toxic epidermal necrolysis.

**Immunological SCAR mechanisms**

After drug stimulation via HLA-encoded MHC proteins, immune mechanisms of SCARs include the activation of drug-specific cytotoxic T cells, inflammatory cells, or regulatory T cells (T-regs) and the differential secretion of inflammatory cytokines.

In Stevens-Johnson syndrome and toxic epidermal necrolysis, drug-specific cytotoxic cells are probably not the sole effector mechanisms of epidermal necrolysis, and their effects might be amplified by massive production of death mediators, altered anti-apoptotic pathways in target cells, or defective negative regulation of drug-specific immune reactions. Inhibition of drug-specific cytotoxic cells by nucleic acid-based blocking agents has been shown. Analysis of blister fluid from patients with Stevens-Johnson syndrome and toxic epidermal necrolysis first identified MHC-I-restricted cytotoxic T cells, some of which had natural killer (NK) cell markers. This analysis also identified various proinflammatory and anti-inflammatory cytokines secreted by cytotoxic T cells, NK cells, keratinocytes, CD1a+CD14+ non-lymphoid dendritic cells, or CD14+CD16+ monocytes. Epidermal cell death results from necrosis and massive T-cell-mediated apoptosis via three described pathways: Fas–Fas ligand interaction, a perforin–granzyme B pathway, and a granulysin-induced pathway. Granulysin has been shown to be a major cytotoxic molecule responsible for extensive keratinocyte necrosis through cytoxic or NK-cell-mediated cytotoxicity without direct cellular contact, whereas Fas–Fas ligand interaction had no detectable effect, and the perforin–granzyme B pathway only a minor one. Increased concentrations of granulysin and interleukin 15 were significantly correlated with severity of Stevens-Johnson syndrome and toxic epidermal necrolysis, and interleukin 15 was significantly associated with mortality. High expression of receptor-interacting protein kinase 3 (RIPK3) in a toxic epidermal necrolysis lesion has suggested that RIPK3 is an essential actor in the programmed death and necrosis of keratinocytes.

Drug-specific T cells, activated in skin and internal organs, mediate DRESS syndrome, and recruitment of HHV6+ peripheral mononuclear cells to damaged skin is required for virus transmission and replication in CD4+ T cells. A high proportion of CD8+ T cells expressing granzyme B was detected in skin samples of patients with severe DRESS syndrome. Involvement of viruses in DRESS syndrome—e.g., when a viral disease is triggered through direct reactivation by the drug or a strong immune reaction (e.g., graft-vs-host disease or organ transplantation)—was not found in in-vitro studies. In a study of patients with DRESS syndrome, circulating CD8+ T cells secreting tumour-necrosis factor (TNF) α and interferon γ were identified and nearly half of the activated circulating CD8+ T cells recognised HHV, whereas CD8+ T-cell visceral or skin infiltrates mainly recognised EBV. Culprit drugs could also trigger EBV replication via patients’ EBV-transformed B lymphocytes.

In AGEP, the identification of dermal cytotoxic CD8+ T-cell infiltrates also suggests neutrophil recruitment and activation through drug-specific T cells via interleukin 8. Increased circulating, interleukin-22-producing, Th17 cells stimulating keratinocyte secretion of interleukin 8 for neutrophil recruitment are reported in patients with AGEP.

To explain the broad phenotypic variability induced by the same drugs, researchers compared Stevens-Johnson syndrome and toxic epidermal necrolysis, DRESS syndrome, and non-SCAR cytokine profiles or levels with type and density of inflammatory cells. Concentrations of granulysin and Fas ligand in serum samples were suggested as predictive factors of phenotype severity and skin detachment in Stevens-Johnson syndrome and toxic epidermal necrolysis, but their clinical relevance needs further assessment. Patients with Stevens-Johnson syndrome or toxic epidermal necrolysis had significantly more proinflammatory cytokines (TNFα, interleukin 6, and interferon γ) and anti-inflammatory cytokines (interleukin 10 and interleukin-1-receptor antagonist) than patients with other cutaneous adverse reactions, including DRESS syndrome. Analyses of immunoglobulin profiles, white blood cell subsets, and lymphocyte subsets also revealed significant differences between patients with Stevens-Johnson syndrome or toxic epidermal necrolysis and those with DRESS syndrome, suggesting the role of an underlying viral infection coinciding with drug exposure. This concept of T-cell readiness to react suggests that drug reactions follow a type of non-drug-specific immune activation such as occurs in a viral infection.

T-reg functions during acute and chronic SCAR stages have also been thought to influence phenotype. By comparison with healthy controls, T-reg frequency in early and late stages of Stevens-Johnson syndrome and toxic epidermal necrolysis did not differ, whereas non-T-reg cell frequency was increased upon resolution of Stevens-Johnson syndrome and toxic epidermal necrolysis. In the acute stage of DRESS syndrome, functional T-regs were dramatically expanded, whereas they were profoundly diminished in Stevens-Johnson syndrome and toxic epidermal necrolysis. By contrast, T-regs became functionally deficient upon resolution of DRESS syndrome, whereas their functionality was restored after Stevens-Johnson syndrome and toxic epidermal necrolysis.

**Assessment of SCARs**

Case assessment relies on the eruption’s clinical appearance (e.g., potentially virus-related or drug-related), how long the eruption has been present, associated symptoms (e.g., fever, pruritus, lymphadenopathy), and the time elapsed between drug intake and SCAR onset.
During the acute stage, SCARs can require intensive care because they can lead to multiorgan failure and fluid loss due to skin damage. Supportive care consists of restoration of haemodynamic equilibrium and prevention of life-threatening complications. Patients with epidermal detachment or erythromelalgia are exposed to increased fluid and protein loss, hypovolaemia, renal insufficiency, thermal dysregulation, and sepsis. Fluid replacement should be started as soon as possible and adjusted daily. The environmental temperature of the patient should be raised to 28°C. Nutritional hypercaloric and hyperproteic enteral feeding of patients with Stevens-Johnson syndrome and toxic epidermal necrolysis is systematically discussed and often initiated through a nasogastic tube. Peripheral venous lines are placed, when possible, in a region of uninvolved skin.

For Stevens-Johnson syndrome and toxic epidermal necrolysis, opioid agonists are used to limit the pain or stress inherent in mucosal or skin-debris removal, necessitating respiratory monitoring. Systematic invasive mechanical ventilation is unnecessary and is associated with high risk of in-hospital death. Anxiolytics can be prescribed for the prevention of post-traumatic stress disorder (PTSD). Antibiotic prophylaxis is not recommended, and the prescription of unnecessary or non-vital medications should be avoided.

**Dermatological care**

Wound care of patients with Stevens-Johnson syndrome or toxic epidermal necrolysis is done daily with antiseptic baths or diluted antiseptic spray. Skin injuries should be avoided to minimise epidermal detachment, so transportation and manipulation of the patient must

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**Management and treatment**

**Principles of symptomatic treatment**

SCAR-management strategies (table 3) are predominantly symptomatic, aimed at avoiding short-term morbidity and mortality and severe long-term sequelae.

For all patients, culprit-drug identification (appendix p 23) and its early withdrawal are the first mandatory steps (figure 2). For Stevens-Johnson syndrome and toxic epidermal necrolysis, early culprit-drug discontinuation is associated with better prognoses, and causative drugs with long half-lives are associated with an increased morbidity risk.

**SCAR management, outcomes, and main sequelae**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Acute-stage mortality</th>
<th>Outcome</th>
<th>Sequelae</th>
<th>Management after resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific</td>
<td>Symptomatic</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SJS and TEN</td>
<td>Drug withdrawal, no RCT-validated</td>
<td>Supportive care strongly recommended, cutaneous and mucous membrane care, enteral feeding, fluid-loss treatment, analgesia, no systemic infection, environmental temperature &gt;28°C, analgolysis</td>
<td>10–40%</td>
<td>Bacterial superinfection, visceral-specific involvement, lung failure</td>
</tr>
<tr>
<td>DRESS</td>
<td>Drug withdrawal, no RCT-validated</td>
<td>Symptomatic, antipyretics</td>
<td>1–10%</td>
<td>Acute organ failure, virus reactivation, relapses</td>
</tr>
<tr>
<td>AGEP</td>
<td>Drug withdrawal, no RCT-validated</td>
<td>Symptomatic, antipyretics, no antibiotics</td>
<td>1%</td>
<td>Recovery</td>
</tr>
</tbody>
</table>

SCAR=severe cutaneous adverse reaction. SJS=Stevens-Johnson syndrome. TEN=toxic epidermal necrolysis. RCT=randomised controlled trial. DRESS=drug reaction with eosinophilia and systemic symptoms. AGEP=acute generalised exanthematous pustulosis. Thalidomide increased mortality in an RCT that was stopped early; the benefits of intravenous immunoglobulins and systemic corticosteroids are still being debated; a single-arm trial† found some benefits of ciclosporin. Follow-up is adapted to disease severity and sequelae, a multidisciplinary approach is mandatory for all cases of SJS and TEN and often necessary for DRESS syndrome. Intravenous immunoglobulin had no benefit; the antiviral ganciclovir provided no clear benefit in case reports.

Table 3: SCAR management, outcomes, and main sequelae.

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Physical examination includes the description of the distribution of SCAR-specific lesions. Cutaneous or mucous membrane involvement in orifices, indicating a severe reaction (external or internal), must be specified. Photos and clinical signs should be collected as often as possible to enable retrospective expert validation of the SCAR. Skin biopsy, including direct immunofluorescence, of blistering eruptions and biological tests to eliminate differential diagnoses, are strongly recommended (table 2).

If Stevens-Johnson syndrome, toxic epidermal necrolysis, or DRESS syndrome is confirmed, management by a referral centre or specialised intensive-care unit is strongly recommended. A diagnosis of Stevens-Johnson syndrome or toxic epidermal necrolysis within 7 days of onset is associated with improved survival compared with a more delayed diagnosis. AGEP is usually a transient onset is associated with improved survival compared with a more delayed diagnosis. AGEP is usually a transient onset is associated with improved survival compared with a more delayed diagnosis. AGEP is usually a transient onset is associated with improved survival compared with a more delayed diagnosis. AGEP is usually a transient onset is associated with improved survival compared with a more delayed diagnosis.
be restricted, as should use of adhesive electrocardiogram electrodes. Petroleum jelly should be applied systematically to all areas of detached skin.\textsuperscript{72} Unlike burn management, large and aggressive skin debridement should be avoided, and might delay re-epithelialisation, because necrotic epidermal sheets act as a natural biological dressing. Topical antimicrobial agents or sulfadiazine cream (containing antibacterial sulphonamides) are not recommended. When necessary, non-adhesive (eg, hydrocellular) dressings are used to cover pressure points, particularly on the back.\textsuperscript{74,75}

During the acute phase, ocular, oral, nasal, genital, or anal mucosae lubrications with emollient are recommended to reduce mucosal adhesion formation and functional sequelae.\textsuperscript{76,77} Mucosal bleeding or erosions are treated with topical anaesthetics, mouthwashes, application of swabs, local administration of adrenaline, and clotting agents.\textsuperscript{78,79} Ocular management relies on inflammatory debris removal with daily saline rinses, and, after topical anaesthesia, removal with a moist cotton bud or smooth blunt instrument. Prophylactic topical antibiotics, topical ciclosporin, or corticosteroids have been used, but were shown to have no benefit in terms of ocular sequelae.\textsuperscript{65} To protect and reduce conjunctival or corneal sequelae, amniotic membrane transplantation has been proposed to prevent eyelid scarring.\textsuperscript{64}

For DRESS syndrome and AGEP, dermatological care mainly relies on appropriate skin moisturisation.\textsuperscript{63} When necessary, mucous-membrane management should be the same as that used for Stevens-Johnson syndrome and toxic epidermal necrolysis.

**Targeted therapeutic approaches**

In parallel with supportive care, therapeutic approaches for patients with Stevens-Johnson syndrome and toxic epidermal necrolysis are still being debated. Most of the information comes from case reports and small, uncontrolled series.\textsuperscript{65,66} The shortage of large randomised controlled trials (RCTs) comparing treatment strategies reflects the rarity of SCARs. Several immunosuppressants or immunomodulatory treatments (eg, corticosteroids,\textsuperscript{87} cyclophosphamide,\textsuperscript{73} calcineurin inhibitors,\textsuperscript{75} anti-TNF therapies,\textsuperscript{88} intravenous immunoglobulins [IVIg]\textsuperscript{80,81} or plasmapheresis) have had controversial results.

Systemic corticosteroids (eg, intravenous methylprednisolone) administered in pulses of 1–2 mg/kg up to 600–1000 mg per day were considered to be a treatment option for many years in some centres. In large case-control study,\textsuperscript{87} which assessed the preventive effect of corticosteroids on Stevens-Johnson syndrome and toxic epidermal necrosis, previous exposure to corticosteroids was associated with a longer disease progression—ie, 2.2 days (95% CI 1.1–3.2) longer—than no previous exposure, with no effect on disease severity or mortality.

IVIg as a treatment for Stevens-Johnson syndrome and toxic epidermal necrolysis has had varying results, some supporting their efficacy and others not.\textsuperscript{82,83–85} A meta-analysis of 17 studies yielded a 19.9% overall mortality rate in patients with toxic epidermal necrosis given IVIg. The pooled odds ratio for mortality from six observational studies comparing IVIg with supportive care was 1.00 (95% CI 0.58–1.75; p=0.99), but IVIg dose (high vs low) did not correlate with mortality in a multivariate analysis.\textsuperscript{91}

In a series of 82 patients with Stevens-Johnson syndrome or toxic epidermal necrolysis that compared corticosteroids with or without IVIg, complication or mortality rates did not significantly differ with the addition of IVIg, whereas hospital stays were significantly shorter for those receiving IVIg.\textsuperscript{86} Furthermore, mortality did not significantly differ in patients receiving IVIg or corticosteroids in comparison with supportive care in a retrospective study.\textsuperscript{86} In a prospective cohort study, no difference with regard to mortality was observed between IVIg and supportive care for patients with Stevens-Johnson syndrome or toxic epidermal necrolysis, suggesting that supportive care alone is optimal.\textsuperscript{86}

Ciclosporin, an anti-apoptotic agent, has also been proposed to inhibit CD8+ T cells, limiting disease
progression after a short-term administration of 3–10 mg/kg. This potential benefit of ciclosporin on mortality of patients with Stevens-Johnson syndrome and toxic epidermal necrolysis, compared with IVIg, was also established in a single centre study of 71 patients. In an uncontrolled series of ten patients with toxic epidermal necrolysis, a single dose of etanercept (an anti-TNF agent) was found to provide prompt healing (in a median of 8–5 days) without complications. Granulocyte-macrophage colony-stimulating factor might be interesting to investigate further in future trials, because preliminary data from two patients with toxic epidermal necrolysis suggest that it had an effect on re-epithelialisation via T-reg mobilisation, expansion of tolerogenic myeloid precursors, immature dendritic cell mobilisation, or enhanced cytolytic functions of NK T cells.

In our clinical experience, neither corticosteroids nor IVIg affect mortality, whereas death occurs more frequently with thalidomide than placebo as shown in an RCT of patients with toxic epidermal necrolysis. Oral ciclosporin (at decremental doses for 10 days) prevented skin-detachment progression in patients with Stevens-Johnson syndrome or toxic epidermal necrolysis; only 38% of patients developed progressive disease, as opposed to 65% of those in the open-label IVIg trial. No severe adverse events, relapse, or deaths occurred in the 29 patients who received oral ciclosporin, even though the SCORE of Toxic Epidermal Necrolysis (SCORTEN) scale—a prognosis score built on seven independent variables (appendix) that can determine a patient’s mortality risk—predicted that three patients would die. No RCTs have been done for DRESS syndrome or AGEP. Pulse or oral corticosteroids have been administered to patients with DRESS syndrome in retrospective series, but no standardised assessment of outcomes has been done. The role of high-dose corticosteroids in DRESS relapses has not been specifically analysed. The use of corticosteroids in patients with DRESS syndrome enhances cytomegalovirus and HHV6 viral load, but not EBV. Despite virus reactivation, antiviral therapy (eg, with ganciclovir) should not be used currently because of both poor demonstrated efficacy in this syndrome and toxic effects. IVIg, which has been used to treat patients with DRESS syndrome, has antiviral and immunomodulatory properties, affecting the innate and adaptive immune system. In a prematurely stopped prospective study, 140 patients with severe DRESS syndrome were given IVIg, but three had severe malaise, one had pulmonary embolism, and four required rescue oral corticosteroid treatment. In our retrospective study on the therapeutic management of DRESS syndrome, patients treated with topical corticosteroids had fewer relapses than those treated with systemic steroids, but this result might be overestimated by the higher severity of DRESS syndrome in patients who received systemic steroids. An RCT comparing superpotent topical corticosteroids with systemic corticosteroids in patients with mild-to-moderate DRESS syndrome is ongoing (NCT01987076).

To limit AGEP progression, drug withdrawal might be sufficient, and topical steroids appeared to be favourable in patients with AGEP and visceral involvement, without requiring systemic corticosteroids, in a retrospective study. As in mild-to-moderate DRESS syndrome, potent or superpotent topical corticosteroids—eg, clobetasol propionate 30 g per day—are widely used for patients with AGEP, although their efficacy has not yet been assessed in an RCT.

Outcome and sequelae

SCARs—mainly Stevens-Johnson syndrome, toxic epidermal necrolysis, and DRESS syndrome—are life threatening and carry a non-negligible risk of severe sequelae (table 3). During the acute stages of Stevens-Johnson syndrome and toxic epidermal necrolysis, visceral involvement (eg, renal failure, intestinal, ocular-specific pulmonary lesions, or sepsis) is the main complication. Respiratory insufficiency in patients with SCARs can result from direct effects of the SCAR on the organs or inhalation of foreign substances leading to superinfection, and pulmonary infection is significantly associated with severe laryngeal lesions caused by Stevens-Johnson syndrome or toxic epidermal necrolysis. During the acute stage, impaired skin barrier function or translocation of gut bacteria might facilitate bacterial colonisation and bloodstream infections. Sepsis is the predominant cause of death attributed to Stevens-Johnson syndrome and toxic epidermal necrolysis. In a study of pregnant women with acute-stage Stevens-Johnson syndrome or toxic epidermal necrolysis, caesarean was the delivery method in 50% of patients; no maternal deaths occurred but fetal outcomes were poor.

Acute-stage mortality ranges from at least 10% for Stevens-Johnson syndrome to around 40% for toxic epidermal necrolysis, with overall in-hospital mortality of 22% in Europe for both conditions. Increased mortality is described in patients with malignancies who develop Stevens-Johnson syndrome or toxic epidermal necrolysis, and several factors contribute to poor prognosis in this population: malnutrition, cancer type, and chemotherapy type. SCORTEN successfully predicts 3-day mortality and the individual risk of death for patients with Stevens-Johnson syndrome and toxic epidermal necrolysis when assessed at admission to hospital. A five-point auxiliary score that does not require laboratory data has also been devised, and might be useful to predict severity of illness in retrospective settings when laboratory data are missing. At 1 year, overall mortality in patients with Stevens-Johnson syndrome or toxic epidermal necrolysis remains high at 34% (95% CI 30–39; 24% [18–29] for Stevens-Johnson syndrome and 49% [36–60] for toxic epidermal necrolysis). Re-epithelialisation of lesions in patients with Stevens-Johnson syndrome or toxic epidermal necrolysis...
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The incidence of long-term DRESS sequelae was 11.5%, and natural chronic course of DRESS and in relapses were associated with severity and prolonged duration of increased anti-HHV6 IgG and anti-HHV6 DNA titres. Routine screening for sequelae is imperative to limit or reduce their burden or impact on quality of life. Cutaneous sequelae (eg, hyperchomatic macules, photosensitivity, telogen effluvium, nail loss, vaginal adhesion bands) and ocular sequelae (eg, photophobia and chronic tearing, eyelid malposition, and punctate keratitis) are the most frequent. Regarding mucous membrane sequelae, in our experience their severity is not related to cutaneous disease severity at the acute stage. During the chronic stage, up to 65% of patients develop late ocular complications—eg, dry eyes or synechial visual loss. In a retrospective study that assessed outcomes 15 months after discharge, acute-stage ocular severity was significantly associated with late complications. Management of ocular sequelae involves treatment for dry eye, cornea inflammation, and refractory ocular disease. To rehabilitate visual function, scleral lenses were successfully used, and minor salivary gland or mucous-membrane grafts have been tested to treat sylmblepharon and dry eyes. Saliva acidity associated with Stevens-Johnson syndrome or toxic epidermal necrosis can cause chronic sialadenitis, tooth decay, and dental atrophy. Male genital synchiae requiring circumcision are often observed, whereas strictures of vaginal mucosa or birth-canal stenosis can complicate spontaneous vaginal delivery and normal sexual intercourse.

With regard to DRESS syndrome, acute-stage mortality ranges from 5% to 10% and is mainly attributed to specific myocardial or pulmonary lesions, and haemophagocytosis. In a cohort study, the overall cumulative incidence of long-term DRESS sequelae was 11–5%, and mainly consisted of autoimmune diseases. If PTSD, anxiety, or depression are reported at a late stage or after remission of DRESS, systematic psychological or psychiatric screening is likely to be needed. Although increased anti-HHV6 IgG and anti-HHV6 DNA titres were associated with severity and prolonged duration of DRESS syndrome, the role of virus reactivation in the natural chronic course of DRESS and in relapses remains hypothetical. Chronic virus activation is suspected of triggering excessive autoimmune responses and inducing autoimmune diseases, such as scleroderma, lupus erythematosus, diabetes, or thyroiditis arising after DRESS remission. Corticosteroids were shown to limit autoimmune diseases in patients with DRESS syndrome assessed retrospectively.

AGEP has a good prognosis and no described sequelae. Medication risk and drug causality When assessing drug causality in a patient with a SCAR, several factors should be taken into consideration: SCAR type, day of symptom onset, drug notoriety, and time since drug intake. The interval between exposure and SCAR onset differs according to the type of SCAR; generally, it is short for AGEP, intermediate for Stevens-Johnson syndrome and toxic epidermal necrosis, and long for DRESS syndrome. For each SCAR, determining the first day of symptoms (index day) is the first step towards identifying a potential causative agent and withdrawing it to assess its role in causing the SCAR prodrome. Additionally, a drug stopped before disease onset should still be suspected if it has a long half-life. At date, the French pharmacovigilance causality score test or the Naranjo algorithm have been the most frequently used worldwide to identify culprit drugs. Additionally, ALDEN (ALgorithm for Drug causality in Epidermal Necrolysis) has been validated to improve individual assessment of a suspected drug’s role in Stevens-Johnson syndrome and toxic epidermal necrosis. This specific algorithm, the scores of which strongly correlated with those in the EuroSCAR case-control study for drugs associated with epidermal necrosis, uses time since drug intake, pharmacokinetics, rechallenge or dechallenge, and drug notoriety to classify drug causality as very unlikely, unlikely, possible, probable, or very probable.

For each phenotype—Stevens-Johnson syndrome, toxic epidermal necrosis, DRESS syndrome, and AGEP—a few drugs are strongly associated with most cases. European case-control studies on SCAR cases yielded a list of potential high-risk drugs; the main ones are listed in the appendix (pp 5–6). European case-control studies on SCAR cases yielded a list of potential high-risk drugs; the main ones are listed in the appendix (pp 5–6). Even though the prevalence of non-drug-induced Stevens-Johnson syndrome and toxic epidermal necrosis is higher in children than in adults, high-risk drugs identified for children were the same as for adults—eg, antibacterial sulphonamides, carbamazepine, phenobarbital, and phenytoin—and no risk has been identified for any vaccines. Paracetamol’s inclusion in the list of culprit drugs might be linked to confounding comedication, because it is prescribed for influenza-like syndromes during the early stage of Stevens-Johnson syndrome and toxic epidermal necrosis. Drug regulatory agencies also frequently publish alerts for drugs undergoing safety monitoring for the risk of SCARs (appendix pp 5–6). Taken together, acute-stage management, hospital stays, and chronic sequelae care mean that the cost of Stevens-Johnson syndrome and toxic epidermal necrosis is high; therefore, benefit–risk analysis is warranted for high-risk drugs. For idiopathic Stevens-Johnson syndrome or toxic epidermal necrosis with no identified culprit drug, drugs in food (eg, phenylbutazone in meat) have been suggested to be a cause. This hypothesis was not confirmed by the concentration of phenylbutazone or its metabolites in plasma in patients with idiopathic or drug-induced Stevens-Johnson syndrome and toxic epidermal necrosis.

Known risk factors associated with SCARs are HIV infection, specific HLA allele and drug combinations, and systemic lupus erythematosus. The strength of the
HIV association might be affected by the severity of the patient’s immunodeficiency or the use of drugs that are high risk for SCARs (eg, co-trimoxazole or abacavir) in this population.109,112 Even though for nevirapine, hypersensitivity predominated in non-HIV-infected patients with high CD4+ counts who were given nevirapine as a prophylactic treatment after HIV exposure.121 For abacavir, in-vitro T-cell reactivity can also occur in drug-naive individuals with the HLA-B*57:01 allele.122 Supported by the results of extensive HLA-B*15:02 screening in Taiwanese neurology clinics, the US Food and Drug Administration recommends genetic testing before prescribing carbamazepine to patients of Asian ancestry from China and southeast Asia.120 HLA screening is thus only relevant to avoid SCARs caused by a few drugs in specific populations.42,111 Relapsing SCARs are mainly a result of re-exposure to the same high-risk medications as caused the initial SCAR, but incomplete relapse of DRESS syndrome has been reported by Picard and colleagues122 following the administration of drugs not previously taken by the patient. These authors hypothesised that a persistent immune stimulation or viral reactivation was responsible for minor DRESS relapses with drugs that were chemically unrelated to the initial causative agent.123 For Stevens-Johnson syndrome and toxic epidermal necrolysis, individual susceptibility was postulated to explain a recurrence risk of 7% after a first episode in a cohort of 708 patients.111 This recurrence risk is overestimated by misclassification of Stevens-Johnson syndrome and toxic epidermal necrolysis, and recurrence solely occurs with causative or relative drugs.114 Nevertheless, there is no particular recommendation about the prescription of other potential high-risk drugs to a patient with a history of SCARs.71 Because relapse can be worse than the initial reaction, and sometimes even fatal, we recommend that patients who have had a SCAR carry an allergy card stating the culprit drug and medication contraindications (including generic and proprietary names of drugs), so that physicians can avoid giving them potential causative drugs, drugs of same structure, and members of the same family of molecules, but not necessarily the entire therapeutic group.

**Culprit-drug tests**

Methods to link a particular drug to a SCAR are scarce, with no standardised strategy, and none is associated with a 100% negative predictive value. To confirm a potential culprit drug, HLA screening might be useful when a strong association between the suspected drug and a particular HLA allele exists. None of the current tests has sufficient specificity and sensitivity to rule out a potential culprit drug when negative, enabling its rechallenge.41

Routine assessment to identify the culprit drug includes establishing the chronology of drug intake and patch testing if several drugs have been taken. Thus far, no standardised protocol of drug quantity or vehicle has been established for patch testing, but is frequently 10% of native drug in petrolatum.120 The allergen-containing patch is usually taped onto the patient’s back for 2 days and then assessed at a minimum of 48 h and 96 h. Patch test specificity and sensitivity vary according to the suspected drug and the SCAR subtype (higher sensitivity for AGEP than for Stevens-Johnson syndrome, toxic epidermal necrolysis, and DRESS).117 Patch tests are reportedly safe, with few reported relapses or severe reactions. Neither prick nor intradermal tests are recommended, despite the intradermal test having higher sensitivity than the prick test, and an oral drug provocation test is definitely prohibited because of the risk of relapse.120 In our experience of DRESS syndrome, tests are done 6 months after the acute stage to avoid relapse.

In-vitro tests can be used to measure peripheral blood mononuclear cell activity in patients with SCARs, with the culprit drug displaying pharmacological activity. For hypersensitivity reactions, two methods to identify the culprit drug have been described: the lymphocyte-transformation test (LTT) and enzyme-linked immunospot assay (ELISPOT). LTT is usually done 1 month after the reaction (5–8 weeks after DRESS onset, and within 1 month of onset of Stevens-Johnson syndrome and toxic epidermal necrolysis), whereas ELISPOT can be done at an earlier stage after SCAR onset.109,110 Neither are done routinely. Although LTT shows promise as a culprit-drug test in patients with DRESS syndrome or AGEP, it has low relevance in Stevens-Johnson syndrome and toxic epidermal necrolysis, even after enhancement of its sensitivity after removal of T-reg CD25+ cells.130,131 ELISPOT has a higher sensitivity (82%) than LTT (50%), and detects drug-specific T cells or identifies the culprit drug via drug-specific interferon-γ, interleukin 4, or granulysin production.132 In patients with Stevens-Johnson syndrome or toxic epidermal necrolysis, the combination of the LTT and ELISPOT, detecting granulysin, granzyme B, and cytokines, have been proposed.109

**Public health and drug-policy issues**

At the population level, the avoidance of SCARs should be considered a high-priority public health and drug policy. A specific focus should be accorded to the following approaches: pharmacogenetic tests to select patients at risk for SCARs in specific subpopulations (table 3); epidemiological studies; pharmacovigilance, including systematic reporting of culprit drugs (including a precise assessment of a drug’s harm potential and benefit–risk ratio) by practitioners to health authorities, drug companies, or independent registries (eg, RegiSCAR); consumer self-reporting of drugs eliciting severe or prevalent cutaneous adverse reactions;115 improvement of drug dictionaries, particularly with regard to the description of drugs associated with Stevens-Johnson syndrome and toxic epidermal necrolysis;144 organisation of experts and referral centres to improve SCAR
management and outcomes; recording of patients’ viewpoints and encouraging the use of patient associations such as Amalyste, which have contributed to improving patients’, health-care providers’, and decision makers’ knowledge of outcomes and sequelae associated with Stevens-Johnson syndrome and toxic epidermal necrolysis; and establishment of specific funds to financially compensate patients with SCARs and improvement of patients’ awareness of these funds.141

For a potentially high-risk drug, several decisions should be considered: (1) withdrawal from the market (eg, chloromezalone was recommended for withdrawal by the European Medicines Agency in 1997 due to its unfavourable benefit–risk relationship); (2) restriction of its use (eg, since June, 2012, minocycline is no longer recommended in France for acne and rosacea but is still allowed to be marketed for other indications);142 (3) changes to the prescription (eg, lamotrigine should be initially given at a low dose and then progressively increased to avoid skin reactions);143 and (4) establishment of a safer alternative agent as first-line therapy (eg, prescription of an isoxazolyl penicillin such as cloxacillin instead of high-risk co-trimoxazole first for severe meticillin-resistant Staphylococcus aureus infection, especially in countries with non-community-acquired strains).144 However, these approaches might not be suitable for some high-risk drugs such as allopurinol, which is widely prescribed. Fewer SCARs would be expected if allopurinol was prescribed only according to its accepted use or guidelines—ie, for gout and kidney stones rather than for asymptomatic hyperuricaemia accompanied by renal or cardiovascular disease, for example, for which an increased risk of hypersensitivity reaction was noted.110

Conclusions

We highlight the difficulty and necessity of early and accurate SCAR diagnosis (figure 2). The expertise of the treating physician is vital in the early diagnosis and specific management of SCARs to prevent or limit long-term sequelae. Physicians should be aware of the potential role of high-risk medication in triggering SCARs, especially when predisposing factors are present.

Contributors

All authors contributed equally to the literature search, analysis, manuscript writing, and critical revision of this Seminar.

Declaration of interests

IV-A has received personal fees from Janssen Cilag for a booklet on cutaneous adverse reactions to telaprevir and educational presentations, Cephalon for review of phosphogluconol in cutaneous adverse reactions, Pinnacle Biologics for expertise regarding cutaneous reactions to amifostine, Boehringer Ingelheim for expert consultancy on severe cutaneous adverse reactions, and Pierre Fabre for educational presentations. PW is a principal investigator of a clinical trial (NCT01412892) on neutrophilinisation funded by Novartis and has received personal fees from Pierre Fabre and Expanscience for his expertise on isoretinoincin for acne. OC has received personal fees from Roche for a trial assessing the preventive role of dicyclene in cutaneous-related anti-EGFR toxicities, and Janssen Cilag for a meeting dedicated to the Centorcor registry for ustekinumab side-effects, as well as personal fees from Sanofi-Aventis, Novartis, Albivie, GlaxoSmithKline, Bailleul, Astellas, and Galderma for being on a scientific board. OC has also received a grant from GlaxoSmithKline and consultant and speaker fees from Bayer. TAD declares no competing interests.

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