Multisystem organ failure secondary to acute generalised exanthematous pustulosis (AGEP) with atypical presentation resembling septic shock

Karen Cravero, Teja Chakrala, Andrew Shychuk

SUMMARY
A woman was admitted for sepsis secondary to cellulitis. After clinical improvement of sepsis, non-follicular small pustules were observed on the trunk, limbs and face while vesicles/bullae and skin exfoliation were noted on upper extremities. Larger systemic manifestations included fever, hypertension and tachycardia. Laboratory results revealed neutrophilic leucocytosis, eosinophilia, mild transaminitis and acute renal failure. Despite treatment for potential sepsis and discontinuation of offending agents, her condition worsened leading to haemodynamic instability and renal failure requiring vasopressor support, intubation and continuous venovenous haemodialysis. Skin biopsy revealed a diagnosis of acute generalised exanthematous pustulosis (AGEP), a rare condition usually caused by antibiotic treatment. The suspected offending drug was clindamycin, with possible combined effects by metronidazole and/or vancomycin. Improvement of skin manifestations were seen within 48 hours of starting systemic steroids. Here, we present an atypical case of AGEP clinically presenting with atypical skin lesions, severe systemic involvement mimicking septic shock, which culminated in multisystem organ failure.

BACKGROUND
Acute generalised exanthematous pustulosis (AGEP) is a rare severe cutaneous adverse reaction afflicting 1–5 patients per million per year. It is seen in both men and women, however, it tends to have a female predominance. A genetic component to susceptibility is suspected, as human leucocyte antigens B51, DR11 and DQ3 are found to be more commonly identified in patients with AGEP. Furthermore, Navarini et al described a higher percentage of IL36 receptor antagonist (IL36RN) mutations in AGEP patients with mucosal involvement.

AGEP typically presents as an abrupt onset of superficial, pin-sized, sterile, non-follicular pustules usually on a background of erythematous and edematous skin. Pustules are often first identified in intertriginous areas, rapidly expanding to the trunk and limbs without involvement of the palms and soles. Systemic manifestations are usually limited to fever (≥38°C) and neutrophilia (≥7.5x10^9/L), while eosinophilia is seen in about 30% of all cases. Mucosal membrane involvement is absent or minimal, and if present, is usually localised to the lips. Rarely, AGEP cases may present with mild hepatic and kidney injury.

AGEP is a type intravenous hypersensitivity reaction caused most commonly by antibiotics, although other drug classes, viral infections and unknown triggers have been reported. Onset of skin manifestation is highly variable, with clinical signs observed in less than 48 hours or up to weeks. In most cases, complete resolution of pustules and fever is achieved within 2 weeks following discontinuation of the offending agent, when skin lesions transition from pustules to diffuse exfoliation which may last for extended periods of time. The above described presentation is by far the most common way AGEP may manifest. Unusual cases have been reported in which AGEP may present with multiple vesicles, large bullae, and diffuse skin desquamation; a presentation more commonly observed with toxic epidermal necrolysis (TEN). TEN is a separate entity with its own unique manifestations and pathophysiology; however, atypical cases of AGEP clinically presenting with overlapping TEN features (appropriately named TEN-like AGEP) have been described in the literature. Presentation of TEN-like AGEP cases are often associated with mucous membrane involvement. They present with skin manifestations observed in both conditions, however, demonstrate AGEP specific histopathology. While most of these cases tend to follow the clinical course of AGEP with full recovery, more severe events can present with signs resembling septic shock. These more severe forms may lead to the requirement of ICU-level care and vasopressor support. Multisystem organ failure may also occur with consequential increase in mortality rates. To the extent of the authors’ knowledge, there have only been four cases reported of TEN-like AGEP progressing to shock and multiple organ dysfunction.

It is important to keep AGEP in the differential when patients develop a septic shock-like picture in association with skin manifestations of pustules and/or bullae shortly after drug administration. Given the rarity of AGEP, it is important for physicians to familiarise themselves with the spectrum of presentations to aid with timely diagnosis and therapeutic interventions. Herein, we describe a case of severe AGEP with overlapping TEN skin manifestation, which rapidly progressed into shock and multisystem organ failure.

CASE PRESENTATION
A morbidly obese African-American woman presented with acute cellulitis superimposed on...
chronic bilateral lower extremity lymphedema. At the time of admission, the patient stated that she had a history of stable lymphedema for over 3 years, but within the last week swelling in both legs increased, leading to severe pain, weakness and difficulty with ambulation. A day prior to presentation, she noticed white drainage from her left lower extremity prompting her to visit her local emergency department. She was found to be tachycardic, hypotensive, and febrile. Laboratory results were significant for an increased white cell count (WCC) (31 x 10^9/L), mild liver dysfunction and acute renal failure. She was given acetaminophen, and intravenous fluids. Blood cultures were collected, and she was started on intravenous ceftriaxone. The patient was transferred to our hospital for further management of sepsis secondary to left lower extremity infection.

On presentation (day 1), she was hypertensive 199/73 mm Hg, tachycardic (103 bpm), with temperature of 37.8 C and saturating 98% on room air. On examination, she was alert and oriented to person, place and time. Her bilateral lower extremities were hyperpigmented, dry, scaly and thickened throughout. Significant bilateral pitting oedema was also present with circumferential induration. Numerous deep fissures, several containing maggots, were present in addition to purulent fluid leakage from the medial and plantar surfaces of both feet. Aside from severe pain and tenderness to palpation bilaterally, the patient denied any other symptoms. Initial labs were notable for an elevated lactic acid, creatinine and white WCC. No fluid collections or other symptoms. Initial labs were notable for an elevated lactic acid, creatinine and white WCC. No fluid collections or thromboses were present on imaging. Full infectious workup was ordered, and the patient was started on empiric antibiotic therapy with intravenous vancomycin, cefepime and metronidazole. On day 2, a one-time dosing of intravenous clindamycin was added for additional gram-positive coverage. Over the next 4 days, the patient’s condition improved with resolution of her symptoms and haodynamic instability. Her WCC steadily continued a downward trend to near-normal levels. Patient also reported decrease in pain in both legs.

On the late hours of day 5, approximately 87 hours after the first dose of clindamycin, the patient became febrile (37.6 C) and her WCC began to rise. Her mentation was normal, and she was deemed stable, not requiring an escalation of care. Repeat blood and urine cultures along with imaging studies were obtained, which revealed no new infections or abnormalities. During the early hours of day 6, patient developed another fever spike (38 C) with significant neutrophilia despite being on broad spectrum antibiotic coverage (intravenous vancomycin and cefepime). At this point, clindamycin and metronidazole were added back to the antibiotic regimen. At approximately 14:00 hours, the patient received a dose of clindamycin, the second one overall since being admitted.

Approximately 1 hour later, a skin lesion was first observed on her right medial thigh. Over the next few hours, her condition quickly deteriorated. She became unresponsive, haemodynamically unstable, which necessitated escalation of care and she was transferred to the intensive care unit for management of possible septic shock. Several hours later, the skin condition worsened. Physical examination revealed numerous scattered small studded pustules over an erythematous and edematous background, with intertriginous predominance. Small vesicles and flaccid bullae were also present in the upper and lower extremities as well as the neck region. Diffuse non-blanching redness was present throughout the body. Erythematous plaques with desquamation, denuded erosions and excoriations were also noted, most significantly under the breasts, and the posterior area of both ears (figure 1). There was no evidence of mucous membrane involvement. During this time, the patient had become encephalopathic requiring intubation for airway protection. Her urine output diminished, she developed metabolic acidosis and she was started on continuous veno-venous haemodialysis (CVVHD).

The next day, her cutaneous examination revealed more prominent pustules, vesicles, and superficial desquamation diffusely throughout her trunk and extremities (figure 2). Dermatology was consulted and a punch biopsy of affected skin was obtained. Clindamycin was stopped for suspicion of AGEP, and systemic corticosteroids was started. Laboratory studies revealed significant neutrophilic leukocytosis, mild eosinophilia, evidence of acute kidney injury, and some liver damage, with mild aspartate transaminase and alanine aminotransferase elevations. Because neutrophilia and eosinophilia are classic characteristics seen in AGEP, a clinical timeframe for WCC is presented in figure 3. Of note, absolute neutrophil levels obtained at time of rash detection, were the highest values to date. Levels of her kidney and liver function, are depicted in figure 4.

INVESTIGATIONS

Extensive laboratory testing was performed throughout the patient’s hospital stay with relevant lab values at different time points of her treatment duration listed in table 1. CT imaging of the abdomen and pelvis revealed subcutaneous fat stranding; however, no soft-tissue gas was observed ruling out necrotising fasciitis. Chest imaging did not show any signs of new infections, no airspace opacities and no pleural effusion.

It is worth mentioning that the only positive blood culture results were obtained from medical records received from Cravero K, et al. BMJ Case Rep 2022;15:e247040. doi:10.1136/bcr-2021-247040
Case report


the institution the patient was originally treated at; which grew Streptococcus spp in both aerobic and anaerobic bottles. An extensive infectious workup performed at our hospital, including multiple blood and urine cultures, was largely negative. A skin punch biopsy was performed on a left proximal thigh lesion (0.3 cm×0.5 cm) and demonstrated collections of sub-corneal neutrophils overlying mild spongiosis, along with perivascular inflammation with infiltration of mostly neutrophils, but also eosinophils and lymphocytes to a lesser degree. Of special significance, no evidence of epidermal necrosis was noted. Figure 5 presents these histopathological findings.

DIFFERENTIAL DIAGNOSIS

The differential diagnosis for the case included AGEP, septic shock, pustular psoriasis, TEN, and Stevens-Johnson syndrome (SJS).

► A comprehensive infectious evaluation was negative. Culture of an open wound did grow Actinobacter iwoffii, a normal skin flora. Lack of an infectious agent along with the progressive deterioration of the patient’s health despite broad antibiotic therapy, made the diagnosis of septic shock less likely.

► Pustular psoriasis is very similar to AGEP in its presentation. AGEP has an acute onset, is a single episode, and

Figure 2 Second day of skin manifestations presented with more diffused and pronounced collarette-shaped desquamation, and erythematous skin in (A) chest, neck and (B) back areas. (C) Erosive lesions in the thigh and abdomen area with pustules coalescing into larger structures.

Figure 3 Absolute WCC at different times during length of clinical presentation. Counts demonstrated significant neutrophilia and mild eosinophilia during time of skin presentation. This figure was created by the authors specifically for inclusion in this manuscript. WCC, white cell count.

Figure 4 Pictorial representation of labs for kidney and liver function during length of clinical presentation. This figure was created by the authors specifically for inclusion in this manuscript.
resolves within 2 weeks; whereas pustular psoriasis is a chronic condition with frequent flare ups and has a slower onset.17 Skin biopsy results in our patient lacked characteristic histological findings seen in psoriasis such as tortuous dilated blood vessels, psoriaform acanthosis, Munro’s microabscesses and increased mitotic figures.17

► TEN and SJS tend to have mucous membrane involvement, a finding which was absent in our patient. The latency period between drug exposure and symptom onset is relatively more prolonged in these conditions. Drugs such as allopurinol and sulfonamides have been found to trigger SJS and TEN, but have not been associated with AGEP.12 18 Finally, both SJS and TEN are characterised by extensive epidermal necrosis.12 Our patient’s physical signs resembled those seen in TEN, but her histological analysis was more suggestive of atypical TEN-like AGEP than TEN itself.

Overall, the temporal relationship of antibiotic initiation to skin eruption is strongly suggestive of drug reaction. When examining a patient with pustules or desquamation, the above-mentioned diagnoses should be considered. In our case, given a high EuroSCAR score and the biopsy results, AGEP was the most likely diagnosis.

**TREATMENT**

Figure 6 is a pictorial representation of all antibiotics and systemic corticosteroids used with their respective time course. Treatments received by our patient at the outside hospital prior to transfer included acetaminophen, 3 L normal saline and 1 g of ceftriaxone. On day 1, after transfer to our hospital, the patient was empirically treated with intravenous metronidazole and cefepime for suspected sepsis. Intravenous vancomycin and one dose of clindamycin were added at later hours of the day. Due to the patient’s clinical improvement, metronidazole was discontinued on day 4, while vancomycin and cefepime remained. During the late hours of day 5, early day six the patients’ WCC and temperature began to rise, and on day 7 she was restarted on metronidazole and clindamycin. One dose of metronidazole was given in the early morning hours, while clindamycin was given in the early afternoon. About 1 hour post clindamycin (second dose overall), and approximately 5 hours after the last metronidazole treatment, skin manifestations were first detected. At this time, the patient became hypotensive, tachycardic with significant alteration in her mental status. Septic shock was suspected, and the patient received intravenous fluid resuscitation, along with vasopressor support for hypotension and continued with empiric broad-spectrum antibiotics (vancomycin, cefepime, clindamycin and metronidazole). Given her pressor requirement, patient’s level of care was upgraded to ICU. She was sedated due to acute encephalopathy, intubated for mechanical ventilatory support and started on CVVH. Dermatology was consulted for

![Figure 5](http://casereports.bmj.com/bcr-2021-247040)

**Table 1** Relevant laboratory results across clinical timeline

<table>
<thead>
<tr>
<th>Test</th>
<th>Reference range</th>
<th>Results</th>
<th>Day 1</th>
<th>Day 5</th>
<th>Day 7</th>
<th>Day 11</th>
<th>Day 15</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leucocyte count x 10^9/L</td>
<td>1–7</td>
<td>29.4</td>
<td>15.4</td>
<td>57.2</td>
<td>50.3</td>
<td>27.6</td>
<td></td>
</tr>
<tr>
<td>Eosinophil count x 10^9/L</td>
<td>0.03–0.46</td>
<td>0</td>
<td>0.28</td>
<td>0.92</td>
<td>0.5</td>
<td>0.83</td>
<td></td>
</tr>
<tr>
<td>Lactate mmol/L</td>
<td>0.3–1.5</td>
<td>2.5</td>
<td>n.a.</td>
<td>3.7</td>
<td>2.2</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Albumin g/dL</td>
<td>3.5–5.2</td>
<td>3.5</td>
<td>2.8</td>
<td>2.7</td>
<td>2.2</td>
<td>2.3</td>
<td></td>
</tr>
<tr>
<td>Aspartate transaminase U/L</td>
<td>0–37</td>
<td>23</td>
<td>15</td>
<td>24</td>
<td>41</td>
<td>37</td>
<td></td>
</tr>
<tr>
<td>Alanine transaminase U/L</td>
<td>0–35</td>
<td>16</td>
<td>10</td>
<td>20</td>
<td>28</td>
<td>47</td>
<td></td>
</tr>
<tr>
<td>Alk phos U/L</td>
<td>33–133</td>
<td>89</td>
<td>67</td>
<td>158</td>
<td>181</td>
<td>171</td>
<td></td>
</tr>
<tr>
<td>Total bilirubin mg/dL</td>
<td>0.0–1.0</td>
<td>1.1</td>
<td>0.7</td>
<td>0.8</td>
<td>1.4</td>
<td>1.4</td>
<td></td>
</tr>
<tr>
<td>Creatinine mg/dL</td>
<td>0.38–1.02</td>
<td>1.21</td>
<td>1.25</td>
<td>2.82</td>
<td>2.24</td>
<td>1.2</td>
<td></td>
</tr>
<tr>
<td>Comments</td>
<td>Admission</td>
<td>Before rash</td>
<td>Rash detected</td>
<td>Rash improve</td>
<td>End of steroids</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

na, not applicable.

![Figure 6](http://casereports.bmj.com/bcr-2021-247040)
her cutaneous manifestations and per their recommendations pending biopsy results, clindamycin was discontinued (day 8). Despite use of topical corticosteroids (triamcinolone 0.1% ointment), the patient’s skin eruption worsened over the next few hours, coinciding with rapid clinical decline, haemodynamic instability and multiorgan involvement with continued requirement of vasopressors.

By day 9, on AGEP confirmation via biopsy, cefepime and vancomycin were also discontinued due to the possibility of these agents contributing to the condition. Instead piperacillin/tazobactam and linezolid were started for three and 4 days, respectively. Systemic corticosteroid therapy with hydrocortisone 50 mg intravenous every 6 hours was also started, which led to a rapid clinical improvement of the skin manifestations within 24–48 hours post-treatment. Antibiotic therapy was transitioned to monotherapy with intravenous meropenem for 4 days after discontinuation of all other antibiotics. Corticosteroids were used for 8 days total, with the last dose given on day 15 of treatment.

OUTCOME AND FOLLOW-UP

Within 48 hours of clindamycin discontinuation and 24 hours after initiation of intravenous hydrocortisone, the rash improved. The patient’s pustules and bullae progressed to extensive desquamation of her skin (figure 7). Systemic steroids were continued for an additional 5 days, and during this time the patient’s antibiotics were discontinued sequentially with her remaining only on meropenem for the final 4 days of therapy. By day 14, vasopressors were no longer required, and the patient was no longer febrile. She exhibited spontaneous movement of her limbs and was responsive to tactile stimuli. On day 15, the patient began responding to simple questions by moving her head; at this point steroids were discontinued. She remained intubated and continued to receive CVVHD for an additional 12 days. By the conclusion of this report, patient had been extubated, and started on intermittent haemodialysis. Both her WCC and liver enzymes were steadily improving; however, her serum creatinine levels and other markers of kidney function were worsening. Her mentation recovered, and to date she is conscious, alert and oriented, without any focal neurological deficits.

DISCUSSION

AGEP is a severe cutaneous adverse reaction (type intravenous hypersensitivity reaction) classically presenting with the acute eruption of hundreds of sterile pustules most commonly arising in intertriginous areas. Pustules are superimposed on a background of erythematous and edematous skin, with systemic manifestation limited to leukocytosis and fever. AGEP cases are traditionally self-limiting, resolving within 2 weeks of discontinuation of the causative agent with a mortality rate less than 5%. Multi-organ involvement is rarely seen, although severe complications have been reported, especially with atypical presentation, such as the case presented here. Atypical AGEP refers to cases with superimposed manifestations that are not characteristically seen. In our patient, cutaneous findings also included vesicles, bullae, and extensive desquamation. Since these signs are usually associated with TEN, these uncharacteristic AGEP cases are referred to as TEN-like. As observed in our patient, these cases tend to be more severe in their presentation, with signs of septic shock, and may even lead to multiorgan failure. In the case described, our patient required intensive care treatment, vasopressor support and use of systemic corticosteroids.

While some AGEP cases have been attributed to viral infections, mercury contact and insect bites, the majority of cases (>90%) result from medication exposure. The most common
Table 2 Drugs most commonly associated with AGEP development

<table>
<thead>
<tr>
<th>Drug class</th>
<th>Specific agents</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antibiotics</td>
<td>Ampicillin, amoxicillin, ceftriaxone, clindamycin, vancomycin, pristinamycin and spironamycin</td>
<td>1 4 8 11 18 23</td>
</tr>
<tr>
<td>Antifungals</td>
<td>Nystatin, terbinafine, fluconazole</td>
<td>27 28</td>
</tr>
<tr>
<td>Anticonvulsants</td>
<td>Carbamazepine</td>
<td>29</td>
</tr>
<tr>
<td>Antimalarial</td>
<td>Hydroxychloroquine</td>
<td>10 30 31</td>
</tr>
<tr>
<td>Antihypertensive</td>
<td>Diltiazem hydrochloride</td>
<td>32</td>
</tr>
</tbody>
</table>

culprits include antibiotics, antifungals and antihypertensives. Table 2 lists specific agents most frequently associated with AGEP development.

Because a patch test was not performed, the offending drug was unable to be identified. Nonetheless, based on the time period between administration of the drug to onset of symptoms (~80 hours), and consequential improvement of skin manifestations on removal of the medication, we hypothesise that clindamycin was the causative agent. Clindamycin is a well described culprit of not only classical AGEP but also TEN-like incidents. 4 8 11 13 23 24 Although we postulate that clindamycin is the offender, we cannot rule out the role that metronidazole, vancomycin and/or cephepine may have played on this patient’s presentation. Interestingly, cases of AGEP presenting as septic shock, as seen in our patient, have been reported with the use of vancomycin. 8 18 23

Although a comprehensive description of the pathophysiology of AGEP is beyond the scope of this report, briefly, AGEP is categorised as a type intravenous hypersensitivity reaction mediated by CD8+ plus T cells. 26 During the initial phase, activated cytotoxic T-cells (referred in the literature as drug-specific T-cells), migrate to the skin and contribute to the development of subcorneal vesicles by inducing keratinocyte apoptosis via granzyme B, perforin, and the Fas ligand mechanism. 2 26 Infiltreation of CD4 + T cells leads to the release of CXCL-8, a potent neutrophil chemotactic agent, causing subsequent recruitment of neutrophils and transformation of vesicles into pustules which are characteristic of AGEP. 2 26 Other compounds released by drug-specific T-cells include interferon-gamma and granulocyte/macrophage colony stimulating factor which further prolong the viability of neutrophils. 2

Diagnosis of AGEP can be made by clinical assessment for which the EuroSCAR scoring system developed by Sidoroff et al5 can be helpful. This system combines histological as well as clinical features to determine the likelihood of an AGEP diagnosis. Description of the rash, its distribution, timing, and presence of other symptoms are all taken into account. The algorithm yields a score between −18 to +12, confirming the diagnosis of AGEP. 5

Table 3 AGEP validation score developed by EuroSCAR (European Severe Cutaneous Adverse Reactions) study group5

<table>
<thead>
<tr>
<th>Variable</th>
<th>Our patient results</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Morphology</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pustules</td>
<td>Typical</td>
<td>+2</td>
</tr>
<tr>
<td>Erythema</td>
<td>Typical</td>
<td>+2</td>
</tr>
<tr>
<td>Distribution/pattern</td>
<td>Typical</td>
<td>+2</td>
</tr>
<tr>
<td>Course</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mucosal involvement</td>
<td>Yes</td>
<td>+1</td>
</tr>
<tr>
<td>Acute onset (&lt;10 days)</td>
<td>Yes</td>
<td>+1</td>
</tr>
<tr>
<td>Fever ≥38°C</td>
<td>Yes</td>
<td>+1</td>
</tr>
<tr>
<td>Polymorphonuclear neutrophils &gt;7000/mm³</td>
<td>Yes</td>
<td>+1</td>
</tr>
<tr>
<td>Histology</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spongiform, subcorneal pustule(s) with papillary oedema, perivascular infiltrates of neutrophils</td>
<td>Yes</td>
<td>+3</td>
</tr>
<tr>
<td>Overall score (−18 to +12)</td>
<td>12</td>
<td></td>
</tr>
</tbody>
</table>

Giving the probability of AGEP diagnosis.

AGEP, acute generalised exanthematosus pustulosis.

Learning points

- Acute generalised exanthematosus pustulosis (AGEP) is a rare, cutaneous drug reaction, classically presenting with small non-follicular pustules primarily visualised in intertriginous areas.
- AGEP can present with atypical skin manifestations mimicking TEN and should be suspected in patients being managed for septic shock associated with a negative infectious workup, along with a deteriorating clinical condition while on anti-infective agents.
- Most cases of AGEP are self-limited with good prognosis; however atypical TEN-like AGEP presentations can be more severe and lead to multigorgan failure.

Table 3 shows the scores allocated to our patient for each category. Clinical suspicion of AGEP can be validated by skin biopsy analysis.

Our case presents several unique factors. First, it is not a typical presentation of AGEP, involving signs that are usually seen in more severe conditions such as TEN. Furthermore, unlike most AGEP cases, our patient had multi-system involvement, with rapid deterioration, manifesting with hypotension, tachycardia, tachypnoea and fever; a picture reminiscent of septic shock. Moreover, while discontinuing the suspected drug, and supportive care is the principal treatment, our case presents an instance in which treatment with systemic steroids was warranted for clinical recovery. To the best of our knowledge, there have been only a few reported cases of AGEP with such severe presentation. It is crucial for physicians to become acquainted with the rarer AGEP manifestations so that recognition may lead to a faster, and accurate diagnosis thus facilitating a more appropriate treatment plan.

Contributors KC contributed in the drafting, editing and revision of the case report. TC contributed in the editing, revision and submission of the case report. AS contributed in the editing and revision of the case report. Consent obtained directly from patient(s). None declared.

Funding The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

Competing interests None declared.

Patient consent for publication Consent obtained directly from patient(s).

Provenance and peer review Not commissioned; externally peer reviewed.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially.
Acute generalised exanthematous pustulosis: an update.

REFERENCES


