AstraZeneca (AZD1222) COVID-19 vaccine-associated adverse drug event: A case report

In the current pandemic of SARS-CoV-2, the fast development of vaccines was mandatory due to many deaths and cases across the world since the beginning of the outbreak.

Questions about the potential immunogenicity of the vaccine and the safety in the short and long term have been raised.1,2 Reports of cutaneous reactions have been made for other vaccines (as small case series following Moderna mRNA-1273 vaccination).3

We report the case of a 37-year-old woman, with no medical history nor allergies, who developed a skin rash 4 days after the vaccination with the AstraZeneca AZD1222 COVID-19 vaccine.

Rash was characterized by plaques of vesicular, vesicular-papular, partially confluent lesions with peripheral halo and necrotic center on thighs, feet, and face, without dermatomal distribution, nor hitching (Figure 1).

She also had arthritis of metacarpophalangeal joints (demonstrated with ultrasound), pitting edema of hands and feet, muscle weakness, and foot neuropathic pain.

We performed three different sites skin punch biopsies of different evolutive types of lesions, with specimens sent for histological, immunohistochemical, and immunofluorescence analysis.

We collected blood samples for ANA, ANCA, dsDNA antibodies, connective tissue disease immunoblot, cryoglobulins.

We started an empirical course of clarithromycin and valaciclovir before steroid infusion and for the following two weeks.

She received a 125 mg infusion of methylprednisone succinate and subsequent oral tapering (Prednisone 25 mg) in 3 weeks until complete suspension. The lesions rapidly disappeared in the first week.

On blood exams CRP elevation in a neutrophilic state (13.3 × 10^9/L leukocytes with 76.3% of neutrophils) was present. Procalcitonin and autoimmunity assay were negative.

Histology showed neutrophilic pustular dermatitis at different evolutive states sometimes with the presence of necrotizing neutrophilic infiltrate with nuclear debris (“nuclear dust”), without any demonstration of vasculitis. Immunohistochemistry showed a prevalence of neutrophils (MPO+) and histiocytes-macrophages (CD68+) with rare T (CD3+) and B (CD20+) cells without plasma cells (Figure 2).

The immunofluorescence was negative, so we classified the skin rash as a Sweet syndrome.4,5

Four weeks after steroid discontinuation, the patient underwent an electromyographic study due to the persistence of foot neuropathic pain, muscle weakness, and fatigue.

Myositis in the tibialis anterior muscle was found without evidence of polyneuropathy.

We repeated auto-antibodies: ANA was positive (1:160 Ac1, Ac 4.5) with borderline positivity of anti-Pm/scl-75 antibodies with aldolase raise (10.3 U/L NV < 7.6).

FIGURE 1 Different lesion types in (A), (C), (E), and (F). Pitting edema in (B) and (D). Arthritis in (B)
We repeated electromyography with the demonstration of moderate severity subacute myositis without muscle damage and polyneuropathy.

We made a diagnosis of polymyositis secondary to an autoimmune reaction following vaccination.

At present time, the patient is scheduled to be treated with IVIg (2 g/kg).

To our knowledge, this is the first reported case of adverse reaction to the AstraZeneca COVID vaccine that had studied in detail the skin and systemic autoimmune reaction.

Development of autoimmune reaction following SARS-CoV-2 infection has been described extensively6,7; however, evidence of autoimmunity following vaccination seems to lack at present. Our case suggests that in predisposed subjects’ vaccination could trigger an autoimmune reaction as the natural infection. Larger and accurate immunological studies are needed to elucidate this interrogative that is of fundamental relevance due to the future mass vaccination.

CONFLICT OF INTERESTS

The authors declare that there are no conflict of interests.

AUTHOR CONTRIBUTIONS

Drafting of the work, biopsy sampling, and patient management: Capassoni Marco. Histological examination and critical review: Ketabchi Sheyda and Cassisa Angelo. Electrophysiological study and critical review: Caramelli Riccardo. Critical review: Molinu A. Anna. Critical review, biopsy sampling, and patient management: Galluccio Felice.

Coordinator and critical review: Guiducci Serena.

Marco Capassoni1,2, Sheyda Ketabchi3, Angelo Cassisa3, Riccardo Caramelli4, Anna A. Molinu5, Felice Galluccio2, Serena Guiducci1,2

1Department of Clinical and Experimental Rheumatology, University of Florence, Florence, Italy
2Department of Rheumatology, Azienda Ospedaliero Universitaria Careggi, Florence, Italy
3Department of Anatomopathology, Ospedale San Giovanni di Dio, Florence, Italy
4Department of Neurophysiopathology, Azienda Ospedaliero Universitaria Careggi, Florence, Italy
5Dermatology, Istituto Ortopedico Toscano, Florence, Italy

Correspondence

Marco Capassoni, Department of Clinical and Experimental Rheumatology, University of Florence, Florence, Italy. Email: marco.capassoni@gmail.com

ORCID

Marco Capassoni http://orcid.org/0000-0003-2209-4450
Riccardo Caramelli https://orcid.org/0000-0002-7110-040X
Felice Galluccio https://orcid.org/0000-0001-7485-471X
Serena Guiducci https://orcid.org/0000-0003-2722-6475

REFERENCES

1. Akinosoglou K, Tzivaki I, Marangos M. Covid-19 vaccine and autoimmunity: Awakening the sleeping dragon. Clin Immunol. 2021;226:108721. https://doi.org/10.1016/j.clim.2021.108721
2. Talotta R. Do COVID-19 RNA-based vaccines put at risk of immune-mediated diseases? In reply to “potential antigenic cross-reactivity
between SARS-CoV-2 and human tissue with a possible link to an increase in autoimmune diseases”. 

Clin Immunol. 2021;224:108665.

3. Blumenthal KG, Freeman EE, Saff RR, et al. Delayed large local reactions to mRNA-1273 vaccine against SARS-CoV-2. 

N Engl J Med. 2021;384(13):1273-1277. https://doi.org/10.1056/NEJMct2102131

4. von den Driesch P. Sweet’s syndrome (acute febrile neutrophilic dermatosis). 

J Am Acad Dermatol. 1994;31(4):535-556. https://doi.org/10.1016/s0190-9622(94)70215-2

5. Su WP, Liu HN. Diagnostic criteria for Sweet’s syndrome. 

Cutis. 1986;37(3):167-174.

6. Caso F, Costa L, Ruscitti P, et al. Could Sars-cornavirus-2 trigger autoimmune and/or autoinflammatory mechanisms in genetically predisposed subjects? 

Autoimmun Rev. 2020;19(5):102524. https://doi.org/10.1016/j.autrev.2020.102524

7. Rodríguez Y, Novelli L, Rojas M, et al. Autoinflammatory and autoimmune conditions at the crossroad of COVID-19. 

J Autoimmun. 2020;114:102506. https://doi.org/10.1016/j.jaut.2020.102506