To the Editor,

STAT1 gain-of-function (STAT1 GOF) mutations underlie chronic mucocutaneous candidiasis (CMC), a phenotypically diverse inborn error of immunity, ranging from isolated infectious susceptibility to complex immune deficiency [1]. The hypermorphic effect of the STAT1 mutation is likely due to reduced STAT3 promoter binding as a result of strong antagonizing type I and II interferon (IFN-I, IFN-II)-driven STAT1 recruitment [2]. Recently, JAK inhibitors, such as ruxolitinib, operating upstream from STATs, were shown to suppress the STAT1 response to IFNs and ameliorate the immune dysregulation [3].

The host immune response to SARS-CoV-2 infection, the cause of COVID-19 disease, is largely governed by the innate viral sensing mechanisms [4, 5]. Following their activation in the infected cells, the upregulation of the downstream transcription factors leads to increased production of IFNs. IFNs act as ubiquitous alarm triggers for the neighboring cells, activating the JAK/STAT pathways to augment the anti-viral defenses [6].

Recently, the most severe forms of COVID-19 were shown to be associated with disrupted IFN-I signaling, either due to inborn genetic errors or the presence of anti-IFN antibodies [7, 8]. Moreover, the early therapeutic enhancement of IFN response may be beneficial in preventing the infection and lowering mortality [9].

Hypothetically, STAT1 GOF patients may be to some degree protected from SARS-CoV-2 infection/severe COVID-19 via the pre-emptive overactivation of IFN-I signaling, despite their otherwise immune deficient status. On the other hand, abnormally augmented IFN response may contribute to the catastrophic cytokine-driven hyperinflammation in delayed stages of COVID-19 [10]. So far, three STAT1 GOF patients were reported to suffer COVID-19 (one on ruxolitinib), all experiencing a mild course [11–13].

The mRNA-based SARS-CoV-2 vaccines activate the mechanisms of innate immunity, resulting in the production of multiple inflammatory mediators, including IFNs, which effectively promote both T and B cell-mediated response and antigen-specific immune memory [14]. The introduction of such agents into an IFN-biased environment in STAT1 GOF might amplify the immune dysregulation-related symptoms. Conversely, STAT1 GOF patients receiving JAK inhibitors, such as ruxolitinib, may have blunted vaccine response, due to the artificially suppressed JAK/STAT pathway. Recently, two STAT1 GOF CMC patients were reported to mount a normal vaccine-specific anti-spike 1 antibody response and one who failed to do so [15, 16].

Here, we report seven adult STAT1 GOF patients (two with novel mutations) with an uneventful course of COVID-19 vaccination, and/or SARS-CoV-2 infection, including two patients receiving JAK inhibitor ruxolitinib.

Results

Seven adult Czech STAT1 GOF CMC patients (previously described p.Y68C, p.A267V, p.M390T and novel p.T288N and p.E29A; cohort characteristics detailed in Supplementary Table 1) received SARS-CoV-2 spike 1 protein-encoding mRNA vaccine (Comirnant®, Pfizer/BioNTech). All seven received two doses, six of them 3 to 6 weeks apart, one patient 4 months apart. Two patients were receiving ruxolitinib at the time of vaccination. No adverse events were
noted during 1–4-month follow-up. The antibody response was determined prior to the vaccination (Enzyme immunoassay COVID-19 RBD by TestLine Clinical Diagnostics, Czech Republic) and 3–5 weeks after the second vaccine dose (Microblot-Array COVID-19 by TestLine Clinical Diagnostics, Czech Republic). Four seronegative patients achieved vaccine-specific receptor-binding domain spike protein (RBD) IgG seroconversion (median 1211.5, range 999–1347 U/ml), which was comparable to vaccinated healthy controls ($N = 100$, median 985.5, range 470–1924 U/ml). Only two patients responded with anti-RBD IgA, compared to 76 out of 100 healthy controls. This may be an important efficacy parameter, as serum IgA was shown to be an early SARS-CoV-2–neutralizing agent [17]. Despite the absence of any previous COVID-19 symptoms, one patient (P5M390T) was found to be S1-spike protein seropositive before vaccination, and one patient (P4A267V, unavailable for testing prior to vaccination) had anti-nucleocapsid, as well as anti-RBD antibodies after vaccination, suggestive of past infections in both patients. One patient did not mount an antibody response after vaccination (P7E29A) (Fig. 1A). Regardless of the serologic status, six out of seven patients (one patient not tested), including the two ruxolitinib receivers, were found to mount a virus-specific (S1-spike protein) cellular immune response (median 1192.5, range 585–4115 mIU/ml), which was comparable to vaccinated healthy controls ($N = 7$, median 1830, range 675–1735 mIU/ml), detected by whole blood IFN-γ release assay (EUROIMMUN SARS-CoV-2 IGRA, Germany) (Fig. 1B). The summary of the humoral and cellular COVID-19-related immune parameters is listed in Table 1.

Within the cohort, two patients were of particular interest. An 18-year-old female PsM390T with CMC and severe pulmonary damage had asymptomatic COVID-19 infection, as evident by anti-S1 seropositivity prior to vaccination. Despite that, the anti-nucleocapsid antibodies were not detectable. This patient has been on ruxolitinib (0.4 mg/kg/day, dose titered to healthy control’s phosphorylated STAT1 level) for the past 6 months; however, it is unknown whether the infection occurred prior or after the JAK inhibitor initiation.

The second patient, a 45-year-old male P7E29A with CMC, severe pneumopathy, and multiple autoimmune phenomena, was vaccinated while COVID-19 naïve and 5 months on ruxolitinib (0.2 mg/kg/day, dose titered to healthy control’s phosphorylated STAT1 level). The patient failed to produce detectable SARS-CoV-2 antibodies. However, this may be attributed to continuing B cell impairment after previous CD20-depletion therapy (rituximab) 18 months prior to vaccination; while the overall CD19 B cells count represented 5.7% of lymphocytes, the B cell pool constituted predominantly of CD27 negative naïve B cells prior to vaccination. In fact, such skewed antibody response to primary antigen exposure post-rituximab has been reported by multiple studies, including, recently, the impaired response to SARS-CoV-2 vaccines [16, 18]. Nevertheless, the patient developed a strong S1-peptide-specific T cell immune response in IGRA, interestingly, the strongest in our cohort (4115 mIU/ml; Fig. 1B).

In summary, while limited by the cohort size and the absence of antibody-neutralization assays, our observations indicate that SARS-CoV-2 mRNA vaccination in
STAT1 GOF is immunogenic and may be safe, even during treatment with JAK inhibitor and after past infection. Furthermore, we affirm that COVID-19 disease may take a mild/asymptomatic course in STAT1 GOF CMC. This data may help guide clinical counseling for patients with STAT1 GOF CMC.

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Author Contribution MB treated the patients, established the hypothesis, and wrote the manuscript. ZP established the hypothesis and co-wrote the manuscript. JH treated the patient and co-organized the sampling. JL carried out the testing and analyzed the samples. AS treated the patients, established the hypothesis, and co-wrote the manuscript. All authors reviewed the manuscript.

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Data Availability On a reasonable request, the data supporting study’s findings are available from the corresponding author.

Code Availability Not applicable.

Declarations Ethics Approval Ethical review and approval was not required for the study on human participants in accordance with the local legislation and institutional requirements.

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