To the editor

Patients with severe combined immunodeficiency (SCID) have an increased incidence of BCG disease following vaccination compared to other infants [1]. BCG disease is classified in local, regional, distant, and disseminated [2]. Disease flares often occur following hematopoietic stem cell transplantation (HSCT). Consequently, management of BCG disease in SCID patients undergoing HSCT remains a challenge, and optimal management is not standardized.

We retrospectively analyzed the clinical course of BCG disease in 36 SCID patients who underwent HSCT at the Department of Pediatrics at the University Medical Center Ulm between 1983 and 2014, focusing on the impact of immune reconstitution and tuberculostatic therapy. All patients had a documented BCG vaccination. The study was approved by the ethical committee of Ulm University. Furthermore, we collected information via a standardized questionnaire about current BCG disease management in 15 international centers associated with the European Society for Blood and Marrow Transplantation (EBMT). Results from the international survey can be found in Table 1 of the supplemental online material.

Main clinical and epidemiological characteristics of the cohort are summarized in Table 1. Although BCG strains were not documented in our patient files, we were able to map the strains using http://BCGatlas.org, which allows identification of most strains according to year of vaccination and location. Not for all, but for 28/36 patients we managed to infer the BCG strain (Table 2, supplemental online material): respectively Copenhagen 1331 (25 children from Germany, Spain, and Latvia), Moreau (1 child from Croatia), Tokyo-172 (1 child from Portugal), Pasteur (1 child from Bosnia). For UAE and Morocco there are no official data available, whereas in Saudi Arabia three strains were in use (Copenhagen 1331, Pasteur, Tokyo-172). The Copenhagen 1331 strain is classified by the WHO as a “strong” one (WHO, Recommendations to assure the quality, safety and efficacy of BCG vaccines, Annex 3, 2013). We then compared time to remission from BCG disease in patients who received Copenhagen 1331 (n = 25) with all others (n = 11).

Surprisingly, time to remission was significantly longer in patients following vaccination with the Copenhagen 1331 strain as shown in Table 2 of the supplements (464 days post HSCT vs 191 days, p = 0.01).

Molecular genetics of SCID variants were determined in 26 patients (72%): IL2RG deficiency and RAG1/RAG2 defects (including Omenn syndrome) were the most frequently reported, followed by Artemis (DCLRE1C), JAK3, ADA, AK2, IL7RA, ZAP70, and CD3ε. We analyzed the distribution of BCG disease subtype on the basis of immunophenotype (T−B+ versus T−B− SCID). There was no difference between these groups as for distribution of disease subtype. There was no statistical significance of BCG disease subtype with respect to the presence of T cells or MFT. Most infants presented with a combination of respiratory tract infections, enteritis, and failure to thrive. Rotavirus was detected in 12 patients, whereas 7 children were infected with cytomegalovirus (CMV). BCG disease was often one of the initial manifestations of immunodeficiency: 24 patients developed symptoms of BCG disease before or concomitantly with SCID diagnosis. In our cohort, most patients who presented with first BCG disease symptoms prior 4 months of age developed disseminated disease, though these results
locoregional No BCG trial therapy was started concomitantly (mostly quinolones and aminoglycosides. Antimycobactericidal drugs were administrated on average for 12 or more months in most patients. The large majority (25/26) of patients received myeloablative conditioning before transplantation: 18 with reduced intensity conditioning (RIC) and/or immunosuppressive drugs. The large majority (25/26) of haploidentical grafts had been CD34 + enriched and T cell depleted via lectin-agglutination and erythrocyte-rosette depletion or by immunomagnetic methods. One patient showed HLA homozygosity for three loci (A, B, and C) and was HLA-identical in the DRB1 locus with his mother; hence, this graft was not manipulated before transplant. Four patients received myeloablative conditioning before transplantation: 18 with reduced intensity conditioning (RIC) and/or immunotherapy (IT), and 14 without conditioning. There were no differences in time to remission from BCG disease at 12 months after HSCT (range: 1–31 months), depending on the clinical course and immune reconstitution. This was in line with the results from our international survey, as tuberculostatic therapy is applied for 12 or more months in most centers (9/15).

HSCT was performed in all patients to cure underlying SCID (Table 3a/3b, supplemental online material). Haploidentical or mismatched familiar donors were available for the majority of children (n = 26), whereas matched familiar and matched unrelated donor could be identified for 7 and 3 patients respectively. A total of 34/36 patients received GvHD prophylaxis via T cell depletion of the graft, and/or immunosuppressive drugs. The majority (25/26) of haploidentical grafts had been CD34 + enriched and T cell depleted. Patients with symptoms at 12 months’ follow-up (n = 10) had all been symptomatic for BCG disease from the beginning; hence, absence of BCG disease symptoms at first medical assessment correlates with a better outcome. However, these results were not statistically significant.

Tuberculostatic drugs were administrated on average for 12 months after HSCT (range: 1–31 months), depending on the clinical course and immune reconstitution. This was in line with the results from our international survey, as tuberculostatic therapy is applied for 12 or more months in most centers (9/15).
developed acute GvHD (15 following haploidentical and 6 following HLA-identical transplantation), which resulted in chronic GvHD in 6 of them. Patients who underwent haploidentical HSCT or were treated with immunosuppressive therapy due to GvHD presented more often with locoregional as compared to disseminated BCG disease, but this finding was not statistically significant. Time to remission from BCG disease did not substantially differ between haploidentical or HLA-identical transplantation. In our international survey, nine centers would administer not only tuberculostatic but also immunosuppressive treatment to better control BCG disease flares. Presence of BCG disease, however, would not influence donor choice, and HLA-identical donors are mostly preferred.

BCG disease worsened in 27/36 patients following HSCT (Fig. 1, supplemental material). Exacerbation occurred either early after HSCT (2–16 days) in 9 patients or concomitantly with T cell immune reconstitution (defined as CD3 + T cells > 500/μL) in 18 patients. In 3 of these late occurrences, the Mantoux test turned positive during disease flare. Early exacerbation, which is rarely described in the literature, occurred mainly after transplantation from HLA-identical donors.

Cure from BCG disease was documented in 27/36 patients or 75% (median time of absence of BCG disease signs and symptoms: 380 days after HSCT, range 14–930). Locoregional disease resolved significantly earlier than disseminated BCGitis (220 and 458 days respectively, p = 0.03).

In the course of the 30 years analyzed, distribution of BCG disease subtype or time to remission from BCG disease remained consistent. It is however worth mentioning that all deaths occurred before 1995.

Our data emphasize the importance of early SCID diagnosis and HSCT to prevent BCG disease outbreak and/or severe BCG disease in patients with a history of BCG vaccination, as previously observed [1, 3, 4]. Although no risk factors for developing severe (i.e., disseminated) BCG disease were identified when analyzing data according to underlying SCID gene variant or immunophenotype, time of treatment plays an important role in management of BCG disease. Patients presenting BCG disease symptoms early in life (< 4 months) seem to have a higher risk for dissemination. This finding needs to be verified in an even larger cohort. Initiating tuberculostatic treatment at the time of SCID diagnosis—even in the absence of BCG symptoms—may protect from BCG complications [1]. Furthermore, our results suggest that tuberculostatic treatment initiated prior HSCT is usually associated with better prognosis 1 year after HSCT. Data about whether preemptive treatment with a single antimycobacterial agent versus combination therapy is indicated is inconclusive. However, there are reports that routine administration of isoniazid alone was not effective in preventing BCG dissemination in BCG-vaccinated SCID patients (Verzegnassi Fiorotte et al., Poster Session ESID Meeting 2020, unpublished data). Careful observation is always recommended in the absence of BCG symptoms [5].

BCG-vaccinated SCID patients who undergo HSCT are at high risk for exacerbation of BCG disease. This may occur concomitantly with immune reconstitution shown by tuberculin test positivity [1], reminiscent of IRIS (immune reconstitution inflammatory syndrome) in HIV-infected individuals started on antiretroviral therapy [2]. In presence of T-replete grafts, exacerbation can occur few days after transplantation. Whether T cell reduction may prevent or dampen inflammatory response in the context of immune reconstitution remains an open question. The use of immunosuppressive drugs to control disease flare can be discussed and is currently adopted in some centers (results from international survey, Table 1 supplemental online material).

In conclusion, HSCT is curative for most SCID patients with BCG disease. Early initiation of tuberculostatic treatment prior HSCT is beneficial. We recommend early combination of antimycobacterial drugs (≥ 2) prior HSCT in BCG-vaccinated SCID babies including for locoregional disease or asymptomatic infants. Moreover, timely HSCT should be encouraged as it is associated with less severe, i.e., locoregional, disease (p = 0.03).

Identifying BCG disease flares, which frequently occur following HSCT, is necessary in order to intensify tuberculostatic therapy; furthermore, adjuvant immunosuppressive therapy may also be useful.

Delaying BCG vaccination is key for preventing BCG disease in undiagnosed SCID babies. With the expanding availability of TREC screening for T cell lymphopenia in newborns, timing of BCG vaccination should be reevaluated by local health authorities, and no longer be administered right after birth.

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**Declarations**

**Conflict of interest** The authors declare no competing interests.

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**References**

1. Marciano BE, Huang CY, Joshi G, et al. BCG vaccination in patients with severe combined immunodeficiency: complications, risks, and vaccination policy. J Allergy Clin Immunol. 2014;133:1134–41.
2. Hesseling AC, Rabie H, Marais BJ, et al. Bacille Calmette-Guérin vaccine-induced disease in HIV-infected and HIV-uninfected children. Clin Infect Dis. 2006;42:548–58.
3. Mazzucchelli JT, Bonfim C, Castro GG, et al. Severe combined immunodeficiency in Brazil: management, prognosis, and BCG-associated complications. J Investig Allergol Clin Immunol. 2014;24:184–91.
4. Bacalhau S, Freitas C, Valente R, et al. Successful handling of disseminated BCG disease in a child with severe combined immunodeficiency. Case Rep Med. 2011;2011: 527569. https://doi.org/10.1155/2011/527569.5pages.
5. Bernatowska E, Wolska-Kusnierz B, Pac M, et al. Risk of BCG infection in primary immunodeficiency children. Proposal of diagnostic, prophylactic and therapeutic guidelines for disseminated BCG based on experience in the Department of Immunology, Children’s Memorial Health Institute in Warsaw between 1980-2006. Centr Eur J Immunol. 2007;32(4):221–5.