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Clinical presentation, immunologic features, and hematopoietic stem cell transplant outcomes for IKBKB immune deficiency

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ABSTRACT

IKBKB immune deficiency is a rare but life-threatening primary immunodeficiency disorder, involving activation defects in adaptive and innate immunity. We present sixteen cases of a homozygous IKBKB mutation (c.1292dupG) in infants characterized by early-onset bacterial, viral, fungal and Mycobacterial infections. In most cases, T- and B-cells were quantitatively normal, but phenotypically naïve, with severe hypogammaglobulinemia. T-cell receptor excision circles were normal, meaning newborn screening by TREC analysis would miss IKBKB cases. Although IKBKB immune deficiency does not meet traditional laboratory based definitions for SCID, this combined immune deficiency appears to be at least as profound. Urgent HSCT, performed in eight patients, remains the only known curative therapy, although only three patients are survivors. Ongoing infections after transplant remain a concern, and may be due to combinations of poor social determinants of health, secondary graft failure, and failure of HSCT to replace non-hematopoietic cells important in immune function and dependent upon IKK/NF-κB pathways.

1. Introduction

In the Canadian province of Manitoba, our group has periodically managed young infants of Northern Cree First Nations (Aboriginal) descent presenting with early-onset and life-threatening viral, bacterial, Mycobacterial, and fungal infections, clinically resembling severe combined immune deficiency (SCID). Infants were initially identified in the 1970s from two small northern Manitoba communities. In later years, infants from the neighboring province of Saskatchewan were also identified. In 2013, our group and collaborators identified the genetic defect in four patients as being due to a homozygous duplication mutation (c.1292dupG) in exon 13 of IKBKB, the gene encoding inhibitor of nuclear factor kappa B kinase subunit beta (IKKβ), also known as IKK2 [1].

The homozygous IKBKB duplication mutation seen in our population results in complete loss of IKKβ expression, a critical component of the canonical IKK-nuclear factor kappa B (NF-κB) pathway [1]. Normally, signaling through tumor necrosis family receptors, toll-like receptors, and antigen receptors on T- and B-cells induces activation of the IKK complex, which consists of the kinases IKKα and IKKβ, as well as the regulatory protein nuclear factor Kappa-B essential modifier (NEMO) [2]. The active IKK complex, and specifically IKKβ, phosphorylates the inhibitory protein IκBα, resulting in release of cytoplasmic NF-κB. Free from the influence of IκBα, NF-κB translocates to the nucleus to induce the transcription of a multitude of immune and inflammatory genes.

Given the central role of the IKK/NF-κB pathway in regulating both

Abbreviations: BCG, Bacillus Calmette-Guerin; CID, Combined Immune Deficiency; CMV, Cytomegalovirus; HSCT, Hematopoietic stem cell transplant; HLH, Hemophagocytic Lymphohistiocytosis; IKKβ, Inhibitor of nuclear factor kappa B kinase subunit beta; NEMO, Nuclear factor Kappa-B essential modifier; PHA, Phytohemagglutinin; RSV, Respiratory Syncytial Virus; SCID, Severe combined immune deficiency; TCR, T-cell receptor; TRECs, T-cell receptor excision circles

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innate and adaptive immune responses, it is not surprising that patients with mutations in genes from this pathway manifest with primary immunodeficiency disorders [3]. Males with hypomorphic hemizygous mutations in IKKBG, which encodes for the NEMO protein and results in reduced NEMO expression, present with a combined immune deficiency that includes enormous allelic, immunologic and clinical heterogeneity [4–9]. Patients with hypermorphic gain of function mutations in NFKBIA [10–12], resulting in reduced degradation of the IκBα protein, have also been shown to exhibit significant combined immune deficiencies. Furthermore, the IKK/NF-κ pathway operates in both hematopoietic and non-hematopoietic cells, leading to the extra-immune findings seen in patients with defects in NEMO and IκBα, such as ectodermal dysplasia and colitis.

Similar to patients with IKBKG and NFKBIA mutations, the first four patients reported by our group with deficiency of IKKβ due to the c.1292dupG IKKB mutation also presented with immune deficiency [1]. Since then, six additional patients from various ethnic backgrounds (including Turkey, Qatar, and the Arabian Peninsula) with different nonsense IKKB mutations have been reported, with all patients presenting with early onset and profound combined immune deficiencies, but typically without findings of ectodermal dysplasia [13–15]. In our initial report, patients had normal to near-normal numbers of T- and B-cells of an almost exclusively naive phenotype, severe hypogammaglobulinemia, and activation defects to a variety of stimuli in a number of immune cells, both innate and adaptive, more suggestive of a global functional defect as opposed to a quantitative lymphocyte deficiency [1].

Herein we describe the clinical presentation, immunologic features, and HSCT outcomes for the largest cohort of infants with IKKβ immune deficiency resulting from complete loss of IKKβ expression published to date.

2. Materials and methods

2.1. Patients and study design

The study protocol was approved by the Health Research Ethics Board of the University of Manitoba. The authors identified all known (genetically confirmed) or highly suspected infants with the IKKβ mutation managed through the Department of Pediatrics, Winnipeg Children's Hospital, CancerCare Manitoba and the Manitoba Blood and Marrow Transplant Program dating to the 1970s. Data collected by retrospective chart review included clinical presentation, immunologic features, and HSCT approaches and outcomes. Although data was complete for patients diagnosed in a contemporary era (after 2000), data availability was variable for some patients before this (1970–2000). Young infants of Northern Cree descent from the known affected communities presenting with a clinical picture consistent with a severe immune deficiency but without genetic confirmation of an IKKβ mutation were considered highly suspected if they also had a close familial relationship to a genetically confirmed infant with the mutation (e.g. sibling, or second-degree relative where there was known consanguinity within the family). We chose to report the decade of diagnosis (instead of year) to provide the reader with perspective on the immunologic investigations and treatments available at the time, while protecting against the potential for inadvertently identifying individuals, given that patients and their families reside in small communities. Detailed family trees documenting consanguineous relationships within families have not been included for the same reason.

2.2. Immunologic and genetic investigations

Patient laboratory values, including absolute lymphocyte count, CD3, CD4 and CD8 T-cell counts, CD19 B-cell counts, CD56 NK-cell counts, and immunoglobulin levels were compared against published, age-adjusted variables [16,17]. Laboratory parameters are described as low, normal, or high with respect to these ranges. Maternal engrafment of T-cells was assessed by analysis of short tandem repeats using the AmpF STR® Profiler Plus™ PCR Amplification Kit from Life Technologies. Chimerism analysis after HSCT was performed on T-cells, B-cells, and myeloid cells labeled with monoclonal antibodies to CD3, CD19, and CD33/66b and separated using an automated immunomagnetic cell separation method with the RoboSep instrument from StemCell Technologies. Due to a potential paucity of cells with each chimerism test, purity of the various lymphocyte and myeloid subsets was not evaluated with each assay. From validation experiments, however, the purity of separated subsets is > 95% with a ± 5% error around lineage specific chimerism results.

T-cell proliferative responses using PHA were performed by 3H-thymidine incorporation assay of peripheral blood mononuclear cells after stimulation with PHA at 1:400–1:800 following a 3-day culture. Percentage of normal PHA response was measured by the stimulation index of the patient relative to the average stimulation index of two concurrently run healthy adult control samples. TREC analysis, when performed at the time of diagnosis (n = 6), was either done locally in our laboratory (n = 3) [18] or at a clinical laboratory (Mayo Clinical Laboratories) (n = 3). Units of measure for TRECgs (copies/μl. whole blood, copies/million PBMCs, copies/million CD3 cells) changed over time.

Sanger sequencing for the c.1292dupG in exon 13 of IKKβ was performed locally on a 3500 Genetic Analyzer™ as per the method developed by Pannicke [1]. Samples containing homozygous G insertions within the region of interest were classified as affected, and confirmed by Sanger sequencing at an independent laboratory.

3. Results

3.1. Clinical presentation of IKKβ immune deficiency

Sixteen patients with genetically confirmed (n = 11) or highly suspected (n = 5) IKKβ mutations were included (Table 1). Genetic confirmation for the five infants with highly suspected IKKβ immune deficiency was not possible since all had died, and DNA was not available (all were born in years before the IKKβ mutation was known). One patient was diagnosed immediately after birth by direct IKKβ mutational analysis of a Guthrie card, as part of a targeted newborn screening project started in November 2013 in two of the heavily affected communities. This patient’s younger sibling was later diagnosed in-utero, due to the known family history, with confirmation of the mutation after birth. Both siblings were placed in protective isolation and proceeded to HSCT by 1–2 months of age. The other fourteen patients presented to health care facilities with severe infections at a median of 2.5 months of age (range: 2 weeks - 4 months). Twelve patients came from the province of Manitoba and four from the province of Saskatchewan. Four patients were previously reported in less detail in our initial report of IKKβ immune deficiency [1]. We included these patients in order to describe additional clinical and immunologic characteristics and provide updates on long-term outcomes.

Details on the number and type of infections present at the time of initial presentation (before HSCT) were complete for twelve and incomplete for two (patient 1 who died of disseminated Mycobacterium bovis-Bacillus Calmette-Guerin (BCG) and patient 13 who died of unknown sepsis). The two siblings identified by newborn screening and family history had no infections at the time of diagnosis of IKKβ immune deficiency and proceeded to urgent HSCT without infections.

For patients not identified by newborn screening, clinical presentation consisted of a multitude of severe bacterial, fungal, Mycobacterial, and viral infections beginning at an early age (< 3 months), usually in association with failure to thrive. Mucocutaneous infection of the oropharynx and perineum with Candida was common (n = 11) with evidence of disseminated Candida infection
| Patient | 1 | 2 | 3 | 4 | 5 | 6 |
|---------|---|---|---|---|---|---|
| Decade of Diagnosis | 1970s | 1970s | 1980s | 2000s | 1990s | 2000s |
| IKBKB Mutation Confirmed | No | No | Yes | Yes | Yes | Yes |
| Direct Family Relationships to Known IKBKB Cases | Affected siblings, and second degree relative with confirmed mutation | Affected siblings, and second degree relative with confirmed mutation | Affected siblings, and second degree relative with confirmed mutation | Affected second degree relatives | Affected half sibling and distant relative | Affected half sibling and distant relative |
| Age at Clinical Presentation | Unknown | 3 months | 2 weeks | 2.5 months | 1 month | 3.5 months |
| Major Infections at Presentation and Before HCT* | Disseminated BCG* | Bacteremia (Klebsiella pneumoniae, E. Coli)* | Recurrent pneumonias | Disseminated BCG* | Disseminated CMV (lungs, blood, urine)* | Disseminated Adenovirus (lungs, urine)* |
| | Otitis Media (E. Coli) | Pneumonia* (polymicrobial gram negatives) | | Candida albicans (oropharyngeal, perineum) | | Bacteremia (Streptococcus pneumoniae, Stenotrophomonas maltophilia) |
| | Tinea capitis | | | | | Bacteremia (E. Coli, Morganella morganii, Staphylococcus aureus) |
| Died Before HCT | Yes | No | No | No | Yes | No |
| Received HCT | No, HCT not available at the time | No, HCT not available at the time | No | No | No, Multi-organ failure present. Not HCT candidate. | No, Two maternal HLA 10/10 identical bone marrow transplants. Yes (Day +122) |
| Died After HCT | N/A | N/A | Yes | Yes (Day +9) | N/A | N/A |
| Age at Death | Infant ( < 4 months) | 14 months | 7 years | 4 months | 5 months | 8 months |

**Table 1**

Clinical presentation and outcome of IKBKB immune deficiency.

| Patient | 7** | 8** | 9 | 10** | 11 |
|---------|-----|-----|---|-----|-----|
| Decade of Diagnosis | 2000s | 2000s | 2000s | 2010s | 2010s |
| IKBKB Mutation Confirmed | Yes | Yes | No | Yes | Yes |
| Direct Family Relationships to Known IKBKB Cases | None | None | Affected siblings with confirmed mutations | Affected siblings | Affected siblings |
| Age at Clinical Presentation | 2 months | 1 month | 1 month | 3 months | 2 months |
| Major Infections at Presentation and Before HCT* | Listeria monocytogenes bacteremia and meningitis * | Bacteremia (E. Coli) | Pneumonia (RSV)* | Bacteremia (Streptococcus hominis, Enterococcus faecalis, Enterococcus faecium, Lactobacillus lactis) | Pneumonia (Rhovirus)* |
| | Thrush | | | Meningitis (Enterococcus species)* | Klebsiella pneumoniae bacteremia, pneumonia and meningitis* |
| Died Before HCT | Yes | No | Yes | No | Yes |
| Received HCT | No, Multi-organ failure present. Not HCT candidate. | No, SCID was suspected but since IKBKB mutation had not yet been described diagnosis was never confirmed. | No | No, SCID was suspected but since IKBKB mutation had not yet been described diagnosis was never confirmed. | No, Multi-organ failure present. Not HCT candidate. |
| Died After HCT | N/A | Yes (2 years) | N/A | N/A | N/A |
| Age at Death | 2 months | 2 years | 15 months | No – Alive 7 years post-HCT | 2.5 months |

**Table 2**

Clinical presentation and outcome of IKBKB immune deficiency.

| Patient | 12** | 13 | 14 | 15 | 16 |
|---------|------|---|---|---|---|
| Decade of Diagnosis | 2010s | 1980s | 1990s | 2010s | 2010s |
| IKBKB Mutation Confirmed | Yes | No | Yes | Yes (NBS) | Yes (in-utero and post-natal) |
| Direct Family Relationships to Known IKBKB Cases | No | Affected sibling with confirmed mutation | Affected sibling | Affected sibling | Affected sibling |
| Age at Clinical Presentation | 3 months | 4 months | 2.5 months | 0 months - Newborn Screening | 0 months – Family History Screening |
| **Note:** | | | | | |

(continued on next page)
### Table 1 (continued)

| Patient | Died Before HCT | Received HCT | Age at Death | Organ Abnormalities |
|---------|-----------------|--------------|--------------|---------------------|
| 12**    | No              | Yes          | N/A          |            |
| 13*     | No              | No           | 11 months    | Disseminated BCG and died. Biopsies (n=2) revealed Mycobacterium bovis in multiple organs in- |
| 14      | No              | Yes          | 4.5 months   |                      |
| 15      | No              | Yes          | 3.5 months   |                      |
| 16      | No              | Yes          | 6 months post-HCT |                      |

**Single asterisks refer to infections that ultimately resulted in death due to multi-organ failure and / or severe neurological injury. Involvement of organs was confirmed at the time of autopsy.**

**Double asterisks refer to patients included in the original publication (Pannicke et al, reference 1).**

### Abbreviations:

- BCG: Bacillus Calmette-Guerin
- CMV: Cytomegalovirus
- HCT: Hematopoietic Cell Transplantation
- HLA: Human Leukocyte Antigen
- HSV: Herpes Simplex Virus
- HPV: Human Papilloma Virus
- NBS: Newborn Screening
- ND: Not Documented Infections
- N/A: Not Available
- RSV: Respiratory syncytial virus

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In the blood (n = 1), urine (n = 1), and cerebrospinal fluid (n = 1) in three. Severe, recurrent gram-negative and gram-positive bacterial infections were frequent and included bacteremia / sepsis episodes with Escherichia coli (n = 7), Pseudomonas aeruginosa (n = 3), Klebsiella pneumoniae (n = 2), Enterococcus species (n = 2), Staphylococcus aureus (n = 2), Stenotrophomonas maltophilia (n = 2), Serratia marcescens (n = 1), Streptococcus pneumoniae (n = 1), Staphylococcus epidermidis (n = 1), Listeria monocytogenes (n = 1), Moraxella osloensis (n = 1), Leuconostoc side lactis (n = 1) and Enterobacter cloacae (n = 1). Intracranial infections were present in six infants and included a combined Staphylococcus aureus and Candida albicans meningitis in one, and Listeria monocytogenes meningitis, Enterococcus meningitis, Klebsiella pneumoniae meningitis, Serratia marcescens intracranial abscesses, and Mycobacterium bovis meningitis with intracerebral abscesses in the other five. Disseminated viral infections occurred with cytomegalovirus (n = 3), adenovirus (n = 2), varicella (n = 1), and herpes simplex virus (n = 1). Severe pneumonia syndromes were documented in eleven as result of unknown pathogens (n = 3), cytomegalovirus (n = 2), respiratory syncytial virus (n = 2), polymicrobial gram negative bacteria (n = 1), adenovirus (n = 1), rhinovirus (n = 1), and parainfluenza (n = 1).

Consistent with standard practice in the two affected communities at the time, four infants received the BCG vaccination after birth, all of whom developed disseminated BCG and died. Biopsies (n = 2) and autopsies (n = 2) revealed Mycobacterium bovis in multiple organs including the lungs, liver, spleen, lymph nodes, thymus, bone marrow, adrenal glands, kidneys, eyes, skin, brain and gastrointestinal tract. Following these deaths, BCG vaccination in the two affected communities was discontinued. More recently, with the advent of targeted newborn screening for the homozygous IKKβ deficiency, one patient had documented organ abnormalities (dysplastic kidney at time of autopsy).

### 3.2. Patient outcomes

Of the sixteen IKKβ immune deficiency patients, eight died of overwhelming infection before HSCT could be performed, and eight proceeded to HSCT. Detailed autopsy reports were available for three who died prior to HSCT. Aside from evidence of infection, all exhibited severe thymic hypoplasia with scant lymphoid cell populations, minimal to no lymph node and tonsillar tissue, and absence of germinal follicles in the spleen.

### 3.3. Immunologic presentation of IKKβ immune deficiency

A range of immunologic tests were performed over time, with the extent and type of testing dependent upon the year of presentation and initial suspicion of an immunodeficiency diagnosis. Extensive immunologic investigations were more likely to occur in the contemporary era (after 2000).

Median total lymphocyte count at the time of diagnosis was $6.94 \times 10^9/L$ (range: 1.52–11.85 \times 10^9/L), with 11/14 patients exhibiting normal (n = 6) to elevated (n = 5) total lymphocyte counts for age. The three lymphopenic infants all had severe infections (disseminated BCG in two and Klebsiella sepsis in one) at the time their initial blood work was performed, along with various lymphocyte subset and hematologic cytopenias (anemia, thrombocytopenia), more consistent with bone marrow suppression from infection. Median CD3 T-cell count (12 patients) was 3298 cells/μL (range: 240–8480 cells/μL), with only one patient having a CD3 T-cell count < 1500 cells/μL.
and one other < 300 cells/μL. As above, in both patients with CD3 T-cell lymphopenia, severe infection was present at the time of assessment. Median CD4 T-cell count (11 patients) was 2490 cells/μL (range: 120–6520 cells/μL) and median CD8 T-cell count (11 patients) was 1660 cells/μL (range: 100–2654 cells/μL), with 6/11 patients having at least one of the CD4 and/or CD8 T-cell count above the normal range for age. Maternal engraftment studies were performed at diagnosis in eight patients, with none showing evidence of maternal engraftment. Median CD19 B-cell count (11 patients) was 750 cells/μL (range: 100–3470 cells/μL), being low in two, normal in eight, and elevated in one. Median CD56 NK-cell count (10 patients) was 240 cells/μL (range: undetectable – 1090 cells/μL), being low in three and normal in seven.

Despite generally normal to elevated T- and B-cell numbers in most IKBKB immune deficiency patients, all infants presenting with infections were severely hypogammaglobulinemic (n = 12), with a median IgG level of 0.75 g/L (range: undetectable – 2.06 g/L). The two infants identified as newborns both had normal IgG levels immediately after birth, likely reflecting maternal placental transfer of IgG.

T-cell proliferative responses were available for 13 patients (Fig. 1). Relative to controls, three had PHA responses < 10%, two had PHA responses between 10 and 30%, and eight had PHA responses above 30% (range: 53% - 146%). No relation was evident between PHA re-

Fig. 1. Phytos hemagglutinin responses at the time of IKBKB diagnosis by patient number. Percent of control is defined as the stimulation index of the patient relative to the average stimulation index of two healthy controls. Corresponding CD3 T-cell count (cells/μL) at the time of diagnosis is under each patient number. Patients 4 and 14 both had disseminated BCG with bone marrow involvement and severe cytopenias, including lymphopenia, which may have impacted PHA response.

3.4. Clinical outcomes following hematopoietic stem cell transplantation

Three patients receiving transplant remain alive 6-months, 6-years and 7-years post-HSCT. Two patients died soon after HSCT (days +9 and +122) from complications related to infections present at the time of transplant. Three others died of new infections acquired post-HSCT, including two (patients 3 and 8) who died of disseminated Mycobacterium avium-intracellulare infection in association with secondary graft failure at 2- and 5-years post-HSCT. The third (patient 15) died of Streptococcus pneumoniae bacteremia and meningitis despite full donor chimerism at 8-months post-HSCT. New clinically relevant infections, including viral, bacterial, fungal, and Mycobacterial infections, were common in patients who survived at least 6-months after transplant, and were still occurring years after transplant despite full and mixed lymphoid and myeloid donor chimerism and normal neutrophil and lymphocyte counts. Such infections included Streptococcus pneumoniae bacteremia (n = 5), meningitis (n = 1), osteomyelitis (n = 1), and septic arthritis (n = 1); Staphylococcus aureus bacteremia, cellulitis, and osteomyelitis (n = 1); Salmonella bacteremia and enterocolitis (n = 1); and Pseudomonas aeruginosa bacteremia (n = 1). Viral infections (e.g. influenza, parainfluenza, varicella, adenovirus) also occurred, with resolution in all patients. One patient developed Mycobacterial tuberculosis pneumonia at 1-year post-HSCT, was successfully treated with anti-tuberculosis medications, and remains a long-term survivor.

3.4.3. Immune reconstitution following hematopoietic stem cell transplantation

All patients achieved normal numbers of CD3+, CD4+, CD8+, CD56+, and CD19+ cells post-HSCT. Immunoglobulin replacement was variable between patients and generally continued between 3- and 12-months post-HSCT. PHA responses were normal (above 50%) for all patients. Despite these findings, specific antibody production to post-transplant vaccinations once immunoglobulin was discontinued was poor. For instance, patients 8 and 10 did not make anti-pneumococcal
Table 2
Hematopoietic stem cell transplant outcomes for IKBKB immune deficiency.

| Patient: | 3    | 4    | 6    | 8    | 10   | 12   | 15   | 16   |
|----------|------|------|------|------|------|------|------|------|
| Decade of Diagnosis | 1980s | 2000s | 2000s | 2000s | 2010s | 2010s | 2010s | 2010s |
| Identified by IKBKB NBS | No | No | No | No | No | No | Yes | Yes |
| Major Infections Present at Start of HCT | HSV | Disseminated BCG, Adenovirus, CMV | No | Disseminated Adenovirus, CMV | No | Parainfluenza Pneumonia | No | RSV Pneumonia and Serratia Marcescens Abscesses |
| Age at First HCT | 19 months | 4 months | 4.5 months | 5 months | 7 months | 8 months | 2 months | 1 month |
| Donor | Father | Half-Sibling | Mother | Unrelated | Brother | Unrelated | Unrelated | 5/6 |
| HLA Match | 6/6 antigen match (HLA-A, B, HLA-C) and non-reactive in mixed leukocyte culture | 6/6 | 10/10 | 6/6 | 10/10 | 5/6 | 5/6 | 5/6 |
| Graft Source | Marrow | Marrow | Marrow | Cord Blood | Marrow | Cord Blood | Cord Blood | Cord Blood |
| Conditioning | MAC - Bu/Cy/ATG | None | None | Tacrolimus / Methotrexate | RIC - Flu/Mel/Camptothecin | Tacrolimus / Methotrexate | Tacrolimus / Methotrexate | Tacrolimus / Methotrexate |
| GVHD Prophylaxis | None | None | None | Tacrolimus / Methotrexate | None | Tacrolimus / Methotrexate | None | None |
| Time to ANC > 0.5 x 10^9/L | N/A | Never Achieved | Primary Graft Failure | +14 | +31 | +54 | +44 | +35 |
| Time to Platelets > 50 x 10^9/L | N/A | Never Achieved | Primary Graft Failure | N/A | N/A | N/A | N/A | N/A |
| Second Transplant or Boost | No | No | Yes. Mother, Bone Marrow, Myeloablative (Bu/Flu/ATG) with neutrophil engraftment day + 12. | No | No | No | No | No |
| CD3 T-Cell Count Post-HCT (x10^9/L) | N/A | N/A | 6.55 | 3.51 | 5.78 | 6.55 | 5.11 | 3.04 |
| CD4 T-Cell Count Post-HCT (x10^9/L) | N/A | N/A | 3.84 | 1.51 | 4.36 | 5.27 | 0.9 | 0.11 |
| CD8 T-Cell Count Post-HCT (x10^9/L) | N/A | N/A | 2.71 | 1.73 | 1.27 | 0.9 | 0.11 |
| CD56 NK cell Count Post-HCT (x10^9/L) | N/A | N/A | 0.61 | 0.22 | 0.13 | 0.12 | 0.11 |
| CD19 B-Cell Count Post-HCT (x10^9/L) | N/A | N/A | 1.57 | 1.62 | 3.26 | 5.27 | 7.98 |
| IgG (g/L) off IVIg | N/A | N/A | 19.2 | 14.2 | 6.53 | 2.06 | N/A (on IVIg) | N/A |
| IgA (g/L) | N/A | N/A | 1.62 | 2.04 | 0.65 | < 0.07 | Not tested | Too early post-HCT |
| Response to Post HCT Vaccinations | N/A | N/A | No response to tetanus, diphtheria (3 vaccinations). | No response to tetanus, diphtheria (4 vaccinations). | No response to tetanus, diphtheria (4 vaccinations). | No response to tetanus, diphtheria (4 vaccinations). | Negative anti-HBs (6 hepatitis B vaccinations). | Negative anti-HBs (6 hepatitis B vaccinations). |
| CD57 | N/A | N/A | 0.9 | 0.11 | 0.11 | 0.11 | 0.11 | 0.11 |
| CD16 | N/A | N/A | 0.1 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 |
| CD11c | N/A | N/A | 0.1 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 |
| CD3 T-Cell Count Post-HCT (x10^9/L) | N/A | N/A | 6.55 | 3.51 | 5.78 | 6.55 | 5.11 | 3.04 |
| CD4 T-Cell Count Post-HCT (x10^9/L) | N/A | N/A | 3.84 | 1.51 | 4.36 | 5.27 | 0.9 | 0.11 |
| CD8 T-Cell Count Post-HCT (x10^9/L) | N/A | N/A | 2.71 | 1.73 | 1.27 | 0.9 | 0.11 | 0.11 |
| CD56 NK cell Count Post-HCT (x10^9/L) | N/A | N/A | 0.61 | 0.22 | 0.13 | 0.12 | 0.11 | 0.11 |
| CD19 B-Cell Count Post-HCT (x10^9/L) | N/A | N/A | 1.57 | 1.62 | 3.26 | 5.27 | 7.98 | 7.98 |
| IgG (g/L) off IVIg | N/A | N/A | 19.2 | 14.2 | 6.53 | 2.06 | N/A (on IVIg) | N/A |
| IgA (g/L) | N/A | N/A | 1.62 | 2.04 | 0.65 | < 0.07 | Not tested | Too early post-HCT |
| Response to Post HCT Vaccinations | N/A | N/A | No response to tetanus, diphtheria (3 vaccinations). | No response to tetanus, diphtheria (4 vaccinations). | No response to tetanus, diphtheria (4 vaccinations). | No response to tetanus, diphtheria (4 vaccinations). | Negative anti-HBs (6 hepatitis B vaccinations). | Negative anti-HBs (6 hepatitis B vaccinations). |
| CD57 | N/A | N/A | 0.9 | 0.11 | 0.11 | 0.11 | 0.11 | 0.11 |
| CD16 | N/A | N/A | 0.1 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 |
| CD11c | N/A | N/A | 0.1 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 |

(continued on next page)
| Patient: | 3 | 4 | 6 | 8 | 10 | 12 | 15 | 16 |
|----------|---|---|---|---|----|----|----|----|
| Decade of Diagnosis | 1980s | 2000s | 2000s | 2000s | 2010s | 2010s | 2010s | 2010s |
| Major Infections in First 6-Months Post-HCT | HSV (stomatitis, perianal, ocular) Thrush | Disseminated BCG, Adenovirus, CMV* | Disseminated Adenovirus, CMV* Enterobacter cloacae bacteremia | Streptococcus pneumoniae bacteremia | Varicella Zoster Recurrent Pneumonias | Disseminated Adenovirus Pseudomonas aeruginosa bacteremia | Candida Diaper Dermatitis | Streptococcus pneumoniae bacteremia Candida Diaper Dermatitis Methicillin-Resistant Staphylococcus aureus bacteremia |
| Major Infections After 6-Months Post-HCT | Salmonella bacteremia / gastroenteritis Disseminated Mycobacterium avium-intracellulare* | N/A | N/A | Streptococcus pneumoniae bacteremia / oesophagitis. Disseminated Mycobacterium avium-intracellulare* | Mycobacterium tuberculosis pulmonary disease. Staphylococcus aureus bacteremia / osteomyelitis / cellulitis HPV-70 Warts Norwagian Scabies | Streptococcus pneumoniae bacteremia / septic arthritis Bordetella pertussis pneumonia Scabies Varicella Zoster | Parainfluenza pneumonia Influenza A pneumonia Candida Diaper Dermatitis Streptococcus pneumoniae bacteremia / meningitis* |
| Graft Versus Host Disease | “Mild” Acute GVHD (skin) | None | None | Grade III acute GVHD (gastrointestinal) | None | None | None | None |
| Died After HCT | Yes | Yes | Yes | Yes | No | No | Yes | No |
| Time of Death Post-HCT | 5-years | Day +9 | Day +122 | 2-years | N/A | 7-years | 6-months | N/A |
| Time of Last Follow Up Post-HCT if Alive | N/A | N/A | N/A | N/A | N/A | N/A | N/A | N/A |

Abbreviations: ANC: Absolute Neutrophil Count; ATG: Antithymocyte globulin; BCG: bacillus Calmette-Guerin; Bu: Busulfan; CMV: Cytomegalovirus; Cy: Cyclophosphamide; Flu: Fludarabine; GVHD: Graft-versus-host disease; HCT: Hematopoietic Cell Transplant; HPV: Human papilloma virus. HSV: Herpes simplex virus; IVIg: Intravenous immunoglobulin; MAC: Myeloablative conditioning; Mel: Melphalan; N/A: Not available; NBS: Newborn screening; ND: Not determined; RIC: Reduced intensity conditioning; RSV: Respiratory syncytial virus.

* Asterisks refer to infections that ultimately resulted in death due to multi-organ failure and/or severe neurological injury. Involvement of organs was confirmed at the time of autopsy.
antibodies despite both conjugated and unconjugated pneumococcal vaccinations, and in the case of patient 8, a Staphylococcus pneumoniae infection. Patient 10 received six combined hepatitis A and B vaccinations and despite this, did not make antibody against hepatitis A virus or hepatitis B surface antigen. Similarly, neither patient made protective titres against tetanus or diphtheria despite being vaccinated after HSCT.

Detailed autopsy reports were available for 2 patients after HSCT. Aside from evidence of infection, both exhibited severe thymic hypoplasia with scant lymphoid cell populations, minimal to no lymph node and tonsillar tissue, and absence of germinal follicles in the spleen. Patient 3, who experienced secondary graft failure by 3-years post-HSCT, developed disseminated Mycobacterium avium-intracellulare and was managed for 1-year with anti-tuberculous medications and splenectomy before death. The spleen was entirely replaced by atypical Mycobacteria. At autopsy, performed 5-years after HSCT, the thymus was small and atrophic, weighing only 2 g, and lymph nodes were not readily apparent.

4. Discussion

This series presents the clinical features, immunologic presentation, and HSCT outcomes for the largest cohort of IKKßB immune deficiency patients published to date. A number of important conclusions can be made.

First, the clinical presentations in our case series indicate that IKKßB immune deficiency, as result of profound defects in innate and adaptive immunity, manifest as exceptional susceptibility to a variety of different but severe infections, including bacteria, viruses, fungi, and Mycobacteria. We believe that IKKßB immune deficiency, although not meeting conventional laboratory-based diagnostic criteria for SCID [19] (due to higher CD3 T-cell counts, moderately decreased to normal PHA responses, normal TRECs, high proportions of naïve CD45RA+ T-cells, and low CD45RO+ T-cells in IKKßB immune deficiency), is as severe an immune deficiency as SCID at a clinical level. We would suggest therefore, that inclusion of IKKßB deficiency under the descriptive category of “combined immune deficiencies generally less profound than SCID” in the most recent 2017 International Union Immunologic Societies classification [20], likely underestimates the severity of the immune deficiency and should be modified in future iterations.

Supporting a more profound immune deficiency are the other six reported patients with IKKßB mutations, who also presented with severe bacterial, fungal and viral infections as young infants. Mycobacterial infection was also prominent, with disseminated BCG noted in four of the six [13–15]. Similar to our cohort, previously reported IKKßB immune deficient patients exhibited normal to elevated absolute T-cell and B-cell numbers, low CD45RO+ memory T-cells, and absent or low class-switched memory B-cells. Immunoglobulins were universally low or undetectable, but proliferative responses to PHA were present and typically normal. Only one patient was noted to have features of ectodermal dysplasia at an older age (18 months) [13], a feature which we have not seen in any of our three post-HSCT survivors. Two of the other reported patients died from overwhelming infection prior to definitive therapy, and four received HSCT, with only one long-term survivor.

Secondly, this data further expands the understanding of primary immune deficiencies involving aberrant IKK/NF-kB signaling. At least 14 patients with germline gain of function mutations in NFKBIA have been reported, with a severe but more variable clinical and immune phenotype that appears dependent upon the genotype (with missense mutations resulting in a more severe clinical phenotype) [21]. Like IKKßB immune deficiency, patients with hypermorphic NFKBIA mutations that result in reduced degradation of IkBα, present with multiple and severe bacterial, fungal and viral infections starting at an early age, typically before 3-months. Infections reported include invasive bacteria such as Klebsiella, Pseudomonas, Haemophilus influenzae, and Staphylococcus aureus, Mycobacteria (including vaccine-strain BCG), as well as chronic mucocutaneous candidiasis, pneumocystis pneumonia, and severe viral infections (rotavirus, norovirus, parainfluenza, RSV, CMV). Similarly, patients with NFKBIA mutations present with normal to elevated T-cell counts, an excess of naïve CD4 and CD8 cells, and normal mitogen proliferative response to PHA in most. Dysgammaglobulinemia (usually with one or more quantitatively abnormal immunoglobulin isotypes), impaired response to vaccine antigens, and when tested, absent or reduced memory B-cells and class switched memory B-cells have also been reported. Unlike patients with IKKßB immune deficiency, however, almost all patients with IkßBα defects show some degree of ectodermal dysplasia.

By comparison, individuals with hemizygous hypomorphic IKKßB mutations with reduced NEMO expression manifest a more variable clinical and immunologic phenotype when compared to patients with IKKßB immune deficiency and hypermorphic NFKBIA mutations [22]. In addition to infectious predisposition, patients with NEMO deficiency may present with autoimmunity (25%) and inflammatory conditions (most notably inflammatory bowel disease in 21%), features not described in either our cohort of IKKßB immune deficient patients nor in those with hypermorphic IkßBα defects. As in patients with IKKßB immune deficiency, patients with NEMO deficiency commonly present with infections due to pyogenic bacteria (87%), Mycobacteria (44%), viruses (21%) and fungal and opportunistic pathogens (10%) [22] The median age at HSCT for a large cohort of NEMO deficient patients was 3.3 years old (range: 4.4 months – 18.8 years), an age that is older compared to our cohort, suggesting NEMO deficiency may be a less severe immune deficiency relative to IKKßB immune deficiency [23]. NEMO deficiency also has a more variable immunologic phenotype, with hypogammaglobulinemia in only 24/41 (59%) and defects in specific antibody production in 18/28 (64%). Similar to IKKßB immune deficiency, most patients with NEMO deficiency have normal to elevated CD4 and CD8 counts, and 91% have normal mitogen induced proliferations [22].

A third important finding of this study is that SCID newborn screening by TREC assays will not detect IKKßB immune deficiency. Given the severe clinical phenotype of IKKßB immune deficiency, a pilot project of targeted newborn screening by direct genetic mutation analysis of Guthrie cards for the IKKßB mutation was developed in select Northern Cree communities in Manitoba. This investigation identified a carrier frequency of 1 in 13.1 newborns, with a predicted mutation homozygosity of 1/686 births [24]. Over 5-years (2013–2018), 760 newborns from two communities were screened, with one infant identified at birth (patient 15) and a subsequent sibling (patient 16) identified in-utero. Both were placed in protective isolation after birth and received HSCT at an early age, before the onset of serious infections, important variables associated with excellent outcomes in SCID HSCT [25,26]. In other populations with founder mutations or high rates of consanguinity, a similar genetic-based newborn screening strategy such as ours might be applicable.

Unfortunately, patient 15 later died of invasive Streptococcus pneumoniae infection in their home community, 8-months following an otherwise successful transplant. This illustrates a complex interplay between what we suspect are ongoing defects in immunity post-HSCT, combined with poor social determinants of health. For familial and cultural reasons, our patients often have strong desires to return home soon after HSCT, and the tertiary care HSCT center is not easily accessible from their remote communities. With the death of patient 15, and with the recent transplant of patient 16, we have instituted a program of housing the family close to the tertiary care center for at least 5 years post-HSCT, as well as regular immunoglobulin replacement and indefinite daily anti-pneumococcal antibiotic prophylaxis.

Much remains unknown about the immune abnormalities in patients with IKKßB immune deficiency, both before and after HSCT. From our previous investigations, the broad immune defects relate to abnormal IKK/NF-kB signaling in T-cells, B-cells and innate immune
cells, resulting in generalized activation defects to a variety of stimuli [1]. With the exception of regulatory T-cells and gamma-delta T-cells, which are absent in IKBKB immune deficiency, other lymphocyte subsets appear to develop normally. Despite thymic hypoplasia in five of our patients who died either prior to or after HSCT, six other patients had normal TREC levels at the time of diagnosis, suggesting no intrinsic defect in T-cell development and maturation within the thymus. Unfortunately, none of the patients who had autopsies performed had detailed TREC analysis, making direct correlation with the under-development of the thymus and TREC levels in a single patient impossible. Detailed T-cell studies were not available for most of our patients pre-HSCT, but when previously studied in one patient (and in one additional patient in this series), the T-cell receptor beta-chain variable region repertoire was polyclonal [1]. We are unable to resolve this apparent contradiction in the autopsy and immunologic findings, and believe it warrants further study. Likewise, when studied in two patients, kappa-deleting recombination excision circles (KRECs) in dried blood spots from newborn Guthrie cards of two patients were found to be normal, suggesting normal B cell development.

In a mouse model involving a hypermorphic mutation of IκBk, NF-κB signaling in non-immune “architectural” cells were shown to be severely disrupted [27]. Although such cells are not derived from hematopoietic stem cells, they are nonetheless integral to lymphoid organogenesis, as in the development of lymph nodes, Peyer’s patches, splenic marginal zones, follicular dendritic cells, and the formation of germinal centers in the spleen, which were absent. Further, the immune defect in this model was not fully corrected by HSCT. It is reasonable to suspect, therefore, that similar architectural cell dysfunction with resultant disruption of the immunologic niche could also play a role in IKBKB immune deficiency. Failure to correct non-immune cells with HSCT could explain the ongoing infectious complications seen years after an otherwise apparently successful, lymphoid engrafted transplant in IKBKB immune deficiency. The absence of splenic germinal centers and lack of tonsils and lymphoid tissues in the autopsies of patients who died post-HSCT, and absent specific vaccine responses despite excellent donor T- and B- cell chimerism, supports that HSCT does not fully correct all aspects of the immune deficiency in IKBKB.

In years past, and in patients who were profoundly ill at presentation, full immune evaluations were not always available to further elucidate the nature of the immune deficiency in this population. At the moment, there are only three survivors in our cohort, for whom detailed T cell studies pre- and post-HSCT (including serial measurements of T cell markers such as recent thymic emigrants, markers of naivety and memory cells and markers of activation or exhaustion) have been limited to date. Moving forward, and with genetic-based newborn screening introduced for this condition in our province, we will now have this opportunity.

Despite these challenges, we believe that IKBKB immune deficiency is still a disorder that requires HSCT. To our knowledge, no individual has been a long-term survivor of IKBKB immune deficiency without HSCT. We are unaware of any reports of hypomorphic IKBKB mutations producing a milder clinical phenotype, allowing clinicians to take a wait and see approach. HSCT for other disorders involving the IκK/NF-κB pathway have been previously described, including for hemizygous hypomorphic IKBKG mutations [23] and hypermorphic NFκBIA mutations [21]. In the cohort of fourteen IκBα gain of function patients reported by Boisson et al., eleven received HSCT, with only five survivors. One patient died without receiving HSCT, and two patients survived without receiving HSCT.

Since most centers will not have genetically based newborn screening for IKBKB immune deficiency, clinicians would likely be faced with an infant presenting with severe life-threatening infections, given that TREC based newborn screening will not detect the disorder. Commercially available next-generation sequencing gene panels for immune deficiency disorders (including many SCID panels) increasingly include IKBKB, which should facilitate diagnosis, although not always in a timely manner before HSCT. In our series, two of the three patients surviving HSCT had serious life-threatening infections before transplant, including a case of multiple Serratia marcescens intracerebral abscesses, where the long-term neurocognitive outcome was initially unclear. We have tended to be aggressive with antimicrobials and other supportive care measures, with the goal of reaching HSCT as quickly as possible. This strategy has resulted in long-term survival of 2/6 infants receiving HSCT with active infections. That being said, we suspect that survival will be significantly improved with current newborn screening strategies, in the absence of pre-HSCT infection.

Unfortunately, and based on this limited dataset, we believe myeloblastic conditioning may be required for engraftment in IKBKB immune deficiency. The largest and most recent study of HSCT in NEMO patients, however, did not find this to be necessarily true [23]. In their cohort of fifteen patients receiving classical myeloablative conditioning and thirteen patients receiving reduced intensity conditioning, the global engraftment rate was 93%, with similar survival rates and secondary graft failure rates, independent of conditioning regimen intensity. Our center currently uses once daily intravenous dosing of busulfan for 4 days (aiming for an area under the curve approximating 4000 μmol/L*min), fludarabine 1.33 mg/kg/dose for 4 days, and rabbit anti-thymocyte globulin 2.5 mg/kg/dose for 3 days. We acknowledge other conditioning regimens might also be efficacious, although caution that late secondary graft-failure appears to be a concern in IKBKB immune deficiency.

Finally, and importantly, clinicians should be mindful that despite apparently reasonable basic laboratory immune reconstitution data for IKBKB immune deficiency patients post-HSCT (e.g. T-cell counts, chimerism studies), late infections, including bacterial and Mycobacterial infections in particular, appeared to be common. Of interest, one of our long-term survivors who was fully engrafted developed primary Mycobacterium tuberculosis pulmonary disease at 14-months post-HSCT. With appropriate therapy, the infection resolved, suggesting HSCT corrected enough of the immune deficiency. In contrast to our post-HSCT patients with IKBKB immune deficiency, Miot et al. found that bacterial infections only recurred at > 6-months post-HSCT for NEMO deficiency in 2/27 patients [23]. We hypothesize that failure of HSCT to improve all aspects of innate and adaptive immune function after transplant, as shown by the number of ongoing serious bacterial and Mycobacterial infections and lack of protective specific antibody responses to vaccinations, could be because HSCT does not correct cells of non-hematopoietic origin important in the immune response that are also dependent upon IKK/NF-κB signaling. A similar pattern of incomplete immune correction and ongoing bacterial and viral infections after HSCT has been reported in a patient with the related autosomal dominant anhidrotic ectodermal dysplasia with immune deficiency due to hypermorphic IκBα defect [28].

In summary, we present a relatively large cohort of a rare primary immunodeficiency disorder, IKBKB immune deficiency, characterized by the presence of naive T-cells, evidence of a functional activation defect, and amenable to HSCT, although with ongoing long-term infectious challenges.

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