Recurrent and Sustained Viral Infections in Primary Immunodeficiencies

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Viral infections are commonplace and often innocuous. Nevertheless, within the population of patients with primary immunodeficiencies (PIDDs), viral infections can be the feature that drives a diagnostic evaluation or can be the most significant morbidity for the patient. This review is focused on the viral complications of PIDDs. It will focus on respiratory viruses, the most common type of viral infection in the general population. Children and adults with an increased frequency or severity of respiratory viral infections are often referred for an immunologic evaluation. The classic teaching is to investigate humoral function in people with recurrent sinopulmonary infections, but this is often interpreted to mean recurrent bacterial infections. Recurrent or very severe viral infections may also be a harbinger of a primary immunodeficiency as well. This review will also cover persistent cutaneous viral infections, systemic infections, central nervous system infections, and gastrointestinal infections. In each case, the specific viral infections may drive a diagnostic evaluation that is specific for that type of virus. This review also discusses the management of these infections, which can become problematic in patients with PIDDs.

Keywords: virus, primary immunodeficiency, morbidity, herpes, papillomavirus, norovirus, enterovirus

INTRODUCTION

Frequent infections are a common reason for physician visits. Distinguishing a pattern or a type of infection that suggests an immunodeficiency as opposed to part of the normal susceptibility to infection can be a challenge. Common causes of recurrent infections are allergies, anatomical contributions, secondary immune deficiency, and an unusual burden of exposures. Primary immunodeficiencies (PIDDs) are much less common and therefore difficult to appreciate during the wealth of infections that are typically seen in a physician’s practice. During the first 5 years of life, children can experience six to eight respiratory tract infections per year. These tend to peak in the winter months and daycare attendance, exposure to smokers, and atopy can increase this frequency significantly (1–4). Respiratory tract infections in adults are somewhat less common; however, three to five respiratory tract infections per year in adults are typical (5). Recurrent sinus infections, pneumonia, and bronchitis are common signs of an immunodeficiency, recognizing that frequent bacterial infections of the respiratory track are often a harbinger of antibody disorders, the most common type of primary immunodeficiency. This review will address recurrent and sustained viral infections for which there are fewer studies to assist the physician in the identification of patients with potential immunodeficiency. This review will address unusual viral respiratory tract infections, systemic viral infections, infections of the brain and meninges, and cutaneous viral infections. Unusual viral infections can be a sign or complication of PIDD. There are several excellent reviews that address the
RESPIRATORY VIRAL INFECTIONS IN PIDD

Respiratory viruses are extremely common in most patients with PIDDs (9, 10). In most cases, they represent nuisance infections that can be a predisposing condition leading to bacterial superinfection. In patients with antibody defects, respiratory infections fall into this category. Although defense against recurrence of respiratory tract viruses is mediated largely by antibody, eradication of an infecting virus is mediated largely by the T cell compartment. Respiratory viral infections are therefore more significant in patients with T cell immune deficiencies. Today, many newborns with severe combined immune deficiency (SCID) are detected by newborn screening; however, this is not true in all parts of the world, nor is it true in all states in the USA. A study by the PIDTC found that 21% of their cohort had a respiratory infection prior to transplant with the most common being parainfluenza followed by RSV, rhinovirus, and influenza (11). Although other types of infections were more common as presenting features in this cohort, respiratory infections were among the least likely to resolve prior to transplant. All viral infections are typically prolonged in patients with T cell defects. However, in SCID, there are no T cells and a simple respiratory virus will progress relentlessly unless a hematopoietic stem cell transplant (HSCT) allows the infant to develop a competent immune system (Figure 1) (11). The specific pathogens to which children with T cell defects are susceptible include all of those common in the general population. RSV, influenza, and rhinovirus are typically the most prevalent during the respiratory season (12). Coronavirus and metapneumovirus have been increasingly recognized as causing respiratory infections. Exposures will dictate the pattern of viral infections in patients with T cell defects. The severity of disease is a result of both the degree of T cell compromise and the nature of the infecting virus. Any patient with prolonged viral infections is at risk for bacterial superinfection. It is not, therefore, uncommon to see a mixed picture of viral and bacterial infections. Additionally, severe T cell defects are associated with compromised antibody production, also contributing to a mixed infection picture.

T cell defects are associated with a generally increased predisposition to viral infections. IRF7 deficiency, in contrast, is associated with a selective susceptibility to influenza (13). It is otherwise uncommon to see an isolated susceptibility to a respiratory virus. The single reported patient had a severe primary infection with influenza associated with poor production of type I and type III interferons.

Diagnostic Approach

There are many specific genetic types of immunodeficiencies associated with T cell deficiency, and the diagnostic considerations are different depending on the age of the patient. Infants with a prolonged or severe respiratory viral infection should be evaluated for SCID, and patients of any age with dysmorphic features or other associated features should be evaluated for chromosome 22q11.2 deletion syndrome (14). Most states now have a newborn screening program to detect SCID. The results can be accessed to identify significant T cell lymphopenia in early infancy. Newborn screening has significantly improved survival of infants with SCID. There are, however, significant T cell disorders not identified by this test, and therefore, T cell disorders still represent a concern in the setting of a prolonged viral infection. T cell enumeration is often the quickest way to screen for T cell defects. The vast majority of T cell deficiency conditions will have...
Management of respiratory tract infections in non-SCID T cell defects is largely supportive with optimization of bronchodilators, antiviral therapy if available and attention to nutrition (12). Management of respiratory tract infections in SCID is highly problematic. There is a race to replace the immune system before the virus can progress to the point of no return. This race is highly dependent on the type of transplant donor, type of conditioning, and type of transplant, but respiratory infections clearly impact the transplant outcome (11). Any adjunctive measure to improve respiratory status should be sought.

SYSTEMIC VIRAL INFECTIONS IN PIDD

Children with severe T cell defects are also susceptible to systemic viral infections. Patients with SCID are extremely susceptible to progressive infection with cytomegalovirus (CMV) as well as other systemic viral infections. Infants with suspected SCID should be protected from exposures such as breast milk, transfusions, potentially infected siblings, live viral vaccines, and caregivers.

There is another circumstance in which susceptibility to gamma-herpes viruses such as CMV and Epstein–Barr virus (EBV) occurs. Table 1 lists conditions in which gamma-herpes virus susceptibility dominates the clinical picture. Patients with cytolytic T cell defects (with or without concomitant NK cell defects) exhibit unique susceptibility to these gamma-herpes viruses (15). In some circumstances, the susceptibility is almost entirely limited to susceptibility to either severe mononucleosis or hemophagocytic lymphohistiocytosis (HLH). HLH is characterized by excessive immune activation and can be diagnosed either by a molecular diagnosis consistent with HLH or clinically when patients meet five out of eight criteria: fever, splenomegaly, cytopenias affecting two or more blood lineages, hypertriglyceridemia and/or hypofibrinogenemia, hemophagocytosis, low/absent natural killer cell activity, hyperferritinemia, and high soluble interleukin-2 receptor levels (16). Any patient presenting with exceptionally severe mononucleosis or HLH should be screened for the HLH defects.

A second phenotype with susceptibility to EBV has a smoldering or even asymptomatic presentation. These patients generally have an increased risk of lymphoma due to chronic EBV, but the manifestations of EBV may be subtle or absent. The X-linked disorder due to deficiency of MAGT1 is associated with a mild susceptibility to other infections but chronic EBV. Similar conditions include deficiencies of CD27, CTPS1, RASGRP1, CD70, and MCM4. The importance in recognizing this group is due to their unpredictable capacity to control EBV and the risk of lymphoreticular malignancies.

Nearly all of the leaky SCID types and the combined immunodeficiencies are associated with an increased risk of CMV and EBV (17, 18). This set of disorders can have a broad phenotype including Omenn’s phenotype, autoimmunity, granulomas, and infections (18–23). In these patients, EBV and CMV can drive progression to malignancy, and they require careful monitoring. Leaky SCID has been defined as T cell lymphopenia (CD3 300–1,500 cells/mm³); functional impairment as defined by proliferative responses, absence of maternal engraftment, and most often having identified hypomorphic mutations in genes associated with SCID.

Diagnosis Approach

When the consideration is prolonged infection with gamma-herpes viruses without HLH, T cell counts and function can be helpful as supporting information but often genetic testing is the quickest approach. For HLH disorders, enumerating the HLH criteria is a useful exercise. CD163 staining of the bone marrow can be a sensitive way to identify active hemophagocytosis, but this is not required for the diagnosis (15, 16, 24). Additional maneuvers are measurement of IL-2R in the serum and CD107a on the surface as a marker for degranulation. HLH can occur without an underlying PIDD, and thus genetic analysis is often central to the management. Nearly always an underlying PIDD will require HSCT as definitive therapy, whereas HLH due to uncommon infections such as Leishmania, certain influenza viruses, and arboviruses will not require HSCT.

Management

Management of systemic viral infections relies on the availability of antiviral compounds. For CMV, therapy is often begun with ganciclovir or valganciclovir (25, 26). Foscarnet may be added if

**TABLE 1 | EBV susceptibility.**

| Phenotype          | Gene defect | Viral susceptibility | Other features                          |
|--------------------|-------------|---------------------|-----------------------------------------|
| EBV viremia        | ITK         | EBV                 | Lymphoma                               |
| EBV viremia        | MAGT1       | EBV                 | Lymphoma                               |
| EBV viremia        | CD27        | EBV                 | Low IgG                                |
| EBV viremia        | CORO1A      | Many viruses        | EBV                                    |
| EBV HLH            | SH2D1A      | EBV                 | Lymphoma, dysgammaglobulinemia, and vasculitis |
| EBV HLH lymphoma   | XIAP        | EBV                 | Hypogammaglobulinemia                   |
| EBV familial HLH   | PRF1, UNC13D | EBV, CMV           | Malignancy, short stature, adrenal insufficient |
| EBV Pigmentary     | LYST, RAB27A | EBV, CMV           | Pigmentary dilution                     |
| dilution with      | AP3B1, BLOC156 | others             |                                         |
| HLH                |             |                     |                                         |
| EBV susceptibility  | Leaky SCID, | Many viral         | Gene dependent                         |
| with broad         | most combined | susceptibilities |                                         |
| infectious         | immunodeficiencies, |                   |                                         |
| susceptibility     | WASP, WIPF1, |                     |                                         |
|                    | PLCG2, PRKCD, |                     |                                         |
|                    | ORAI1, STIM1, |                     |                                         |
|                    | IKBKG, CASP8, |                     |                                         |
|                    | STAT1 GOF,   |                     |                                         |
|                    | DOCK8, GATA2 |                     |                                         |

GOF: gain of function; LOF: loss of function; EBV, Epstein–Barr virus; HLH, hemophagocytic lymphohistiocytosis; CMV, cytomegalovirus; SCID, severe combined immune deficiency; IgG, immunoglobulin G.
the virus is resistant or progressive in spite of adequate ganciclovir (27). Bone marrow toxicity from ganciclovir may also require a change to foscarnet. EBV in some cases is treated with rituximab to eliminate one important reservoir of virus (28). When HLH is present, a systemic approach to stabilize the patient and treat the underlying inflammation is essential (29). Risks and benefits of antiviral therapy must be carefully weighed as all approaches can have significant adverse events. Management decisions are often impacted by subsequent transplant strategies.

**CHRONIC VIRAL SKIN INFECTIONS IN PRIMARY IMMUNODEFICIENCY**

Cutaneous manifestations are common in PIDD. As many as two-thirds of the patients have cutaneous manifestations at some point. Atopy, infection, and inflammatory lesions have all been described, and there may be interplay between the features (30). Awareness of common skin infections is important both to aid in the early diagnosis and also in the treatment of potentially life-threatening infections that can begin in the skin. Bacterial infections are one of the most common findings in PIDD. For example, folliculitis, abscesses, and impetigo are typical in neutrophil defects. Similarly, a significant subset of PIDD diagnoses is associated with fungal infections. These can be seen both in T cell defects as well as defects of the myeloid compartment. Chronic mucocutaneous candidiasis is most often due to defects that affect the Th17 cell production or function. These diseases generally do not overlap those with a susceptibility to cutaneous viral infections. One exception is STAT1 gain-of-function (GOF) mutations that render patients susceptible to a broad range of cutaneous infections. Viral infections of the skin are not nearly as common but are much more suggestive of PIDD. Severe herpes infections and papillomavirus are particularly characteristic of PIDD and can become the most notable feature in a patient. Chronic herpes virus and papillomavirus, in turn, predispose to cutaneous carcinoma and surveillance becomes important for this evolution. In this section, we will provide a brief synopsis about the individual disorders associated with susceptibility to papillomavirus.

**Papillomaviridae**

There are more than 200 strains of human papillomavirus (HPV). The diverse strains have variable malignant potential and tissue tropism. HPV causes warts in the general population with an incidence of cutaneous warts (Figure 2) that range from 1 to 12% (31–33). School age children have been estimated to have a cutaneous warts prevalence of over 40% (34). While sexually active women 20–24 years of age have a prevalence of genital papillomavirus of nearly 50% (35), clinically significant genital warts occur in only 5% of women (36). Genital HPV infection has been associated with malignant transformation, leading to the development of the first vaccine intended to prevent cancer. Worldwide, 5% of cancer is caused by HPV (37). Nearly all T cell disorders can be associated with increased susceptibility to warts; however, there is a small group of PIDDs that have warts as a cardinal feature (38). In these patients, the warts are recurrent, severe, and resistant to therapy. In most cases, the specific papillomaviruses are identical to those in the general population, and the appearance of the warts is the same as in the general population. In epidermodysplasia verruciformis (EV), however, there is a broader susceptibility to HPV types. In addition, the cutaneous manifestations are often atypical. Table 2 compares the features of the different conditions described below.

**DOCK8 Deficiency**

Patients with DOCK8 deficiency have a complex combined immunodeficiency secondary to disrupted cytoskeletal rearrangement (39). This includes an inability to properly assemble the immune synapse that fosters the signaling cascades required for lymphocyte memory differentiation (40). Lymphocyte migration through tissues is also compromised, contributing to the susceptibility to cutaneous infections (39). T cell counts are typically low for age, and there is impaired memory differentiation that may be progressive with age (41). Clinical features resemble that seen in Wiskott–Aldrich syndrome (WAS) including low IgM.
and elevated IgE and IgA. Patients have significant atopy, infections, impaired specific antibody responses and poor memory B cell responses, increased rates of malignancy, elevated IgE, and eosinophilia and are highly susceptible to cutaneous viral infections. Early in life, the atopic manifestations may dominate while the infectious susceptibility evolves. Skin infections are most often caused by HPV (including increased risk of malignant transformation of skin lesions), herpes simplex virus (HSV), molluscum contagiosum virus, and varicella zoster virus (42). HSCT has been shown to be curative (43).

**Epidermodysplasia Verruciformis**

The term EV refers to a group of disorders in which patients are susceptible to beta-HPV with severe diffuse warts, and there is a striking increase in the rate of skin carcinomas (44). *EVER1/TMC6 and EVER2/TMC8* inactivating mutations cause autosomal recessive (AR) EV. *EVER* proteins are intracellular zinc transporters, and mutations lead to altered cell activation and a more permissive environment for HPV. Interestingly, the only infectious susceptibility is to HPV. Most patients present in childhood, but the appearance of the warts can lead to misdiagnosis as seborrhea or tinea versicolor. The warts are generally worse on sun-exposed skin for reasons that are not clear. The specific HPV types are not necessarily those seen in the general population as wart-associated. The therapy in EV is usually local control since the susceptibility does not relate to hematopoietic cell dysfunction; there is no role for HSCT.

**LCK Deficiency**

*LCK* deficiency causes atypical EV with CD4 T cell deficiency as well as recurrent pneumonia and severe warts complicated by non-melanoma skin cancer (45). To date, only a single patient with *LCK* deficiency has been described, leading to uncertainty in the full spectrum of infectious susceptibility. Therapy is not clear for the same reason, but HSCT would be expected to be curative.

**CARMIL2/RLTPR**

Three Norwegian families have been identified with increased viral cutaneous infections (warts and molluscum contagiosum) as well as dermatitis and pneumonia. The four affected family members were found to share a single variant in this gene, with some evidence for a role in T cell activation (46).

**RHOH Deficiency**

Mutations in the *RHOH* gene (an atypical Rho GTPase) cause susceptibility to EV-type HPV strains, due to alterations in T cell activation and homing (47). Naïve T cell counts are low, and there is poor skin homing of T cells, with an increase in effector memory T cells in the setting of altered T cell receptor signaling. *RHOH* deficiency infectious susceptibility was largely limited to HPV in the two siblings identified. In mice, the defect was correctable by transfer of wild-type bone marrow, suggesting that this is a potential treatment.

**IKBKGD Deficiency (NEMO)**

Hypomorphic mutations in the central kinase of the canonical NFκB pathway lead to increased susceptibility to warts (48). This PIDD is associated with fine sparse hair, dental defects, and a broad susceptibility to infections including opportunistic infections. The spectrum of phenotypes is among the broadest among PIDDs. Lymphedema, osteopetrosis, susceptibility to pneumocystis and other fungi, mycobacterial susceptibility, and viral susceptibility call all be associated with *IKBKGD* deficiency. Inflammatory bowel disease is also common. Cutaneous viral infections are common in this population but are not usually the dominant feature. Among the PIDD in this section, this is a disorder where the benefit of HSCT is unclear because HSCT outcomes have been poor.

**GATA2 Deficiency**

GATA2 mutations yield a complex set of phenotypes. Patients tend to have monocytopenia and low NK and B cell counts.

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**TABLE 2 | Cutaneous viral infections in primary immunodeficiencies.**

| Viral family | Virus | Increased susceptibility in which PID | Other features |
|-------------|-------|--------------------------------------|---------------|
| Papillomaviridae | HPV | Ataxia telangiectasia; DOCK8; EV (EVER1, EVER2, RHOH; LCK); GATA2; Idiopathic T cell lymphopenia; Netherton syndrome; STK4/MST1; WHIM (CXC4R); WILD, CARMIL2/RLTPR, Clouston’s syndrome | EV warts are often flat, appearing as actinic keratosis or seborrhea-like lesions and can have increased susceptibility to unusual HPV strains. No other infectious susceptibility | DOCK8, GATA2; also include susceptibility to HSV. Progressive lymphopenia seen |

**Herpesviridae**

| Virus | Increased susceptibility in which PID | Other features |
|-------|--------------------------------------|---------------|
| HHV8/KSHV | DOCK8; NEMO; STAT1 GOF; STK4; CXC4R; Wiskott–Aldrich syndrome (WAS) | Susceptibility to mycobacteria |
| HSV | DOCK8; GATA2; NEMO; STAT1 GOF; STK4; CXC4R; Wiskott–Aldrich syndrome (WAS) | DOCK8, NEMO, STAT1 GOF: broad infectious susceptibility CXC4R: pancytopenia, abnormal neutrophils WAS: thrombocytopenia, eczema |
| VZV | DOCK8; GATA2; STAT3 GOF; IFNGR1; RHOH; STAT1 GOF; STK4 | DOCK8, NEMO, STAT1 GOF: broad infectious susceptibility CXC4R: pancytopenia, abnormal neutrophils WAS: thrombocytopenia, eczema |

**Poxviridae**

| Virus | Increased susceptibility in which PID | Other features |
|-------|--------------------------------------|---------------|
| MCV | DOCK8; GATA2; IKBKGD; STAT1 GOF; STK4; CXC4R; CARMIL2/RLTPR | DOCK8, IKBKGD, STAT1 GOF: broad infectious susceptibility CXC4R: pancytopenia, abnormal neutrophils WAS: thrombocytopenia, eczema |
| Orf virus | STAT1 GOF | Broad infectious susceptibility |

GOF, gain of function; LOF, loss of function; HPV, human papilloma virus; HSV, herpes simplex virus; EV, epidermodysplasia verruciformis; WHIM, warts, hypogammaglobulinemia, infections, and myelokathexis.
and have susceptibility to HPV, among other infections (49). The laboratory defects are progressive with age and correlate with increasing infection burden. Clinical features include lymphedema, risk of malignancy, and pulmonary alveolar proteinosis. Infections include fungal infections and cutaneous viral infections. The cutaneous viral infections can be the most prominent feature but there is a wide-ranging phenotypic heterogeneity (50, 51). The cutaneous viral infections and genital HPV are associated with a high rate of malignant transformation.

**STAT1 GOF**

Patients with STAT1 GOF mutations can present with dramatic autoimmunity, usually including enteropathy and endocrinopathies, or they can have a picture with a pronounced infectious susceptibility. Susceptibility to candida is common, but cutaneous viral infections can also be problematic (52, 53). The viral infections can become more problematic as immune suppression is used to control the enteropathy.

**Netherton Syndrome**

Patients with Netherton syndrome (secondary to SPINK5 mutations) demonstrate congenital ichthyosis and have higher susceptibility to EV-associated HPV strains (53, 54). Although Netherton syndrome can have a mild humoral immune deficiency, the susceptibility to warts seems to be a result of disrupted local response to infection (55). The warts are generally controllable with local measures.

**MST1 Deficiency**

Serine–threonine kinase 4 (STK4), encoded by a gene called MST1, affects the FOXO1 transcription factor and thus impacts T cell lifespan and has been shown to increase susceptibility to HPV infections (56), as well as causing B cell lymphopenia. The phenotype also includes susceptibility to EBV and cutaneous viral infections.

**Warts, Hypogammaglobulinemia, Infections, and Myelokathexis (WHIM) Syndrome**

Warts, hypogammaglobulinemia, infections, and myelokathexis syndrome is caused by an autosomal dominant (AD) GOF mutation in CXCR4, which leads to increased susceptibility to HPV (57, 58). Varying degrees of pancytopenia can be seen, and hypogammaglobulinemia occurs in many (but not all) patients. Warts can be the most prominent susceptibility, but neutropenia and hypogammaglobulinemia can drive diverse infectious susceptibilities. COPD and cutaneous carcinoma have been observed (59). WHIM can be treated with plerixafor or topical control measures for the warts.

**WILD Syndrome**

WILD (warts, depressed cell-mediated immunity, primary lymphedema, and anogenital dysplasia) is also correlated with severe warts without a known genetic etiology (60). Of note, this diagnosis does not lead to EV-defining HPV strain infections. A recent study demonstrates a case of a patient whose warts improved after quadrivalent HPV vaccination (61).

**Clouston Syndrome**

An ectodermal dysplasia syndrome with alterations in hair and nails (not generally in teeth) which can present with eccrine syringofibroadenomatosus that is reminiscent of EV and is associated with HPV infection (62).

**Other Settings**

Idiopathic CD4 T cell lymphopenia is associated with increased risk of cutaneous warts (63), and SCID patients post-HSCT can have an increased risk of warts, especially with specific underlying mutations (64, 65).

**Management**

Management of cutaneous warts typically progresses from low level removal approaches involving topical therapy (freezing, electrosurgery, curettage, laser, chemical softening, and cantharidin) to immune stimulants (imiquimod and antigens), bleomycin, and antiviral therapy including cidofovir (66). Genital warts are treated with conceptually the same approach. Management of genital warts also includes special considerations for pregnancy, partners, and screening for malignancy (67–69). Interferon a2 and GM-CSF have been used successfully for papillomavirus (70, 71). In severe cases, therapy can be very unsatisfying. HSCT can result in prompt eradication, though JAK3 mutations and common γ chain SCID can have significant warts post-HSCT, which may be in part due to poor NK cell function (64). Topical cidofovir has recently been shown in a case report to be effective in some of these patients (72). In one case, regression was observed after papillomavirus vaccination (61).

**Herpesviridae**

The Herpes family of DNA viruses includes nine viruses pathogenic for humans; CMV, EBV, HHV6a/b, HHV7, HSV1, HSV2, and KSHV, not all of which have skin tropism. Almost any type of T cell dysfunction is associated with an increased frequency of cutaneous HSV. The PIDD with a high likelihood of chronic or disfiguring cutaneous herpes are DOCK8, GATA2, NEMO/IKBKG, STK4, WAS, STAT1 GOF, and WHIM (CXCR4 mutations) as described above in association with warts. These same syndromes can manifest with severe primary varicella and an increased frequency and severity of zoster (73). Varicella and zoster can also be seen in settings with altered proportions of T cell subsets, as in STAT3 LOF deficient patients (AD-hyper IgE syndrome) (74) or cytokine signaling defects as seen in IFNγR1 mutations (75). Thus, susceptibility to herpes viruses occurs in a broad range of T cells defects and is often part of a complex of susceptibility to many cutaneous viral infections. Herpes viruses can be particularly persistent and cause significant morbidity and mortality. Carcinoma is a feared consequence of recurrent or persistent infection and can be difficult to distinguish from persistently infected skin.

**Management**

Management includes acyclovir, valacyclovir, or fampiclovir as initial options. IL-2 therapy has been used successfully in WAS (76). In WAS, it specifically improves NK cell function (77). Type
I and type II interferons have been used successfully in model systems (78, 79). Small studies of non-PIDD populations have supported its use in patients (80–83). Management should focus on prevention of recurrences and healing of cutaneous lesions. Malignant transformation relates in complex ways to persistence of infection.

**Poxviridae**

Poxviridae are double stranded DNA viruses. Molluscipoxvirus (i.e., molluscum contagiosum virus or MCV), orthopoxvirus (i.e., smallpox), parapoxvirus, and yatapoxvirus are the four genera that can infect humans. Susceptibility to poxviridae tends to be associated with susceptibility to other cutaneous viral infections. Therefore, many of the disorders described above have an increased susceptibility to poxviridae. Molluscum contagiosum in the most common pox virus infection in humans and in the general population is a self-limited infection with minimal residua (84). Severe molluscum contagiosum can be seen in DOCK8 deficiency, STK4 deficiency (56), GATA2, NEMO, STAT1 GOF, and WHIM syndrome. Molluscum can be a significant issue for patients with WAS, as described in the Herpesviridae section. In addition, Orf virus, found in pasture animals, was found in a single patient with STAT1 GOF (85). Similar to herpes virus susceptibility described above, molluscum can be seen with nearly any T cell defect and typically arises in that setting as part of a susceptibility to many cutaneous viruses.

**Management**

Initial management of molluscum in a PIDD patient is control of spread through curettage, topical therapy such as salicylate, cantharadin, or immune stimulation with imiquimod (86, 87). Other topical approaches have also been used successfully. Retreatment 2–4 weeks later is often required. If addressed early, spread may be controlled and the outbreak contained. For diffuse disease or if addressed early, spread may be controlled and the outbreak contained. For diffuse disease or disease that spreads despite all attempts at control, type I interferon (interferon a2) has been suggested (88, 89). Intralosomal immunotherapy with live antigen has been promoted but is contraindicated in PIDD patients with T cell defects (90, 91).

**Persistent Vaccine-Stain Rubella Infection**

Three studies have identified persistent vaccine-strain rubella in patients with moderate T cell defects (91–93). Most of the patients have had ataxia telangiectasia but a wide range of PIDD diagnoses have been seen. Generally, the patients have had sufficient T cell function to be leading relatively normal lives and the immune deficiency might not even be recognized at the time of the MMR vaccine administration. The manifestations have been largely cutaneous granulomas although chronic inflammation at other sites has been observed (91). Persistence of virus due to compromised T cell control and acquisition of mutations that may further impact clearance is the proposed mechanism.

**Diagnostic Considerations for Patients with Cutaneous Viral Infections**

The above conditions are derived from defects in T cell, NK cell, and local tissue immunity. It is therefore nearly impossible to systematically screen for gene defects related to cutaneous viral infection susceptibility. A reasonable start is to define T cells both quantitatively and functionally. If that is unrevealing and the phenotype suggests a PIDD, then whole exome sequencing may be appropriate.

**GASTROINTESTINAL (GI) VIRAL INFECTIONS**

Chronic diarrhea (>6 weeks) is a frequent finding in PIDD patients. Given that the etiologies of chronic diarrhea in immunodeficient patients can be diverse, it is important to first distinguish if the diarrhea is infectious, malabsorptive, or inflammatory in nature as there are multiple types of PIDD that can present with autoimmune enteropathy or inflammatory bowel disease (94–96). PIDD patients are susceptible to multiple types of GI pathogens, and this section will focus on GI viruses. The concerted action of both the innate and adaptive immune system is necessary for viral clearance (97, 98). Therefore, there are a number of combined immunodeficiency phenotypes that result in susceptibility to GI viral pathogens, which manifest as prolonged illness as well as prolonged asymptomatic viral shedding.

**Norovirus**

Persistent infection with norovirus resulting in prolonged viral shedding and symptomatic disease has been noted in patients with SCID and various secondary immunodeficiency states (99). In a series of pediatric PIDD patients, it was the most frequently isolated virus at 20.6%, with patients with SCID, major histocompatibility complex II deficiency, CD40L deficiency, and agammaglobulinemia represented in this series (100). Norovirus shedding can be prolonged in the stool of patients who were immunosuppressed following infection and norovirus can be part of multiple infections in the GI tract (101). In CVID, norovirus infection has been linked to development of severe enteropathy with prolonged viral carriage over the course of years (102). In several patients in this series, clearance of norovirus resulted in normalization of the GI enteropathy.

A concern about norovirus is the great difficulty in public health containment. A patient with PIDD who is shedding for a prolonged period of time is not only themselves at risk but also places those around at risk. Norovirus is a common pathogen in the general population, and exposures are therefore common. Norovirus is spread from the moment of illness to several days after clinical recovery. Both vomit and feces can spread virus. The virus lives on surfaces for up to 20 days, and alcohol-based cleaners are not completely effective. Patients with chronic norovirus should use bleach to clean surfaces. Vigorous hand washing with soap and water is also effective.

**Hepatitis C**

In the early 1990s, the United States FDA recommended that hepatitis C positive donors be excluded from the plasma donor pool resulting in loss of neutralizing hepatitis C viral antibodies from IVIG (103). Hepatitis C virus (HCV) infection was subsequently reported from several countries, which was due to the
presence of contaminating HCV virus from the small numbers of seronegative HCV positive donors (103–106). The severity of hepatitis seen in immunodeficient patients was variable, but subsets of patients with primary hypogammaglobulinemia were observed to have a more severe course of hepatitis, which in some cases was rapidly fatal (103, 105, 106). Younger age and early treatment with IFN were associated with better overall outcomes (103, 106). Following adoption of PCR screening for HCV and viral inactivation processes with solvent–detergent or pasteurization there have been no subsequent reports of IVIG-associated HCV since 1996 (107). This cautionary tale supports surveillance of PIDD patients who have risk factors for HCV: blood product exposure, including IV drug use, infants born to HCV positive mothers, high-risk sexual behavior, shared personal items with potential blood exposure among HCV positive individuals. Today, therapy for HCV should improve outcomes compared to the cohort in the early 1990s.

Other GI Viruses

Adenovirus, enterovirus, and rotavirus have been isolated from single PIDD patients with chronic diarrhea, and the true incidence is not known (100). Chronic rotavirus infection has been described in patients with SCID and agammaglobulinemia (108). In patients with immunodeficiency, rotavirus can be poorly contained within the GI tract. On investigation at autopsy, active rotavirus replication has been identified in the liver and kidney of patients with SCID, complete DiGeorge syndrome, and acquired-immunodeficiency syndrome, illustrating poor control of viral replication in the setting of profound immunodeficiency (109). An important consideration is that SCID has been associated with susceptibility to the live rotavirus vaccine (110). Indeed, vaccine-strain illness is cleared only after immune reconstitution (110, 111).

CVID and agammaglobulinemia can rarely have prolonged asymptomatic shedding of vaccine-strain polio following immunization with live-attenuated oral polio vaccine, which can pose risk to other immunocompromised members of the community (112–114). Additionally, central nervous system (CNS) infection can occur in agammaglobulinemia (see below). These are the main reasons that live polio vaccination is no longer used in the USA.

Management

Care for chronic GI viral infections in PIDD is primarily supportive: optimizing hydration and nutrition. Orally administered immunoglobulin G (IgG) has been demonstrated to be effective as a therapy for chronic infectious diarrhea in antibody deficient patients (115, 116). Oral IgG survives passage through the stomach and is bioavailable (116). Antiviral therapies are untested; however, they could be considered in severe disease.

VIRAL INFECTIONS OF THE CNS IN PIDD

Viral infections of the CNS confer significant morbidity and mortality in the general population (117–119). Therefore, they are not often considered to be indicators of primary immunodeficiency. There are two circumstances where an infection of the CNS is often associated with a primary immunodeficiency: atypical herpes simplex encephalitis and CNS enteroviral disease.

Herpes Simplex Encephalitis

Herpes simplex encephalitis in the general population is most typically seen in newborns and is typically caused by herpes simplex type 2. Infection occurs at the time of delivery and infants present in the second week of life with agitation, obtundation, or seizures. Adults can develop herpes simplex encephalitis (120). Underlying immune compromise can be a risk factor for adult-onset herpes simplex encephalitis, and today HIV is the most common associated condition in adults. Herpes simplex encephalitis outside of the neonatal period may therefore suggest an immunodeficiency. Among PIDD, defects in the toll-like receptor pathway are most strongly associated with this infection (Table 3) (121). Approximately 5% of children with herpes simplex encephalitis have defects in the toll-like receptor pathway (122). Patients with these defects may present in childhood or adulthood, and some patients with just keratitis have been described (123). Recognition is important because therapy can be tailored if the defect is known. Surveillance and prevention of relapses is important. Several of these defects are inherited in an AD fashion, and therefore recognition of these PIDD is critical not only for management of the patient but also surveillance for other family members. A population study suggested that there may be additional defects inherited in an AR fashion yet to be defined (123). A key consideration is that the described toll-like receptor pathway defects are due to loss of local control in the CNS. Antibody and T cell responses are normal, and indeed, local mucosal recurrences are uncommon. As a consequence, testing of the hematopoietic cells is not revealing typically.

Enteroviral Meningoencephalitis

Viral meningoencephalitis due to a prolonged infection with enterovirus is strongly suggestive of a specific class of PIDD. Enteroviruses are the most common cause of viral meningitis in the general population manifesting as acute onset headache with gradual resolution over days to a few weeks. In patients with agammaglobulinemia, manifestations are quite different (124). These children typically present with regression of developmental milestones. Ataxia or clumsiness may be noted by parents or on examination. Features early on are subtle, and the slow progression can lead to efforts at mitigation with physical therapy or behavioral strategies. In a patient with a known humoral immune deficiency, the index of suspicion should be high and a workup should not be delayed if there are clear neurologic signs or symptoms. CNS infection in patients with agammaglobulinemia has a
very poor prognosis. There can be other phenotypes associated with enteroviral disease in patients with agammaglobulinemia; however, CNS infection is the most common. Dermatomyositis and hepatitis have been described and have progressed in some cases to CNS infection. Treatment for enteroviral disease includes high dose immunoglobulin and when available, drugs directed at enterovirus.

A unique subset of CNS enteroviral infections occurs in either SCID or agammaglobulinemia with live-attenuated polio vaccine. Wild-type polio, occurring in three serotypes, has been nearly eradicated. Even early on, it was recognized that the live-attenuated vaccine could cause disease (125) and that patients with hypogammaglobulinemia could excrete virus for years (126, 127). Currently, circulating wild-type polio is seen only in Afghanistan and Pakistan although virus can be isolated in sewage from other countries supporting ongoing risk for immunodeficient individuals (128). Vaccine-associated poliomyelitis can be due to infection of an immune deficient individual and spread to the CNS or to revertants of vaccine-strain virus (129, 130). In the latter case, even normal hosts can have overt paralytic disease. Vaccine-associated poliomyelitis can appear as acute flaccid paralysis or with a meningoencephalitis in immunodeficient individuals. The prognosis has generally been poor (131).

Diagnostic Approaches

Testing for defects related to herpes simplex encephalitis often involves genetic sequencing although functional analyses are available on a research basis. Table 3 lists the currently recognized genetic causes of susceptibility to herpes simplex encephalitis.

The diagnosis of enteroviral meningoencephalitis in PIDD patients requires a specific description. In a patient with agammaglobulinemia detection of enterovirus is surprisingly difficult. PCR analysis of cerebrospinal fluid or stool (less specific) should be performed. However, it is not unusual for children with agammaglobulinemia and suggestive clinical features to require a brain biopsy for diagnosis. The biopsy tissue can be tested for enterovirus by PCR. In a patient who presents with CNS enteroviral infection, identification of an immune deficiency is critical because of the prognostic implications. The strong association of CNS enteroviral disease with agammaglobulinemia supports a strategy that begins with enumeration of peripheral blood B cells by flow cytometry. Only if that is negative and there are no other secondary immune deficiencies should alternatives such as CD40L or CVID be sought. A reasonable secondary screen would be to measure immunoglobulin levels and responses to vaccines.

Management

Management of herpes simplex encephalitis requires specific antiviral approaches as well as attention to seizures, increased intracranial pressure, and a comprehensive intensive care approach. Acyclovir delivered intravenously is the cornerstone of management. One should consider a prolonged course of oral therapy after initial management that could include oral acyclovir or valacyclovir because the relapse rate is high in these toll-like receptor pathway defects. The role of steroids is controversial. One small study of immune competent children supported the use of beta-interferon (132), and it could be argued that interferons specifically mitigate the underlying defect in the toll-like receptor pathway disorders.

Management of CNS enteroviral disease in agammaglobulinemia has recently been reviewed (124). In the USA, antiviral drugs are not available, but pocapavir is under study and may become available. High dose IVIG has been proposed as therapeutic, but survival rates remain dismal and functional outcomes are poor.

SUMMARY

Viral infections are a common cause of morbidity in patients with PIDDs. They can be a clue to the diagnosis when persistent or unusually severe and can represent a significant management challenge.

AUTHOR CONTRIBUTIONS

MR, SH, and KS collectively conceived and wrote the manuscript.

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