Case Report

Limbic Encephalitis following Allogeneic Hematopoietic Stem Cell Transplantation

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A woman with myelodysplastic syndrome (MDS) was treated with allogeneic hematopoietic stem cell transplantation (allo-HSCT). 65 days after the transplantation, she developed fatigue and central neurological symptoms. Clinical workup including magnetic resonance imaging (MRI) and cerebrospinal fluid (CSF) examination revealed findings suspicious for limbic encephalitis (LE), successfully treated with intravenous immunoglobulins and intravenous corticosteroids. Although a rare complication after allo-HSCT, physicians should be aware of neurological symptoms that develop throughout the transplantation course.

1. Introduction

The use of allogeneic hematopoietic stem cell transplantation (allo-HSCT) is increasing in most countries [1], since this treatment is becoming more reliable and more accessible. The treatment, on the other hand, is associated with a relatively high incidence of serious, and in worst case fatal, complications. Furthermore, neurological complications are rare, but a dreaded complication of allo-HSCT [2]. Reports support that a persistent neurological impairment occurs in many of those patients who suffer from neurological complications after allo-HSCT [3]. Herein, we present a patient who developed a serious neurological complication after allo-HSCT.

2. Case Presentation

A 49-year-old previously healthy woman, with no history of mental illness or neurological abnormalities, was diagnosed with myelodysplastic syndrome (MDS) with fibrosis. She was initially treated with 5-azacitidine, while preparing for an allo-HSCT. Allo-HSCT was performed with reduced intensity conditioning (RIC) regimen with fludarabine, treosulfan, and anti-thymocyte globulin (ATG), before transplant with a matched unrelated donor (MUD), with both recipient and donor being positive for Epstein–Barr virus (EBV) and cytomegalovirus (CMV) IgG. She received cyclosporine A from the day before allo-HSCT and methotrexate the 1st, 3rd, and 6th day after allo-HSCT as prophylaxis towards graft versus host disease (GVHD). Already at day +21, she developed a generalized exanthema diagnosed as toxic epidermal necrolysis treated with intravenous immunoglobulin (IVIG) and methylprednisolone. The biopsy of the skin showed no signs of GVHD. She had a late engraftment, but at day +28, she had trilineal hematopoiesis and 99% donor chimerism and she showed no signs of acute GVHD (aGVHD). However, by day +65, she developed fatigue and altered mental level with confusion and seizures. She had low fever but normal C-reactive protein (CRP). The seizures were dominated by
contractions lasting only seconds in face, neck, arm, and leg. However, an electroencephalogram (EEG) did not show seizure. Magnetic resonance imaging (MRI) demonstrated bilaterally increased signal intensity of the amygdala and hippocampus (Figures 1(a)–1(c)). Investigation of the cerebrospinal fluid (CSF) revealed increased number of leukocytes, $42 \times 10^6/L$ (reference < 3), mononuclear pleocytosis, increased protein level 1.93 g/L (reference 0.15–0.50), and serum like IgG bands in isoelectric focusing of CSF. She was initially treated with broad-spectrum antibiotic and antiviral intravenous treatment, but these were discontinued as investigations for underlying viral or bacterial causes were negative, including human herpes virus-6 (HHV-6) in cerebrospinal fluid. Furthermore, as demonstrated in Table 1, 17 different standard onconeural and encephalitis autoantibodies were not detected in serum or CSF and antinuclear antibodies were not detected in serum. A computer tomography (CT) scan with intravenous contrast of the thoracic, abdominal, and pelvic regions had recently been performed with no signs of other malignancy. The clinical and radiological presentation were considered to be an autoimmune limbic encephalitis (LE) following allo-HSCT, due to cognitive impairment, seizures, MRI findings, and pleocytosis in the CSF. Accordingly, she was treated with intravenous corticosteroids, methylprednisolone 1000 mg for 5 days, and intravenous immunoglobulins for 5 days, total of 2 g/kg. She had an initial improvement, but the symptoms relapsed after 15 days, and the treatment was repeated and supplemented with tapering dose of per oral prednisolone for several weeks. The patient’s symptoms went slowly in regression, and control MRI after four weeks was normal (Figures 2(a)–2(c)).

3. Discussion

The classical presentation and clinical findings of LE include rapidly progressive short-term memory loss, psychiatric symptoms, and seizures, combined with MRI findings of temporal lobe involvement, and CSF inflammatory abnormalities including detection of autoantibodies [4]. Our patient highlights the possibility of this rare complication post allo-HSCT and is a reminder that detection of autoantibodies in the CSF is not a prerequisite for the diagnosis of LE. Subacute development of short-term memory deficits is often typical for LE, although it was probably overlooked in the present case by other symptoms such as headache, irritability, sleep disturbance, delusions, hallucinations, and seizures [4]. Thermal sensations, pилоeection, and paroxysmal dizziness might be symptoms resulting from focal seizure activity in LE [4]. Faciobrachial dystonic seizures are especially helpful diagnostically as they are nearly pathognomonic in a subtype of LE. A normal EEG during seizures should not eliminate the diagnosis, as coincident epileptiform discharges are seen only in the minority [4, 5]. Our patient did not have an EEG with typical findings that could correlate to the seizure, but an EEG with generalized low frequency activity supporting a diffuse central nerve system dysfunction.

LE is an inflammatory disease involving the mesial temporal lobes and the limbic structures, amygdala, and hippocampus. MRI often reveals abnormal high signal intensity on FLAIR and T2 weighted images in the mesial temporal lobes [6]. Typical MRI signal was found in our case. However, sometimes MRI may be normal, and in such cases, 18-fluorodeoxyglucose (18F-FDG) PET imaging has been reported to typically reveal mesial temporal lobe hypermetabolism [7].

Post-transplant acute/subacute LE has been described as a complication of allo-HSCT, occurring relatively early in the post-transplant period. Risk factors have been identified as heavily pretreating conditioning including ATG and corticosteroids treatment [8].

Neurological complications after allo-HSCT can be classified according to etiology or clinical presentation as listed in Table 1. All autoantibodies tested in serum and spinal fluid in our patient (all were below reference values).

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<th>Autoantibodies in serum</th>
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<td>Anti LGI 1, anti CASPR2, anti gaba R b1/2, anti DPPX, anti-GluR type AMPA½, anti GluR type NMDA, anti Tr, anti Zic4, anti GAD 65, antiSOX1, anti Ma2, antiMa1, anti CRMP5, anti Amphiph, anti Yo, anti Ri, anti Hu</td>
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Table 1: All autoantibodies tested in serum and spinal fluid in our patient (all were below reference values).

Figure 1: (a–c) Coronal FLAIR-MRI scan reveals slight enlargement and increased signal intensity of the amygdala and hippocampus bilaterally.
The etiology of LE could be divided in infectious and non-infectious causes, where the former often has been related to HHV-6 reactivation [9]. However, without evidence of infectious origin, immunological mechanisms should be considered. When first-line therapy with steroids, immunoglobulins, and/or plasma exchange fails, one converts to second-line immunotherapy. Second-line immunotherapy includes usually monoclonal antibodies (mAbs) directed at B-cells such as the anti-CD20 antibody rituximab [10]. Our patient responded to corticosteroid and immunoglobulin therapy, although it had to repeated. We suggest that a rapid and circumstantiated diagnosis of autoimmune encephalitis, herein limbic encephalitis, could anticipate specific treatments and influence the rate or the extent of long-term disabilities [3].

### Data Availability

No data were used to support this study.

### Consent

Written informed consent was obtained from the patient before publication.
Conflicts of Interest

The authors declare that they have no conflicts of interest.

Acknowledgments

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