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Case study

2	FUSARIUM INFECTION AFTER ALLOGENEIC STEM CELL
3	TRANSPLANTATION WITH PROLONGED NEUTROPENIA
4	
5	Abstract:
6	Fusarium is an opportunistic fungal pathogen which is emerging as a significant cause of
7	morbidity and mortality in the immunocompromised host (1). This disease can be localized,
8	focally invasive or disseminated, when two or more noncontiguous sites are involved. We
9	present a case of disseminated fusariosis in a patient with prolonged and profound neutropenia
10	after the third allogeneic stem cell transplantation for acute lymphoblastic leukemia.
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12	Keywords: stem cell transplantation, fusarium infection, neutropenia

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14 Case:

A 22-year-old male applied to university hospital outpatient clinics for fever and fatigue. He 15 was diagnosed as T cell acute lymphoblastic leukemia and induction chemotherapy was 16 started as methotrexate (1 $gram/m^2$, on day 1) and cytarabine (3 $gram/m^2$, on days 2 and 3). 17 During the maintenance therapy, relapse has occurred and induction chemotherapy was 18 started. Induction therapy including cyclophosphamide $(300/m^2, \text{ on days 1 to 3})$, vincristine(2 19 mg per day, on days 4 and 11), doxorubicin (50 mg/m², on day 4) and dexamethasone (40 mg 20 21 per day, on days 1 to 4 and days 11 to 14) yielded to complete remission. Therefore, after 13 22 months of the diagnosis, an allogeneic hematopoietic stem cell transplantation (allo-HSCT) 23 was planned. We searched his family but couldn't find full match donor and donor selection 24 was made among from his one related brothers with two-human-leucocyte mismatched 25 antigens. Because of HLA mismatches, autologous back-up stem cells were removed for potential graft failure. The conditioning regimen consisted of total body irradiation (TBI) 26 2x200 cGy on days -6,-5,-4 and cyclophosphamide 60 mg/kg on days -3, and -2. CD34+ 27

peripheral stem cells on a number of 4.6×10^6 /kg were collected from the donor together with T cell depletion (4 log).

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31 During the allogeneic hematopoietic stem cell transplantation (HSCT) period, 32 immunosuppressive medications (cyclosporine), antibiotics (carbapeneme, aminoglicoside, 33 teicoplanin, fluconasole, caspofungin) and filgrastim were given, as a HSCT protocol. 34 Altough we expected for an engraftment after four weeks of treatment, we did not observe neutrophil or platelet engraftment signs. We therefore assessed this clinical picture as a 35 36 primary graft failure. Therefore, after one-month of the first HSCT, a second allogeneic-HSCT from the same donor without conditioning regimen was performed (5.02 $\times 10^{6}$ CD34+ 37 38 cells /kg).

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After 10 days of the second HSCT, while he was receiving broad-spectrum multi 40 41 antibiotherapy (meropenem, teicoplanin, caspofungin, acyclovir) due to intractable fever over 38 ^oC , and immunosuppressive therapy (cyclosporine and mycophenolate mofetil) for graft 42 43 versus host disease (GVHD) prophylaxis, a target-shaped, hyperemic, painful but not itchy 44 skin lesion, 2-cm. diameters appeared on the right hand palm (Figure 1a). Laboratory examination showed white blood cell: $0x10^{9}/L$, hemoglobin: 8,6 g/dL, platelet: $5x10^{9}/L$. 45 Cultures taken regularly did not show any positive results. Blood, urine cultures and skin 46 47 biopsy were taken from the patient and empirical oral voriconazole (200 mg two times per 48 day) was started. The biopsy and blood cultures yielded fusarium. (Figure-2a, 2b). Persistent 49 fever and extended skin lesions involving whole body, confirmed disseminated character of 50 the infection under voriconasole therapy (figure-1b,1c). Then, a combination antifungal 51 therapy with liposomal amphoterisin-B (LAmB) (5mg/kg) and intra-venous voriconazole 52 (loading dose, 6 mg/kg/day, followed by 4 mg/kg/day intravenously every 12 h) was initiated. 53 The graft failure as well as persisted, that obligated the patient for another (third) HSCT. The donor, was selected this time the other HLA two-mismatched younger seventeen-year-old 54 55 brother. T-cell depletion was not preferred in order to prevent a loss in CD 34+ cell number in the graft (total number of 3.9×10^6 CD34+ cells/kg were infused after 55 day of the first 56 transplant). The conditioning regimen included fludarabine (30 mg/m² on days -4,-3 and -2) 57 and cyclophosphamide (800 mg/m^2 on days -4,-3 and -2). Antifungal and antibiotic therapies 58 59 went on the whole transplantation period without breaking. Two weeks after the third 60 transplantation, neutrophil engraftment occurred and the skin lesions related with fusarium 61 improved remarkably (90 percent) and the fever regressed. After neutrophil engraftment his

62 clinical condition improved to ECOG 1. The skin biopsy and blood cultures did not yield 63 fusarium. Four weeks after the third transplantation grade 2 GVHD of skin and 64 gastrointestinal tract have occured. Corticosteroid, mycophenolate mofetil, cyclosporine were 65 given as GVHD therapy. Due to GVHD presentation we could not discharge the patient. The patient went on ambisome for 32 days and voriconasole for 81 days after fusarium infection. 66 67 Also he received granulocyte-colony stimulating factor (Neupogen 48 MU) 91 times for 4 68 months due to profound neutropena, 72 units erythrocyte suspension and 53 units platelet 69 suspensions transfused because of severe anemia and trombositopenia. 6 months after the 70 first transplantation, although neutrophil engraftment has occurred, the patient did never 71 achieve a successful platelet engraftment and has died due to massive gastrointestinal 72 hemorrhage.

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74 **Discussion:**

Fungi belonging to the genus Fusarium are ubiquitously present in soil, air, and water and are parasites of numerous plants. In humans, these microorganisms usually cause superficial or subcutaneous infections such as keratitis or onychomycosis, but they may cause severe disseminated infections in immunocompromised patients (1).

79 Invasive or disseminated fusariosis is a rare but severe complication in hematological diseases 80 (2). Patients with compromised immune function are at high risk for invasive fusariosis, 81 particularly in the setting of prolonged and profound neutropenia and/or severe T cell 82 immunodeficiency (3). Among patients with hematologic malignancy, the infection 83 predominates during periods of neutropenia, typically among patients with leukemia receiving 84 induction chemotherapy (4). Invasive fusariosis also occurs with an increased frequency 85 among HSCT recipients higher than the autologous recepients(5). Marcio nucci et. al. 86 presented that the incidence of fusariosis among HSCT recipients varies according to the type 87 of transplantation. It is lowest among autologous and highest among the allogeneic.(6). The 88 prognosis of fusariosis is directly related to the patient's immune status, with high death rates (mortality reaches to 80 to 90%) in patients with persistent immunodeficiencies (4). We 89

90 described an adult allo-HSCT patient with disseminated fusariosis treated with early 91 combination of voriconazole and LAmB. The patient underwent to the third allo-HSCT from 92 another younger brother and we report a unique case report in this respect. The duration of the 93 neutropenia was 68 days and engraftment occurred after the third allo-HSCT. The patient's 94 clinical condition improved after neutrophil engraftment and did not die due to infection or 95 complications related with infection.

96 In severely immunocompromised patients, two characteristics suggest the diagnosis of 97 disseminated fusariosis: the presence of skin lesions (either cellulitis at sites of skin 98 breakdown caused by trauma or onychomycosis, or metastatic lesions) and mold growing 99 from blood cultures. Skin biopsies should be performed in all immunocompromised patients 100 with suspicious skin lesions, and should be sent for both histopathology and microbiology 101 studies. Blood cultures should also be obtained (7).

The optimal treatment strategy of patients with fusariosis remains unclear because of the lack of clinical trials and the critical role that immune reconstitution plays in the outcome of this infection. Successful outcomes have been reported with various antifungal agents including amphotericin B deoxycholate , liposomal amphotericin B (4), amphotericin B lipid complex, and the triazole antifungals, voriconazole (8) and posaconazole . Combination antifungal therapy has also been described in single case reports and a retrospective study (9). We presented treatment varieties and demographic features on table 1.

A lipid formulation of amphotericin B (3 to 5 mg/kg IV once daily) is usually the preferable first-line therapy. A combination of a lipid formulation of amphotericin B and voriconazole (6 mg/kg IV every 12 hours for two doses, followed by 4 mg/kg IV every 12 hours) is often used because of the variable susceptibility of Fusarium spp to antifungal agents and the need to ensure that at least one active antifungal agent is given.

114 Because a recovering immune system is essential for the successful outcome of fusariosis, 115 every effort should be made to enhance immunity; this includes decreasing the dose of immunosuppressants when possible and the use of adjunctive immunotherapy such as 116 117 granulocyte or granulocyte-macrophage colony-stimulating factors (G-CSF or GM-CSF), G-CSF-stimulated granulocyte transfusions, or interferon-gamma adjunctive therapies. The 118 119 efficacy of these therapies for fusariosis has not been established (10). 120 **References:** 121 122 1. Nucci M, Anaissie E. Fusarium infections in immunocompromised patients. Clin Microbiol

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	Gulsum Tezcan	Isabelle	Marta	Erturk et al.
	et. al.	Durand-Joly	Stanzani et.	(Reported Case)
		et. al.	al.	(F)
Underlying	ALL	ALL	AML	ALL
condition				
Underlying	_	+	_	_
condition not in				
complete remission				
Type of stem cell	Allo-HSCT	-	Allo-HSCT	Allo-HSCT
transplantation				
Conditioning	Busulfex,	-	Fludarabine,	TBI,Cyclophospha
regimen	Etoposid,		BCNU,	mide, and
C C	Cyclophosfamid		Alcaran	Fludarabine and
				Cyclophosphamide
				(for the second
				HSCT)
Immunosuppressive	ATG +		ATG,	Cyclosporine and
Agents	Mycophenolate		Cyclosporin	Mycophenolate
	Mofetil		А,	Mofetil,Steroid)
			Methotrexate,	
Culture	Blood	Skin	Skin,	Skin,
	Synovial fluid		Blood	Blood
GVHD	-	-	-	+
Graft failure	+	-	-	+ (first and second
				HSCT)
Duration of	95	15	30	68
neutropenia (day)				
Antibiotherapy for	LAmB +	LAmB +	LAmB +	LAmB +
fusariosis	Voriconasole	Voriconasole	Voriconasole	Voriconasole
		(Voriconasole		
		after		
		dissemination)		
Death because of	-	-	-	-
fusariosis				
Disseminated	+	+	+	+
Infection				
Number of HSCT	2	-	1	3

Table 1. Demographic and clinical characteristics of HSCT recipients with fusariosis

- 151 ALL: Acute Lymphoblastic Leucemia, AML: Acute Myeloid Leucemia, ATG: Anti-
- thymocyte Globuline, TBI: Total Body Irradiation, GVHD: Graft Versus Host Disease,
- 153 HSCT: Hematopoetic Stem Cell Transplantation, LAmB: Liposomal Amphoterisin B
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Figure-1a

Figure-1b

Figure -1c

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- Figure-1a: A target-shaped, hyperemic, painful but not itchy skin lesion, 2-cm. diametersappeared on the right hand palm
- **Figure-1b and 1c:** Extended skin lesions involving whole body, confirmed disseminated
- 161 character of the fusarium infection

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Figure 2b

Figure-2a and 2b: The biopsy and blood cultures yielded fusarium