# Recurrent Peripheral Facial Paralysis Revealing Behçet s Disease: A Case Report

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#### RecurrentPeripheral Facial ParalysisRevealingBehçet'sDisease: A Case Report 1

#### 2 Abstract

- We report the case of a 21-year-old female patient presenting with recurrent peripheral facial 3
- paralysisassociatedwithoropharyngealaphthosis and diffuse arthralgia.
- Immunologicaltestingrevealed HLA-B51 positivity, suggestingBehçet's disease. The patient's 5
- condition improvedwith a combination of corticosteroidtherapy, colchicine, 6
- functionalrehabilitation, and symptomaticmeasures.

#### 8 **Key Words**

Behçet'sdisease - Recurrent facial paralysis - Neuro-Behçet - HLA-B51

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### Introduction

- Behçet'sdisease (BD) is a chronic, relapsing, multisysteminflammatorydisordercharacterized 12
- by recurrent oral and genitalulcers, ocularinvolvement, and variable systemic manifestations. 13
- It is classified among the variable vesselvas culitides, as it can affect both arteries and veins of all 14
- calibers. Although the etiologyremainsunclear, 15 exact
- interactionbetweengeneticpredispositionparticularly HLA-B51 positivityand environmental or 16
- 17 infectious triggers issuspected.
- Neurologicalinvolvement, termedneuro-Behçet's disease, occurs in approximately 5-10% of 18
- 19 cases and represents one of the mostsevere complications. It canmanifest as
- 20 parenchymallesions involving the brainstem or as vascular complications such as
- cerebralvenousthrombosis. involvement, however.
- system 21 Peripheralnervous
- remainsexceptional. 22
- 23 Peripheral facial paralysisis a commonneurological disorder, most often idiopathic or post-viral
- in origin. Its recurrences hould prompt investigation for an underlying systemic disease. We 24
- 25 report here an unusualpresentation of Behçet's diseaser evealed by recurrent peripheral facial
- 26 paralysis in a youngwoman, emphasizing the importance of multidisciplinaryevaluation and
- 27 early recognition of atypical forms.

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## Case Report

- 31 Ms. M. M., a 21-year-old womanwith no significant medical history, experienced a right-
- sidedperipheral facial paralysisthreemonthsearlier, whichhad been attributed to idiopathic 32
- 33 (cold-induced) Bell'spalsy. Her condition hadimprovedundercorticosteroidtherapy and
- physicalrehabilitation. 34
- 35 Threemonthslater, shepresentedagainwith a recurrence of the same facial deficit.
- Clinicalexaminationrevealed a complete right-sidedperipheral facial paralysis (House-36
- 37 Brackmann grade IV), with no involvement of othercranial
- examinationidentified a single aphthousulcerlocated on the right tonsillarfossa. The patient 38

alsoreportedmigratorypolyarthralgia, mainlyaffecting the knees and wrists, withoutswelling or 39 40 local inflammatorysigns. Laboratoryworkupshowed: normal CBC, erythrocytesedimentation rate (ESR) of 28 mm in 41 42 the first hour, nearly normal C-reactive protein (CRP), and positive HLA-B51. Brain MRI 43 wasstrictly normal, showing no brainsteminvolvement or inflammatorylesions. 44 Ophthalmologic examination revealed no uveitis or retinal involvement. 45 Discussion 46 Peripheral facial paralysisis one of the mostcommonreasons for ENT consultations. It isusuallyidiopathic or post-viral. However, recurrence of such an episodewithina short period 47 in a young patient should prompt consideration of a systemicetiology. 48 49 In this case, the presence of oropharyngealaphthosis, polyarthralgia, and HLA-B51 positivityled 50 to the diagnosis of Behçet'sdisease. 51 chronicsystemicvasculitispredominantly affects youngadults and isclassicallycharacterized by 52 triad of oral aphthosis, genitalaphthosis, Neurologicalinvolvement in Behçet's disease is rare but severe, occurring in approximately 5-53 10% of cases, and maybeeitherparenchymal or vascular. In our observation, the absence of 54 system 55 brain MRI lesionsruled out central nervous involvement suggestedinflammatoryperipheral 56 damage to the facial nerve. 57 This case highlights the importance of acomprehensive approach and multidisciplinary followup in patients presenting with recurrent facial paralysis. 58 The patient wastreated with colchicine (1 mg/day) to control systemic inflammation, low-dose 59 oral corticosteroids to reduce facial nerve edema, vitamin B12 to support nerve regeneration, 60 and artificialtears for ocular protection. Functionalrehabilitation sessions were prescribed to 61 acceleraterecovery of 62 Clinicalevolutionwas favorable aftertwoweeks, with progressive recovery of facial movement 63 and disappearance of joint pain. No recurrence was observed after three months of follow-up. 64 65 66 67 Conclusion 68 69 Behçet's disease should be considered in any case of recurrent facial paralysis associated with 70 oral aphthosis. Early recognition of thisetiologyallows for targeted management and prevention of severeneurological or ocular complications. This case underscores the 71 importance of aholisticclinicalapproach and close collaboration between ENT specialists, 72 73 internists, and ophthalmologists in the management of suchatypical presentations. 74

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