SUDDEN ONSET OF DOUBLE ELEVATOR PALSY: A RARE CLINICAL ENTITY
Sundip Shenoy¹, Anitha Maiya², R. Jayaram³, Venita J. Noronha⁴, Vasanth Kumar⁵

HOW TO CITE THIS ARTICLE:

ABSTRACT: Double elevator palsy (DEP) currently known as Monocular Elevation Deficit is a rare condition characterized by restricted elevation of one eye in all positions of up gaze. Clinically it presents as a unilateral restriction of superior rectus (SR) and inferior oblique (IO). We present a case report of DEP in 3 year old boy who presented with sudden onset of DEP in right eye in our OPD. Very little information on acquired causes of DEP is available. Diagnosis, differential diagnosis and management are discussed and then the literature is reviewed.

KEYWORDS: DOUBLE ELEVATOR PALSY (DEP), MONOCULAR ELEVATION DEFICIT (MED), UPGAZE PALSY.

INTRODUCTION: Double elevator palsy (DEP) is a rare condition which was first described by White in 1942 and later by Dunlap.¹ In DEP there is apparent paralysis of the elevators of one eye, the superior rectus and inferior oblique resulting in restricted elevation. DEP is actually a misnomer because true paralysis of the elevator muscles is seen only one-fourth of cases and mostly only one of the elevator muscles may be involved in certain cases. Hence DEP is better known as Monocular Elevation Deficit (MED)².

CASE REPORT: A 3 year old school going male child presented to our clinic with complaints of sudden onset of downward vertical displacement of right eye of 15days onset with chin elevation. His birth history was normal with normal Apgar score and no history of squinting (intermittent / manifest) in past 3 years. No history of stress, fever, trauma, measles or whooping cough or patching of eye. He is immunised upto age. Developmental milestones were normal. His ocular, medical and surgical history was unremarkable. No history of any systemic diseases or association with any congenital syndromes. Family history was negative.

General examination was normal. Old photographs were normal and showed no deviation of either eye.

On ocular examination, Visual acuity in both eyes was noted to be 6/6 OU, not a spectacle wearer. Pupils were round regular and reactive. No nystagmus was noted. He presented with a head posture, mainly chin up position and head tilt to right. Visual field was normal by confrontation test. Cover uncover test revealed hypotropia of right eye seen in primary gaze associated with pseudo ptosis with brow elevation. No variation in lid position was seen during the course of the day. Neurological examination revealed intact Bell's phenomenon indicating supranuclear lesion in DEP.

Extra ocular muscle movement revealed a restriction of movement of right eyeball in up gaze (supraduction /supraversion) both in adduction and abduction pointing towards elevator deficit (Right SR and right IO) (PHOTO 2 and 6). All other movements were normal except for excessive movement of eyeball down, revealing right inferior rectus overaction (PHOTO 7). Buckling and tethering of central portion of lower lid was not seen which implies Inferior rectus restriction is
absent. Krimsky and PBCT was done with base up prisms which confirmed vertical deviation of 10 pd. Worth’s 4 dot test revealed no diplopia. There was no restriction of movement in left eye. Near point of convergence and accommodation was normal. FDT was negative.

Anterior segment was otherwise normal with normal fundus evaluation. Intra ocular pressure was normal. Atropine refraction revealed normal retinoscopic studies

**Investigations:**

Neurological examinations was normal.

CT of the brain and orbit revealed normal study.

**DISCUSSION:** In DEP, there is unilateral limitation of elevation in all horizontal orientations of the eye. When the patient fixates with the nonparetic eye (left), the paretic eye (right) will take a hypotropic position and the upper lid may be slightly ptotic. Fixation with the paretic eye (right) will cause a hypertropia of the nonparetic eye (left), and ptosis may disappear, provided the levator palpebrae is not involved (PHOTO 5). Patient may also complain of frequent upshoots of normal eye. Anatomical improbability that both superior rectus and inferior oblique are involved by single lesion suggests long standing SR palsy as the primary event with later spread of comitance leading to inferior oblique involvement. FDT is done to confirm any mechanical restriction of movement and was negative. Often there is chin up position to achieve binocular single vision. Bell’s phenomenon is usually preserved and differentiates between supra nuclear and infra nuclear lesions. No risk of post-operative corneal exposure.

There are mainly two forms; Congenital and Acquired and has also been reported in twins\(^3\). Among the acquired forms one needs to exclude pineocytoma, polycythemia vera, vascular events and metastatic events. Possible differential diagnosis of DEP include blow-out fracture of the orbital floor, congenital or acquired fibrosis, endocrine myopathy, myasthenia gravis, skew deviation, Parinaud’s syndrome, Browne syndrome, heavy eye syndrome, pineocytoma and 3rd nerve palsy.

(DIFFERENTIAL DIAGNOSIS OF DEP/MED)

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<tbody>
<tr>
<td>1</td>
<td>BROWNS SYNDROME</td>
<td>Absence of elevation in adduction, FDT positive</td>
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<td>2</td>
<td>ORBITAL FLOOR FRACTURE</td>
<td>Monocular limitation of elevation with numbness of infra orbital region and enophthalmos, FDT positive on elevation, X ray and CT confirmatory</td>
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<td>3</td>
<td>THYROID OPTHALMOMOPATHY</td>
<td>Acquired limitation in elevation with eye signs and IR involvement</td>
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<td>4</td>
<td>MYOSITIS</td>
<td>Limited EOM with signs of inflammation, FDT Positive</td>
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<td>5</td>
<td>CONGENITAL MUSCLE FIBROSIS</td>
<td>Bilateral ptosis, hypotropia, up gaze limitation</td>
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<td>6</td>
<td>HEAVY EYE SYNDROME</td>
<td>Seen in myopia</td>
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**TABLE 1A: RESTRICTIVE CAUSES**
CASE REPORT

### 1. 3rd NERVE PALSY
- Ptosis, eye in abducted position, dilated pupil

### 2. MYASTHENIA GRAVIS
- Diurnal variation of ptosis, eye signs positive

### 3. CHRONIC PROGRESSIVE EXTERNAL OPHTHALMOPLEGIA
- Deafness, ptosis, retinal degenerations, Ophthalmoplegia

<table>
<thead>
<tr>
<th>TABLE 1B: INFRA-NUCLEAR CAUSES</th>
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<tbody>
<tr>
<td>1. PARINAUD'S/ DORSAL MID BRAIN SYNDROME</td>
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<td>2. SKEW DEVIATION</td>
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<td>3. SUPRANUCLEAR PARESIS OF MONO OCULAR ELEVATION</td>
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### Table 1C: Supranuclear Causes
- **Von Noorden** attributed DEP to long-term paralysis of the superior rectus (SR) in the presence of a normally functioning inferior oblique (IO) muscle, indicating that the motor signs may be due to SR paralysis alone. Ziffer et al classified DEP into three subgroups, using scleral search coil technique for evaluating saccadic velocities into groups of IR restriction, complete or incomplete SR paralysis and supranuclear palsy.

**INVESTIGATIONS:**
1. **MRI/CT:** Neurological and squint evaluation was discussed. CT normal

**TREATMENT:** There are different surgical treatments for DEP which reflects the various causative mechanisms. Results are seen only in improvement of eye position in primary gaze with minimal improvement in elevation.

   Treatment is generally based on carrying out the forced duction test (FDT).

   If FDT is positive, inferior rectus recession is carried out and most patients recover satisfactorily.

   In the absence of IR restriction, (forced duction test negative) surgical treatment comprises of the KNAPP PROCEDURE. Here the entire tendon of both the medial and lateral rectus muscle is transferred to the side of superior muscle insertion. In transposition, as the 4 anterior ciliary arteries are sacrificed, a 6 month adaption time should be given before operating the third rectus to prevent anterior segment ischemia. In the absence of IR restriction, results are good. If the IR is restricted, the muscle has to recessed before or after transposition.

   If vertical deviation in primary position is large, then IR recession with SR resection is done. Recently, tucking of IR of normal eye is also done to remove residual hypotropia and pseudoptosis that may persist after the Knapp Procedure.

   Therefore, the surgical procedure should be individualized for each case based on the etiology and mechanisms of involvement.
SUMMARY: A case of sudden onset of Monocular elevation deficit /DEP (right eye) in a 3 year old boy with chin elevation and head tilt to the right is described with various etiologies and management.

PHOTO 1: HEAD POSTURE (CHIN ELEVATION AND HEAD TILT TO RIGHT)

PHOTO 2: SUPRAVERSION SHOWING RESTRICTED UPGAZE ONLY IN RE

PHOTO 3: DEXTROVERSION

PHOTO 4: LEVOVERSION SHOWING OVERACTION OF SUPERIOR OBLIQUE-RE

PHOTO 5: PRIMARY GAZE SHOWING HYPOTROPIA CORRECTED WITH PARETIC EYE FIXING (RIGHT EYE) AND CORRECTION OF PSEUDOPTOSIS

PHOTO 6: SUPRADUCTION RE SHOWING RESTRICTION OF UPGAZE
PHOTO 7: INFRADUCTION RE SHOWING OVERACTION OF IR

PHOTO 8: ADDUCTION RE

PHOTO 9: NORMAL ABDUCTION RE

PHOTO 10: NORMAL SUPRADUCTION LE

PHOTO 11: NORMAL INFRADUCTION LE
REFERENCES:

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