Management of congenital muscular torticollis Under one year of age
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Summary:
Background and study aim:
Torticollis is the postural deformity of head and neck. Congenital Muscular Torticollis (CMT) is a postural deformity of head and neck detected at birth or shortly after birth, primarily resulting from unilateral shortening of Sternocleidomastoid muscle (SCM). In neonates and infants, patient may cure conservatively by physiotherapy but surgery is the treatment of choice for children and adolescents. Here we show our experience regarding management of congenital muscular torticollis with physiotherapy.

Patients and Methods:
This is an observational descriptive study. Verbal consent from parents was taken. Patients of congenital muscular torticollis with other disease or other congenital anomaly were excluded from study. Twenty patients of congenital muscular torticollis were treated. The cases were enrolled between Nov' 2005 to Oct’ 2008 in Bangabandhu Sheikh Mujib Medical University, Gonosasthaya Somaj Vittik Medical College Hospital, ZH Sikder Women's Medical College Hospital, Shaheed Shurawardy Medical College Hospital. Neonates and infants were treated conservatively with physiotherapy and non responsive cases were referred for surgery.

Results:
Patients age range from 5 days to 1 year of which eleven were females and nine were males. Sternocleidomastoid muscle (SCM) was shortened in all cases (12 on right side and 8 on left side). Of 20 patients 6 neonates, rest 14 infants within 1 year age. Out of 20 neonates and infants 17 were cured conservatively with physiotherapy and rest 3 were referred for surgery. Conclusion: Most of the patient of congenital muscular torticollis can be treated conservatively during infancy.

Key words: Congenital Muscular torticollis (CMT), Wryneck, Conservative Management, Manual Stretching.

Introduction:
Torticollis is the postural deformity of head and neck which may be acquired or congenital. There are various types of congenital torticollis. Congenital Muscular torticollis (CMT) is a postural deformity of head and neck detected at birth or shortly after birth, primarily resulting from unilateral fibrosis & shortening Sternocleidomastoid muscle. The shortening of Sternocleidomastoid muscle (SCM) results in traction of mastoid process toward the sternoclavicular joint. The head is therefore rotated & tilted toward the involved Sternocleidomastoid muscle. The condition is some times called "Wryneck". Torticollis is also known as twisted neck. The incidence of Congenital Muscular torticollis (CMT) is one in every 300 live births. Plagiocephaly may co-exist in 80-90% of children with CMT. Though impairment of Sternocleidomastoid muscle (SCM) function is the most frequent cause of CMT but torticollis could also be result from other underlying disorders. The exact pathophysiology and etiology of sternocleidomastoid impairment in CMT still unknown. In Congenital Muscular torticollis (CMT) patient's head...
remain tilted toward the involved side. Child and adolescents patient can't look forward. If the patient wants to look other sides he/she have to rotate whole body as he / she can't move the neck. Gradually patient develops facial hemihypoplasia which results in flattening and under development of the malar eminence, downward displacement of the eye, ear and angle of mouth on the effected side. It also hampers the development of facial skeletons.

Children with Congenital Muscular torticollis (CMT) can be assigned to one of three groups

1. Children with a palpable swelling or pseudo tumor of the sternocleidomastoid,

2. Children with sternocleidomastoid (SCM) tightness but no tumor,

3. Children with all features of muscular torticollis without muscle tightness or tumor. Diagnosis can do by history, physical examination & clinical progression but in some cases ocular, neurological evaluation and radiological investigation of cervical spine is necessary.

Management depends upon the age of patient. Patient of below 1 year of age, treatment is conservative - Physiotherapy and above one year of age - treatment is surgery. Within one year of age 69 to 91 % patient may cure conservatively with physiotherapy.

Conservative management of infants with torticollis consists of positioning, gentle range of motion, and strengthening through activation of head and trunk muscles as the infant gains control of upright postures. Manual stretching is the most common form of treatment for CMT. Proper stabilization and hand placement is vital for the success of each stretch; however, all child/parent pairs will not be comfortable with the same method of stretching or the same stretch positions. The severity of the torticollis, the age of the child, the tolerance of the child for handling, and the parent's ability to carry out the exercise program will determine the method of stretching. When performing stretching exercises, the position of the head and neck in flexion versus extension will impact the effectiveness of the stretch.

Materials and methods

This is an observational descriptive study. Verbal consent from parents was taken. Patients of congenital muscular torticollis with other disease or other congenital anomaly were excluded from study. Data was collected by self administered pre-tested semi-structured questionnaire. Twenty cases of Congenital Muscular Torticollis (CMT) were treated from Nov' 2005 to Oct' 2008 in Bangabandhu Sheikh Mujib Medical University, Gonosasthya Somaj Vittik Medical College Hospital, ZH Sikder Women's Medical College Hospital, Shaheed Shurawardy Medical college Hospital Bangladesh were included in the study. Patients were diagnosed clinically. All the patients present with short sternocleidomastoid muscle. There were no facial hemihypoplasia or any other associated abnormalities. All the cases were treated conservatively with physiotherapy as out patient - none was admitted. Physiotherapy is simple - manual stretching of affected sternocleidomastoid, frequent movement of head on opposite direction of head position, lied lateral position on effected side. Patient's mother was trained accordingly. Physiotherapy was given at home by mother.

Results:

Total 20 patients were selected. Range of age was from 5 days to 1 year. Eleven were female and nine male. SCM was shortened in all cases (12 on right side and 8 on left side) Right: Left = 1.5:1. Among them neonates 6 and infants 14 within 1 year of age. There was no associated anomaly. Fifteen patients had the history of normal vaginal delivery, one patient had the history of forceps delivery and four had the history caesarian operation. Among the normal vaginal delivery patients, Six have prolonged labor of which four was first issue. (Table-I).

Table-1: Mode of delivery

<table>
<thead>
<tr>
<th>Mode of delivery</th>
<th>No of Patients</th>
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</thead>
<tbody>
<tr>
<td>Caesarian Section</td>
<td>4</td>
</tr>
<tr>
<td>Normal Vaginal Delivery</td>
<td>15</td>
</tr>
<tr>
<td>Forceps (First Born Baby)</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>20</td>
</tr>
</tbody>
</table>

None can give the history of breech delivery. Antenatal history was uneventful. There is no family history of torticollis in any patient. Out of 20 neonates and infants 17 cured conservatively with physiotherapy (Table-II).

Table- II: Treatment Result

<table>
<thead>
<tr>
<th>Outcome of physiotherapy</th>
<th>No of patient treated</th>
<th>% of treated patient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cured</td>
<td>17</td>
<td>85%</td>
</tr>
<tr>
<td>Failed</td>
<td>03</td>
<td>15%</td>
</tr>
<tr>
<td>Total</td>
<td>20</td>
<td>100%</td>
</tr>
</tbody>
</table>

Total 03 failed with physiotherapy due to non compliance with physiotherapy and were referred for treated
surgically.

Discussion:

Congenital Muscular Torticollis (CMT) patient's head always remain tilted to the affected side, they can't look forward, can't rotate the head and neck as they want. They have to move the whole body to see the side objects. Untreated cases develop mandibular hypoplasia, facial hemihypoplasia, skeletal deformity of face, etc which leads to permanent disfiguration. So Congenital Muscular Torticollis (CMT) patients suffer both functional and cosmetic problems. Though in literature, the incidence of CMT is 1 in every 300 live births\(^4\), but we have no data about CMT in Bangladesh. The optimal management of Congenital Muscular Torticollis (CMT) has been urged for many years. Most agree that physiotherapy is the mainstay of the treatment. In literature success rate of physiotherapy is 69-91 %. In our study it is 85% which is within the limit of international studies.

Surgery is usually reserved for patients whose conditions were persistent beyond the age of one year or failed with physiotherapy or when cervical function and facial deformities are unacceptable\(^11\).

Movement of the head and neck depends upon the synergistic and antagonistic activities of the different cervical muscles, such as splenius capitis, trapezius, platysma, the longus coli, the longus capitis, rectus capitis, etc\(^15\). So, in absence of Congenital Muscular Torticollis (CMT) in one side will not interfere with the movement of head and neck.

In our study, Right : Left = 1.5:1 which corresponds with the study of others\(^16\). There are records of family history of CMT in 3.6%\(^16\) but we have no such finding which also corresponds with the study of others\(^3\). Facial asymmetry and plagiocephaly are common, though not invariable associated anomaly with CMT. Both believed to be secondary to the Torticollis. Reported co-existence of hip dysplasias with CMT varies from 0.6% to 20%\(^16\) but we have no such condition or any other associated anomalies like others\(^17\). Though in reported literature recurrent torticollis is about 3%\(^19\) but we have no such record.

Conclusion

Early detection and initiation of physical therapy is related to improved outcomes and less need for surgical treatment of the SCM. Repositioning is a required element of early management of torticollis. Most of the patient of congenital muscular torticollis can be treated conservatively during infantile period.

Beyond the age of one year and failed cases surgery is the treatment of choice. Limitation of this study is small sample size. Further study with large sample size is advocated. To find out the burden of CMT or CMT with other congenital abnormality prevalence study can done in community.

References:
