"TUBEROUS SCLEROSIS COMPLEX" - WITH RARE MULTISYSTEM INVOLVEMENTS

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ABSTRACT

BACKGROUND
Tuberous Sclerosis represents a genetic disorder of hamartoma formation in many organs, particularly the skin, brain, eye, kidney, and heart. It is now recognized that about half the families are linked to 9q34 (TSC1) and the other half to 16p13 (TSC2).

KEYWORDS
Tuberous Sclerosis, Rare Systemic Involvements, Topical Treatment with Sirolimus for Adenoma Sebaceum.


CASE REPORT
Patient 27 yrs., female, unmarried, not working residing at Usmanabad, hailing from Maharashtra came with the chief complaint of:
1. Raised skin lesions all over the face and chest since age of 5 yrs.
2. Outgrowing lesions from both the great toe since age of 5 yrs.
3. Bleeding from nose since 3 months
4. Watery discharge from right ear since 3 months
5. U/L headache on right side since 8 days
6. Pain in jaw since 8 days.

Pt. was apparently alright till the age of 5 years, then suddenly her mother noticed raised lesion on the nose which gradually spread to both side of face over the time. She also noticed similar lesion on the chest.

Patient also complaint of outgrowing lesion from the great toe, which keeps growing and then sheds off by itself. Patient took treatment for the same lesion which was satisfactory and the lesion recurred later.

On further inquiry, patient revealed that 9 months back she was admitted for the pain over both flanks, nausea, vomiting, headache with generalized weakness since 2 months, for which USG ABD. Was done, which was S/O multiple renal angiomyolipomas.

Then 3 months back she developed bleeding from nose every 2-3 days watery discharge from ears and then she also developed U/L right side headache and pain in jaws.

No h/o seizures.
No h/o mental retardation.
No h/o trauma.

Cutaneous Examination
Multiple skin coloured hyperpigmentation, dome shaped papules distributed B/L symetrically over the nose, nasolabial fold, cheeks, chin forehead, eyelid measuring 1-5 mm in size. (Adenoma sebaceum, cutaneous angiofibroma) [Fig. 1, Fig. 2].

Contrast Enhanced CT Abdomen and Pelvis Reveals,
- Multiple renal angiomyolipomas bilaterally. [Fig. 14]
- Lymphangiomyomatosis in the visualized lung bases. [Fig. 17]
- Patchy sclerosis at most places in visualized axial skeleton. [Fig. 15 & 16]
- Mild hepatomegaly with incidental right ovarian cyst and pelvic congestion. [Fig. 18]

CT PNS reveals,
- Changes of right maxillary sinusitis with relatively hyperdense contrast suggest possibility of, Desicated secretions, areas of haemorrhage, secondary infections. [Fig. 11].
- Subependymal calcified tubers with possible subependymal giant cell ependymoma in the region of foramen of Monro. [Fig. 10].

Plain MRI Brain Reveals,
- Areas of subependymal calcification or tubers seen involving frontal horns, right occipital horn largest of size 9 x 8 mm. [Fig. 13]
- Patchy area of altered signal intensity of size 11 x 11 mm on left side involving base of tongue. [Fig. 12]
  - 2D echo/Doppler study- Normal
  - Normal LV EF
  - No pulmonary artery hypertension
  - ECG- Normal
  - EEG: normal, no epileptiform activity.
  - Chest X-ray: NAD
Biopsy
Biopsy of shagreen patch showed increased collagen in dermis with H&E stain and Verhoeff-Van Gieson stain demonstrates increased red collagen and sparse black elastic fibers.

Figure 1

Figure 2

Figure 3

Figure 4

Figure 5
Nodular lesion in the region of foramen of Monro on the left suggestive of Subependymal giant cell astrocytoma. Areas of subependymal calcification in relation to the right occipital horn.
Figure 12. Patchy area of altered signal intensity of size 11 X 11 mm on left side involving base of tongue

Figure 13

Figure 14

Bilateral enhancing mass lesions with fat attenuation areas within at most places in an intra-renal, perinephric and subcapsular locations suggestive of angiomyolipomas.

Figure 15

Figure 16. Patchy sclerosis was noted at most places in visualized axial skeleton on bone windows

Figure 17. Lung bases covered in the CT ABD study showed multiple small well defined thin walled cystic lesions in both lungs suggestive of lymphangioleiomyomatosis
DISCUSSION

Tuberous sclerosis complex is an autosomal dominant neurocutaneous disorder in which multisytem hamartomas are seen. It is caused by the mutations in two genes: Tuberous sclerosis complex 1 on 9q34.3 that encodes the protein hamartin, and tuberous sclerosis complex 2 on 16p13.3 that encodes the protein tuberin. Normally, these suppressor proteins inhibit the activation of mammalian target of rapamycin (mTOR). Any mutation in these genes lead to the dysregulation of this inhibitory process, leading to uncontrolled cell growth, proliferation and protein synthesis that results in multi-organ hamartomas. This disorder has an incidence of 1/6000 births, and mostly affects the skin, kidneys, heart, brain and lungs.

The cutaneous manifestations of tuberous sclerosis complex include angiofibromas, forehead fibrous plaques, hypomelanotic macules or confetti-like lesions, periangual fibromas and shagreen patches. Facial angiofibromas are small, symmetrical, pink to red papules which coalesce to form plaques. They are composed of blood vessels and fibrous tissue, and are seen in approximately 80% of patients, making it the most frequent cutaneous finding of this disease. According to the recommendations of the “International Tuberous Sclerosis Complex Consensus Conference 2012,” they are considered as one of the major diagnostic criteria. Although benign in nature, they can bleed or obstruct the nasal openings. They cause great psychosocial burden to patients. Therapeutic options included cryotherapy, electrocoagulation, dermabrasion, laser surgery and podophyllotoxin application.

Epilepsy is seen in almost all mentally retarded patients and in 70% of those with average intelligence. It usually begins in infancy or early childhood, thus often preceding the skin lesions by many years. Ocular signs occur in 50% of cases but may be hard to detect. Retinal phacomas are seen as white streaks along the vessels or as small, rounded tumours near the disc. Cardiac and renal tumours are often asymptomatic unless by reason of their size or site.

Cardiac rhabdomyomas, detected by echocardiography, occur in over 50% of infants. Renal involvement includes angiomyolipomas, a benign tumour of the renal parenchyma, renal cyst. Pulmonary changes are rare and seldom cause symptoms, but if extensive can result in increasing dyspnoea and recurrent spontaneous pneumothorax.

**Major Criteria**
1. Facial angiofibroma / forehead plaque.
3. Shagreen patches
4. Cortical tuber (Brain).
5. Subependymal nodules (Brain).
6. Subependymal giant cell astrocytoma (Brain).
7. Lymphangioleiomyomatosis (lung).
8. Renal angiomyolipoma.
9. Cardiac rhabdomyoma.
10. >3 hypomelanotic macules.
11. Multiple retinal nodular hamartoma.

**Minor Criteria**
1. Multiple random pits in enamel.
5. Bone cysts.
6. Cerebral white matter radial migration lines.
7. Retinal achromatic patch.
8. Confetti skin lesions.
9. Multiple renal cysts.

We also studied another 11 patients to evaluate the effectiveness and tolerability of topical sirolimus for facial angiofibromas in patients with tuberous sclerosis complex. We investigated the effect and safety of topical 0.1% sirolimus, which was obtained by crushing sirolimus tablets and mixing it with petrolatum. The patients were asked to apply the cream to one side of their face, and Vaseline to the other side. The effect of topical sirolimus was evaluated using the “facial angiofibroma severity index.”

There was a significant improvement in the redness and extension of the tumours on the sides to which the active ingredient was applied. Some side effects such as itching and irritation occurred in three patients, which were treated with topical hydrocortisone cream.

Topical sirolimus appears to be a promising, fairly well tolerated treatment, for facial angiofibromas, in patients with tuberous sclerosis complex. Although its efficacy diminishes with time, repetitive usage is effective.

**REFERENCES**


