Port Wine Birthmark Presentation

Port Wine birthmarks (capillary vascular malformations) of the face are the hallmark of Sturge-Weber Syndrome.

Wounds in Port Wine Birthmarks
- Port Wine birthmarks are comprised of a massive excess of capillary blood vessels. While these vessels have low flow, wounds in port wine birthmarks may bleed significantly more than normal skin.
- Compression bandages are important before and after suturing wounds in Port Wine birthmarks.
- Local anesthesia is the same as for any wound.
- Extensive electrocau-ogulation of bleeding dermal blood vessels should not be done in order to avoid cutaneous necrosis.
- Deeper wounds require dermal suturing to obtain perfect dermal alignment. Resorbable 4.0 sutures should be used.
- Cutaneous sutures should not be bono-resorbable, 4.0 or 5.0 and removed in 7-10 days.
- Patients should wear a compressive bandage for 24 hours following suturing.
- Patients should be seen in follow up the next days.

Pyogenic Granulomas in Port Wine Birthmarks
- Pyogenic granulomas are neovascular growth that are seen in normal skin, but also occur commonly in port wine birthmarks.
- They have a very friable surface and bleed frequently.
- Local anesthesia is the same as for any skin injury.
- The simplest and best single treatment for pyogenic granulomas is superficial removal and then cautery or laser photocoagulation of the bleeding papule.
- Bleeding from the subsequent wound is best controlled by pressure and gentle electrocau-ogulation.
- Recurrence of the lesion can occur requiring re-treatment.
- Excisional surgery will lead to a larger scar and should be performed by physicians with experience in treatment of pyogenic granulomas.
- For lesions less than 5 mm, pulsed dye laser treatment if available, is simple and may be effective.

Other Dermatological Conditions of the Skin Involved with a Port Wine Birthmark
- Common skin conditions such as eczema, acne, impetigo and the majority of skin diseases may be treated the same as in patients without Port-Wine birthmarks.

The information contained herein is specific to the United States. We recognize that emergency procedures in other countries may differ.
**Neurologic Presentation**

The major neurological features of patients with Sturge-Weber syndrome include seizures, headache, focal neurologic deficits and cognitive and psychological impairment.

### Seizures

- **Seizures** develop in 70 to 80% of patients with Sturge-Weber syndrome.
- The risk of developing seizures is highest in the first two years. 75% of those with seizures have the initial event during the first few months of life often in conjunction with hemiparesis or other focal deficits.
- The natural course of Sturge-Weber syndrome depends on the presence, persistence and resistance to treatment of seizures. Onset of seizures before 1 year of age, and the occurrence of episodes of status epilepticus predict an unfavorable outcome.
- Seizures are usually focal motor or secondarily generalized tonic clonic seizures, but infantile spasms, myoclonic seizures and atonic seizures may also occur.
- Many patients have frequent and repeated seizures, but some children have only occasional seizures. In some patients, seizures may cluster.
- EEG studies document decreased amplitude and frequency of electrocerebral activity over the affected hemisphere.
- Diffuse, multiple and independent spike foci are commonly present.
- MRI of brain with gadolinium enhancement is the preferred imaging modality to evaluate the extent and severity of intracranial involvement. Brain MRI may be occasionally negative during the first few months (or even years) of life despite showing abnormalities subsequently.
- When a child presents with a new-onset seizure, a neurologic consultation should be sought. Routine studies including blood glucose, electrolytes, serum calcium and magnesium and CBC should be obtained to evaluate for precipitating factors. EEG and neuroimaging studies such as CT or MRI scan of brain should also be obtained. Aggressive antiepileptic therapy should be established from first seizure.
- Arrest of status epilepticus is of utmost importance. Immediate management includes assessment of vital functions, establishment of intravenous lines, blood sampling for investigation of possible metabolic or infectious causes and IV benzodiazepines such as lorazepam. IV benzodiazepines may be repeated if necessary if seizures are not controlled within minutes of giving benzodiazepines, IV Sphenotoin or phenobarbital should be given in adequate doses. For resistant status epilepticus, close EEG monitoring and supervision by specialized medical and nursing personnel in intensive care unit are required. General anesthesia with IV barbiturates or other agents, neuromuscular blockade and respiratory support may be needed.
- Approximately half of the patients achieve complete control with standard antiepileptic agents and a significant proportion of the remaining patients achieve partial control.
- In medically-refractory cases, focal cortical resection or hemispherectomy can improve seizure control and may prevent cognitive decline.

### Headaches

About half of the patients complain of headaches. Migraine represents the most frequent type of headache. Hemiplegic headaches commonly occur after epileptic seizures. In a few patients headache may be due to glaucoma. Headaches should be managed as in other persons. However due to increased risk of focal neurologic deficits, vasoconstrictor agents such as triptans and ergots should probably be avoided. When a patient presents with severe headache for the first time, neurologic consultation should be obtained to establish a diagnosis. Neuroimaging and cerebrospinal fluid analysis should be performed if unexpected complications such as intracranial bleeding and infection are suspected.

- Cerebrospinal fluid analysis may exclude subarachnoid hemorrhage or central nervous system infection.
- Neuroimaging studies are indicated for recurring headaches that do not fulfill the strict definition of migraine or other primary headache disorder. For recurring migraine-type headaches, preventive therapy may be needed.

### Focal Neurologic Deficits

- **Hemiplegia** occurs in at least one-third of the cases and is localized to the side opposite to the facial Port-Wine birthmarks.
- Hemiplegia commonly first appears after an episode of seizure and may become more severe with the recurrence of seizures.
- Hemiplegia is present in majority of patients, either alone or in association with hemiparesis.
- Transient hemiplegias not following an epileptic attack and sometimes accompanied by migraine-like headaches are observed in many cases of Sturge-Weber syndrome. These hemiplegic episodes are apparently not epileptic nature and may be a consequence of vasomotor disturbances within or around the angioma.
- Intracranial hemorrhage is quite rare but has been documented in a few cases.
- When a child presents with acute focal neurologic deficit, determination of the cause is essential in guiding therapy.
- Low dose aspirin has been used by some physicians theoretically to prevent recurrent venous thromboses that are supposed to cause stroke-like episodes and focal deficits. However, lack of controlled clinical trials and the clinical arbitrariness of the syndrome make it impossible to determine whether aspirin is helpful.

### Ophthalmic Presentations

Ophthalmic abnormalities are part of the Sturge-Weber syndrome (SWS). The information below is designed to offer Emergency Room physicians insight into the common eye findings unique to SWS and the special ocular problems that patients with SWS may experience.

#### Common (Typical) SWS Eye Findings

- **Port-Wine birthmark involving the eyelids**
- **Differentiate from erythema by lack of tenderness, presence of sharply demarcated edges often respecting the midline, and lack of warmth to touch.**
- **Red eye (prominence of conjunctival and episcleral vessels).**
- **Differentiate from ocular inflammation by lack of discharge, lack of pain, clearly visible tiny vessel complexes, tortuous and telangiectatic in appearance and absence of symptoms.**
- **Retina and red reflex appearing redder than usual or than in the uninvolved contralateral eye.**
- **Due to choroidal hemangiomia (choroid lies underneath retina).**
- **Differentiate from retinal hemorrhagia by absence of distinct blood spots and appearance of diffuse redness.** Particularly obvious in unilateral SWS when compared to unaffected eye.

#### Common SWS Eye Problems

- Lid swelling which most often occurs on waking.
- Differentiate from cellulitis by history of long recurrent period, recurrence, absence of fever, absence of tenderness, lack of true erythema (see above), lack of warmth to touch, and resolution within hours with no treatment.
- **Glaucoma - present, especially in children less than 5 years old, with ipsilateral enlarged eye, photophobia, tearing, cloudy cornea, decreased red reflex, and enlarged optic nerve cup.**
- All signs and symptoms not required for diagnosis.
- Requires urgent referral to ophthalmologist if any suspicious signs or symptoms occur.
- **Glaucoma in older children usually present with no symptoms or signs except enlarged optic nerve cup but may have signs of LV pressure increases mentioned above.**
- **Intracocular pressure significantly higher than in the contralateral uninvolved eye.**
- Be sure to check vision in both eyes, with glasses on and/or with pinhole to get best corrected visual acuity measurement.
- **Visual field defects especially in children with seizures and those who have had hemispherectomy, most common is one side hemianopsia.**
- **Decreased peripheral vision common either due to brain involvement or glaucoma.**
- Retinal detachment and other complications from choroidal hemangiomia are very rare.
- **Refer to ophthalmologist if new discovery of decreased vision in one eye and absent or white (leukocoria) pupillary “red flux test”.**

#### Special Considerations

- All newborn babies with SWS and Port-Wine birthmarks affecting the eyelids and/or brain involvement should see an ophthalmologist in the first few weeks of life. Children with Port-Wine birthmark involving lids may be at life long risk of glaucoma and require periodic eye examination, often under anesthesia or sedation in the first years of life until able to be done awake.
- Any child who has had prior eye surgery and presents to the ER with a red eye (above baseline), pain, or any other ocular concern (including conjunctivitis) should be urgently referred to their ophthalmologist.
- **Children with glaucoma are often on multiple medications. In general:**
  - **Red Top Drops = pupil dilators (symphatomimetics or parasympathomimetics).**
  - **Green Top Drops = pupil constrictors (parasympathomimetics).**
  - **Purple Top Drops = alpha agonists.**
  - **Orange Top Drops = carbonic anhydrase inhibitors.**
  - Others: topical prostaglandin analogues, antibiotics, steroids and other anti-inflammatory drugs may be in use.
  - **May also be using oral steroids or oral carbonic anhydrase inhibitors for eye problems.**
- **Note that some bottles may contain more than one agent (combination preparation).**
- **Children with SWS on certain alpha agonists (e.g. brimonidine) may be more prone to sedative side effects that can be acute and dramatic. In infants or children with severe SWS, bradycardia, apnea, and hypothermia may also be seen even if only one drop given.**
- If a child has had a glaucoma drainage tube implant (also called seton or tube), there may be a visible whitish square at the edge of the cornea in one quadrant. This is tissue used to cover the plastic tube. If a child has had glaucoma surgery he or she may have a cystic appearing mass (bleb), either vascular or avascular, in one or more quadrants beyond the edge of the cornea. This is the expected area of fluid drainage.