Incontinentia Pigmenti

BS G,1 Pai GS,1 Pai AH,1 Vinekar AS,2 Pai HS,3 Noronha T,1 Fernandes MS1

1Department of Dermatology
K.S.Hegde Medical Academy
Deralakatte, Mangalore

2Pediatric Vitreoretina service
Narayana Nethralaya, Bangalore

3Alpha Eye centre, Mangalore
India

Corresponding Author
Girisha B.S
Department of Dermatology
K.S.Hegde Medical Academy
Deralakatte, Mangalore
India
Email: drgirishbs@gmail.com

Citation

ABSTRACT

Incontinentia pigmenti is an X-linked dominant condition characterized by cutaneous lesions associated with developmental defects of the eye, skeletal system and central nervous system. We report a case of incontinentia pigmenti in a 30 day old female infant who presented to us with skin eruptions over bilateral upper limbs, lower limbs and trunk since birth. She had linear verrucous plaques and vesicles distributed along the Blaschko’s lines in addition to macular hyperpigmentation in a linear and whorled pattern involving the concerned areas. On ophthalmological examination, proliferative retinopathy in the right eye was noted.

KEY WORDS
Incontinentia pigmenti, proliferative vitreoretinopathy

INTRODUCTION

Incontinentia Pigmenti (IP) also called Bloch-Sulzberger syndrome is a rare X-linked dominant genodermatosis affecting mostly female patients and is usually lethal to male fetus in utero. The gene affected is NEMO or IKK gamma gene located on Xq28. It is a multisystem disorder which can affect skin, teeth, eyes and central nervous system. The skin lesions can be accompanied by ocular, dental and neurological manifestations.1,2

CASE REPORT

A 30-day-old baby girl delivered at term of a non consanguineous marriage by a caesarian section was brought to our hospital with skin eruptions all over the body since birth. The indication for caesarian section was a previous caesarian section. The weight of the baby at birth was 3200gms. She had fluid filled lesions on the left arm at birth, which progressed to involve the left forearm, right arm, right forearm, trunk, lower legs and back subsequently within a few days as noted by her mother. These fluid filled lesions would flatten after a few days without rupturing and would crust. She had also noticed hyperpigmented linear streaks over the abdomen from birth. There were no history of any episodes of seizures, feeding difficulties or delay in milestones. Family history is unremarkable. Her four year old sibling was not affected. Mother had regular antenatal checkups. There was no history of any miscarriage. On examination of skin, hyperpigmented verrucous plaques were arranged linearly over the upper and lower extremities (Fig 1 and 2). A few vesicles were present over the upper limb and back. Reticulate hyperpigmentation was present over bilateral thighs. A few tense bullae were present over the soles (Fig 2). Face was spared. Whorled pattern of hyperpigmentation was present over the abdomen (Fig 3). Scalp, hair, nails and oral mucosa were normal. Systemic examination was within
normal limits: central nervous, respiratory, per abdomen
and cardiovascular systems were normal. Ophthalmological
examination showed proliferative vitreoretinopathy of the
right eye (Fig 4 and 5). Routine blood investigations were
within normal limits. Biopsy of a vesicle showed spongiosis
with numerous eosinophils in the epidermis with a few
dyskeratotic cells. Patient was referred to higher centre for
further evaluation and treatment of her retinal problem,
where she underwent laser photo coagulation.

DISCUSSION

Incontinentia pigmenti is a genodermatosis that can affect
skin, teeth, eyes and central nervous system. It has an
X-linked dominant inheritance. Clinical manifestations
vary from subtle dental and cutaneous findings to severe ophthalmological and neurological involvement.
Cutaneous involvement is present in nearly all the cases
and is often the first sign. Cutaneous features are divided
in to four stages. They are vesiculobullous stage, verrucous
stage, stage of pigmentation and stage of atrophy. These
stages can classically evolve in succession or may occur
currently.3

Our patient had overlapping features of vesiculobullous
stage, verrucous stage and stage of pigmentation when
presenting to us. She did not have any family history
of any skin and dental disorders. On her routine ocular

examination proliferative vitreous retinopathy was
detected. No neurological deficits were observed.

Ocular manifestations are seen in 25-77% patients with IP.1
They account for severe manifestations of IP and are often
associated with neurological deficits. Ocular anomalies in IP
can be subdivided into retinal and nonretinal manifestations.
The characteristic retinal manifestation of IP is proliferative
retinopathy that consists of peripheral retinal vascular
nonperfusion, preretinal neovascularisation, preretinal
or vitreous hemorrhage and infantile tractional retinal
detachment.1,4 Demonstrable retinal lesions may occur
from neonatal period up to one year of age. The nonretinal
manifestations include strabismus, crossed eyes, optic
nerve atrophy, conjunctival pigmentation, iris hypoplasia,
nystagmus and uveitis.1

Prognosis for normal vision is considered good if there is no
evidence of retinal lesions in first year of life. However the
course of established lesions is unpredictable. It ranges from
slow progression over many years to rapid deterioration
and blindness.1 Hence ophthalmic examination should
form an integral part of routine examination of patients
with IP. Abnormalities if detected in the neonatal period
can be promptly treated and blindness can be prevented.

In our patient the proliferative vitreoretinopathy of the
right eye was detected at a very early stage and was
referred to a tertiary eye care centre where it was treated
Case Note

by laser photocoagulation. She is also on regular follow up to recognise progression of her retinal lesions as well as neurological deficits. Early detection and treatment of retinal lesions in incontinentia pigmenti can preserve vision.

REFERENCES


