Introduction

Xeroderma pigmentosum, or XP, is an autosomal recessive genetic disorder of DNA repair in which the ability to repair damage caused by ultraviolet (UV) light is deficient. In extreme cases, all exposure to sunlight must be forbidden, no matter how small; as such, individuals with the disease are often colloquially referred to as *Children of the Night*.

Multiple basal cell carcinomas (basaliomas) and other skin malignancies frequently occur at a young age in those with XP. In fact, metastatic malignant melanoma and squamous cell carcinoma are the two most common causes of death in individuals with XP. This disease involves both sexes and all races.

Xeroderma Pigmentosum (XP)

Xeroderma pigmentosum, which is commonly known as XP, is an inherited condition characterised by an extreme sensitivity to ultraviolet (UV) rays from sunlight. This condition mostly affects the eyes and areas of skin exposed to the sun. Some affected individuals also have problems involving the nervous system.

The signs of XP usually appear in infancy or early childhood. Many affected children develop a severe sunburn after spending just a few minutes in the sun. The sunburn causes redness and blistering that can last for weeks. Other affected children do not get sunburned with minimal sun exposure, but instead tan normally.
By age 2, almost all children with XP develop freckling of the skin in sun-exposed areas (such as the face, arms, and lips); this type of freckling rarely occurs in young children without the disorder. In affected individuals, exposure to sunlight often causes dry skin (xeroderma) and changes in skin colouring (pigmentation). This combination of features gives the condition its name, Xeroderma Pigmentosum.

People with XP have a greatly increased risk of developing skin cancer. Without sun protection, about half of children with this condition develop their first skin cancer by age 10. Most people with XP develop multiple skin cancers during their lifetime. These cancers occur most often on the face, lips, and eyelids. Cancer can also develop on the scalp, in the eyes, and on the tip of the tongue.

Studies suggest that people with XP may also have an increased risk of other types of cancer, including brain tumours. Additionally, affected individuals who smoke cigarettes have a significantly increased risk of lung cancer.

The eyes of people with XP may be painfully sensitive to UV rays from the sun. If the eyes are not protected from the sun, they may become bloodshot and irritated, and the clear front covering of the eyes (the cornea) may become cloudy. In some people, the eyelashes fall out and the eyelids may be thin and turn abnormally inward or outward. In addition to an increased risk of eye cancer, XP is associated with noncancerous growths on the eye. Many of these eye abnormalities can impair vision.

About 30 percent of people with XP develop progressive neurological abnormalities in addition to problems involving the skin and eyes. These abnormalities can include hearing loss, poor coordination, difficulty walking, movement problems, loss of intellectual function, difficulty swallowing and talking, and seizures. When these neurological problems occur, they tend to worsen with time.

Researchers have identified at least eight inherited forms of XP: complementation group A (XP-A) through complementation group G (XP-G) plus a variant type (XP-V). The types are distinguished by their genetic cause. All of the types increase skin cancer risk, although some are more likely than others to be associated with neurological abnormalities.

Incidence of Xeroderma Pigmentosum in South Africa
The National Cancer Registry (2014) does not provide any information regarding the incidence of Xeroderma Pigmentosum in South Africa.

Causes of Xeroderma Pigmentosum (XP)
Xeroderma pigmentosum is an autosomally recessive inherited disease, which means that one must inherit two recessive XP genes (one from each parent). If one’s parents are only carriers of the XP trait (each have one XP gene and one normal gene), they will not show signs or symptoms of the disease. By having the two XP genes this causes one to have an extreme sensitivity to UV light and as
a result experience a range of signs and symptoms of XP. At least eight different gene abnormalities or complementation groups have been described in different families (XPA to XPG) resulting in varying disease severity.

Essentially, the signs and symptoms of XP are a result of an impaired DNA repair system. In people who do not have XP, cell damage from UV light is mended by the DNA repair system. However, people with XP have a defect in this repair system and any damaged cells from UV light remain unrepaired, leading to cancerous cells or cell death.

**Symptoms of Xeroderma Pigmentosum (XP)**
The following symptoms may appear:

- Sunburn that does not heal after just a little bit of sun exposure
- Blistering after just a little bit of sun exposure
- Spider-like blood vessels under the skin
- Patches of discoloured skin that get worse
- Crusting of the skin
- Scaling of the skin
- Oozing raw skin surface
- Discomfort when being in bright light (photophobia)
- Skin cancer

**Differential Diagnosis**
There are other causes of photosensitivity, e.g., congenital erythropoietic porphyria.

Other genetic conditions with photosensitivity due to defective DNA repair - eg, Cockayne’s syndrome, the XP-CS complex, trichothiodystrophy (TTD), the XP-TTD complex, cerebro-oculo-facio-skeletal (COFS) syndrome and the UV-sensitive syndrome. (Patient Info).

**Treatment of Xeroderma Pigmentosum (XP)**
Children with this condition need total protection from sunlight - even the light coming through windows or from fluorescent bulbs is dangerous.

When these children go out in the sun, they need to wear special protective clothing.

Use high SPF protection sunscreen and very dark, UV sunglasses. The doctor may prescribe medicine to help prevent certain precancerous growths from becoming skin cancers.
Many patients with Xeroderma Pigmentosum may die at an early age from skin cancers. However, if a person is diagnosed early, does not have severe neurological symptoms or has a mild variant, and takes all the precautionary measures to avoid exposure to UV light, they may survive beyond middle age.

Patients with Xeroderma Pigmentosum and their families may face many challenges in daily living. Constant educating and reminding of the need to protect oneself from sunlight is paramount to the management of Xeroderma Pigmentosum.

Genetic Counselling and Risk to Relatives
Inheritance of Xeroderma Pigmentosum (XP) is autosomal recessive. If parents are considering further pregnancies, prenatal diagnosis is often possible. Where XP is suspected, siblings should be protected from UV light until XP can be excluded.

Recent investigations of heterozygotes with one of four XP genes (XPA, XPC, ERCC2, or ERCC5) have reported an increased risk of skin cancer, lung cancer, or altered response to certain chemotherapeutic agents.

The South African Xeroderma Pigmentosum Society Contact Details
The XERODERMA PIGMENTOSUM SOCIETY is a Non Profit Company incorporated in South Africa on January 13, 2010. Their business is recorded as In Business. The activity is registered as NON PROFIT SUPPORT GROUP TO RELIEVE THE NEEDS OF PERSONS WITH XERODERMA PIGMENTOSUM (XP) AND RELATED CONDITIONS AS WELL AS FAMILY SUPPORT. It is not part of a group. The company was incorporated 6 years ago.

Xeroderma Pigmentosum Society

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Sources and References Consulted or Utilised

DermNet NZ
http://www.dermnetnz.org/systemic/xeroderma-pigmentosum.html

Genetics Home Reference

Medline Plus

Patient Info
https://patient.info/doctor/xeroderma-pigmentosum-pro

Stage III XP
http://www.dermrounds.com/photo/xeroderma-pigmentosum-3rd-1?context=album&albumid=1980062%3AAlbum%3A8589

Wikipedia
https://en.wikipedia.org/wiki/Xeroderma_pigmentosum

Xeroderma Pigmentosum Boy

XP
https://www.google.co.za/search?q=Xeroderma+Pigmentosum&espv=2&biw=1517&bih=714&source=lnms&tbm=isch&sax=KNGDVYmG8aGR7aajpD4&ved=0CAAYUoAQ&amp;dpr=0.9#imgdii=TPCITPMX4NXWIFM%3A%3BTPCITPMX4NXWIM%3A%3BTPCITPMX4NXWIFM%3A%3BPeoNtnmVUwIbyM%3A&imgc=TPCITPMX4NXWIFM%3A%3BPeoNtnmVUwIbyM%3A&imgrc=TPCITPMX4NXWIFM%3A%3BPeoNtnmVUwIbyM%3A&imgrc=TPCITPMX4NXWIFM%3A%3BPeoNtnmVUwIbyM%3A&imgrc=TPCITPMX4NXWIFM%3A%3BPeoNtnmVUwIbyM%3A