Xeroderma Pigmentosum-A Rare Genodermatosis: Overview of Literature

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Abstract

Xeroderma pigmentosum is a rare genodermatosis, autosomal recessive in nature in which excessive ultraviolet radiation causes skin, ocular, neurological, and oral lesions along with development of cutaneous and internal malignancies at an early age. There is no definitive cure for the disease. Avoidance of ultraviolet radiation, use of protective clothing, sunscreens, oral retinoids, 5-fluorouracil and regular consultations with dermatologists, ophthalmologists, neurologists and dentists forms an important part of the treatment protocol. This paper aims to throw light on the etiopathogenesis, clinical features and treatment modalities of this life threatening disease. There is also a special mention on the oral manifestations and dental health considerations of the rare disorder.

Keywords: Xeroderma pigmentosum; Ultraviolet radiation; Cutaneous lesions; Dental health considerations

Introduction

Xeroderma pigmentosum (XP) is a rare genetic autosomal recessive disease marked by extreme photosensitivity, hyperpigmentation and premature ageing of the skin, along with the development of cutaneous and internal malignancies at an early age. There is a defect in the nucleotide excision repair (NER) mechanism which results in a defective repair of DNA damaged by ultraviolet (UV) radiation [1]. XP is widespread in all continents and all racial groups are affected. As the disorder has an autosomal recessive inheritance, both males and females are affected. The reported incidence rate (estimates made in the 1970’s) was 1:250,000 [2] and 1:20,000 in the USA and Japan respectively, with a rare occurrence in India [3]. Recent survey analysis in Western Europe suggested an incidence of approximately 2.3 per million live births [4]. As consanguinity is highly prevalent in North Africa and the Middle East, the incidence rate of XP in these areas is substantially higher. The disease is categorized into eight forms based on the clinical aspects and the patterns of the molecular defect [5]. The diagnosis of XP can be established by studies performed in specialized laboratories. These studies include cellular hypersensitivity to UV radiation and chromosomal breakage studies, complementation studies, and gene sequencing to identify the specific gene complementation group [6]. Early and rigorous photoprotection with sun avoidance, sunscreens and appropriate clothing forms a part of treatment plan. DNA repair enzymes and oral retinoids are currently under clinical trials and show a hope for the future [7,8].

Discussion

Xeroderma pigmentosum (XP) is a rare autosomal recessive, progressive, degenerative disease which is related to photosensitivity, cutaneous pigmentation, abnormal DNA repair, various neoplasms and occasionally, neurologic degeneration [9].

Xeroderma pigmentosa (XP) was described by Dr. Moriz Kaposi in 1870, a dermatologist in Vienna [10]. The term “xeroderma or parchment skin” was initially described and in 1882, the term “pigmentosum” was added to emphasize the striking pigmentary abnormality [11].

Etiopathogenesis

Exposure to UV radiation

Ultraviolet (UV) irradiation is composed of UV A spectrum and UV B spectrum, where UVB plays an important role in the etiology of XP. UV irradiation causes photoproducts in DNA, chiefly cyclobutane pyrimidine dimers (CPD) and 6-4-pyrimidine-4-pyrimidone, which further brings about cell death, mutagenesis, carcinogenesis and cellular ageing [12]. XP is an autosomal recessive disorder which results from mutations in any of the eight genes. These genes restore the DNA damage induced by UV radiation by a process known as nucleotide excision repair (NER) [13]. XP patients have mutations in one or more NER genes, and cause molecular defects in cellular DNA repair mechanisms and hypersensitivity to UV radiation. As a result, the accumulation of unrepaired UV induced DNA damage occurs which either facilitate cell death, contributing to accelerated skin ageing, or cellular transformation resulting in the development of malignancies [14-16].

Many XP patients with tumors show mutations in the p53 gene, indicating that p53 mutations are characteristic of UV exposure [17].

Consanguinity

Consanguinity has been implicated as an etiological factor. This has been reported to varying degrees of up to 92.8% in XP patients in Libya [18]. Other studies reported from Egypt, Pakistan, and Nigeria etc. have a high incidence of XP [19]. However, a case report by Stephanie Christen et al. showed a family where the uncle and nephew were affected by XP [20].

Drugs and Chemicals

A number of DNA-damaging agents apart from UV radiation have been implicated to cause a hypersensitive response to XP cell [21] (Table 1).

Clinical Features

Most clinical features are subject to amount of exposure to sunlight, the complementation group, the mutation, and some unknown factors.

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Hence XP patients present with varied clinical manifestations. Factors aggravating the skin lesions include- Sunny weather, outdoor living, fair skin, smoking, poor accessibility of diagnostic amenities, resulting in delayed diagnosis, and improper shield from sunlight. These factors ultimately cause cutaneous pigmentary changes, cutaneous cancers, and early death [22].

**Cutaneous lesions**

Extreme photosensitivity is usually the first sign in about 60% of cases [23]. Lentigines (freckle-like pigmentation) in sun-exposed areas are seen in remaining 40% XP patients as early as two years of age. These pigmented areas are seen on the nose, zygoma, and forehead and followed by the sides of the neck, sparing the area under the chin [22] (Figure 1). Photophobia is often present [24].

Lack of sunlight avoidance causes skin ageing, making it dry, rough, and atrophic. With subsequent sun exposure, lentigines increase in number and darken. They then become difficult to distinguish clinically from flat, pigmented seborrhoeic warts, which also proliferate and become warty with time. Small, hypo-pigmented macules are commonly seen amongst the lentigines and give rise to the characteristic mottled hyper-pigmented and hypo-pigmented appearance known as salt and pepper pattern of skin [21,25] (Figure 2). Atrophic, hypo-pigmented patch is often seen on the skin of the nose in these patients. Telangiectasia can be a late manifestation.

In the general population, Dark-skinned individuals usually have a lower incidence of skin cancer compared to light skinned people. This most probably occurs due to the photo protective properties of melanin [26].

Excessive sun exposure usually results in areas of hyper- and hypopigmentation, followed by skin ageing, warty growth, melanocyte and keratinocyte malignancy, and ultimately multiple basal cell carcinomas and invasive squamous cell carcinomas and melanomas [22,23]. The risk of cutaneous basal cell or squamous cell carcinoma or melanoma is increased to a 1000-fold in XP patients under 20 years of age [21,27]. In XP, the median age of onset of non-melanoma skin cancer was 8 years, in contrast to 60 years in the general population [28]. Usually, patients with the most severe repair defects would show the most extreme sunburn reactions and the highest incidence of skin cancer. Paradoxically, patients with acute sunburn reactions develop fewer skin cancers than those who do not. An early diagnosis of the former group and a disinclination of this group to go out in sunlight without protection is likely to play an important role.

**Ocular lesions**

Approximately 40% of XP patients present with ocular lesions like photophobia and blepharoconjunctivitis, which are the most common. Erythema, pigmentation, atrophy, malignant change, telangiectasias, loss of lashes, and chronic blepharitis are less common ocular lesions [29]. Ectropion of the lower eyelid and symblepharon are some symptoms that may occur due to atrophic scarred skin [30]. Concomitant opacification, neovascularization, pterygia, and band keratopathy are common, and bandshaped nodular dystrophy and squamous cell carcinomas have also been reported. Keratitis, edema, corneal ulceration and perforation may occur due to lower lid loss [31]. Conjunctival involvement may result in conjunctivitis, pinguecula, symblepharon, melanosis, and tumors developing from the interpalpebral zone of the limbus [29]. Squamous cell carcinomas, malignant melanomas and limbal stem cell deficiency have been often reported. Cases of iritis, stromal atrophy, pigment abnormalities, and, occasionally melanoma have been reported [32]. Fundus abnormalities are less frequently seen because the posterior segment is protected from UV damage by the cornea and lens [33].

**Neurological lesions**

Neurological problems are often seen in approximately 20-30% of the patients. The onset of neurological symptoms usually begins at the

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**Table 1: Drugs and chemicals.**

<table>
<thead>
<tr>
<th>Drugs</th>
<th>Carcinogens</th>
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<tbody>
<tr>
<td>1. Psoralens plus long wavelength UV radiation (PUVA)</td>
<td>1. Plant toxin-Aflatoxin</td>
</tr>
<tr>
<td>2. Chlorpromazine</td>
<td>2. Alkylation agents-benzo (a) pyrene</td>
</tr>
<tr>
<td>5. Anthramycin</td>
<td>5. Phenanthrene derivatives</td>
</tr>
<tr>
<td>6. Cisplatin and Carmustine</td>
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**Figure 1:** Hyper pigmented macules interspersed with patchy hypopigmentation over face, ear, eyes & forehead.

**Figure 2:** Rough, dry, scaly skin with hyperpigmented macules over the truncal area, back, hands and feet.
age of two and may occur up to middle age [23,26]. The neurological abnormalities include isolated hypo-reflexia, mental retardation, sensorineural deafness, spasticity, or seizures [34].

Loss of high-frequency hearing, is the most common neurological abnormality while the most severe neurological deficits are seen in DeSanctis-Cacchione syndrome [21,35]. The incidence of central nervous system tumors (CNS) is also ten times higher than in the normal population. Some common CNS tumors include, astrocytomas, medulloblastomas, glioblastomas, and malignant schwannoma [31].

Oral lesions

Oral manifestations are rarely seen and only about 4% patients exhibit these symptoms [27]. Some common oral manifestations associated with XP are leukoplakia, erythroplakia, Actinic cheilitis and SCC of the tip of the tongue and lips [25]. The precancerous and cancerous lesions of the tip of the tongue are sites that are seldom affected in the normal population and are likely to be induced by UV radiation [24,26,36]. SCC of the tip of the tongue in XP patients usually occurs in individuals younger than 20 years of age and progresses slowly [26]. Actinic cheilitis is a potentially malignant lesion that affects the lower lip of white patients who are frequently exposed to the sun. Multiple labial plasty result in areas of fibrosis. Stretching of these fibrous areas causes pain when the patient opens the mouth for eating, speaking, breathing, and oral hygiene procedures. These patients usually have poor oral hygiene habits and a high rate of dental plaque, caries, and periodontal disease [37]. Cases of chronic desquamative gingivitis, fissured tongue, geographic tongue and keratoacanthoma have also been reported [21,38] (Figures 3-5).

Treatment

Excessive sun exposure results in irreversible skin damage in XP patients. Treatment protocol aims at minimizing the sun exposure and may include avoidance of being outdoors and use of sunscreen and dark ointments. Avoiding sun exposure for long duration may alter the calcium ion concentration and affect bone growth and development in infants and young children. Dietary supplementation with vitamin D, vitamin A and nicotinamide or zinc sulfate may minimize these side effects [39]. Prophylactic treatment with anticancer drugs, including isotretinoin or fluorouracil can reduce the incidence of skin cancer in XP [40]. Enzymatic treatment and gene therapy have shown good effects [39,41,42]. Surgical resection is effective in patients who have developed tumors.

Dental considerations

Important dental health considerations in XP patients include-

1. Use of a UV light meter. The patient may be exposed to a variety of light sources in the dental office, including overhead lights, view boxes, dental lamps, fiber optic lights, computer screens and dental curing units. If the UV light meter shows a reading above 0 nm/cm² for UV light, the use of that unit should be contraindicated.

Although light-emitting diode curing lights emit a wavelength of only 450 nm, [43] the degree of biological damage resulting from such exposure is not sufficiently understood. Restorative materials such as glass ionomers offer a good substitute to resin sealants and composite restorations for treating XP patients [43]. Curing light filters may become ineffective or worn out over time, and allow transmittance of wavelengths in the UVB range (290-320 nm), which are known to have substantial carcinogenic effects on normal epithelium [44,45]. A regular clinical examination is usually a must for timely detection of premalignant or malignant lesions. Furthermore, establishment of protocols for prophylaxis and topical fluoride application, as well as the use of chlorhexidine digluconate 0.12%, aims at the homeostasis of the oral environment. Mouthwashes with high alcohol concentration are usually avoided as this is associated with an increased risk of oral cancer development in these patients [46]. Regular dental procedures, such as dental extraction, restoration and rehabilitation pose a challenge to the dentist due to difficulty in accessing the oral cavity.

Periodic oral examinations are important and usually include complete inspection and palpation of soft tissues (lips, tongue and oral mucosa). Radiation therapy is used to treat the head and neck tumors and are known to cause radiation induced caries and permanent damage to salivary glands. Meticulous oral hygiene procedures such as proper brushing, flossing and fluoride rinses are usually advised for XP patients, as these patients do not maintain routine dental appointments [43,47].

Conclusion

Xeroderma pigmentosum is a autosomal recessive disorder characterized by UV induced damage to the skin, eyes and nervous system tumors (CNS) is also ten times higher than in the normal population. Some common CNS tumors include, astrocytomas, medulloblastomas, glioblastomas, and malignant schwannoma [31].
tissue. Oral lesions are rarely associated with the disorder and include leukoplakia, erythroplakia and carcinomas. There is no cure for the disease, however, clinical trials are in progress to understand the etiopathogenesis and the development of an effective treatment protocol.

References
