THE CORNEA IN MEASLES

PROEFSCHRIFT

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To Cora,

Oscar, Reinout, Muriel,

Nico-muti and all his little
African brothers and sisters
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1 - Introduction

1.1. Blinding eye complications after measles: “Post-Measles-Blindness”

In several developing countries blindness after measles is a considerable public health problem. Many patients, who have lost useful vision through leukomata, adhaerent leukomata or phthisis bulbi, relate this with an episode of measles when they were young. For example, in a survey in Western Kenya of two Schools for the Blind, I found a prevalence of 30% for corneal blindness, and the majority of these children blamed the measles. (unpublished data; cfr Sauter 1976).

It seemed unjustified to me, to connect the corneal blindness after measles with a single, specific pathogenetic mechanism, e.g. Vitamin A dependant keratomalacia. For this reason the more comprehensive term “Post-Measles-Blindness” (PMB) is preferred. This indicates the relation in time between measles and the subsequent blindness, and permits study of the problem without prejudice.

In addition to a high prevalence in blindness statistics (§ 2.7) a high incidence of PMB is currently reported in studies of measles from developing countries. Kimati and Lyaru (1976) reported 5 cases with corneal involvement leading to blindness out of 624 patients with measles, admitted to the Mwanza Regional Hospital, Tanzania. Ophthalmological details are not available. 44 Out of 2,376 (1.9%) measles patients, reported by Morley, Martin and Allen (1967) in East-Africa, developed permanent ocular damage. In this study ocular damage was associated with a high mortality (cfr Animashaun 1977).

Similar reports are published from West-Africa: 31 out of 2,164 patients (1.4%) with destruction of one or both eyes (Morley 1969) and 27 cases of corneal blindness out of 2,772 measles-patients (1.0%) in Lagos, Nigeria (Animashaun 1977).

These statistics make it likely, that in at least some African countries in development, somewhere around 1% of all children with measles will sustain permanent, severe ocular damage of corneal origin.

Ocular complications in measles are not necessarily localized in the cornea. Retrobulbar neuritis (Srivastava and Nema 1963) and retinitis (Bücklers 1969; Haydn 1970; Regensburg and Henkes 1976) are described, but in developing countries these are rare compared to the corneal complications. In this study only the corneal involvement with measles will be taken into account.

1.2. Pathogenesis of Post-Measles-Blindness

In the literature on PMB 3 possible pathogenetic pathways can be identified: infection, malnutrition and treatment. In chapter 2 a detailed review of the literature will be given. In this section only the outlines will be given, as far as they are needed to define the purpose of this study.
a. Infection

Occasionally a coarse punctate keratitis is mentioned as a sign of measles (Trantas 1903; Thygeson 1959); severe corneal damage, however, is seldom attributed to the measles virus itself: Frederique, Howard and Boniuk (1969) are the exception.

A herpetic keratitis was seen early in the measles infection (Sauter 1976) and Whittle et al. (1979) were able to culture herpes simplex virus from corneal ulcers after measles.

An unspecified bacterial superinfection is, however, considered as a more important possibility by many authors (Gaud 1958; Armengaud et al. 1961; Quené 1964; Benezra and Chirambo 1977).

No report on fungal infections of the cornea in connection with PMB was found.

b. Malnutrition

Protein Energy Malnutrition (PEM) and morbidity from measles go hand in hand: measles runs a more severe course in malnourished children (Scheifele and Forbes 1973) and overt kwashiorkor is a frequent complication of measles (Kimati and Lyaruu 1976; Alleyne et al. 1977). The deep disturbance of the protein metabolism in PEM might in itself be a reason for the necrosis of the cornea (Moore 1957; McLaren 1963; Kuming and Politzer 1967).

Most attention however has been given to the disturbance of Vitamin A metabolism in measles. The intake of Vitamin A and its precursors is reduced with grossly reduced food intake. The absorption is diminished because of diarrhoea. The serum levels of retinol and Retinol Binding Protein (RBP) are lowered because of infection and fever (Moore 1957; Morley, Woodland and Martin 1963; Arroyave and Calção 1979; Axton 1979). This reduced availability of Vitamin A, alone, or in combination with PEM, is held responsible for a Vitamin A dependant keratomalacia in the wake of measles (Oomen, McLaren and Escapini 1964). In this view, measles is only the trigger to produce a manifest keratomalacia in a previously only marginally deficient child.

c. Treatment

In the ophthalmological literature originating in Africa much attention is given to the possible role of traditional medicines. Vivid descriptions of their use can be found (Phillips 1961).

They are widely in use in Africa (Ayanru 1974; Kokwaro 1976; Chirambo and Benezra 1976; Maina-Ahlberg 1979). Some of the vegetable materials in use as ocular medicines can cause corneal damage. (Crowder and Sexton 1964; Cordero-Moreno 1973). The significance attached to their use is controversial and varies from "nonsense" (McManus 1968) to "the most important" cause of Post-Measles-Blindness (Phillips 1961).
In tropical countries childhood morbidity and mortality are considerable. Measles is not the only disease severely afflicting the under-fives in underprivileged conditions. It shares its bad reputation with malaria, pneumonia, gastro-enteritis, whooping cough and tuberculosis. But still, of these diseases measles especially is extremely frequently mentioned as a factor connected with corneal blindness, originating in childhood.

This clinical observation suggests that the cornea is intrinsically involved with measles: measles might directly affect the cornea.

The previously mentioned pathogenetic pathways take this connection insufficiently into account and the question remains: why does Post-Measles-Blindness occur? To answer this question it is first of all necessary to know what happens to the cornea in the acute stage of measles.

1.3. Lack of knowledge about the cornea in early measles

The question about corneal involvement in measles may look selfevident, but remarkably little attention has been paid to this subject: the literature is very scanty and gives only few ophthalmological details.

In 1903 Trantas (Constantinople) describes for the first time a punctate keratitis as a sign of measles. The existence of this keratitis is — among others — confirmed by Thygeson (1959), but very few detailed descriptions are to be found.

Even less attention has been given to the early corneal complications of measles.

Also in the limited amount of literature on the corneal signs of the acute stage of measles no connection is made with possibly blinding complications. The statement of Quéré (1964) “Ophthalmology has neglected measles” still holds true.

1.4 Purpose and content of the present study

The involvement of the cornea in the acute stage of measles is the subject of the present study. The best study on the measles-keratitis now available is still the one by Trantas in 1903. It seems worthwhile therefore to study this self-limiting keratitis with the investigative tools now available. The attention paid to this keratitis is above all warranted by the possible occurrence of blinding complications: the existence of Post-Measles-Blindness (1% in developing countries) is the incentive for this study.

It is hoped that by the study of the early corneal signs of measles, some data relevant for the pathogenesis of PMB can be obtained, which hopefully might have implications for the prevention of PMB.

In the background of every infection, the nutritional status of the child is the most important, not to say the decisive factor for the final outcome of the disease. It is even said that “a community is malnourished so long as children
die of measles” (King et al. 1972). In the literature PMB is also invariably connected with Protein Energy Malnutrition (P.E.M.) (Rodger 1959; Oomen, McLaren and Escapini 1964). Complications are associated with PEM, but the possibility exists that also the severity of physical signs is influenced by the nutritional status (O’Donovan 1971; Dossetor, Whittle and Greenwood 1977). In this study much attention is therefore given to the possible association of corneal disease with the nutritional status.

In the literature on measles the terms signs, symptoms and complications are used indiscriminately to describe anything that is observed on the cornea. For the purpose of this study physical signs and symptoms are defined as any phenomenon related to the multiplication of the measles virus and the normal reaction of the healthy body against this viral invasion. Anything beyond this interaction constitutes a complication. According to these definitions the Koplik’s spots and the rash are signs, buccal ulcers are complications.

In chapter 4 a detailed description of the conjunctival and corneal signs of measles will be given and in chapter 6 it will be demonstrated that these signs are independent of the nutritional status of the affected children.

It is, however, in no way the purpose of this study to deny the association of blinding complications after measles with the nutritional status (i.e. Protein Energy Malnutrition and/or Vitamin A Deficiency). Nor will it be possible to derive from this study any conclusion — positive or negative — about the prevalence of Vitamin A Deficiency in the population. The subject of this study is measles.

This study was mainly done in a 150 bed rural mission hospital in Western Kenya. Here in a two year period 248 children, acutely ill with measles were followed by daily slitlamp examination. They all had an anthropometric assessment of their nutritional status. Serum samples were analysed in Nairobi (400 km away); conjunctival specimens for immunofluorescence and electron-microscopy were sent to The Netherlands.

A rural mission hospital and a governmental Provincial Hospital are hardly equipped for research purposes. But, because of the motivation for this study (PMB in under-privileged conditions) it had to be done somewhere in the bush. This gave much difficulty in getting things organised, and done. This work would never had been carried out if I hadn’t had the enthusiastic support and cooperation of so many others. I owe them many thanks.
2 – Review of literature

2.1. The epidemiology of measles

In developing countries measles is a public health problem of considerable importance, because of its high prevalence in combination with a high mortality rate.

In Kenya measles is an endemic disease, with an epidemic outbreak every 2(-3) years (cfr Morley, Woodland and Martin (Nigeria) 1963; Voorhoeve et al. 1977) Nearly all children get measles, and, compared to Europe, at a relatively early age (8–36 months). This is probably caused by the overcrowding of the African households, resulting in a higher infection load at this early age (Leary and Obst 1966).

The mortality for measles is high. In hospital statistics the case fatality rate is 8.6% (Bwibo 1970) to 25% (Morley 1969) and is usually presented to be about 15–20%. In field studies however the mortality rate is understandably lower: 6.8% in Nigeria (Morley, Woodland and Martin 1963), 6.5% in a Kenyan study (Voorhoeve et al. 1977). The West African saying “count your children after the measles” needs no further explanation.

The mortality can vary considerably in different epidemics (Muller et al. 1977). The lower age of infection is accompanied by a higher mortality (Bwibo 1970; O’Donovan 1971; Cruickshank, Standard and Russell 1976) and is highest in the age-group of 1–2 years (Voorhoeve et al. 1977). According to many studies this higher mortality could be caused by the higher frequency of undernutrition in the lower age groups (Leary and Obst 1966; Morley, Martin and Allen 1967; O’Donovan 1971; Hendrickse 1975; Dossetor, Whittle and Greenwood 1977; Muller et al. 1977).

Also the morbidity from measles is considerable: the accompanying stomatitis, laryngotracheitis and gastroenteritis may be severe, whereas bronchopneumonia, otitis media, and Protein Energy Malnutrition are frequent complications (Leary and Obst 1966; Bwibo 1970; Kimati and Lyaruu 1976). It has been mentioned already in §1.1. that 1% of the measles children sustain permanent ocular damage.

Early this century the same situation existed in Western Europe. For example, in 1908 in Glasgow 5.8% of the under-fives infected with measles died (Morley, Woodland and Martin 1963). In Europe measles has now become a benign disease (in Great Britain the case fatality rate is 1:10,000 (Cruickshank, Standard and Russell, 1976) because of the socio-economic development, rather than the introduction of modern medicine. The differences in the epidemiology of measles, in Europe and developing countries, can be explained on non-geographical grounds. These differences are associated with socio-economic and cultural differences, not with different climates or different strains of the measles-virus. There are therefore no reasons to consider “Tropical measles” a separate entity. The use of this
term is to be dropped (cfr Cruickshank, Standard and Russell 1976; Muller et al. 1977; Benezra and Chirambo 1978).

2.2. The extra- and intracellular morphology of the measles virus

The measles virus belongs — together with the viruses of rinderpest and canine distemper — to the group of pseudomyxoviruses. They share a common morphology and some cross-immune reactions (Wilterdink 1979).

The virion measures 120–300 nm. (1 nanometer = $10^{-9}$ m) on electron-microscopical examination, but only 100 nm. in filtration experiments. It is polymorph in shape and occasionally anomalous and long filamentous forms can be seen.

Pseudomyxoviruses have a lipoprotein envelope, on which numerous spikes, 10 nm. in length, are visible. The envelope itself has also a thickness of 10 nm. (fig. 2.2.a). A unit of nucleocapsid of measles virus consists of RNA combined with a protective protein. About 2000 of these units are wound into a single helix, where the RNA as the carrier of genetic material is surrounded and protected by the protein. The diameter of this helix is 18 nm., the pitch of the helix is 4.5 nm. (fig. 2.2.b).

The overall length in electronmicroscopical specimens, stained with negative contrast, is approximately 1000 nm. Inside the lipoprotein-envelope the nucleocapsid helix is haphazardly folded. (Hall and Martin 1975; Wilterdink 1979).

The measles virion makes contact with the receptor sites at the surface of

Figure 2.2.a. The morphology of the complete virion, the appearance of the extra-cellular measles-virus
the host cell. The envelope merges with the cell membrane and the viral nucleocapsid is taken into the cytoplasm. With the use of the metabolism of the host cell, RNA re-duplication commences with the formation of a viral messenger-RNA. Later on newly formed RNA is combined with the protecting proteins to form the helical nucleocapsid. These nucleocapsids are located in inclusion bodies, visible 16–20 hours after the infection in the cytoplasm, 96–120 hours after the infection inside the nucleus (Morgan and Rapp 1977).

The newly formed nucleocapsid induces a change in the cellular membrane, visible as a higher density on electronmicroscopic examination. In a "budding process" the altered cellular membrane, with newly induced haemadsorption properties, engulfs the nucleocapsid and subsequently forms the envelope. The now complete virions are set free into the intercellular fluids, and continue on their way to the next cells (Nakai and Imagawa 1969; Wilterdink 1979).

The virion is thus the extracellular, the nucleocapsid helix the intracellular morphological appearance of the measles virus.

2.3. The pathogenesis of the measles infection

The first contact with the highly infective virus is at the mucous membrane of the respiratory tract. Also the conjunctiva might act as a portal of entry for the measles infection (Papp 1954). If not inactivated by mucus or specific secretory IgA antibodies (Dawson 1976) the virus enters the ciliated columnar epithelium and a small focus is located in the oro-pharynx, where it is hardly ever detected. Where the conjunctiva is the portal of entry, a "conjunctivitis of infection" may be the clinical result. During the primary viraemia, 2–6 days after the infection, the virus is transported intracellularly inside the formed elements of the blood. Macrophages withdraw most of the virus and
debris of infected cells from the blood, and an extensive proliferation of virus follows in the reticulo-endothelial system in the tonsils, spleen, liver, bone-marrow and other lymphoid tissues. The second viraemia starts 10 days after the infection, with proliferation of the virus inside the leucocytes. Neutralizing antibodies appear 14 days after the infection, at the time of appearance of the rash. The rash is the expression of immunological defense: in cases with severely impaired cell-mediated immunity a measles-infection may run its course without a rash (Burnet 1968; Scheifele and Forbes 1973; cfr Cruicksrank et al. 1974; Morgan and Rapp 1977; Wilterdink 1979).

The formation of lymphoid giant cells (Finkeldey-Warthin cells) is a characteristic cytopathogenetic effect of measles virus. The demonstration of their presence in tissues or secretions can be a valuable diagnostic sign of prodromal measles. They are however present in only half of all measles cases (Roberts and Bain 1958).

The virus is eliminated by humoral and cell-mediated immune responses: in most cases it will not be possible to culture the virus from the blood later than 36 hours after the outbreak of the rash.

The conjunctiva is involved in this pathogenetic process in two ways: as a possible portal of entry for the infection and the development of a characteristic conjunctivitis in the prodromal stage of measles. The potential importance of the conjunctiva as a portal of entry for measles has been demonstrated by Papp (1954, 1956, 1957), who found experimentally that, in the contact with measles patients, protection of the eyes with either goggles or anti-measles-convalescent serum dropped into the eyes, prevented the infection. When the primary focus of the measles infection is located in the conjunctiva, a “conjunctivitis of infection” may be the result (Goodall, in Grist 1950; Robbins 1962).

The superficial layer of the lamina propria of the conjunctiva is made up of lymphoid tissue. Also in this tissue, a multiplication of the measles virus takes place after the first viraemia, and a manifest conjunctivitis is the clinical result.

This is the conjunctivitis, characteristic for the prodromal stage of measles.

2.4. Ocular signs and complications in measles

2.4.1. The conjunctivitis of infection

When the conjunctiva is the portal of entry for the measles infection, a “conjunctivitis of infection” may be the result. (Herrman 1914; Goodall cited in Grist 1950; Robbins 1962). The practical importance of this observation is limited and bears no relation with the subject of the present study.

2.4.2. The conjunctivitis in prodromal measles.

A catarrhal conjunctivitis of variable extent is a characteristic sign of measles
(cfr Gemert, Valkenburg and Muller 1977) and has a high diagnostic value. In accordance with the pathogenesis of the measles-infection, this conjunctivitis has a subepithelial localization.

The conjunctival epithelium can be involved too. Occasionally lesions comparable to Koplik's spots can be observed. When localized at the caruncula or the semilunar fold they are of diagnostic value (Bonamour 1953; Gaud 1958; Nataf, Lépine and Bonamour 1960; Fedukowicz 1978). Very rarely vesicles with a contagious content are found in the conjunctival epithelium (Bonamour 1953b).

Azizi and Krakovsky (1965) observed in the majority of their measles cases — on slitlamp examination and using fluorescein 2% eyedrops — lesions of the conjunctival epithelium, continuous with similar lesions in the corneal epithelium. These epithelial lesions were localized in the palpebral slit. They were strictly epithelial, without subepithelial infiltration or other signs of inflammation. These authors state explicitly that this keratoconjunctivitis is a sign, not of a complication, of measles.

These same lesions were observed by Sauter (1976), their presence could be demonstrated with the use of the vital stains Rose Bengal and Lissamine Green. Because of these staining properties Sauter (1976) considered these epithelial lesions as signs of Vitamin A deficiency.

For the purpose of the present study it will be very important to know whether these lesions are of viral origin or induced by Vitamin A deficiency. Much attention will therefore be given to the immunofluorescence and electronmicroscopy of conjunctival biopsies (§ 5.1 and § 5.23) and the significance of the vital stains Lissamine Green and Rose Bengal (§ 3.4).

Complications of this conjunctivitis occur. An (unspecified) bacterial superinfection is possible; a combination with a diphtheric conjunctivitis is a rarity (Gaud 1958; Nataf, Lépine and Bonamour 1960).

Occasionally a phlyctenular keratoconjunctivitis is seen (Gaud 1958).

### 2.4.3. The epithelial keratitis in measles

In 1903, Trantas (Constantinople) described for the first time a coarse punctate, strictly epithelial keratitis as a sign of measles. This keratitis occurs at the time of the rash, is usually bilateral, gives remarkably little subjective symptoms and heals without sequelae.

The existence of this keratitis is mentioned by some other authors (Cosmöttatos 1908; Armengaud et al. 1961; Quéré 1964; Franken 1974; Sauter 1976), but only a few give some more details (Florman and Agatston 1962; Azizi and Krakovsky 1965). This keratitis is supposed to be caused by the measles virus (Trantas 1903; Jones 1960; Thygeson 1961; Casanovas 1976; Morgan and Rapp 1977).

There is considerable discongruence as regards the incidence of this keratitis:

- 4%: Armengaud et al. (1961) Senegal
- 10%: Quéré (1964) Senegal
- 30%: Sauter (1976) Kenya
- 71%: Lagruelet and Bard (1967) Upper Volta
- 76%: Trantas (1903) Turkey.

Thygeson (1961, USA) even states that this keratitis can be observed in every measles patient, provided he is seen early enough in the disease. These differences in incidence can at least be partially explained by the differences in examination technique: the incidence is necessarily higher in longitudinal studies (Trantas, Thygeson) than when it concerns snapshot visits (Sauter).

Not all authors agree upon the duration of the keratitis, most agree that it vanishes within days, without sequelae. Azizi and Krakovsky (1965) mention a duration of some weeks, whereas in the description of Florman and Agatston (1962) it may take the cornea some months to clear up.

It was already mentioned (§2.4.2) that Azizi and Krakovsky (1965) observed a continuity between the lesions in the corneal and conjunctival epithelium.

Central exfoliations of the corneal epithelium of larger size than the lesions of the coarse punctate keratitis of Trantas, were described by Förster and Berger (1892, cited in Trantas, 1903). This paper was not available in its original form. No other description of these central exfoliations could be traced.

2.4.4. Corneal ulcers, a complication of measles

The development of stromal ulcers is a complication of measles. These corneal ulcers have no particular morphological characteristics and are therefore indistinguishable from most other causes of corneal ulceration.

Usually a rim of normal corneal tissue is still present at the limbus. Perforation is common, the end result of these ulcers is a more or less dense leucoma, adhaerent leucoma or phthisis bulbi.

The leucomata are preferentially localized at the lower half of the cornea. Two explanations for this occurrence at the 6 o'clock position are given. Phillips (1961) (Zambia) mentions that when traditional medicines that hurt are applied to the eye, the eye is turned away (upward) as forcibly as possible and only the lower part of the cornea is exposed to the possibly noxious substances.

Sandford-Smith and Whittle (1979) (Nigeria) state that exposure and drying of the exposed cornea is an important pathogenetic factor in the cause of corneal ulceration after measles.

In this respect it may be of importance that Oomen (1961) distinguishes two different clinical forms of xerophthalmia: an acute total liquefaction of the cornea with extrusion of the contents of the eye and subsequent phthisis bulbi, and a more localized quiet perforation in the lower (or nasal) half of the cornea, resulting in a descemotocele and subsequently an adhaerent
leucoma with the retention of at least some useful vision. This last form suggests the presence of a localizing factor.

However, much controversy exists as far as the pathogenetic interpretation of the ulceration is concerned. As mentioned earlier (§1.2) 3 main pathogenetic mechanisms can be identified: infection, malnutrition and treatment.

a. Infection

During a measles epidemic in Haiti, Frederique, Howard and Boniuk (1969) observed 25 children with corneal ulcers. In 14 patients the ulcers perforated, in 7 of them bilaterally. No signs of previous Vitamin A deficiency were present: no nighblindness, no xerosis, no Bitot’s spots. Moreover, they mention ‘‘— the absence of any prior epidemic of nutritional keratomalacia with corneal perforation’’. Three eyes were examined histologically. The most prominent feature was the presence of multinucleate and syncytial cells in the epithelium, like those commonly seen in measles (cfr Scheifele and Forbes 1973). No specific changes in the corneal stroma were described. They concluded that the measles virus is to be held responsible for the corneal ulcers. Probably it is no coincidence that all 25 children were described as “markedly malnourished”.

Bacterial superinfection is another possible cause of corneal ulceration. Armengaud et al. (1961) observed 14 corneal ulcers in 416 consecutive cases of measles. These ulcers developed from the fifth day onwards after the rash in an area of punctate keratitis. The authors consider secondary bacterial infection responsible for the ulcers (Thygeson 1957; Rodger 1959; Quéré et al. 1967; cfr Lagraulet and Bard 1967). No complications were seen, however, when topical antibiotics were administered routinely (Quéré 1964; Lagraulet and Bard 1967; Quéré et al. 1967; Muller et al. 1977).

A bacterial infection of the xerotic cornea (i.e. in cases of corneal xerophthalmia) might be another potential cause of severe corneal damage (Kuming and Politzer 1967; Sullivan, McCulley and Dohlman 1973).

Sandford-Smith and Whittle (1979) demonstrated the possibility of a viral superinfection (with the herpes simplex virus) in cases of corneal ulceration after measles.

The possibility exists that fungal infections are initiated by the instillation of traditional African medicines (Phillips 1961).

b. Malnutrition

If malnutrition is considered as the main cause of corneal ulceration after measles, a distinction is to be made between Vitamin A Deficiency (= xerophthalmia) and Protein Energy Malnutrition (PEM).

Xerophthalmia after measles is considered to be a frequent event (Oomen, McLaren and Escapini 1964; ten Doesschate 1968; Oomen J.M.V. 1971; Franken 1974; Sauter 1976). Van Manen (1938) noticed in Indonesia that an epidemic of measles is followed in its wake by an epidemic of xerophthalmia.
Xerophthalmia after measles occurs in the majority of cases in children without any previous clinical sign of Vitamin A deficiency (Oomen 1961). The diminished intake of Vitamin A and its precursors and the decreased bio-availability of retinol are the causes of this xerophthalmia occurring "out of the blue". Also a lack of the B-vitamins can be a cause of corneal disease. The "nutritional corneal dystrophy" in P.O.W.'s (Spyratos 1949; Petzetakis 1950; Alleyne et al. 1977) and the "malnutritional keratoconjunctivitis" (Blumenthal 1950, 1960) are only mentioned here because of their connection with malnutrition. They have probably no relation to measles.

On the other hand, it is to be remembered that in nearly all cases with corneal complications encountered in the literature, the patients are invariably described as (markedly) malnourished. This has even led to the suggestion that keratomalacia might be a matter of PEM and not primarily of Vitamin A deficiency. (Yap-Kie-Tiong 1956; Arroyave et al. 1961; Reddy and Srikantia 1966; Venkataswamy 1967; Kuming and Politzer 1967; MacManus 1968; Emiru 1971; Baisya et al. 1971).

The observation that routine mass distribution of Vitamin A capsules (like in San Salvador and Indonesia) does prevent the minor forms of xerophthalmia (nightblindness, xerosis and Bitot's spots) but not the keratomalacia (Sommer, Faich and Quesada 1975; Pirie 1976), is an argument in the same direction.

c. Treatment

It has been mentioned in § 2.4.4.a that no corneal complications were observed when topical antibiotics were routinely applied to the eyes in cases of measles.

In contrast to the supposedly beneficial effect of these preparations, the use of topical traditional medicines could quite well be an important cause of blindness after measles. "During the prodromal stage of measles – irritant peppers and toxic substances are instilled in the eye as part of "traditional treatment". Corneal burns, ulceration, perforations and secondary infection lead to blindness." (Ayanru 1974.)

"Used in eyes, rendered more susceptible by existing pathology, the mechanical abrasive action of the powdered medication, the toxicity, acidity or alkalinity of the liquid preparations and the introduction of pathogens and fungi by the grossly unhygienic methods of preparation, must destroy countless corneas and eyes every year." (Phillips 1961.) (cfr McGlashan 1969; Jamieson 1970; Osuntokun 1975; Benezra and Chirambo 1977).

Strictly speaking, no proof is available about their harmful effects to the cornea, but "circumstantial evidence" (Agatha Christie) about their possible deleterious side-effects will be presented in a separate paragraph.
2.5. Depression of serum proteins, cell-mediated immunity and serum retinol in malnutrition

Serum proteins in malnutrition

Kwashiorkor and marasmus are the two extremes of the clinical spectrum of Protein Energy Malnutrition (PEM). Traditionally marasmus is supposed to be caused by the lack of food, providing the necessary energy, whereas kwashiorkor is attributed to a lack of protein. To explain the pathogenesis of kwashiorkor and clinical and epidemiological differences between kwashiorkor and marasmus, the concept of "disadaptation" was introduced (Waterlow and Payne 1975; Alleyne et al. 1977). In this view the marasmic child is chronically malnourished and uses its limited external energy sources in combination with katabolic mechanisms to maintain its biochemical equilibrium.

This adaptation mechanism fails in kwashiorkor (McLaren 1974; Waterlow and Payne 1975; Bailey 1975; Eddy 1977). The balance between requirement and input is acutely disturbed (weaning, infection) and the child doesn’t get the time to develop protective adaptive mechanisms, with all the clinical consequences of this failure. This concept explains why in marasmus the biochemical changes are minimal, whereas in kwashiorkor a profound biochemical disturbance exists (Whitehead, Coward and Lunn 1973).

One of the effects of the metabolic disturbance in kwashiorkor is the decrease in protein synthesis. The first proteins to be affected are some transport proteins with a rapid turnover, of which Retinol Binding Protein (RBP) is an example. The RBP level in the serum is therefore regarded as a fast reacting indicator of the impairment or improvement of the nutritional status (Ingenbleek et al. 1972, 1975a–b; Shetty et al. 1979).

The synthesis of serum albumin is also impaired, but at a much slower rate. The serum albumin is considered as a good, but slow, reacting indicator for the nutritional status. When the serum level of this protein drops below 3 gr%, biochemical deterioration sets in (Whitehead, Frood and Poskitt, 1971; Hay, Whitehead and Spicer, 1975). It is to be remembered however, that also measles itself is a cause for a lowering of the serum albumin level (Poskitt 1971).

The estimation of serum albumin and serum RBP therefore gives some information about the nutritional status of the child.

Immunological disturbance in malnutrition.

In general, malnutrition raises the susceptibility for infections (Emiru 1971), this holds especially true for bacterial infections (Scrimshaw, Taylor and Gordon 1959).

In malnutrition the infection, once established, also runs a more severe course. Mortality and morbidity increase considerably with increasing malnutrition (Geddes and Gregory 1974; Orren et al. 1979). This is caused by a
depression of the cell mediated immunity in kwashiorkor and marasmus (Sellmeyer et al. 1972). This depression of cell mediated immunity might be more severe in kwashiorkor.

The same applies to measles in combination with malnutrition. In malnourished children a prolonged excretion of Finkeldey-Warthin cells in nasal secretions was found: 12 days compared to 3 days in "normal" measles (Scheifele and Forbes 1972).

The delay in, or qualitative impairment of, the cell-mediated-immunity could easily be the cause of a bigger virusload and therefore of viral complications (Enders et al. 1959; Scheifele and Forbes 1972; Whittle et al. 1979).

In cases of very severe depression of cell mediated immunity (caused by malnutrition or immunosuppression by other mechanisms) measles can run its course even without the development of a rash (Burnet 1968; cfr Kimura, Tosaka and Nakao 1975). A malnourished child can therefore have a measles infection which stays undiagnosed, because of the lack of a rash.

Malnutrition not only deeply influences the course of the measles infection, but is in itself also a common complication of measles. Measles is an important reason for the "disadaptation" and can provoke an overt malnutrition in previously borderline malnourished children (Kimati and Lyaruu 1976; Alleyne et al. 1977). So it was found in Uganda, that 26% of the admissions for kwashiorkor had an infection with measles less than six weeks before the admission (Hay, Whitehead and Spicer 1975).

Serum retinol

Traditional treatment of measles (and the general illness of the child) includes a reduction in the intake of all food, including Vitamin A and its precursors. Moreover, the diarrhoea of measles interferes with the absorption. Vitamin A is, however, stored in large quantities in the liver and a diminished intake is probably not the cause of an acute deficiency (Moore 1957; Morley, Woodland and Martin 1963).

In all measles cases the serum retinol level is lowered, but this is possibly an aspecific effect of the fever because of increased metabolism (Moore 1957).

In cases of Protein-Energy-Malnutrition (PEM), the Retinol-Binding-Protein (RBP) is also lowered due to a reduction in the synthesis of apo-RBP, the protein part of the molecule. Since retinol is not present in its free form in the serum, a low serum retinol can therefore be caused by either a lack of retinol or a failing protein synthesis or both (Muto et al. 1972; Arroyave et al. 1961).

This means that a low serum retinol is therefore not automatically an indicator for Vitamin A Deficiency, in the same sense that a reduced RBP is not automatically a proof for the presence of PEM. Much caution is therefore needed in the interpretation of these biochemical estimations.
2.6. Traditional ocular medicines

The use of traditional medicines is widely spread all over Africa (Rodger 1959; Phillips 1961; Imperato and Traoré 1969; Kokwaro 1976; Chirambo and Benezra 1976). They are used in practically any condition, and are also frequently combined with western medicine (Maina-Ahlberg 1979).

In the literature several substances, instilled into the eyes, could be traced. They were in use as a treatment for "sore eyes" in general, or measles eyes more specifically.

- Powders: powdered cowrie shell (McLaren 1960a)
  powdered sugar candy (Holmes 1959; Chirambo and Benezra 1976)
  soot (Phillips 1961)
- Minerals: copper stone (Phillips 1961)
  copper sulphate (Holmes 1959)
- Foodstuffs: honey (Imperato and Traoré 1969)
  breastmilk (Ayanru 1978)
  orange juice (Imperato and Traoré 1969)
  egg yolk (Maina-Ahlberg 1979)
- Plants: a great many varieties (Kokwaro 1976)
- Miscellaneous: cow's urine (Animashaun 1977)
  aspirin tablets (Renkema, personal communication)

It will be obvious that powders may damage the cornea for mechanical reasons, and on a damaged cornea infections easily supervene.

Much damage will be caused by the physical and chemical properties of the preparation used: alkalinity or acidity, temperature, osmolarity. Moreover, the medicines can be toxic: copper may etch the cornea, resulting in abrasions and leucomata (Duke Elder and MacFaul 1972) and many "fresh" cases of traditional treatment look like acid or alkali burns (Chirambo and Benezra 1976). In many cases this may result in extensive scarring and symblepharon, whereas in other cases the cornea melts away totally with prolapse of the uvea (Chrambo and Benezra 1976). Also the impossibility of dosing the working principle of these preparations (Okoth, personal communication) is an important factor in the pathogenesis of the deleterious side effects of traditional medicines.

Most traditional medicines are of vegetable origin. Powders, decoctions, extracts and ashes derived from roots, tubers, stems, leaves, twigs and flowers are in use. Kokwaro (1976) mentions 75 plants used as eye medicines in East Africa. Of these, 18 are used in cases of conjunctivitis; measles is not mentioned specifically. Most probably many more are to be found (cfr Kerharo and Bouquet 1948; Rodger 1959).

For the explanation of possible negative effects of these medicines the emphasis is on the toxic substances found in these plants, e.g.: oxalates, saponines, steroid-like substances and cyanogenic glucosides.
The crushed leaves of *Oxalis corniculata* L. are, under the local names of Awayo (Luo) and Nandwa (Abaluyha), used to cure infected eyelids (Kokwaro 1976). *Oxalis* spp. however are known to contain large quantities of oxalates (Lewis and Elvin-Lewis 1977) what can be the cause of corneal ulcers (Duke-Elder and MacFaul 1972).

Many vegetable extracts contain saponins,* which may have a “deleterious action on the cornea, causing —, in concentrated solutions, chemosis, ulcerative keratitis and opacification” (Duke-Elder and MacFaul 1972). These saponins are present, among others, in Euphorbiaceae. An accidental contact with the sap of *Euphorbia tirucalli* L. (pencil tree) or *E. lactea* (candelabra cactus) can cause a considerable kerato-conjunctivitis.

In experiments, especially dogs are susceptible to this toxic effect (Crowder and Sexton 1964; Cordero-Moreno 1973). In this context it is however remarkable that the juice from the leaves and stem of *E. hirta* L. is used as a traditional eye medicine (Kokwaro 1976).

Moreover, some saponins have steroid properties or are steroid precursors. They are present — among others — in *Dioscorea* spp, *Polygala* spp and *Smilax* spp (Lewis and Elvin-Lewis 1977). Of these, the leaves of *Dioscorea astericus* Burk. *Polygala persicariifolia* DC, *P. stenopetala* Klotzsch and *Smilax kraussiani* Meisn are used as traditional eye medicines in East Africa (Kokwaro 1976). The presence of steroids explains their fame as anti-inflammatory agents, but is of course not always beneficial (cfr herpes simplex virus and measles, § 2.4.4.a).

Ilusa is the local Abaluyha name for *Ageratum conyzoides* L., of which the juice from the leaves is used to treat sore eyes (Kokwaro 1976). It contains however a cyanogenic glucoside (Lewis and Elvin-Lewis 1977) and HCN is known to reduce the repair activity of damaged corneal epithelium (Buschke 1949). Also *Cassia mimosoides* L., around Kakamega used as eyedrops, under the name of Masambu or Koinyama, contains cyanide.

It is also to be remembered that in cases of measles these preparations are used in severely ill children, with a reduced resistance, whose corneae are already damaged by a viral keratitis. A deleterious effect of these traditional medicines is therefore quite conceivable.

In general, the application of these preparations causes considerable pain. Logical thinking would therefore discard their use as harmful, but in primitive thinking the pain is considered as a proof of the strength of the medicine, and its use can therefore be continued, despite the apparent lack of therapeutic success (Phillips 1961). The use of these medicines might also explain the common localization of corneal ulcers at the 6 o’clock position on the cornea (Phillips 1961).

The action of the traditional medicines is of course not always only

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*Saponins are neutral amorphous glucosides of vegetable origin, with considerable surface action. They act strongly on cellular membranes (Bakker 1940, Duke-Elder and MacFaul 1972).*
detrimental. The anti-inflammatory effect of some saponin-containing species has been mentioned before. Another example is *Aloe barbadensis*, the extract of which contains chrysophanic acid, which is beneficial for the skin (Lewis and Elvin-Lewis 1977). It may also be the working principle in aloe extracts for the healing of corneal ulcers (Mortada et al. 1976). On the other hand, chrysophanol is supposed to cause a chemicotic conjunctivitis, keratitis and ulcers (Duke-Elder and MacFaul 1972). This difference might also be a matter of dosage. Too little is known about the effect of traditional medicines, their chemistry, toxicology and pharmacology. More attention is to be given to their use, especially in otherwise unexplainable conditions.

2.7. Statistics on Post-Measles-Blindness

In *Europe* Post-Measles-Blindness (PMB) is quantitatively unimportant. At a low overall blindness rate of 0.055% in the Netherlands, only 1% is related to measles. Corneal and posterior segment diseases are evenly represented (Schappert-Kimmijser 1959). The same was found in England (Fraser and Friedmann 1967) and Sweden (Lindstedt 1969).

In most blindness statistics from developing countries morphological and etiological criteria are mixed up and one has to be extremely careful to try to avoid a bias in the interpretation of these single sided blindness statistics.

In *Asia* the accent is on Vitamin A deficiency. In Serawak (Indonesia) 15 out of 103 patients, blinded below the age of 21 years, lost useful vision through keratomalacia "triggered" by measles. The same was found in 30% of the blind patients in Vietnam and South Korea (Oomen, McLaren and Escapini 1964).

Ten Doesschate (1968) describes in great detail the causes of blindness around Surabaya (Indonesia), in 675 blind children 285 cases of keratomalacia were found. Only in 5 children measles was considered to be the principal cause of blindness, but in a considerable percentage of the keratomalacia cases measles was the predisposing factor.

*Africa*

Rodger (1959) reported on the causes of blindness in North Nigeria, North Ghana and The Cameroons. Of the 193 blind children below the age of 10 years, 43 were blinded by measles; 17 were blinded by Vitamin A deficiency.

140 blind children were examined at the University Eye Clinic, Ibadan, Nigeria; 20 of them were blinded by measles (Olurin 1970). In 41 of the 116 enucleated eyes of children below the age of 10 years the corneal necrosis was associated with measles (Olurin 1973).

Also in Malawi 12% of childhood blindness is attributed to measles. One third of the inmates of the "Schools for the Blind" lost eyesight below the age of 3 years due to measles (Chirambo and Benezra 1976; Benezra and Chirambo 1977).
The figures from Zambia are even higher: of 686 blind persons, 64% blame measles as the cause of their blindness (McGlashan 1969).

Phillips (1961), also reporting from Zambia, states that even as much as 80% of blindness is caused by measles. He adds, however: “It is my firm conviction, that... African traditional medicines are the overwhelming cause of these lesions.”

On the contrary, Blumenthal (1954) considers measles in South Africa a negligible cause of blindness. The major cause of blindness (214 out of 895) is - what he calls - “malnutritional keratitis”. This “discrete colliquative necrosis” (McLaren 1960b) is supposed to be caused by lack of the B-Vitamins (Blumenthal 1950, 1960). From these statistics it is evident that PMB is a considerable problem, surely, in Africa.

Kenya

Some statistics on the causes of blindness in Kenya are available. Sauter (1976) surveyed the “Schools for the Blind” and found that 352 of the 749 inmates (47%) had lost their eyesight due to xerophthalmia in “approximately 60%” of the cases in connection with measles. This high prevalence of corneal blindness in “Schools for the Blind” is in concordance with my own findings. Roughly one third of the childhood blindness, as found in these schools, is of corneal origin, one third relates to cataracts (frequently combined with microphthalmus) and one third is attributable to other, mainly congenital or neurological, causes.

These figures, however, don’t give an indication of the overall prevalence of Post Measles Blindness, for which population-based surveys are needed.

The first survey for the causes of blindness in Kenya was conducted by Calcott (1956). 1093 Blind persons were examined: 183 showed a panophthalmitis, 182 corneal ulcers or their sequelae were encountered. No etiological diagnosis is given.

A number of random-sample surveys were done in several ethnic groups in Kenya.

Sinabulya (1976) examined 895 persons who had lost a total of 138 eyes, only one eye was lost in connection with measles. This study was done in Machakos, where also the “Joint Project Machakos” of the Medical Research Centre is going on. In this last study, with its emphasis on Mother and Child Care, not a single eye was lost because of measles. In this respect it is to be noted that Machakos, as far as medical care is concerned, is to be considered as a modern area (Muller et al. 1977; cfr Morley, Woodland and Martin 1963). Of the other surveys the numbers are not yet available.
3 - Patients and methods

3.1. Places and time

The provincial hospital at Kakamega is the referral centre for the district hospitals in Western Province, Kenya (see map 3.1). Specialist medical care is available only at this central facility.

The Provincial Ophthalmologist however, has his location at the St. Elizabeth Hospital Mukumu, a 150 bed Mission Hospital, 12 km South of

Map 3.1. Western Province, Kenya and some nearby district headquarters. Western Province and Siaya District were regularly visited on safaris by the Mobile Eye Unit no. 5. The other places were more or less frequently visited by the ophthalmologist alone, for consultations. The asterix indicates the location of the St. Elizabeth Hospital Mukumu
Kakamega. The eye department (founded and run by the Professor Weve Foundation) has at its disposal a 28 bed in-patient facility and a large outpatient department.

The "Kenya Society for the Blind" runs a nationwide, therapeutic, mobile eye service, and one of their "Mobile Eye Unit"'s has its home base at this hospital. With the Mobile Eye Unit, regular safaris were made in Western Province and the Siaya District, which together covers an area with over 2 million inhabitants. Most of the routine work on these safaris was done by the Ophthalmic Clinical Officer, and his assistants: a driver and a dresser.

Most of the patients to be described in this study were seen in the isolation ward of the St. Elizabeth Hospital. A few patients at the Kakamega Provincial Hospital were included in the clinical trial.

Of the patients seen anywhere on our safaris, only those with late complications after measles were included.

The study started June 1975; the last patient to be included in this study was seen in February 1978. In Kenya measles is an endemic disease, with an epidemic outbreak every 2–3 years. In the period of the study we had an epidemic in the second half of 1976. Moreover, in November 1976 a large measles vaccination campaign was held in the area of this study, a total of 18,000 doses were given and this may also have been instrumental in the reduction of the number of measles patients in 1977.

3.2. The patients

Groups of patients: diagnosis of measles and treatment

Table 3.2.a gives a summary of the patients included in this study.

A preliminary pilot study involved 99 measles children at the St. Elizabeth Hospital, examined in a very irregular way. The most important results were that daily slitlamp examination was necessary, and that all data had to be correlated to the day of first appearance of the rash.

<table>
<thead>
<tr>
<th>Time</th>
<th>Place</th>
<th>No.</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pilot Study</td>
<td>6-'75–4-'76 SEH</td>
<td>99</td>
<td>§ 4.1-4.2</td>
</tr>
<tr>
<td>Mukumu I</td>
<td>6-'76–11-'76 SEH</td>
<td>148</td>
<td>§ 4.3.2</td>
</tr>
<tr>
<td>Clinical trial</td>
<td>3-'77–10-'77 SEH</td>
<td>51</td>
<td></td>
</tr>
<tr>
<td>Mukumu II</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kakamega</td>
<td>KPH</td>
<td>49</td>
<td></td>
</tr>
<tr>
<td>Safaris + Mukumu</td>
<td>1-'77–2-'78 any</td>
<td>9</td>
<td>§ 4.4</td>
</tr>
</tbody>
</table>

SEH = St. Elizabeth Hospital; KPH = Kakamega Provincial Hospital.
In this study four groups of patients will be described. The most important group is "Mukumu I" including 148 patients seen at the time of the measles epidemic in 1976. These patients will be described in detail in §4.1.

The statistical analysis regarding measles keratitis and nutritional status is done on this group only. For statistical reasons it is not possible to combine the different groups of our patients.

The 100 patients in the groups "Mukumu II" and "Kakamega" formed a clinical trial regarding the effectiveness of Vitamin A capsules 200,000 IU and (or) tetracycline eye-ointment, in the prevention of corneal complications in measles. This trial failed because of the limited number of patients. Only the patients who developed complications will be described.

During the safaris I used to check on all children admitted to the hospitals and dispensaries we visited. This means that every year about 2,000 children were examined, apart from those seen at the regular eye clinics. In 1977, 7 patients with late complications after measles were found on safaris. They are included in §4.4.

**Diagnosis of measles**

All children seen at the out-patients department with measles were admitted, except when the mother refused admission. I did not interfere actively with this policy. The diagnosis of measles was made on clinical grounds by the attending (pediatric) clinical officer or physician and this clinical diagnosis was, ipso facto, the reason for admission to the isolation ward and inclusion into this study.

The demonstration of a specific IgM is proof of a recent infection (Wilterdink 1979). A total of 118 serum samples were examined for the presence of anti-measles IgM; 106 were positive, 12 negative.

The 12 patients with negative IgM estimations are tabulated in table 3.2,b.

<table>
<thead>
<tr>
<th>R</th>
<th>W/A</th>
<th>Keratitis classification</th>
<th>Measles vaccination</th>
<th>Tentative explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>−1</td>
<td>64</td>
<td>II</td>
<td>+</td>
<td>too early</td>
</tr>
<tr>
<td>−1</td>
<td>81</td>
<td>III</td>
<td>−</td>
<td>too early</td>
</tr>
<tr>
<td>0</td>
<td>87</td>
<td>neg</td>
<td>−</td>
<td>too early</td>
</tr>
<tr>
<td>0</td>
<td>76</td>
<td>II</td>
<td>−</td>
<td>too early</td>
</tr>
<tr>
<td>0</td>
<td></td>
<td>I</td>
<td>−</td>
<td>too early</td>
</tr>
<tr>
<td>+1</td>
<td>main</td>
<td>V</td>
<td>−</td>
<td>malnutrition</td>
</tr>
<tr>
<td>+2</td>
<td>66</td>
<td>III</td>
<td>−</td>
<td>malnutrition</td>
</tr>
<tr>
<td>+2</td>
<td>93</td>
<td>III</td>
<td>−</td>
<td>?</td>
</tr>
<tr>
<td>+3</td>
<td>90</td>
<td>neg</td>
<td>−</td>
<td>no measles</td>
</tr>
<tr>
<td>+4</td>
<td>86</td>
<td>neg</td>
<td>−</td>
<td>no measles</td>
</tr>
<tr>
<td>+6</td>
<td>58</td>
<td>I</td>
<td>?</td>
<td>no measles/main.</td>
</tr>
<tr>
<td>+7</td>
<td>76</td>
<td>V</td>
<td>−</td>
<td>?</td>
</tr>
</tbody>
</table>

Table 3.2,b. Patients with negative anti-measles IgM in relation to the time of outbreak of the rash (R), nutritional status (W/A = Weight for Age), the keratitis classification (§3.3), vaccination against measles and the tentative explanation for the negative IgM.
In 5 cases the sample was probably taken too early, i.e. before or on the day of the outbreak of the rash. In 2 or 3 cases the possibility of immunosuppression or delay in the development of measles-antibodies, due to Protein Energy Malnutrition, exists, whereas in 2 or 3 other cases the eruption might have been something other than measles. For 2 cases no explanation can be given. This outcome once more proves the validity of the clinical diagnosis of measles (Gemert et al. 1977) and makes it more than likely — especially in epidemics — that we really are dealing with measles patients.

Sex, age and time of admission. (Mukunu I)

During the epidemic of 1976, 152 children, all Luhya's by tribe, were admitted to the isolation ward of the St. Elizabeth Hospital. Three children were excluded from this study, because they had developed measles more than 10 days previously. One well-nourished child with a measles contact failed to develop a measles rash and was therefore also excluded from the study.

The remaining 148 children were 78 girls and 68 boys; in 2 cases the sex was not recorded (for numerical data, see appendix). The age of these children varied from 5 months to 14 years, median age 21 months. The age distribution is given in table 3.2.c.

The time of admission in relation to the day of outbreak of the rash is given in table 3.2.d.

The conclusion to be drawn from tables 3.2.c and 3.2.d is that our measles patients are, as is usually seen in Africa, quite young when they contract measles. Moreover, they are admitted to the hospital early after the outbreak of the rash, probably because the mothers recognize the seriousness of the disease.

Treatment of measles patients

From the beginning of the study it was decided not to interfere with the treatment as prescribed by the attending clinical officer or physician.

All children received antibiotics (mostly penicilline) and anti-malaria

<table>
<thead>
<tr>
<th>Age in months</th>
<th>0–12</th>
<th>13–24</th>
<th>25–36</th>
<th>37–48</th>
<th>49+</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>30</td>
<td>49</td>
<td>31</td>
<td>18</td>
<td>20</td>
<td>148</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Day of admission</th>
<th>-3</th>
<th>-2</th>
<th>-1</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5+</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>1</td>
<td></td>
<td></td>
<td>3</td>
<td>17</td>
<td>58</td>
<td>26</td>
<td>22</td>
<td>14</td>
</tr>
</tbody>
</table>
treatment systemically. Fluids were given in large quantities, when necessary in intravenous-drips or subcutaneously. Phenergan syrup or other cough mixtures and nasal drops were given for symptomatic relief. The oral ulcers were painted with gentian violet. Occasionally children needed to be nursed in steam tents. The treatment also included the administration — routinely — of extra vitamins, in relatively low dosage. The children daily received an average of about 1,000 IU Vit A extra (together with B1, B2, B6, B12, C, nicotinamide, Ca Panthotenate). When the condition of the children permitted the intake of solid food, they had the customary posho (maize) with mboga (green leafy vegetable).

Out of the 148 children in the “Mukumu I” group II children died.

In 84 cases the attending clinical officer (or doctor) prescribed tetracycline eye-ointment, because of the conspicuously red aspect of the eyes. This treatment was accepted and handled as a clinical trial as to the effect of tetracycline eye-ointment on the keratitis. We only interfered with this regimen when ocular complications developed.

3.3. Ophthalmological examination

The ophthalmological examination consisted of a daily examination with a hand held slitlamp (KOWA). Vital stains were used to enhance the visibility of the epithelial lesions: fluorescein 1% was used for the lesions of the cornea, Rose Bengal 1% (colour index 45440) (RB) and, later on, Lissamine Green 1% (colour index 44090) (LG) were used to stain the lesions of the conjunctival epithelium.

A lot of confusion exists about the etiological interpretation of the staining conjunctival lesions. According to Norn (1970, 1973) RB and LG stain degenerating and dead cells in the epithelium, irrespective of the cause of the lesions. On the contrary Sauter (1976) claims that "— vital staining by 1% Rose Bengal or 1% Lissamine Green is a safe, sensitive, specific, simple and cheap method for — early — detection of cases of conjunctival xerosis (X-1A), both in Health Centres and in large-scale field surveys." (pg 194). This claim could not be confirmed (Kusin, Soewondo and Parlindungan Sinaga, 1979; Sommer, 1980).

The use of the word "specific" is the cause of this controversy. For the subject of this study it is of critical importance how the staining conjunctival lesions are etiologically interpreted. Why I adhere to Norn’s opinion, will be explained in §3.4.

In contrast to RB and LG, fluorescein stains "spaces": when an epithelial defect exists, fluorescein diffuses into the intercellular spaces (Norn 1970). Fig. 3.3. (see colour plate I) gives a good example of this differential staining in a case of herpetic keratitis. (= Pat Chw P 172, § 4.4.)

In the examination of the cornea a quantitative scoring system was used to evaluate the correlation measles-keratitis and nutritional status. The number
of lesions on the cornea was counted. If no lesions were present, this was considered negative, 1–5 lesions was positive, 6 lesions or more +++. The observations of both corneas during the first 5 consecutive days were totalled. The maximum to be reached was therefore 20++. The "keratitis score" was classified as follows:

<table>
<thead>
<tr>
<th>extent and duration of corneal lesions</th>
<th>classes of keratitis score</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 +</td>
<td>I</td>
</tr>
<tr>
<td>1 and 2 +</td>
<td>II</td>
</tr>
<tr>
<td>3–5 +</td>
<td>III</td>
</tr>
<tr>
<td>6–8 +</td>
<td>IV</td>
</tr>
<tr>
<td>9–20+</td>
<td>V</td>
</tr>
</tbody>
</table>

When in the first 5 days, one observation day was missing, an interpolation between the previous and following day was used. No "keratitis score" was calculated when the period of observation had been less than 5 days. In some tables a "negative" keratitis score is mentioned instead of a keratitis score class I. This means that the observation period was less than 5 days and for that reason — by definition — no classification of keratitis score could be made.

The ophthalmological examination was — because of the purpose of the study and the generally bad condition of the children — limited to a slitlamp examination. No attention was given to the posterior segment of the eye.

The nearest microbiological laboratory from which reliable results could be obtained was some hundreds of kilometers away. To culture viruses or bacteria was therefore not possible.

3.4. The significance of vital staining of the conjunctiva with Lissamine Green or Rose Bengal

*Lack of specificity for the detection of Vitamin A deficiency*

Rose Bengal (Colour Index 45440) and Lissamine Green (Colour Index 44090) are in a 1% solution in use for the examination of the conjunctival epithelium. Lissamine Green has the advantage over Rose Bengal of a better visibility against a reddish background and hurts less on instillation. They stain mucus and devitalized cells, irrespective of the cause of the cellular damage (Passmore & King 1955; Norn 1970, 1973). The positive staining with Rose Bengal and Lissamine Green nearly always (except, among others, in measles) takes the form of micropunctate lesions. When present in small numbers, they have no pathological significance (Kronning 1954; Norn 1964, 1970, 1973; Lansche 1965). They occur in pathological quantities in keratoconjunctivitis sicca (M. Sjögren) (Kronning 1954; Passmore & King 1955), in traumata of the conjunctival epithelium (mechanical or toxic), sometimes
around Bitot's spots. In 1976 Sauter claimed that a positive staining with Rose Bengal and Lissamine Green was a specific and reliable sign of Vitamin A deficiency. This claim has not been confirmed.

Vijayaraghavan et al. (1978) studied under field conditions the usefulness of the Rose Bengal test for the detection of Vitamin A deficiency. They found a considerable number of false positive children: i.e. children who are positive in the dye test, but fail to show any -- clinical or biochemical -- sign of Vitamin A deficiency.

Kusin et al. (1977) demonstrated the presence of a considerable number of false negatives. Moreover, in a clinical trial, the treatment with a massive dose of Vitamin A (200,000 iU) failed to protect the children against the subsequent development of a positive dye test (Kusin, Soewondo and Parlindungan Sinaga 1979). These findings were confirmed by Sommer (1980) and he reaches the conclusion that the staining with Lissamine Green is useless in the detection of Vitamin A deficiency.

All this work has been done in Asia (India and Indonesia). Unaware of this work, I paid a lot of attention to the value of Lissamine Green test in its relation to Vitamin A deficiency. Early 1976 I did a survey in the Shikusa Borstal Institution, Kenya, where among the inmates clinical signs of Vitamin A deficiency were rather frequent. In a statistical analysis (W. Gemert, Medical Research Centre, Nairobi, Dpt. of the Royal Tropical Institute, Amsterdam, The Netherlands) it was found that the incidence of Bitot's spots correlated significantly with the time spent in prison. No such correlation existed between the Lissamine Green staining alone (i.e. without the presence of Bitot's spots) and the length of stay in prison (see table 3.4.a.).

Also a doubly masked clinical trial was done. Half the inmates got 200,000 iU Vitamin A, the other a placebo. After one month all inmates were re-examined for the presence of clinical signs of Vitamin A deficiency and the Lissamine Green test. The results are given in table 3.4.b.

It appeared that 20% of all boys, who had been negative for L.G. at the initial survey, became positive, whether they got Vitamin A or a placebo. Also, 18% of the boys, initially positive, became negative after the placebo, whereas 37% became negative after Vitamin A. This difference is statistically significant. The only conclusion can be that some epithelial lesions staining with Lissamine Green react to the administration of Vitamin A.

Because of these results I reached the conclusion that Lissamine Green is not a test for Vitamin A deficiency which later was confirmed by others.

**Lissamine Green in measles**

Around the outbreak of the rash Lissamine Green and Rose Bengal staining lesions were observed in the bulbar conjunctiva (§ 4.1.2, figs. 4.1.2.a and b). They had a particular morphology, quite different from the micropunctate lesions around Bitot's spots and in xerosis. They disappeared without any
Table 3.4.a. The incidence of Bitot's spots and Lissamine Green positive staining, in relation to the time spent in the Borstal Institution. The incidence of Bitot's spots and the duration of the stay are significantly correlated, no association exists between the time spent in prison and the incidence of Lissamine Green positive staining. Treatment and were seen in all patients who came under observation before the outbreak of the rash (table 4.1.2.c).

Moreover, the studies mentioned earlier in this section don't substantiate

Table 3.4.b. The staining with Lissamine Green before and after the treatment with Vitamin or a placebo. The administration of a high dose of Vitamin A reduces significantly (compared to placebo) the incidence of Lissamine Green positive staining (compare Groups A and C, \( P = 0.015 \)). The administration of Vitamin A does not prevent the conversion from negative to positive staining in 20% of pupils (compare Groups B and D, \( P = 0.40 \)). At least a very considerable proportion of the Lissamine Green staining appears therefore to be independent of the Vitamin A status.

<table>
<thead>
<tr>
<th>Staining at initial survey</th>
<th>Staining at control survey</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>After Vitamin A</td>
</tr>
<tr>
<td></td>
<td>LG +  LG -</td>
</tr>
<tr>
<td>Group A: LG +</td>
<td>63% 37%</td>
</tr>
<tr>
<td>Group B: LG -</td>
<td>20% 80%</td>
</tr>
<tr>
<td>Group C: LG +</td>
<td>79% 21%</td>
</tr>
<tr>
<td>Group D: LG -</td>
<td>18% 82%</td>
</tr>
<tr>
<td></td>
<td>LG +  LG -</td>
</tr>
<tr>
<td></td>
<td>After placebo</td>
</tr>
</tbody>
</table>

n = 389
the specificity, claimed by Sauter, for Lissamine Green to detect conjunctival xerosis.

For these two reasons, the peculiar morphology and natural history of the conjunctival lesions in measles, and the lack of etiological specificity for Lissamine Green there are no grounds to follow Sauter’s opinion that these lesions are specifically caused by Vitamin A deficiency.

In this study LG and RB will therefore only be considered as vital stains for the detection of epithelial lesions, without ascribing any precise etiological connotation, unless proven by other techniques.

In § 5.1 it will be demonstrated that with the use of immunofluorescent techniques these lesions reveal their viral nature.

3.5. Assessment of the nutritional status

When the nutritional status of a measles child is to be compared with the severity of its measles keratitis, the nutritional status of the child must be measured quantitatively: a qualitative clinical impression is useless for this purpose, so anthropometric or biochemical methods are to be used.

On admission, the nurses of the isolation wards collected data regarding age, sex, vaccinations and measles-contacts of the children. As soon as feasible the nurses took height and weight of the admitted children. Height was taken with the children lying on the bed, using an ordinary tape ruler; weight was taken with ordinary baby weighing scales. The sometimes critical condition of the children and the application of drips and steam-tents did not permit the use of Salter scales. Occasionally measuring was, in any event, impossible.

Weight, height and age, are, in any combination, used to assess the nutritional status.

- Weight for age (W/A): the weight is expressed as a percentage of the ideal weight for the age of the child. Although it is an American standard, the Harvard standard is generally used as reference. The “Road to Health” is based on this standard. (Ministry of Health, Kenyan Government.)

- Height for Age (H/A): the height is expressed as a percentage of the ideal height for the age of the child. Stunting would be indicative of malnutrition of long duration.

- Weight for Height (W/H): the weight is expressed as a percentage of the ideal weight for the height of the child. This index is extremely useful when the age of the child is not exactly known. Moreover, it would be a good indicator of the actual nutritional status of the child (Alleyne et al. 1977).

Height, weight and age can mathematically be combined in single indices, which are then supposed to be independant of the age of the child (Rao and Singh 1970; Dugdale 1971; McLaren and Read 1975). They are, however, only applicable to a limited age group (up to 5 years), moreover, our data were not always complete enough to use these indices.
To evaluate the nutritional status biochemically, as soon as possible after admission, a venous bloodsample was drawn, spun down, and the serum was stored at \(-20^\circ\text{C}\). These samples were transported to the Medical Research Centre (Nairobi), where all immunological and biochemical estimations were done (Miss H.L. Ensering and Miss M.M. van Rens).

For the detection of specific anti-measles IgM a direct immunofluorescent technique was used. Vero cells (Green monkey kidney cells), in tubes with coverslips were inoculated with the Schwarz measles vaccine strain. When, during inoculation at \(37^\circ\text{C}\), a Cyto-Pathogene Effect, typical for measles, appeared, these coverslips were washed three times with Phosphate Buffered Saline (PBS), and fixed in acetone for 20 minutes at a temperature of minus \(20^\circ\text{C}\) and kept at \(4^\circ\text{C}\). Three coverslips were inoculated simultaneously with either the serum to be examined, diluted 1:10 in PBS, or a standard IgM positive serum or a standard IgM negative serum, but with specific anti-measles IgG.

After incubation at \(37^\circ\text{C}\) for 45 minutes and washing three times with PBS, the Fluorescein-Iso-Thiocyanate-Conjugate (FITC)-anti-human-IgM-goat-serum was applied. After incubation and washing with PBS the coverslips were mounted and checked for the presence of fluorescence under the fluorescence microscope. Only a strong fluorescence was read as positive.

The results of the 118 anti-measles IgM estimations are given in §3.2.

99 Serum samples were estimated for the presence of serum retinol and \(\beta\) carotene. The values found were far too high to be reliable, which was probably caused by autofluorescence because of improper storage. These results will therefore not be used.

In 103 sera RBP was estimated, the albumin content in 100 samples. For both estimations a Manchini Radial Immuno-Diffusion Test was used.

Glassplates, 100 \(\times\) 100 \(\times\) 1.5 mm, were coated with 1\% Agar Noble and dried for 15 min at \(100^\circ\text{C}\). A solution of 2\% agarose in Manchini-buffer was mixed with commercially available antiserum (Hoechst) at \(53^\circ\text{C}\): 0.8 cc of anti-Retinol-Binding-Protein serum, or 0.3 cc of anti-albumin-serum was added to 15 cc of the agarose solution. This mixture was poured out on the glassplates, 2.5 mm large holes were punched in this gel with a gel-puncher. Each hole was filled with 5 \(\mu\)l, either of a standard serum in routine-dilutions, or the serum to be tested in a 1:150 dilution.

The plates were left (strictly horizontally) for 2–3 days in which the proteins diffused into the agarose-gel, where they precipitated with the specific antiproteins. The size of the precipitation-ring is a measure for the quantity of protein present in the serum to be tested. The value can be calculated by comparison with the precipitation rings of the standard serum.

In this technique, the normal values (derived from Europeans) are:

- **Albumin**: 3500–5500 mgr%
- **Retinol Binding Protein**: 3–6 mgr%

The results will be presented in §6.2.
3.6. Biopsies and specimens for pathology, electronmicroscopy and immunofluorescence

After consent of the attending parent was obtained, paired conjunctival biopsies were taken from 10 children: one biopsy came from the Lissamine Green positive area in the exposed part of the bulbar conjunctiva, one biopsy — serving as a control — from the non-staining conjunctival tarsi.

For conjunctival biopsies, the conjunctiva was anaesthetized with a few drops of 0.4% descaine (Novesine ®) topically. The conjunctiva was lifted with a non-toothed forceps and a small piece was cut off with a sterile de Wecker scissor. For obvious reasons it was not possible to obtain corneal biopsies.

Four pairs of conjunctival biopsies were fixed in glutaraldehyde according to Sabatini, and prepared for electronmicroscopy (Drs. G.F.J.M. Vrensen and J.J.L. van der Want) at the Netherlands Ophthalmic Research Institute, Amsterdam.

Five pairs of conjunctival biopsies were taken from children admitted to the measles-ward of the Kenyatta National Hospital (Head of the department Dr. M.L. Oduori), and — stored in B.M. Eagles with 2% Foetal Bovine Serum — airmailed on dry ice to Dr. F.E. Nommensen (Dpt. of Virology, Erasmus University, Rotterdam) for investigation with immunofluorescent techniques.

One pair of conjunctival biopsies was taken from a measles child with conjunctival xerosis and prepared for light microscopy.

One child, with a corneal necrosis after measles, came for evisceration of the affected eye. One corneal specimen was sent to Prof. Dr. W.A. Manschot, Inst. of Pathology, Erasmus University, Rotterdam; a second specimen was examined by electron-microscopy.

In 2 children who developed complications (one central abrasion and one an ulcer) and died, we were allowed to remove the corneae. They were sent for electronmicroscopy.

3.7. Representativeness of the patient samples

In this study nearly all the children examined were hospitalized. This might easily become an important cause for selection of the patients to be examined:

(a) Only the more malnourished children are seen, because they are supposedly more susceptible to infections and, in the case of malnutrition, infections run a more severe course.

(b) The admission of children to a hospital is always to be paid for. For this purpose a government hospital is much cheaper than a Mission hospital. It is therefore quite conceivable, that in the St. Elizabeth Hospital only children of more well-to-do families are admitted and hence these children might be in a better nutritional state than the average population.
Table 3.7.a. The percentage of children, age 13–48 months, below the indicated level of Weight for Age. The first group is a random sample survey of 229 children, included in the Rural Kenyan Nutrition Survey (1977). The second group are the 72 children, age 13–48 months of the measles patients in Mukumu I. In the column with 'weight correction' the weightloss because of measles (6%: Morley, Woodland and Martin 1963) has been taken into account.

<table>
<thead>
<tr>
<th>Weight for Age</th>
<th>Rural Kenyan Nutrition Survey</th>
<th>Measles patients</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Weight correction</td>
<td></td>
</tr>
<tr>
<td>&lt; 70%</td>
<td>2%</td>
<td>0%</td>
</tr>
<tr>
<td>&lt; 80%</td>
<td>7%</td>
<td>10%</td>
</tr>
<tr>
<td>&lt; 90%</td>
<td>27%</td>
<td>26%</td>
</tr>
<tr>
<td>n = 229</td>
<td></td>
<td>n = 72</td>
</tr>
</tbody>
</table>

Table 3.7.b. Comparison of Height for Age of 229 children, age 13–48 months, seen in the Rural Kenyan Nutrition Survey and the 73 measles patients, age 13–48 months, in measles patients group Mukumu I.

<table>
<thead>
<tr>
<th>Height for Age</th>
<th>Rural Kenyan Nutrition Survey</th>
<th>Measles patients</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 80%</td>
<td>2%</td>
<td>1%</td>
</tr>
<tr>
<td>&lt; 90%</td>
<td>21%</td>
<td>26%</td>
</tr>
<tr>
<td>n = 229</td>
<td></td>
<td>n = 73</td>
</tr>
</tbody>
</table>

Table 3.7.c. Comparison of the Weight for Age of the children, seen in the Provincial Hospital Kakamega, and the St. Elizabeth Hospital Mukuma, during 1977. No statistically significant differences were present.

<table>
<thead>
<tr>
<th>Weight for Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 60%</td>
</tr>
<tr>
<td>Kakamega</td>
</tr>
<tr>
<td>Mukumu I</td>
</tr>
</tbody>
</table>

4 43 43 90

ad (a) In 1977, the Kenyan Ministry of Finance and Planning published the results from the "Rural Kenyan Nutrition Survey". Table 3.7.a shows that the 148 children in the most important group of our patients (Mukumu I) are a representative sample of the children in Western Province as far as their Weight for Age is concerned.

The similarity is especially good when the fact is taken into account that children with measles loose weight, because of measles. In West Africa an average weightloss of 6% was found.
(Morley, Woodland and Martin 1963) and in the column with
"weight correction" this figure is used.

Table 3.7.b shows the concordance for Height for Age in our
patients and the sample of the Rural Kenyan Nutrition Survey.

(a) Table 3.7.c shows a comparison of the Weight for Age of the
children in the Provincial Hospital Kakamega and the children in
the St. Elizabeth Hospital. The numbers are small, but in our
material, the suggestion that children in the St. Elizabeth Hospital
are in a better nutritional status, compared to the children in the
Governmental Hospital, cannot be substantiated.

This leads to the conclusion that our measles patients form a group of
children representative for those in Western Province.
4 - Clinical description of Ocular signs and Corneal complications of Measles

4.1. Ocular involvement in measles

The clinical description of the ocular lesions in measles is derived from the daily slitlamp-examination of the 148 children in "Mukumu I", during the measles epidemic in 1976. In the well-known catarrhal conjunctivitis of prodromal measles a distinction could be made into two different disease-entities: a subepithelial conjunctivitis and an epithelial conjunctivokeratitis. The former occurs early and is localized in the subepithelial tissue, mainly of the tarsal, and less of the bulbar, conjunctiva, whereas the latter occurs some days later, is strictly epithelial and is localized in the exposed parts of the bulbar conjunctiva.

4.1.1. Subepithelial conjunctivitis

A catarrhal conjunctivitis was a constant, but in extent variable, feature of the prodromal and early exanthematous stage of measles. The palpebral conjunctiva was always inflamed, and in at least half of the cases an inflammatory reaction of also the bulbar conjunctiva was conspicuous without separating the eyelids. The discharge varied from watery to, later on, mucoid. A sometimes frankly purulent discharge suggested bacterial involvement. In some cases a slight oedema of the conjunctiva at the inner canthus existed, subsiding within a few days. No lesions comparable to Koplik's spots were observed in the conjunctiva.

5-7 Days after the outbreak of the rash this conjunctivitis had disappeared. Subconjunctival haemorrhages occurred in some 20% of the cases. Follicles were visible in the lower fornix or at the border of the superior tarsal plate in 10-20% of the cases.

4.1.2. Epithelial conjunctivokeratitis

Conjunctiva

With the use of Lissamine Green or Rose Bengal the presence of epithelial lesions in the conjunctiva could be demonstrated. These lesions were otherwise not visible. They were localized at the nasal part of the bulbar conjunctiva, a little less frequently at the temporal side, but nearly always in the exposed parts of the conjunctiva in the interpalpebral fissure (fig. 4.1.2a: see colour plate II). Only in 3 (of the 148) cases were these lesions observed at the palpebral conjunctiva. In these cases also their number was very limited.

The shape of these lesions varied a great deal. Usually they were more or less round, but occasionally they were multi-angular, circular or horse-shoe shaped (fig. 4.1.2.b: see colour plate II), their size varied from 0.2-0.4 mm.

The number of these "measles spots" varied considerably. On some
The incidence of Lissamine Green staining conjunctival epithelial lesions and keratitis, in relation to the day of outbreak of the rash (R). The data before R are drawn with interrupted lines to indicate the lower level of reliability (compared to data after R) because of the limited number of observations of these early cases. Occasions only a single spot was present, whereas in other cases they could be numerous and give the impression to conflate to a single, large, intensely staining area on the conjunctiva.

The presence of the "measles spots" was dependent on the stage of the disease. All 6 children examined in the prodromal stage of the measles showed these lesions. From the day of the outbreak of the rash the incidence of these lesions dropped, to reach a level of only 11% at the fourth day after the rash. No "measles spots" were seen in the conjunctiva after the eighth day (fig. 4.1.2.c).

These "measles spots" were present independently of the presence of subepithelial vasodilatation.

The "measles spots" had a life-span of one to only a few days and disappeared spontaneously without any treatment. The conjunctival epithelium with "measles spots" is usually not water-repellent: conjunctival xerosis was only observed in 4 cases; two times a capsule of Vitamin A (200,000 iU) was given, the 2 other children received no treatment. In all four patients the conjunctivae returned to normal within a few days.

The shape, the distribution, and the size of these lesions is characteristic for measles; their appearance differs from other conditions in which staining of the conjunctival epithelium occurs. I never observed them in diseases other than measles.

The impression exists that in some children who could be observed for a
long period, the quantity of pigment in the exposed conjunctiva increased during the observation period.

In the literature only two papers could be traced where conjunctival epithelial lesions are mentioned in connection with measles. Azizi and Krakovsky (1965) mentioned conjunctival epithelial lesions, without elaborating their statement. Sauter (1976) mentions a coarse punctate staining of the conjunctival epithelium in measles and ascribes this finding not to measles but to Vitamin A deficiency.

Our findings don't confirm Sauter's opinion (cfr §3.4). Moreover, the morphology of the conjunctival epithelial lesions is very typical for measles. This in itself is an argument against any specificity claimed for the staining with Lissamine Green or Rose Bengal. This fact once more confirms the conclusion of §3.4: Lissamine Green stains epithelial lesions of the conjunctiva, irrespective of the cause.

Conjunctival epithelial lesions of a typical morphology in the prodromal stage of measles are therefore to be considered as a sign of measles.

**Cornea**

Usually the "measles spots" in the conjunctival epithelium had their largest extent around the day of outbreak of the rash (R). Sometimes the whole of the nasal and temporal exposed conjunctiva stained brilliantly green. The lesions did not stay restricted to the conjunctival epithelium, but the same lesions developed at the corneal side of the limbus, in continuity with the lesions in the conjunctival epithelium (fig. 4.1.2.d: see colour plate II).

The epithelial conjunctivitis crosses over to the cornea to give a coarse punctate keratitis. In one case, where I had taken a biopsy from the conjunctiva, only the part of the cornea adjacent to the site of the biopsy stayed free from the keratitis. This confirms the clinical impression that the measles keratitis develops from the epithelial conjunctivitis.

The appearance in size and shape of the lesions in the conjunctival and corneal epithelium was identical. In the cornea the lesions were visible without vital staining. Here they appeared as greyish, slightly opaque, epithelial lesions, strictly confined to the epithelium: Bowman's membrane and the stroma were not involved.

The epithelial keratitis progressed from the limbus toward the centre of the cornea (fig. 4.1.2.e: see colour plate II).

At the time when the conjunctival epithelial lesions started to disappear, the corneal epithelium at the limbus also began to regain its normal aspect. The superficial, coarse punctate keratitis now occupied the mid-periphery of the corneal epithelium (fig. 4.1.2.f: see colour plate II). The centre of the cornea was the last part to become involved. In many cases a punctate keratitis centrally in the cornea was the last manifestation of this measles conjunctiva-keratitis (fig. 4.1.2.g: see colour plate II).

This conjunctivo-keratitis was sometimes observed in restricted forms.
Several times the keratitis stopped at the corneal side of the limbus and did not progress towards the centre. On other occasions only a limited number of lesions were observed, and only one or a few spots developed during a 10 day observation. The time of appearance of the lesions is variable, not the sequence. The limbal lesions appeared from $R-1$ to $R+5$, the central lesions from $R+2$ to $R+9$ to 11. $R$ (like elsewhere in this study) indicates here the day of outbreak of the rash. No lesions were seen after day $R+11$.

The keratitis heals without sequelae. In 7 of the 248 cases however, macro-erosions were seen, directly attributable to the keratitis. They will be dealt with in the next paragraph. No significant differences were observed between right and left eyes.

This keratitis was observed in 115 of the 148 patients (76%). In fig. 4.1.2.c also the incidence of this measles-keratitis is given in relation to the day of outbreak of the rash ($R$). The incidence is highest at day $R+1$: 65% of the measles children show at that day a keratitis. From that day onwards the incidence decreases. No keratitis was observed after day $R+11$.

Comparison of the graphs in fig. 4.1.2.c shows that the conjunctival epithelial lesions occur earlier than the corneal lesions, as was to be expected from the clinical description.

The finding that the incidence of the keratitis is lower than the incidence of the conjunctival epithelial lesions, and the observation that even an extensive measles keratitis at the corneal side of the limbus can be stopped there, suggests the existence of a kind of protective mechanism.

Probably the development of neutralizing antibodies is the cause of this phenomenon, it is, however, not possible to answer this question from the available data.

4.2. "Exaggerated signs" and early corneal complications

In the previous paragraph the ocular signs of measles in 148 measles patients were described. A distinction was made between a prodromal subepithelial conjunctivitis and a strictly epithelial conjunctivo-keratitis. Both are signs of measles.

In this paragraph the early corneal sequelae, exceeding the normal keratitis, occurring during the initial stay in the hospital will be described. Out of the 248 children, seen daily in the Provincial Hospital, Kakamega and the St Elizabeth Hospital, Mukumu, 7 developed macro-erosions of the cornea. The pathogenesis of these macro-erosions is essentially the same as for the smaller lesions of the measles-keratitis. Their "exaggerated" size might quite well be relevant for the purpose of this study. For this reason they are described here separately and more extensively.

3 out of the 248 children developed exposure ulcers because of the inability to close their eyes. An exposure ulcer is to be considered as a real, early corneal complication of measles.
In the pilot study and in safari patients the same phenomena were observed: 5 macro-erosions and 2 exposure ulcers. These will not be described here.

Those complications to be described in § 4.4 are considered late complications, when they developed after the acute stage of measles, but within a time span of 3 months after the rash. The period of 3 months was chosen arbitrarily to be to some extent sure of the relationship with measles. Late complications were less frequently seen than the early ones. They are far more difficult to trace: the mortality in complicated cases of measles is higher, and in the bush medical help is seldom searched for. The restriction that the complication had to be seen within 3 months after the rash greatly reduced the number of patients with late complications (9) who could be included in this study.

No late or severe ocular complications were seen in the 248 children in Kakamega and Mukumu, and no permanent ocular damage was observed in these children, apart from a single patient with minimal corneal nebulae.

4.2.1. Central corneal macro-erosions

7 out of the 248 patients – 4 boys and 3 girls – developed a central corneal erosion: in the central part of the cornea the epithelium was shed, without involvement of Bowman’s membrane or the stromal tissue. The limbus always stayed free. Fig. 4.2.1.a (see colour plate III) is a good example of a corneal erosion after measles.

In 3 cases the erosions were bilateral, in 4 cases only one eye was affected. The erosions developed at day R + 1, R + 1, R + 1, R + 1, R + 8, R + 8 and R + 9.

In some cases the erosions developed in an area of active measles-keratitis, of which the patient presented in Fig. 4.2.1.b (see colour plate III) and c is a good example.

The treatment consisted of tetracyclin 1% eye ointment topically, along with padding of the affected eye(s). The children also had a capsule of 200,000 iU Vitamin A. The erosions healed within 1–5 days.

Especially in some safari-patients it was observed that the corneal erosion was localized in the exposed part of the cornea: some children were so debilitated that closure of the eyelids became a problem. Probably this is a transition to the exposure ulcers to be described in the next paragraph.

Patient M 245 (fig. 4.2.1.d.) is an example of such an erosion.

This 9 months old girl developed a corneal erosion at the six o’clock position on the first day after outbreak of the rash. She was treated with tetracycline eye-ointment, 6–10 times daily. The ever-present mother was instructed to keep the eyelids closed. No pad and bandage were applied, because of the extremely bad condition of the child. She also had a capsule of 200,000 iU Vitamin A. The cornea healed within 3 days without sequelae.
Figure 4.2.1.c. The same patients (M 149) as fig. 4.2.1.b. The course of the measles keratitis and the subsequent corneal erosion is given here in full detail. The cornea was sent for electronmicroscopy (fig. 5.3.b and c)
In table 4.2.1.e. the data regarding age, sex and systemic complications of the 7 measles patients with corneal erosions are tabulated.

<table>
<thead>
<tr>
<th>Patients number</th>
<th>Sex</th>
<th>Age in years</th>
<th>Complications</th>
<th>IgM</th>
</tr>
</thead>
<tbody>
<tr>
<td>M 61</td>
<td>M</td>
<td>1.2/12</td>
<td>Meningitis</td>
<td>+</td>
</tr>
<tr>
<td>M 113</td>
<td>F</td>
<td>3</td>
<td>Pneumonia</td>
<td>+</td>
</tr>
<tr>
<td>M 149</td>
<td>M</td>
<td>1.6/12</td>
<td>Virus Pneumonia</td>
<td>-</td>
</tr>
<tr>
<td>M 223</td>
<td>M</td>
<td>1.9/12</td>
<td>Pneumonia</td>
<td>nd</td>
</tr>
<tr>
<td>M 241</td>
<td>M</td>
<td>7/12</td>
<td>Pneumonia</td>
<td>-</td>
</tr>
<tr>
<td>M 245</td>
<td>F</td>
<td>9/12</td>
<td>-</td>
<td>nd</td>
</tr>
<tr>
<td>K 19</td>
<td>F</td>
<td>2.2/12</td>
<td>-</td>
<td>nd</td>
</tr>
</tbody>
</table>

nd = not done

4.2.2. Exposure ulcers

3 children out of the total group of 248 patients developed real exposure ulcers because of their inability to close their eyes. Fig. 4.2.2.a (see colour
Table 4.2.2.c: The sex, age, general complications and IgM estimation in 3 patients with exposure of the cornea. Patients M 7 and M 207 died. The cornea of patient M 207 was available for microscopic examination.

<table>
<thead>
<tr>
<th>Patients number</th>
<th>Sex</th>
<th>Age in years</th>
<th>Complications</th>
<th>IgM</th>
</tr>
</thead>
<tbody>
<tr>
<td>M 7</td>
<td>M</td>
<td>4</td>
<td>Malnutrition</td>
<td>+ later on -</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>+ TB Meningitis</td>
</tr>
<tr>
<td>M 117</td>
<td>M</td>
<td>5</td>
<td>--</td>
<td>+</td>
</tr>
<tr>
<td>M 207</td>
<td>M</td>
<td>26/12</td>
<td>Dehydration</td>
<td>nd</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Malnutrition</td>
<td></td>
</tr>
</tbody>
</table>

nd = not done

Plate III) shows a typical example of such an exposure ulcer. Pat M 7, a 4 year old boy developed exposure erosions, 3 days after outbreak of the rash, in both eyes. These erosions reacted favourably to treatment. Three weeks later an exposure ulcer suddenly appeared in the right eye. The next morning the boy died.

A 2½ year old boy was admitted, 9 days after the outbreak of the rash, with an exposure ulcer at the six o’clock position in his right eye. The next morning the boy died. The cornea was available for study under the electron-microscope.

Pat M 117, a 5 year old boy, developed a typical exposure at day R + 5, four days after his admission. In fig. 4.2.2.b the clinical course of his exposure ulcers is depicted in more detail. The corneae healed within 5 days with the development of corneal nebulae.

The data regarding age, sex, IgM estimations and systemic complications are given in table 4.2.2.c.

When the patients with the corneal macro-erosions are compared with the patients with exposure ulcers, clinically two differences are apparent:

(a) The patients with corneal erosions have or had a more severe measles-keratitis than the children with exposure ulcers.

(b) The children with exposure ulcers were generally more ill than the children with the central erosions.

These observations suggest that the corneal erosions are a direct progression of the measles-keratitis. This view is confirmed by the findings in the electron-microscope. (§5.3).

On the other hand, the exposure ulcers appear to be more correlated with the severity of the systemic disease than with the measles-keratitis.

Exposure ulcers are therefore supposedly not a specific sequelae of the measles-keratitis, in contrast to the erosions.

It goes without saying that in some patients both factors — viral keratitis and exposure — can cooperate to cause corneal damage.
Figure 4.2.2.b. The course of an exposure ulcer in a 5 year old boy with measles
4.3. The prophylactic value of tetracycline eye-ointment 1% and Vitamin A 200,000 iU

4.3.1. Measles-keratitis

In the first 148 patients we didn’t interfere with the treatment as prescribed by the attending doctor. So 84 patients were prescribed tetracycline ointment because of the red aspect of their eyes.

At first sight it may seem confusing that the treated patients had a more severe keratitis than the non-treated children (average keratitis score 7 compared to 4). Probably this will only mean that the extent of the subepithelial conjunctivitis and the severity of the epithelial keratitis are associated.

In the clinical trial (Mukumu II and Kakamega) evaluation of tetracycline ointment as a prophylactic for the measles-keratitis was attempted. The numbers, however, were too small to allow for a statistical analysis. Anyhow, here no gross differences between the treated and non-treated eyes were observed.

It must also be said that it would be very unlikely that tetracycline ointment could have a preventive value against the viral measles-keratitis.

The limited numbers in our clinical trial also don’t allow for any statement about the prophylactic value of Vitamin A, as far as the measles-keratitis is concerned.

4.3.2. Corneal erosions and exposure ulcers

In the whole group of 248 measles children 10 patients with corneal disease exceeding the common measles-keratitis were observed: 7 corneal erosions and 3 exposure ulcers. None of these corneal problems occurred in children who had received tetracycline eye-ointment. It is to be remembered that of the 148 children in Mukumu I, 84 children had received eye-ointment: generally they had a more severe keratitis, but did not develop corneal erosions. Since corneal erosions develop as an extension of the measles-keratitis, this is a surprising finding. In Mukumu II and Kakamega in every child one eye was protected with tetracycline eye-ointment. Both corneal erosions which were seen to develop during the observation-period occurred in unprotected eyes. The details of these 10 patients are given in fig. 4.3.2.

The same protective value of tetracycline eye-ointment 1% was observed in the safari-patients and no erosions or exposure ulcers were observed in children who had received eye-ointment. In many dispensaries the prescription of eye-ointment is a common routine measure for measles.

These observations demonstrate the prophylactic value of tetracycline eye-ointment in the prevention of erosions and exposure ulcers in early measles.

In table 4.3.2. one patient is to be found who developed a corneal erosion 3 days after the administration of 200,000 iU Vit A (fig. 4.2.1.a, see colour plate III).
Figure 4.3.2. The occurrence of early corneal complications in children with measles in correlation with the time after outbreak of the rash (R) and the prescribed treatment. No erosions or exposure ulcers were seen to develop in eyes treated with tetracycline eye-ointment.

This single observation does not allow a statement about the prophylactic value of Vitamin A in the prevention of corneal disease, because of the limited number of patients in the trial groups.
4.4. Late ocular complications

On our safaris and at the open Eye Clinic Mukumu, 9 patients were seen with late ocular complications after measles.

Late complications are defined as complications occurring after the acute stage of the measles, but within 3 months after the rash.

As always with work originating from the bush of developing countries, many interesting data are missing and follow-up examination is nearly always lacking. The available data regarding sex, age, time of outbreak of the rash and clinical appearance of the cornea are tabulated in table 4.4.a.

Pat Tamb VA 120. (fig. 4.4.b: see colour plate I).

This 6 year old boy was seen 2 weeks after the measles rash had come out. At the time of first examination he had a perforation at the 6 o’clock position in the cornea of the left eye. Traditional herbal medicines had been applied by his mother since one week. It was impossible to retrieve details about the treatment. The boy belonged by tribe to the Elgeyo’s. According to Mulder

Table 4.4.a. Patients with late complications after measles.

<table>
<thead>
<tr>
<th>Number</th>
<th>Sex</th>
<th>Age in years</th>
<th>Rash</th>
<th>Clinical aspect of the cornea</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tamb VA 120 (Fig. 4.4.b)</td>
<td>M</td>
<td>6</td>
<td>2 wk</td>
<td>Perforation at 6 o’clock</td>
<td>TM +</td>
</tr>
<tr>
<td>Muk P 258 (Fig. 4.4.c-d)</td>
<td>F</td>
<td>2.8/12</td>
<td>2 wk</td>
<td>RE: panophthalmitis LE: adherent leucoma</td>
<td>cornea EM TM + Fig. 5.3 d-h.</td>
</tr>
<tr>
<td>Tamb P 273 (Fig. 4.4.e)</td>
<td>M</td>
<td>1.4/12</td>
<td>3 wk</td>
<td>Hypopyon ulcer</td>
<td>TM +</td>
</tr>
<tr>
<td>Chw P 172 (Fig. 3.3)</td>
<td>F</td>
<td>4</td>
<td>4 wk</td>
<td>Herpetic keratitis</td>
<td>TM +</td>
</tr>
<tr>
<td>Tamb VA 28 (Fig. 4.4.f-g)</td>
<td>F</td>
<td>8</td>
<td>4 wk</td>
<td>RE: perforated cornea LE: Phlyctenular keratitis</td>
<td>TB</td>
</tr>
<tr>
<td>Muk VA 7</td>
<td>M</td>
<td>2.6/12</td>
<td>4 wk</td>
<td>Healing corneal ulcer</td>
<td></td>
</tr>
<tr>
<td>Bus P 158</td>
<td>F</td>
<td>1.2/12</td>
<td>5 wk</td>
<td>Herpetic keratitis</td>
<td></td>
</tr>
<tr>
<td>Sia VA 16 (Fig. 4.4.h)</td>
<td>M</td>
<td>7/12</td>
<td>5 wk</td>
<td>Central corneal necrosis</td>
<td></td>
</tr>
<tr>
<td>Bung VA 83 (Fig. 4.4.i)</td>
<td>F</td>
<td>1.6/12</td>
<td>8 wk</td>
<td>Corneal staphyloma</td>
<td></td>
</tr>
</tbody>
</table>
(personal communication) it is unthinkable that an Elgeyo mother will ever serve a meal to her family consisting of only "posho" (maize) without "mboga" (green leafy vegetable of any kind). This excludes more or less Vitamin A deficiency, of which also no other symptoms or signs were present.

It is remarkable that also the mother had an adherent leucoma in the left eye, which — according to her — had developed in connection with measles.

Pat Muk P258. (fig. 4.4.c and d: see colour plate I).
This 2 year 8 month old, severely malnourished girl was seen at the open Eye Clinic Mukumu, 2 weeks after the rash. The right eye was lost because of a panophthalmitis. An evisceration was performed, a purulent hyalitis was found. The corneal-scleral rim was preserved for light and electron-microscopy. On electron-microscopical examination possibly measles virus was detected in the keratocytes (fig. 5.3.h). The left eye showed a descemetocele, with incarceration of iris.

Pat Tamb P273. (fig. 4.4.e: see colour plate I).
This Elgeyo boy, 1 year and 4 months of age, was seen 3 weeks after the outbreak of the rash. The cornea of the right eye showed a small paracentral ulcer, and a secondary iritis with hypopyon was present. A smear was taken from the ulcer and Gram stained. No bacteria were seen in sufficient amounts to be blamed as the cause of the ulceration. The attending mother admitted the application of traditional medicines. It has been mentioned already that the application of herbs frequently gives rise to an iritis (Duke Elder and MacFaul, 1972). After treatment with atropine ointment, chloramphenicol ointment and pad and bandage, the ulcer healed and the hypopyon disappeared. 200,000 iU Vitamin A were also given.

Pat Chw P172 (fig. 3.3: see colour plate I).
Luhya girl, 4 years of age. Herpetic keratitis, 4 weeks after the measles. Traditional medicines had been applied.

Pat Tamb VA28. (fig. 4.4.f and g: see colour plate I).
This 8 year old Kalengi girl was seen for the first time 4 weeks after the rash. The girl made a lamentable impression; she was extremely photophobic and cried continuously. The scalp was covered with impetigo, bald areas alternated with the remaining red hair. The submandibular lymphnodes were swollen and tender. The chest X-ray was very suggestive for pulmonary tuberculosis. In the left eye several phlyctae were visible; the cornea of the right eye had perforated at the six o'clock position. The child was treated with systemic antibiotics and tuberculostatics. The reaction to treatment was good. Four months later the child was seen again in consultation. The perforation in the right eye had healed with an adherent leucoma. The cornea of the left eye was clear.

Pat Sia VA16. (fig. 4.4.h: see colour plate I).
This 7 month old boy was seen 5 weeks after the outbreak of the rash. The cornea of the left eye showed a central necrosis. The perforation was plugged off by the lens, the anterior chamber was virtually non-existent. The child
was lost for treatment and follow-up. No Bitot’s spots were seen. The child was breastfed, supplemented with cow’s milk.

Pat Bung VA 38. (fig. 4.4.i: see colour plate I).
This 18 month old girl was seen 2 months after the measles. The child was extremely well fed. The right eye showed a nearly total corneal staphyloma. No Bitot’s spots were observed, and she did not complain about night blindness.

Pat Muk VA 7.
This 2½ year old Luhya boy was seen with a corneal ulcer, 4 weeks after the measles rash. The ulcer had not perforated and was in transition to a corneal leucoma. No data regarding his nutritional status are available.

Pat P 158.
A 14 month old Luhya girl was seen with an herpetic keratitis, 5 weeks after the measles rash. No other data are available.

These late complications after measles are much more varied than the early ones. In some cases it was not possible to make a distinct etiological diagnosis, but supposedly the application of traditional medicines, tuberculosis and herpes are important etiological factors.

The nutritional data of these patients are given in table 6.4.
50

5 — Immunofluorescence-, light- and electron-microscopy of conjunctival biopsies and corneal specimens

In the previous paragraphs a clinical description was given of the subepithelial conjunctivitis and the epithelial conjunctivo-keratitis as signs of measles. This chapter will deal with the anatomical features of these conjunctival and corneal lesions.

Immunofluorescence for measles virus still gives positive results in skin biopsies up to 4 days after the outbreak of the rash, whereas measles virus can only be cultured exceptionally after 36—48 hours. Moreover, storage of specimens at 4°C for 48 hours does not influence the quality of the immunofluorescence tests in cryostate sections (Olding-Stenkvist and Bjorvatn, 1976). This means that immunofluorescence tests are less vulnerable than culturing techniques. A major advantage of immunofluorescence, important in view of the clinical differentiation made in the two conjunctival signs of measles, is the possibility to localize the virus.

Localization of virus by the electronmicroscope is a far more elaborate enterprise, but allows more detailed insight in the pathological processes occurring at the cellular level.

5.1. Immunofluorescence of conjunctival biopsies

5.1.1. Technique

After thawing, the specimens (§3.6) were taken out of the transport medium and snapfrozen in isopentane, chilled by liquid nitrogen. Cryostate sections of 1 μ thickness were cut on a Slee microtome.

The slides were coded and independently prepared and examined in two different laboratories: Dr. F.E. Nommensen of the Department of Virology and Dr. F.J.W. ten Kate of the Institute of Pathology, both at the Erasmus University, Rotterdam. Both laboratories used essentially the same techniques and materials.

The slides were air dried and stored at −70°C. The air dried specimens were fixed in acetone at −20°C for 10 minutes. After air drying at room temperature the specimens were incubated for 45 minutes with commercially available F.I.T.C. (Fluorescein Iso Thiocyanate Conjugated) anti-measles-goat-serum (Microbiological Associates Inc. Walkersville, Maryland, USA) in an optimal 1:40 dilution. After washing in phosphate buffered saline, the slides were mounted in buffered glycerol 80% (pH 7.4) and examined under a Zeiss fluorescence microscope with incident ultraviolet light.

To assess the specificity of this technique, control experiments were done with the following results.
(a) Lissamine Green had no toxic effect on a Vero cell culture, (Green Monkey kidney cells) and did not influence the auto-fluorescence of these cells.

(b) Lissamine Green had no affinity to cells infected with a Schwarz measles strain in vitro.

(c) The undiluted conjugated serum did not stain respiratory syncytial virus, para-influenzavirus type 1, 2 and 3, mumps virus, influenza virus A and B, herpes simplex virus type I and adenovirus (not typed).

(d) Some slides were incubated with F.I.T.C. antiserum against adenovirus or herpes implex virus type I. Both experiments were negative.

These controls make it very likely that the fluorescence present in our slides demonstrates specifically the presence of measles virus antigen.

It was also tried to do immunofluorescence tests on Epon embedded material, after acid elution (Fulton and Middleton 1975), but all attempts failed to give positive results.

### 5.1.2. Results of the immunofluorescence tests on 5 conjunctival biopsies

The data regarding the children from whom the biopsies were taken are tabulated in table 5.1.2.a.

Culturing of the measles virus from these biopsies was tried, but all cultures stayed negative. This is not surprising in view of the time after the rash at which the biopsy was taken. In a single case (no. 3) a positive culture might have been possible, but probably here also the hazards of long-range transportation and delay have influenced the negative outcome.

Together with the biopsies a venous bloodsample was drawn to assess the presence of anti-measles IgM. The technical difficulties appeared unsurmountable and no IgM estimations were performed on these sera.

Fig. 5.1.2.b (see colour plate III) demonstrates the presence of measles virus antigen in the subepithelial tissue.

In fig. 5.1.2.c (see colour plate III) an example of the immunofluorescence of the conjunctival epithelium is given. The fluorescent activity is localized in the cytoplasm and demonstrates specifically the presence of measles virus antigen.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Sex</th>
<th>Age in years</th>
<th>R</th>
</tr>
</thead>
<tbody>
<tr>
<td>KNH 1</td>
<td>M</td>
<td>2</td>
<td>+ 3</td>
</tr>
<tr>
<td>KNH 2</td>
<td>M</td>
<td>6/12</td>
<td>+ 5</td>
</tr>
<tr>
<td>KNH 3</td>
<td>M</td>
<td>4</td>
<td>+ 1</td>
</tr>
<tr>
<td>KNH 4</td>
<td>F</td>
<td>1.1/12</td>
<td>+ 4</td>
</tr>
<tr>
<td>KNH 5</td>
<td>M</td>
<td>8/12</td>
<td>+ 4</td>
</tr>
</tbody>
</table>

Table 5.1.2.a. Sex, age of the children and time of biopsy taking in relation to the day of outbreak of the rash (R). These biopsies were sent to the Erasmus University Rotterdam for examination with immunofluorescent techniques.
Table 5.1.2.d. The distribution of antimeasles immunofluorescence in 5 paired conjunctival biopsies, staining and non-staining with Lissamine Green (LG + and LG -), performed by 2 different laboratories (Lab A = Dpt. of Virology; Lab B = Dpt. of Pathology, Erasmus University Rotterdam). In the Lissamine Green positive epithelial lesions measles antigen was present. In other biopsies viral activity was found subepithelially, as manifestation of the subepithelial conjunctivitis of prodromal measles.

<table>
<thead>
<tr>
<th></th>
<th>Epithelium</th>
<th>Subepithelium</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Lab A</td>
<td>Lab B</td>
</tr>
<tr>
<td>Pat 1</td>
<td>LG +</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>LG -</td>
<td>-</td>
</tr>
<tr>
<td>Pat 2</td>
<td>LG +</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>LG -</td>
<td>-</td>
</tr>
<tr>
<td>Pat 3</td>
<td>LG +</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>LG -</td>
<td>?</td>
</tr>
<tr>
<td>Pat 4</td>
<td>LG +</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>LG -</td>
<td>-</td>
</tr>
<tr>
<td>Pat 5</td>
<td>LG +</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>LG -</td>
<td>-</td>
</tr>
</tbody>
</table>

All results of the direct anti-measles immunofluorescence tests of both laboratories are tabulated in table 5.1.2.d.

From this table two conclusions are to be drawn.

(a) An excellent concordance exists between the results of both laboratories. It is to be remembered that these results were obtained on coded specimens, which makes the next conclusion even more reliable.

(b) In the specimens taken from Lissamine Green positive areas the measles virus is located in the epithelium; in the Lissamine Green negative areas no virus could be detected in the epithelium. In many of the LG negative areas viral activity was found in the subepithelial lymphoid tissue.

These findings fit into the previously described clinical distinction between a subepithelial conjunctivitis and an epithelial conjunctivo-keratitis.

It is of great importance that the viral nature of the conjunctivo-keratitis in measles could be proved by these observations and this may have far reaching consequences for our thinking about the pathogenesis of Post-Measles-Blindness.

5.2. Light- and electron-microscopy of conjunctival biopsies

5.2.1. Techniques

The 5 paired conjunctival biopsies and the corneal specimens (§3.6) were fixed in a mixture of glutaraldehyde and paraformaldehyde (Peters 1970).

For light-microscopy small pieces were embedded in paraplast; 7 μm
sections were stained with either periodic acid Schiff (PAS) or with hema-
toxylinesin (HE).

For electron-microscopy small blocks were postfixed for 1 hour in os-
miumtetroxyde (Palade 1955), dehydrated in graded series of ethanol and
subsequently embedded in Epon 812. Ultrathin sections were made on a
Reichert OmU 3 ultramicrotome, stained with uranylacetate and leadcitrate
and studied in a Philips EM 400.

In order to experience the normal appearance of measles virus under the
electronmicroscope, a measles infected Vero cell culture (Green Monkey
Kidney cells) served as a control. After centrifugation this cell culture was
treated in the same way as the conjunctival biopsies. Moreover, some thick
sections (0.5 \( \mu \)m) were investigated by a combined application of scanning
and transmission electronmicroscopy (STEM).

5.2.2. Light microscopy of conjunctival biopsies

The data regarding the patients from whom the biopsies were taken are
tabulated in table 5.2.2.a.

Epithelium

In many places the epithelium shows no abnormality: the cells are regularly
arranged, occasionally a goblet cell is seen between light and dark stained
apical cells. In many basal cells the cytoplasm contains melanosomes, arranged
at the apex of the nucleus.

In specimens, taken from Lissamine Green positive areas, the intercellular
spaces seem to be widened, and islands of epithelium are seen where cells are
desquamated (fig. 5.2.2.b., c and e); at some places hardly any epithelium
is left (fig. 5.2.2.d.)

These areas of desquamation showed a patchy distribution and were
bordered by areas of normal epithelium (fig. 5.2.2.e).

Some epithelial giant cells were observed.

In a single specimen, from a 1 year old girl (Bung P 265) with a con-
junctival xerosis (but with no other signs and symptoms of Vitamin A de-
ficiency), was also found a proliferation of the basal epithelium, with the

Table 5.2.2.a. Data regarding sex, age and day after outbreak of the
rash (R) at which conjunctival biopsies for pathology and electron-
microscopy were taken from 5 measles patients.

<table>
<thead>
<tr>
<th>Pat nr.</th>
<th>IOI nr.</th>
<th>Sex</th>
<th>R</th>
<th>Age in years</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>M 223</td>
<td>77141</td>
<td>F</td>
<td>-2</td>
<td>1.9/12</td>
<td></td>
</tr>
<tr>
<td>M 234</td>
<td>77143</td>
<td>F</td>
<td>+1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Bung P 179</td>
<td>77142</td>
<td>F</td>
<td>+2</td>
<td>1.9/12</td>
<td>xerosis</td>
</tr>
<tr>
<td>Bung P 265</td>
<td>7841</td>
<td>F</td>
<td>+2</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Bus P 216</td>
<td>7839</td>
<td>M</td>
<td>+3</td>
<td>10/12</td>
<td></td>
</tr>
</tbody>
</table>
Figure 5.2.2.b. Conjunctiva: the central part of the epithelium (between arrowheads) shows loss of intercellular cohesion. Dyskeratotic and necrotic cells are seen. In the subepithelial tissue infiltration by mononuclear cells is present. Hematoxylin-Eosin × 100. Pat Bus P 216, Negative PA EUR 28092

Figure 5.2.2.c. In the same patient, now at a higher magnification, the loss of intercellular adhesion is demonstrated by the presence of intercellular clefts (arrowhead). Hematoxylin-Eosin × 125. Negative PA EUR 28092
Figure 5.2.2.d. In this conjunctival biopsy (Pat Bung P 265) severe oedematous changes are present in the subepithelial tissue. In some places (arrowheads) all epithelium has vanished: the basement membrane is left bare. Hematoxylin-Eosin × 100, Negative PA EUR 28092

Figure 5.2.2.e. In this conjunctival biopsy the desquamation of the epithelium is seen to take place: on the left hand side the epithelium is desquamating, whereas on the right hand side the epithelium is normal. Hematoxylin-Eosin × 40, Pat Bus P 216, Negative PA EUR 28092
Figure 5.2.2.f. The same patient as the previous fig. The desquamating epithelium is seen at a higher magnification (×100)

Figure 5.2.2.g. Conjunctival biopsy from a xerotic area: focal loss of coherence of epithelial cells with keratinization of superficial layers. The subepithelial tissue shows severe oedema. Pat Bung P265, Hematoxylin-Eosin ×125. Negative PA EUR 28093
formation of pegs below the basement membrane. This was also the only specimen with signs of keratinization of the conjunctival epithelium (fig. 5.2.2.g).

Inclusion bodies were not observed in any of the specimens.

**Substantia propria**

In most cases the substantia propria underlying the epithelium showed little signs of inflammation. In other biopsies a severe oedema (fig. 5.2.2.d) is present. A moderate vasodilation is seen in other specimens. Occasionally an infiltration by mononuclear cells is observed. No inclusion bodies or giant cells were found.

### 5.2.3. Electron microscopy of conjunctival biopsies

**Epithelium**

Many areas with normal epithelium are observed. The epithelial cells show the characteristic ultrastructural features, the microvilli extend from the apical cell membrane, the nuclear chromatin is condensed at the nuclear border.

In other places degenerating cells with pycnotic nuclei are found. The cytoplasm of the apical cells seems relatively empty: inside the cytoplasm mitochondria, ribosomes and endoplasmic reticulum are unevenly distributed. A prominent feature is the presence of dilated intercellular spaces. The lateral cell membrane shows its typical deep indentations, studded with desmosomes, which seem to link the cells together, even when the intercellular spaces are strongly dilated (fig. 5.2.3.a). Occasionally a goblet cell is seen (fig. 5.2.3.b).

**Substantia propria**

The collagen fibrils are irregularly oriented and in some cases clumped together to form amorphous structures. In between this collagen many cells are observed, mainly plasma cells, lymphocytes and macrophages; occasionally polymorphonuclear leucocytes are seen (fig. 5.2.3.c). The cytoplasm contains an often swollen endoplasmic reticulum. Multivesicular bodies and bristle coated vesicles (which resemble lysosomal structures) are seen in great numbers.

In the cytoplasm cytoplasmic aggregates and accumulations of filamentous material are present (fig. 5.2.3.d). These strands measured 20 nm in thickness and resemble the strands found in measles infected vero cells (fig. 5.2.3.e and f)(cfr fig. 2.2.b). This finding suggests that the granular material inside the cells is of a viral nature.
Figure 5.2.3.a. Electron microscopy of conjunctival epithelium. The apical cell border of the superficial cells shows characteristic microvilli. The cytoplasm of the cells differs markedly in electron density, melanin granules (Me) are frequently observed. The intercellular spaces are dilated (arrow). Pat M 234, Negative IOI 79088, magnification 6900 x.

Figure 5.2.3.b. Electron micrograph of a goblet cell (G) between epithelial cells with the adjacent part of the stroma. In the cytoplasm, numerous spherical globules with different electron densities are seen. Pat M 234, Negative IOI 79058, Magnification 6900 x.

Figure 5.2.3.c. Electron micrograph of a polymorphonuclear leucocyte in the conjunctival stroma. The cytoplasm contains numerous vesicles of lysosomal nature: large autophagic vacuoles (AV) and electron-dense bodies (EDB). The nuclear lobes (NUC) are indicated with arrowheads. Pat Bung P 265, negative IOI 79117, magnification 5900 x.
Figure 5.2.3.b.

Figure 5.2.3.c.
Figure 5.2.3.d. Electron micrograph at high magnification of a lymphocyte in the conjunctival stroma. Dark mitochondria (Mi) and a swollen endoplasmic reticulum (ER) are seen in the cytoplasm. Note the fine filamentous material, which may be viral nucleocapsid (NC). Pat M 234, Negative IOI 79023, magnification 15040 x
Figure 5.2.3.e. Scanning transmission electron micrograph of a cultured vero cell, infected with measles vaccine. The nucleocapsid (NC) is clearly visible. A virion (arrowhead) is seen with the fuzzy coat. The cellular membrane shows in several places a slight increase in cell coat (arrows) material. Negative IOI 79304, magnification 28600 X
Figure 5.2.3.f. Scanning transmission electron micrograph shows part of a cultured cell after infection with measles vaccine. The cytoplasm is densely packed with nucleocapsid aggregates (asterisk) and some degenerated membranous structures (arrow) are found. Negative IOI 790496, magnification 50400 x.
5.3. Light and electron microscopy of corneal specimens

Three corneal specimens were available for study under the microscope. The data regarding the patients are given in table 5.3.a.

**Epithelium**

In all 3 specimens the corneal epithelium showed essentially the same features as found in the conjunctival epithelium. At some places the epithelial cells show microvilli at the apical surface, covered by a fine thin filamentous layer, whereas in other parts the epithelium has diminished in thickness, and at some places even only some cell fragments on Bowman's membrane are left (fig. 5.3.b. and c). The intercellular spaces are again widely dilated. The anterior part of the epithelium only shows few cytoplasmic organelles: mitochondria, rough endoplasmic reticulum and Golgi apparatus are scarcely found.

**Stroma**

No abnormalities were observed in Bowman's membrane. In most cases the corneal collagen is normally arranged in bundles with uninterrupted periodicity. No signs of collagen breakdown were found.

Marked changes were found in the cornea with the perforation. Fig. 5.3.d gives the light microscopy. The scar of the corneal perforation is visible, with incarcerated iris pigment. An electronmicroscopical specimen taken from the newly formed scar-collagen, demonstrates the irregular arrangement of the collagen fibres. (fig 5.3.e.)

In the area of the microscopically unaltered cornea the collagen bundles have retained their normal periodicity. Here however many abnormal keratoocytes are found. The cytoplasm of these cells contains swollen mitochondria and large vacuoles and fat inclusion bodies. Celllysis is frequently present, together with disruption of cytoplasmic membranes. Occasionally fine filamentous material, which shows great similarities to the viral strands in the cultured vero cells, can be seen. (fig 5.3.f, g and h)

---

**Table 5.3.a. Data regarding sex, age and time after the outbreak of the rash (R) of the patients with corneal complications after measles, whose corneae were available for pathological examination.**

<table>
<thead>
<tr>
<th>Patient</th>
<th>IOI</th>
<th>Sex</th>
<th>Age in years</th>
<th>R</th>
<th>Corneal condition</th>
<th>Figure</th>
</tr>
</thead>
<tbody>
<tr>
<td>M 149</td>
<td>7702</td>
<td>M</td>
<td>1.6/12</td>
<td>+11</td>
<td>Central macro-erosion</td>
<td>4.2.1.b-c,</td>
</tr>
<tr>
<td>M 207</td>
<td>77139</td>
<td>M</td>
<td>2.6/12</td>
<td>+10</td>
<td>Exposure ulcer</td>
<td></td>
</tr>
<tr>
<td>Muk P258</td>
<td>7838</td>
<td>F</td>
<td>2.8/12</td>
<td>+14</td>
<td>Corneal perforation</td>
<td>4.4.c and d.</td>
</tr>
</tbody>
</table>
Figure 5.3.b. Electron micrograph of the corneal epithelium. The number of epithelial cell-layers is reduced. The apical cell surface is lacking microvilli, numerous cytoplasmic vesicles can be observed. The basal lamina is intact. Pat M 149, Negative IOI 78418, magnification 6160 x
Figure 5.3.c. Electron micrograph of the corneal epithelium of a patient with a corneal erosion: most of the epithelial cells have disappeared, only some disrupted membranes are found. Bowman's membrane (BM) is lightly wrinkled. Pat M 149, Negative IOI, magnification 10,560 x

Figure 5.3.d. Scar of corneal perforation with incarcerated iris pigment epithelium. This patient was seen 2 weeks after the outbreak of the measles rash. The 2½ year old girl was severely malnourished. The clinical impression "panophthalmitis" was confirmed on operation (fig. 4.4.c and d). Pat Muk P258, Hematoxylin-Eosin x 16, Neg PA EUR 28093
Figure 5.3.e. Electron micrograph of the scar of the perforation. A lymphocyte with numerous cytoplasmic electron dense vesicles is found in between the irregularly arranged collagen bundles. Pat Muk P 258, Negative IOI 79088, magnification 6900 X

Figure 5.3.g. The same electron dense material as seen in the previous figure, now at a higher magnification. Pat Muk P 258, Negative IOI 8000257, magnification 22,000 X
Figure 5.3.f. Electron micrograph of the normal corneal stroma of the perforated eye. The collagen bundles are regularly oriented. A large keratocyte is observed. In between the pseudopod-like protrusions of the keratocyte fine electron dense material is seen. Pat Muk P 258, Negative IO1 800256, magnification 78 0 x
5.4. Discussion

The immunofluorescence tests on the conjunctival epithelium demonstrated the presence of measles antigen. Also the patchy distribution of these lesions is in concordance with the clinical findings. This leads to the conclusion that in the clinically observed epithelial lesions the measles virus plays a prominent role. No explanation can be given for the preferential localization of these lesions in the exposed parts of the bulbar conjunctiva.

No giant cells or inclusion bodies were observed. It is to be realized,
however, that inclusion bodies and giant cells need some days to develop (Czajkowsky and Heneen 1976). Probably the time between the first appearance of these lesions and the taking of the biopsy was just too short for such a development. For the same reason it need not be surprising that in the electronmicroscope no viral strands were found, whereas viral antigen was observed in the cytoplasm by immunofluorescence. Immunofluorescence is a far more sensitive technique than light or electronmicroscopy. Further information could well be given by the use of ferritin labeled antiserum on specimens for electronmicroscopy.

The involvement of the conjunctival stroma in the measles infection is — at the time when biopsies were taken — of a much longer duration. This gives the time to develop the morphological correlates of the viral activity: viral strands were seen in the electronmicroscope, together with positive immunofluorescence.

Here also, no giant cells were found. It might be possible that the significance of measles giant cells in "normal" measles is exaggerated. Also Olding-Stenkvist and Bjorvatn (1976) reported a low incidence (cfr Roberts and Bain 1958).

In conclusion: the findings of the immunofluorescence and light- and electronmicroscopy fully support the distinction clinically made between the two conjunctival signs of measles: an early prodromal subepithelial conjunctivitis and an epithelial conjunctivo-keratitis occurring at the time of the rash.

One conjunctival biopsy was (on purpose) taken from a (non-staining) xerotic area. In the light microscopy keratinization of the superficial epithelium and the development of a rete peg were observed. Both changes are considered specific for a Vitamin A deficiency, as found in an experimental animal study by Pfister and Renner (1978).

These changes occurred only in this one specimen, selected on clinical grounds, as with a conjunctival xerosis measles-patient. The microscopy is totally different from the one seen in measles: measles conjunctivitis and Vitamin A deficiency are two, totally different, entities.

The conjunctival and corneal epithelium showed the same pathological changes: a diminished epithelial adhesiveness leading to patchy desquamation of the epithelium. In my opinion this is also to be considered as the cause of the high incidence of corneal erosions in measles-keratitis (cfr Khodadoust et al. 1968; Fogle et al. 1975).

Inside the keratocytes of the perforated cornea a granular material was found. It shares some morphological characteristics with viral RNA strands, and it is very attractive to suppose that this material indeed represents measles virus. In my opinion it is not allowed to state this positively. A study of the cornea with ferritin-labeled antiserum is badly needed.
6 - The nutritional status of the children with measles-keratitis and corneal complications

In the literature not only complications of measles are consistently associated with malnutrition, but also the signs of measles are more severe in malnourished children (§2.5).

For this study the same might apply and the question arises whether the ocular signs of measles (i.e. the measles-keratitis) are more severe in children in a worse nutritional state. A positive association would be very significant in view of the fact that blinding complications of measles occur more frequently in malnourished children.

In order to test the hypothesis of a possible association between the measles-keratitis and nutritional status a quantitative classification for the keratitis was devised (§3.3). This classification was compared with age, sex, history of immunization and anthropometric and biochemical parameters of the nutritional status. In the next paragraphs the tables will be given. In all tables the $\chi^2$-test was used for the statistical analyses.

6.1. Measles-keratitis and age, sex and history of immunization

In table 6.1.a the age of 126 measles patients and the classification for measles-keratitis are compared. No association exists.

In table 6.1.b the sex of the measles children and their keratitis classification are compared. Again, no statistical association was detected. In table 6.1.c the immunization history against measles is compared with the measles-keratitis. These two data are not associated. It is to be remembered however that the data regarding vaccination have only very limited value: the memory of the mother may be at fault (measles vaccination doesn't leave a specific scar like cowpox and BCG) and the vaccination itself may be of a bad quality:

<table>
<thead>
<tr>
<th>Age in months</th>
<th>Keratitis classification</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I</td>
<td>II</td>
</tr>
<tr>
<td>0-12</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>13-24</td>
<td>7</td>
<td>12</td>
</tr>
<tr>
<td>25-36</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>37+</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>total</td>
<td>25</td>
<td>30</td>
</tr>
</tbody>
</table>

$x^2_{12} = 4.01 \quad p = 0.98$
in developing countries it is very difficult to maintain the “cold chain” necessary for the transport of the vaccine to keep its effectivity. Anyhow, no association was found between the measles-keratitis and sex, age and immunization history. These factors therefore don’t have to be taken into account in the following statistical analysis of possible association of the measles-keratitis and nutritional status.

6.2. Measles-keratitis and nutritional status

To assess the nutritional status of our measles children anthropometric and biochemical parameters were used (§3.5). In table 6.2.a the Weight for Age, in table 6.2.b the Height for Age, in table 6.2.c the Weight for Height is compared with the keratitis classification. No statistical association was found between the measles-keratitis and these anthropometric parameters of nutritional status.

The same applies to the biochemical parameters of malnutrition: Serum Albumin and Retinol Binding Protein. The numbers are given in table 6.2.d and 6.2.e.

In conclusion: no statistical association could be detected between the extent and severity of the early virus related features of measles-keratitis and nutritional status in this group of 148 children.
Table 6.2a. The Weight for Age of 105 measles patients and the classification for measles-keratitis. No association was observed

<table>
<thead>
<tr>
<th>Keratitis classification</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
<th>V</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 60%</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>61–80%</td>
<td>8</td>
<td>7</td>
<td>13</td>
<td>12</td>
<td>12</td>
<td>52</td>
</tr>
<tr>
<td>&gt; 81%</td>
<td>11</td>
<td>13</td>
<td>8</td>
<td>4</td>
<td>12</td>
<td>48</td>
</tr>
<tr>
<td>total</td>
<td>20</td>
<td>22</td>
<td>22</td>
<td>17</td>
<td>24</td>
<td>105</td>
</tr>
</tbody>
</table>

\[ \chi^2_a = 9.33 \quad 0.30 < p < 0.50 \]

Table 6.2b. The Height for Age of the measles patients and the measles-keratitis. The extent of measles-keratitis was independent of the nutritional status – measured as Height for Age – of the measles children

<table>
<thead>
<tr>
<th>Keratitis classification</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
<th>V</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 90%</td>
<td>7</td>
<td>11</td>
<td>4</td>
<td>5</td>
<td>8</td>
<td>35</td>
</tr>
<tr>
<td>&gt; 91%</td>
<td>13</td>
<td>12</td>
<td>17</td>
<td>10</td>
<td>16</td>
<td>68</td>
</tr>
<tr>
<td>total</td>
<td>20</td>
<td>23</td>
<td>21</td>
<td>15</td>
<td>24</td>
<td>103</td>
</tr>
</tbody>
</table>

\[ \chi^2_b = 4.07 \quad 0.30 < p < 0.50 \]

Table 6.2c. The Weight for Height of the measles patients and the measles-keratitis. No association exists between the weight for height of the measles children and the extent of the measles-keratitis

<table>
<thead>
<tr>
<th>Keratitis classification</th>
<th>I</th>
<th>II</th>
<th>III</th>
<th>IV</th>
<th>V</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 90</td>
<td>7</td>
<td>8</td>
<td>9</td>
<td>6</td>
<td>13</td>
<td>43</td>
</tr>
<tr>
<td>91–100</td>
<td>4</td>
<td>4</td>
<td>5</td>
<td>8</td>
<td>2</td>
<td>23</td>
</tr>
<tr>
<td>&gt; 101</td>
<td>7</td>
<td>10</td>
<td>6</td>
<td>2</td>
<td>9</td>
<td>34</td>
</tr>
<tr>
<td>total</td>
<td>18</td>
<td>22</td>
<td>20</td>
<td>16</td>
<td>24</td>
<td>100</td>
</tr>
</tbody>
</table>

\[ \chi^2_c = 12.03 \quad 0.10 < p < 0.20 \]

6.3. The nutritional status of 10 children with early corneal complications

10 out of the 248 measles patients developed more severe corneal disease: 7 corneal erosions, 3 exposure ulcers. The data regarding the nutritional status of these patients are tabulated in table 6.3. For reasons of comparison, of all parameters the median values in our sample are given.

A statistical analysis of this table is self-evidently not possible, because of the limited number of patients.
Table 6.2.d. The classification for measles-keratitis and the serum albumin content in 100 measles patients. (normal value: 3,500–5,500 mg %). No association is observed.

<table>
<thead>
<tr>
<th>Serum albumin in mg %</th>
<th>Keratitis classification</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I</td>
<td>II</td>
</tr>
<tr>
<td>≤ 2,500</td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td>2,501–3,000</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>&gt; 3,000</td>
<td>8</td>
<td>11</td>
</tr>
<tr>
<td>total</td>
<td>20</td>
<td>22</td>
</tr>
</tbody>
</table>

χ² = 7.65  0.30 < p < 0.50

Table 6.2.e. The measles-keratitis and the content of Retinol Binding Protein in the serum of 103 measles patients. (normal value: 3–6 mg %). No association was found between the serum RBP and the extent of the measles keratitis.

<table>
<thead>
<tr>
<th>RBP in serum in mg %</th>
<th>Keratitis classification</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I</td>
<td>II</td>
</tr>
<tr>
<td>≤ 1.2</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>1.3–2.2</td>
<td>10</td>
<td>11</td>
</tr>
<tr>
<td>&gt; 2.3</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>total</td>
<td>20</td>
<td>22</td>
</tr>
</tbody>
</table>

χ² = 5.41  0.70 < p < 0.80

Table 6.3. Anthropometric and biochemical parameters for the nutritional status of 10 patients with corneal erosions and exposure ulcers. For comparative reasons the median values of the patients Mukumu I are given. The age is given in years; W/A = Weight for Age; H/A = Height for Age; W/H = Weight for Height; RBP = Retinol Binding Protein, normal value: 3–6 mg %; Alb = Serum Albumin, normal value: 3,500–5,500 mg %

<table>
<thead>
<tr>
<th>Pat</th>
<th>Age</th>
<th>W/A</th>
<th>H/A</th>
<th>W/H</th>
<th>RBP</th>
<th>Alb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Central corneal erosions</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M 61</td>
<td>1 2/12</td>
<td>93</td>
<td>92</td>
<td>109</td>
<td>&lt; 1.2</td>
<td>1995</td>
</tr>
<tr>
<td>M 113</td>
<td>3 2/12</td>
<td>78</td>
<td>91</td>
<td>91</td>
<td>1.3</td>
<td>2715</td>
</tr>
<tr>
<td>M 149</td>
<td>1 6/12</td>
<td>80</td>
<td>92</td>
<td>90</td>
<td>2.4</td>
<td>3390</td>
</tr>
<tr>
<td>M 223</td>
<td>1 9/12</td>
<td>70</td>
<td>91</td>
<td>80</td>
<td>2.4</td>
<td>3015</td>
</tr>
<tr>
<td>M 241</td>
<td>7/12</td>
<td>83</td>
<td>83</td>
<td>82</td>
<td>2.5</td>
<td>2425</td>
</tr>
<tr>
<td>M 245</td>
<td>9/12</td>
<td>80</td>
<td>99</td>
<td>91</td>
<td>2.7</td>
<td>3128</td>
</tr>
<tr>
<td>K 19</td>
<td>2 2/12</td>
<td>71</td>
<td>87</td>
<td>87</td>
<td>2.7</td>
<td>3128</td>
</tr>
<tr>
<td>Exposure ulcers</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M 77</td>
<td>4</td>
<td>63</td>
<td>93</td>
<td>72</td>
<td>1.2</td>
<td>&lt; 1500</td>
</tr>
<tr>
<td>M 17</td>
<td>5 6/12</td>
<td>78</td>
<td>91</td>
<td>98</td>
<td>1.7</td>
<td>2790</td>
</tr>
<tr>
<td>M 207</td>
<td>2 6/12</td>
<td>80</td>
<td>93</td>
<td>95</td>
<td>1.5</td>
<td>2680</td>
</tr>
<tr>
<td>Median value</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 9/12</td>
<td>80</td>
<td>93</td>
<td>95</td>
<td>1.5</td>
<td>2680</td>
<td></td>
</tr>
</tbody>
</table>
This table 6.3 however contains some noteworthy details.

(a) The children with corneal erosions seem to be the younger ones. This would be remarkable since the erosions are to be considered as effects of keratitis, but the keratitis itself is independent of the age.

(b) Exposure ulcers occur preferably in children in a bad condition and are accompanied by a high mortality. This finding is not unexpected and confirms the clinical impression.

(c) Of patient M 149 two serum samples were taken: the first on admission at day R + 3, the second at day R + 9 when the corneal erosion developed. In both samples serum albumin was normal, albeit lower in the second one. The serum RBP had however dropped dramatically. This confirms again the value of fast reacting transport proteins as indicators of the nutritional status, not in a static anthropometric sense, but as a dynamic situation.

This is therefore a good example of the nutritional “disadaptation” in the wake of measles. Also, in both samples the IgM was negative, possibly again an example of a disturbed protein synthesis.

6.4. Nutritional status of patients with late corneal complications

In this study 9 patients with late corneal complications were seen. This number was low, because of the strict criteria used for inclusion into this study. Also a presumably higher mortality in children with corneal complications has to be taken into account.

The data regarding tribe, corneal appearance and the nutritional status (as far as available) are tabulated in table 6.4. Self-evidently, no statistical analysis of these data will be possible.

It is however remarkable, that only a limited number of data are indicative for malnutrition.

6.5. Discussion

Measles-keratitis

The lack of association between measles-keratitis and nutritional status is a rather surprising finding. The connection between malnutrition and impaired defense mechanisms might easily result in a higher virusload and therefore give rise to a more severe (because viral) keratitis.

A false negative result however is not to be excluded. Two possibilities exist to explain such a false negative result.

a. Underrepresentation of malnutrition

It has been described already (§3.7) that our patients form a representative
<table>
<thead>
<tr>
<th>Sex</th>
<th>Age in years</th>
<th>Tribe</th>
<th>Corneal appearance</th>
<th>TM</th>
<th>Nutritional status</th>
<th>Fig.</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>M</td>
<td>6</td>
<td>Elgeyo</td>
<td>Descemetocoele</td>
<td>+</td>
<td>W/A 82% H/A 94%</td>
<td>4.4.b</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>2 8/12</td>
<td>Luhyo</td>
<td>Perforation + secondary infection</td>
<td>+</td>
<td>W/A 61% H/A 76%</td>
<td>4.4.c-d</td>
<td>virus in corneal stroma</td>
</tr>
<tr>
<td>M</td>
<td>1 4/12</td>
<td>Elgeyo</td>
<td>Hypopyon ulcer</td>
<td>+</td>
<td>RBP: 3.2 Alb 2460</td>
<td>4.4.e</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>4</td>
<td>Luhyo</td>
<td>Herpetic keratitis</td>
<td>+</td>
<td>W/A 74% RBP: 2.7</td>
<td>3.3</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>8</td>
<td>Kalengin</td>
<td>Phlyctenular keratitis</td>
<td>Perforation</td>
<td>RBP: 1.9 Alb 1620</td>
<td>4.4.f-g</td>
<td>tuberculosis</td>
</tr>
<tr>
<td>F</td>
<td>2 6/12</td>
<td>Luhyo</td>
<td>Healing corneal ulcer</td>
<td>?</td>
<td>Alb 3090</td>
<td>4.4.f-g</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>1 2/12</td>
<td>Luhyo</td>
<td>Herpetic keratitis</td>
<td>?</td>
<td></td>
<td>4.4.i</td>
<td></td>
</tr>
<tr>
<td>M</td>
<td>7/12</td>
<td>Luo</td>
<td>Corneal necrosis</td>
<td></td>
<td>W/A 77% Breast-feeding</td>
<td>4.4.h</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>1 6/12</td>
<td>Luo</td>
<td>Corneal staphyloma</td>
<td></td>
<td>overfed</td>
<td>4.4.i</td>
<td></td>
</tr>
</tbody>
</table>

TM: Traditional medicine was applied; W/A: Weight for Age; H/A: Height for Age; RBP: Retinol Binding Protein in serum; Alb: Albumin in serum in mg%. 
group of the children in Western Province. Moreover, our patients are comparable to other published measles patients, for example the group of Gupta and Singh from Tanzania (1975).

<table>
<thead>
<tr>
<th>Weight for Age</th>
<th>Mukumu I</th>
<th>Gupta-Singh</th>
</tr>
</thead>
<tbody>
<tr>
<td>≤ 60%</td>
<td>5%</td>
<td>7%</td>
</tr>
<tr>
<td>61–80%</td>
<td>50%</td>
<td>37%</td>
</tr>
<tr>
<td>≥ 81%</td>
<td>45%</td>
<td>56%</td>
</tr>
</tbody>
</table>

This confirms again, that in our group the patients are generally not in a worse or better nutritional status than the general population.

b. Sample size

The possibility exists, that our sample is too small to detect a positive association, for which purpose a much larger sample might be needed. But also then, a positive outcome is — of course — not guaranteed.

The question remains however, whether all this effort would be worthwhile. In the present study no association between the nutritional status and the self-healing measles-keratitis was found.

On theoretical grounds it may be expected, that all children with severe Protein Energy Malnutrition have a viral measles-keratitis. This would result however in only a little higher incidence of this keratitis in severely malnourished children compared to a group of well-nourished children. This difference can therefore only be detected in a very large sample. On the overall incidence of the measles keratitis this effect is minimal.

**Corneal erosions**

In the electron-microscope it was shown that the corneal erosions are direct extensions of the measles-keratitis. This finding confirmed the clinical impression. Since however the measles-keratitis is independent of the nutritional status of the affected children (§6.2) the same will probably apply to the corneal erosions. The data presented in table 6.3 point in the same direction.

If these erosions play a role in the pathogenesis of Post-Measles-Blindness — and I think they do — they constitute an important corneal factor, present in at least 4% of our patients, independently of their nutritional status.

**Late complications**

The data regarding the nutritional status of the children with late corneal complications have only a limited value for two reasons:

(a) A sometimes considerable time has elapsed since the acute stage of the measles.
(b) These patients are most probably a very selected sample of the children with late ocular complications. Usually these complications are associated with malnutrition and therefore with a higher mortality. The children presented here, survived the measles, possibly because of a relatively good nutritional status. In my opinion no conclusion whatsoever is to be drawn from this table.
7 — Discussion and Conclusion

7.1. The pathogenesis of Post-Measles-Blindness

7.1.1. Measles.

The incentive for this study was the existence of Post-Measles-Blindness: in several developing countries 1% of all children with measles sustain permanent ocular damage of corneal origin. Surprisingly, little was known about the corneal signs of early measles. In this study it was demonstrated that at least 76% of the examined measles-children had an early viral keratitis. In the following paragraphs this observation will be taken as the most logical starting point for the thinking about the etiology of Post-Measles-Blindness.

It was demonstrated that the nutritional status had no influence on the extent of this keratitis.

Our group of patients is representative for the children in Western Province and I have no reason to believe that another group of patients from this area would show this measles-keratitis to a substantially different extent.

The matter remains whether this keratitis is a peculiar finding for our group of patients, or that it is allowed to extrapolate our findings to all children with measles. There are some arguments in favour of this extrapolation, but much work is still to be done to justify this assumption.

(1) Trantas, working in Turkey (1903) found an incidence of 75% for this keratitis. The nutritional status is not mentioned.

(2) Thygeson (1959), reporting from the USA, states that probably all children have a viral epithelial keratitis in measles.

(3) Sauter (1976), reported a 30% incidence for measles-keratitis, in a cross sectional study in well-nourished children. This is comparable with 75% in a longitudinal study.

In this study it was found that this measles-keratitis — as a sign of measles — is independent from the nutritional status of the affected children. Also Thygeson’s statement points in the same direction. This means that quite possibly no important differences will exist for the incidence of this keratitis in rich or developing countries.

7 out of 188 children (4%) with measles-keratitis developed central corneal erosions. For clinical reasons and because of the supporting electron microscopical findings, this is to be considered as a direct effect of the measles-keratitis, directly related to measles-virus replication in the corneal epithelial cells.

This means, that the majority of measles children, have — intrinsically to the measles infection — a cornea more open to complications. In 4% of these children even macro-erosions were found, a finding which in my opinion might play an important role in the pathogenesis of Post-Measles-Blindness. Whether the lower age at which these occur plays a role, remains to be investigated.
7.1.2. Corneal ulcers and collagenase

The development of corneal ulcers requires the breakdown of stromal collagen, by endogenous or exogenous collagenase (Itoi et al. 1969; Berman 1978). The pathogenesis of Post-Measles Blindness must therefore probably be centered around the presence of collagenase. Pirie, Werb and Burleigh (1975) have been able to demonstrate this presence of collagenase in cases of Vitamin A dependent keratomalacia.

In measles several potential sources for collagenase might be available. Because of epithelial damage (measles, traditional medicines, bacterial or viral superinfection, Vitamin A deficiency or exposure) endogenous collagenase may be activated (Burda and Fisher 1960; Berman 1978; Van Horn et al. 1978). Another potential source of endogenous collagenase are the leucocytes, activated in cases of toxic or bacterial keratitis (Rowsey et al. 1976).

Bacterial infections easily supervene on a damaged cornea and — e.g. pseudomonas aeruginosa — can release a potent exogenous collagenase (van Horn et al. 1978). This means that in measles-keratitis several sources for collagenase might be available. Whether the pathological changes found in the keratocytes (§ 5.3) have any significance in this respect is unclear.

7.1.3. Malnutrition

According to common clinical opinion Post-Measles-Blindness occurs preferentially in connection with malnutrition.

The most important, dynamic, aspect of acute malnutrition is the catabolic deterioration of the metabolism. As part of the total "disadaptation" especially the protein metabolism is severely disturbed: the emphasis is on a decreased synthesis of serumproteins, and a suppression of the immune-system. Malnutrition enhances infection: in malnourished children a prolonged excretion of measles virus from the nasal mucosa was found, also (bacterial) complications are more frequent.

It was also found (Vasantha 1969) that in cases of kwashiorkor the biochemical equilibrium between the different forms of collagen in the skin shifted towards the more soluble tropocollagen. Under normal circumstances no tropocollagen is to be found in the cornea (Holt and Kinoshita 1973).

It is now very attractive to speculate, that the same mechanism might contribute to the pathogenesis of Post-Measles-Blindness: measles and its sequelae provide collagenase, whereas malnutrition brings the collagen in a more soluble state which facilitates its disintegration.

This theory could at least to some extent explain why the more severe corneal complications are connected with malnutrition. This may be an attractive hypothesis, but remains an object of much speculation.

7.1.4. Vitamin A deficiency

For some people Post-Measles-Blindness is identical to Vitamin A dependent
keratomalacia. For this reason measles is frequently described as a mere trigger for the development of keratomalacia. From this study it appears however, that measles in itself might at least be an extremely important cause, because of its intrinsic and direct effects on the cornea. Also malnutrition, because of its effects on the immune system and proteinsynthesis is an undeniable factor in the pathogenesis of Post-Measles-Blindness.

From this study, no answer as to the significance of Vitamin A deficiency for the pathogenesis of PMB can be given. It must be stated however, that to dismiss measles as merely a trigger for keratomalacia does injustice to the significance of the intrinsic involvement of the cornea in measles and the possible pathogenetic implication of this observation (cfr Oomen and ten Doesschate 1973).

7.2. The prevention of Post-Measles-Blindness

The previous paragraph on the pathogenesis of Post-Measles-Blindness gives several starting points for the prevention of Post-Measles-Blindness: vaccination for the prevention of measles, topical treatment of the cornea and general measures for improvement of the nutritional status.

7.2.1. Measles vaccination

The “International Agency for the Prevention of Blindness” accepted at its first World Assembly (1978, Oxford) that measles-immunization-programmes should be developed for the prevention of anterior segment blindness, especially in Africa (IAPB 1980). Even when measles is considered only as a trigger for keratomalacia, this is an important policy statement. This recommendation of the IAPB gets however its full weight when the association between measles and the cornea is realized: in my opinion measles vaccination is mandatory (cfr Olurin 1970). To be effective the “cold chain” must be maintained from factory to patient. In Kenya this appeared possible (Lema 1975).

7.2.2. Topical treatment of the cornea

In §2.6 several possibilities for the topical treatment of corneal conditions along traditional African lines were collected. In most instances a deleterious effect of these preparations on the cornea is likely. A considerable number of cases with potential PMB will be saved from losing eyesight when these preparations are withheld.

Also in this study it was found, that treatment of the cornea with eye ointment, prevented the occurrence of early complications. It is my conviction, that this beneficial effect is primarily caused by the ointment base: the fatty base acts as a lubricant and prevents the newly formed epithelium from being scraped off: the development of erosions is avoided. Its beneficial effect in cases of (potential) exposure will be self-evident.

That the ointment contained an antibiotic, may have been coincidental:
no other ointment, apart from steroid containing preparations, was available. It might have theoretical advantages to add a collagenase-inhibitor, like CaEDTA or cysteine (Berman 1978), to the ointment, but this is probably more of theoretical than of practical importance. The readily available, good and effective tetracycline eye ointment, which is nearly always in stock serves the purpose as well.

In conclusion: in underprivileged conditions every child with measles must be treated with topical application of eye-ointment, frequently and as long as he is ill from measles. Some traditional african eye medicines seem to be harmful and their application should be avoided.

7.2.3. Improvement of the nutritional status

It will be self-evident that children acutely ill with measles must be treated with adequate fluids, food and extra vitamins, to compensate for their deteriorated nutritional status.

Improvement of the nutritional status of the whole population, by nutrition education, improvement of horticulture and agriculture, mother and child clinics and other measures is probably the most important factor to improve “Public Health”. This will also reduce the severity of measles and reduce substantially the incidence of Post-Measles-Blindness.

This is the hardest, but on the long run, the most effective way for a permanent prevention of Post-Measles-Blindness.

7.3. The measles-keratitis in immunosuppression

In cases of severe immunosuppression, when no rash develops, viral complications of measles occur, and probably also the viral measles-keratitis runs its normal course. We were able to observe at least two, markedly malnourished children, admitted because of a viral pneumonia. They had a typical measles-keratitis, but no rash was observed.

Another child, a 7 month old boy, was admitted with a classical keratomalacia, Bitot’s spots in the right eye, and corneal necrosis of the left eye. Clinically no signs of a measles infection were apparent, or known to the mother. On laboratory examination however a positive anti-measles IgM was found, as proof of a recent infection.

These observations may have some practical consequences:

(a) In children with malnutrition, who contract a measles infection, the cornea is even more in danger than in well nourished children but now without the warning given by the rash.

It also offers the possibility that in cases of spontaneous keratomalacia, triggered by a feverish disease, the corneal necrosis is in fact initiated by a measles infection, undiagnosed because of the lack of a rash.
It might therefore be worthwhile to make a careful immunological study of children with "spontaneous" keratomalacia, in view a possible role of undiagnosed measles in these cases of keratomalacia.

(b) Immunosuppression can also be a consequence of the therapy for e.g. leukemia. Here also the possibility of a rashless measles infection exists. The existence of a viral conjunctivo-keratitis is then of high diagnostic value. (cfr Haltia et al. 1978).

7.4. Measles and herpes simplex keratitis

Much attention has been given to a possible connection between measles and a herpes simplex infection of the cornea.

Sauter (1976) observed in 2% of early measles a dendritic keratitis. In 480 early measles cases however, I never observed a herpetic infection. Careful slitlampexamination and the different clinical course always excluded the existence of a dendritic keratitis.

In cases with late complications the situation might be different. Two cases of herpetic keratitis — with their characteristic appearance and clinical course — were observed. It is remarkable, that both occurred a long time after the outbreak of the rash. It remains a matter of speculation in how far the use of (steroid containing?) traditional medicines in at least one case could play a role.

This observation confirms a report from Nigeria: Sandford-Smith and Whittle (1979) were able to culture the herpes virus from some corneal ulcers after measles.

In conclusion: the herpes simplex virus probably plays a role in the causation of late corneal complications.

7.5. Conclusion

The incentive for this study was the existence of Post-Measles-Blindness in developing countries: around 1% of all children with measles sustain permanent ocular damage of corneal origin.

In this first, longitudinal, study on the corneal effects of measles it was found, that the majority of our patients experienced a viral keratitis, as a sign of measles. This keratitis is independent of the nutritional status. In my opinion this keratitis is an important factor in the pathogenesis of Post-Measles-Blindness. Other important factors are malnutrition and the topical treatment given. Post-Measles-Blindness is therefore caused by an interaction of three factors: infection, malnutrition and treatment, each of which gets a different accent in individual cases. The conclusion from this study must be, that the measles infection with its effects on the cornea is much more important than is generally accepted. Moreover, more emphasis should
be on what happens in the cornea: the measles infection, traditional and modern medicines, exposure.

With a little exaggeration my opinion regarding the pathogenesis of Post-Measles-Blindness can be summarized as follows: the existence of the measles-keratitis explains why Post-Measles-Blindness exists, the epidemiology of malnutrition explains why it happens in underprivileged populations.

For the prevention of Post-Measles-Blindness all factors involved in its pathogenesis should be attacked: measles vaccination, protection of the cornea with ointment, avoiding the application of some harmful traditional medicines, and improvement of the nutritional status, will all contribute their part in the reduction and eventual eradication of Post-Measles-Blindness.
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Summary

Introduction

In many developing countries measles is a very serious disease with a considerable morbidity and a high mortality (5 to 10%). An important complication is "Post-Measles-Blindness" which occurs in around 1% of the children affected with measles.

Protein Energy Malnutrition and/or Vitamin A Deficiency are commonly held responsible for this Post-Measles-Blindness. Also a bacterial superinfection is frequently mentioned. The significance of the routinely used traditional African ocular medicines is controversial.

None of the mentioned pathogenetic mechanisms explains why this corneal blindness occurs preferentially after measles and is only incidentally mentioned in connection with other diseases of childhood like malaria, meningitis, whooping cough, chickenpox or tuberculosis.

No good explanation for the epidemiological association between measles and corneal blindness is available, but it could only be self-evident to suppose that measles itself might do something directly to the cornea. This might initiate a chain of events, ultimately leading to corneal blindness. "Ophthalmology has neglected measles" (Quéré 1964) and it is the purpose of this study to fill this gap partially. Hopefully the results will be useful for the prevention of Post-Measles-Blindness.

Methods

The most important part of this investigation is the longitudinal study as to the involvement of the cornea in early measles. This work was done in a 150 bed Mission Hospital during a measles epidemic in the second half of 1976. 148 children, admitted to the isolation ward, under the clinical diagnosis of measles -- in many cases the diagnosis was confirmed by a positive anti-measles IgM in the serum -- were daily examined with a hand held slitlamp-microscope. Further ophthalmological examinations were unfeasible in these mostly severely ill children.

For the demonstration of epithelial lesions in the conjunctiva and cornea the vital stains Rose Bengal, Lissamine Green and fluorescein were used. Rose Bengal and Lissamine Green stain damaged and degenerated cells, whereas fluorescein stains intercellular spaces, accessible to the dye in case of a damaged epithelium. Conjunctival biopsies and some corneae were sent to The Netherlands for pathological examination and immunofluorescence tests.

Ocular signs of measles

The conjunctiva is involved in the measles in two ways: first, in the prodromal fase virus multiplication takes place in the subepithelial lymphoid tissue, resulting in the characteristic conjunctivitis, and second, more or less
concomitant with the rash characteristic strictly epithelial lesions appear in
the exposed parts of the bulbar conjunctiva. The size of these lesions is 0.2–0.4 mm and they have a limited lifespan of only some days. They can only be
made visible with the use of vital stains.

These lesions appear to cross over the limbus to occupy the cornea, at
first the periphery, later on the corneal centre. In the meantime the con-
junctival lesions have healed. The last corneal lesions were observed 11 days
after the outbreak of the rash. This measles-keratitis has, because of its
strictly epithelial character, no permanent sequelae.

This measles-keratitis was observed in 76% of the 148 children. In 4% of
the children with keratitis, the smaller lesions merge into larger macro-
erosions, which also heal without permanent sequelae with appropriate

treatment.

In some other children – mainly very severely ill – exposure ulcers were
observed. Exposure is also a factor in the pathogenesis of some of the earlier
mentioned macro-erosions.

The clinical picture of the measles-keratitis suggests, that it is caused by
the measles virus. Immunofluorescent tests and electronmicroscopical exa-
imination of 10 conjunctival biopsies and 1 cornea demonstrated the presence
of measles virus in the epithelium and keratocytes of the corneal stroma.

Measles-keratitis and nutritional status

Complications of measles are generally ascribed to Protein Energy Malnu-
trition, but the possibility exists that also signs of a disease are in their extent
and/or severity dependent on the nutritional status. To test this last possibility
also in the case of measles-keratitis a quantitative scoring system for the
measles-keratitis was devised. The outcome was compared with anthropo-
metric and biochemical parameters of the nutritional status. In our material
no association could be demonstrated between the extent of the measles-
keratitis and the nutritional status of the affected children. This finding is
one reason more to suppose that the occurrence of the measles-keratitis will
not be limited to children in developing countries: probably measles-keratitis
is a sign of measles in the majority of all children with measles.

The pathogenesis of Post-Measles-Blindness

Post-Measles-Blindness is in my opinion caused by a cooperation of three
factors: measles, malnutrition and treatment. In this study we demonstrated
that at least 76% of all children with measles have a measles-keratitis, in 4%
of them leading to a corneal macro-erosion. To the keratitis are added the
problems of exposure. This makes the cornea more open to secondary in-
fecion and further complications.

In this respect it is remarkable, that in the corneal stroma of an eye, lost
due to panophthalmitis after measles, on electronmicroscopical examination
particles were found, resembling viral strands. The significance of this isolated
finding is at this moment unclear.
An acute Protein Energy Malnutrition (= kwashiorkor) is frequently caused by measles. This causes an extreme biochemical disturbance, resulting in an impaired immunological defence system and a diminished protein synthesis. This last factor might cause a destabilisation of the corneal collagen. The cornea is in danger by both mechanisms.

The significance of traditional African ocular medicines is controversial. The opinions in this respect vary from “nonsense” to the “most important cause” of Post-Measles-Blindness. I am personally convinced that in some cases they add an important factor to an already endangered cornea.

It was also demonstrated, that the frequent application of tetracycline prevents the development of early corneal complications.

In my opinion the measles-keratitis is the explanation for the existence of Post-Measles-Blindness, whereas the epidemiology of malnutrition explains why it occurs in underprivileged populations.

Prevention

The prevention of Post-Measles-Blindness must take all these factors into account: a combined attack by measles vaccination, topical protection of the cornea and improvement of the nutritional status will eventually eradicate Post-Measles-Blindness.
Samenvatting

Inleiding

In menig ontwikkelingsland is mazelen nog steeds een ernstige kinderziekte. Samen met andere infectieziekten, zoals malaria, menigitis, kinkhoest, gastroenteritis en tuberculose, is het een belangrijke oorzaak voor de hoge kindersterfte. Vrijwel altijd speelt daarbij ondervoeding (Protein-Energy-Malnutrition) een belangrijke rol.

Ook blindheid op de kinderleeftijd is in veel ontwikkelingslanden een aanzienlijk probleem. Bij de niet-aangeboren blindheid is deze veelal corneaal gelokaliseerd en het is opmerkelijk, dat in een groot percentage der gevallen een verband met mazelen wordt aangegeven; de andere kinderziekten worden steeds incidenteel als veroorzaker van blindheid vermeld.

Als uiteindelijke oorzaak van deze corneale blindheid na mazelen wordt veelal een Vitamine A deficientie beschouwd. Een bacteriële superinfectie is een andere vaak genoemde oorzaak. De rol van potentieel toxische Afrikaanse geneesmiddelen is evenwel meer omstreden.

Een goede verklaring voor het epidemiologische verband tussen mazelen en corneale blindheid ontbreekt. De gebruikelijke verklaring is, dat door de mazeleninfectie bij een te voren latent ondervoed kind een Vitamine A gebrek manifest wordt; mazelen is dan de aanleiding tot een Vitamine A afhankelijke keratomalacie.

Hiernee is nog niet verklaard waarom dit juist bij mazelen zou gebeuren. Het zou daarom voor de hand liggen te veronderstellen, dat mazelen zelf een direct effect heeft op de cornea, wat dan een eerste fase zou kunnen zijn van een tot blindheid leidende corneale complicatie. Onder- en/of wanvoeding spelen daarbij waarschijnlijk een belangrijke rol. In deze studie kan slechts een klein facet van de onderlinge samenhang tussen mazelen en voedingstoestand ter sprake komen.

Methodiek

Dit proefschrift is het verslag van een longitudinaal onderzoek naar de betrokkenheid van de cornea bij mazelen. Het belangrijkste deel van het onderzoek werd in de tweede helft van 1976 verricht in een 150 beds Missie ziekenhuis in West-Kenya. Tijdens de toen heersende mazelen-epidemie werden 148 kinderen opgenomen op de isolatieafdeling. De diagnose mazelen werd op klinische gronden gesteld en bij een groot aantal van hen bevestigd door een positieve test op anti-mazelen IgM in het serum.


Om laesies in het epitheel van conjunctiva en cornea aan te tonen werd gebruik gemaakt van de vitale kleurstoffen Bengaals Rood (later Lissamine Groen) en fluoresceine. De eerste kleuren beschadigde epitheelcellen,
fluoresceine daarentegen diffundeert in de intercellulaire ruimten bij een aanwezig epitheel defect.

**Oogheelkundige symptomen van mazelen**

De conjunctiva bleek op twee verschillende manieren mee te doen met de mazelen:

(a) Tijdens de prodromale fase heeft een subepitheliale virusvermenigvuldiging plaats, in de lymfoïde laag van de conjunctiva, wat de oorzaak is van de voor de prodromale mazelen kenmerkende conjunctivitis.

(b) Min of meer samenvallend met het uitbreken van de mazelen rash, traden kenmerkende laesies van het epitheel op, vaak zonder dat zich daar ter plaatse ontstekingsverschijnselen als vaatverwijding voordeden.

De conjunctivale epitheliale laesies waren gelokaliseerd in de lidspleet, waren 0,2 à 0,4 mm groot en hadden een kenmerkende vorm. Hun levensduur was slechts enkele dagen. Deze laesies waren alleen zichtbaar te maken door het gebruik van vitale kleurstoffen.

In de volgende dagen verplaatsten deze laesies zich over de limbus naar de cornea, waarbij veelal het centrum van de cornea het laatst was aangedaan.


Deze mazelen-keratitis werd gezien bij 76% van de 148 kinderen. Bij 4% van de kinderen met keratitis conflueerden de kleine laesies tot grotere macro-erosies. Ook deze erosies genazen zonder corneale littekens met de gebruikelijke conservatieve behandeling.

Alles duidde erop dat de mazelen-keratitis een virale keratitis was. Dit kon door laboratorium onderzoek worden bevestigd. In conjunctiva biopten kon door middel van immunofluorescentie mazelen antigeen worden aangetoond in het conjunctiva epitheel. In conjunctiva biopten, bewerkt voor de electronen microscoop, werd subepitheliale virus aangetroffen in lymfoeyten. De ontwikkeling van de gewone mazelen keratitis tot een macro-erosie kon, zowel bij pathologisch anatomisch als electronen microscopisch onderzoek van conjunctiva biopsen en een mazelen cornea, aannemelijk worden gemaakt door het aantonen van sterk verminderde cellulaire adhaesie.

Dit alles leidde tot de conclusie, dat een subepitheliale conjunctivitis en een epitheliale conjunctivo-keratitis beschouwd dienen te worden als symptomen van mazelen.

**Mazelen-keratitis en voedingstoestand**

Symptomen van mazelen zijn in hun ernst en omvang, evenals de complicaties, gecorreleerd met de voedingstoestand van de mazelen patientjes. Om deze
mogelijke correlatie voor de mazelen-keratitis (als *symptoom* van mazelen) na te gaan, werd een quantitatief scoringsysteem voor de keratitis ontworpen. Deze scores werden vergeleken met enkele biochemische en anthropometrische parameters voor de voedingstoestand. Er kon evenwel in ons materiaal geen verband worden vastgesteld tussen de omvang van de mazelen-keratitis en de voedingstoestand.

*De pathogenese van blindheid na mazelen*

Blindheid na mazelen wordt veroorzaakt door een interactie van de 3 factoren: mazelen, ondervoeding en behandeling.

a. *Mazelen*. In deze studie konden we aantonen dat tenminste 76% van de kinderen een mazelen-keratitis doormaakte, terwijl 4% van alle kinderen een verdergaande beschadiging van de cornea (hetzij een erosie, hetzij een uitdrogings-ulcus) vertoonde. Alleen al hierdoor is de cornea ontvankelijker voor het optreden van secundaire infecties en verdere complicaties. Bij electronen-microscopisch onderzoek van een cornea, afkomstig van een oog, verloren na mazelen, werden in de keratocyten partikels aangetroffen, welke mogelijk virus-partikels zouden kunnen zijn. Deze bevinding was totaal onverwacht en op dit moment is de betekenis nog onduidelijk.

b. *Ondervoeding*. Bij het ontstaan van blindheid na mazelen speelt ondervoeding een zeer belangrijke rol. Een acute ondervoeding (kwashiorkor) kan geëvenereerd worden door mazelen. Deze geeft een extreme verstoring van biochemische processen, wat o.a. aanleiding kan zijn tot vermindering van de korra, afkomstig van een oog, verloren na mazelen, werden in de keratocyten partikels aangetroffen, welke mogelijk virus-partikels zouden kunnen zijn. Deze bevinding was totaal onverwacht en op dit moment is de betekenis nog onduidelijk.

c. *Lokale behandeling van de cornea*. Er zijn argumenten om aan te nemen, dat sommige inheemse Afrikaanse geneesmiddelen een schadelijk effect op de cornea zouden kunnen hebben. Sommigen vinden dit een onbelangrijke factor, terwijl anderen met stelligheid beweren dat het de belangrijkste oorzaak is van blindheid na mazelen. Ik ben er van overtuigd, dat sommige van deze geneesmiddelen een belangrijke schadelijke factor extra zijn.

Indien kinderen met mazelen echter consequent een oogzalf met een antibioticum kregen toegediend, traden aanmerkelijk minder complicaties op: zelf beschermde de cornea.

*Preventie van blindheid na mazelen*

Alle bovengenoemde factoren lenen zich als aangrijpingspunt voor de voorkoming van blindheid na mazelen: mazelenvaccinatie, verbetering van de voedingstoestand, het vermijden van sommige schadelijke handelwijzen en de bescherming van de cornea met zelf, zullen een drastische vermindering van de blindheid na mazelen te zien geven.
Résumé

Introduction

A l’opposé de la situation observée en Europe et aux États-Unis la rougeole a des conséquences beaucoup plus graves dans les pays du Tiers-monde. Les symptômes sont souvent beaucoup plus graves et les complications sont plus fréquentes et la mortalité est élevée (5 à 10%). La raison, pour laquelle je me suis attaché à ces recherches, est la prévention de la cécité cornéenne consécutive à la rougeole qui se produit environ chez 1% des enfants atteints.

On attribue la plupart du temps à l’origine de cette cécité cornéenne une sous-alimentation et un manque de vitamine A; on accorde aussi souvent comme cause une surinfection bactérienne. Le rôle, joué par des médicaments indigènes, à usage fréquent, est très discutable.

Cependant, il est à remarquer que cette forme de cécité se déclare après une rougeole alors qu’un rapport de cause à effet est rarement signalé dans les maladies telles que malaria, méningite, coqueluche, variole ou tuberculose dans les pays, couvrant le Tiers-monde. Il manque à cette relation épidémique une interprétation valable.

On pourrait être tenté d’établir que la rougeole a un effet, direct sur la cornée et déclenche une série de réactions consécutives lesquelles pourraient finalement amener à la cécité. Jusqu’à présent, on a fait très peu de recherches dans le sens de relation directe entre la rougeole et ses complications sur la cornée.

Le but de ces examens longitudinaux est une clarification d’idées à propos des effets de la rougeole sur la cornée, dans l’espoir que ces données pourront contribuer ainsi à la prévention de la cécité après la rougeole.

Méthode

La partie la plus importante de nos recherches — c’est à dire, une étude longitudinale consacrée à la relation de la cornée dans la phase aiguë de la rougeole — fut faite dans un hôpital missionnaire de 150 lits au Kenya-Ouest, pendant la seconde moitié de l’année 1976.

Pendant l’épidémie de rougeole, 148 enfants furent hospitalisés et mis en quarantaine. Le diagnostic de la rougeole a été établi par des symptômes cliniques et pour un grand nombre de ces cas, confirmés par un test positif à l’aide d’un IgM antirougeole dans le sang. On a observé quotidiennement ces 148 enfants grâce à une lampe à fente à main.

Il se révèle impossible de faire chez ces enfants gravement atteints un examen oculaire plus conséquent. Pour donner la preuve des lésions dans l’épithélium conjonctival et cornéen, on a fait l’emploi de colorants vitaux, Rose Bengé, plus tard Lissamine Verte et de fluorescène.

Les premiers teintent les cellules épithéliales lésées à l’opposé de la fluorescène qui se diffuse dans les espaces intercellulaires en présence d’un épithélium déficient.
Les symptômes oculaires de la rougeole

Il est apparu pendant ces recherches que la conjonctive était atteinte de deux façons différentes à compter avec un déroulement normal de la maladie; d'abord pendant la phase prodromale, il se crée une multiplication du virus dans le tissu lymphoïde de la conjonctive qui se manifeste cliniquement sous la forme d'une conjonctivite; en second lieu, simultanément à l'apparition du "rash" l'épithélium conjonctival laisse apparaître des lésions caractéristiques en l'absence de symptômes infectieux locaux, comme par exemple, une dilatation vasculaire.

Sur ce dernier symptôme va se centrer toute notre attention. Les lésions conjonctivales épithéliales nommées ici plus-haut, sont localisées dans la fente palpebrale; elles mesurent 0,2 à 0,4 mm. et elles ont une morphologie caractéristique avec une durée, limitée à quelques jours. On parvient à rendre ces lésions visibles grâce à l'emploi de colorants vitaux.

Ces lésions qui se déplacent les jours suivants vers la cornée, vont traverser de limbe pour se terminer à maintes reprises dans le centre de la cornée; 12 jours après l'éruption du "rash", on ne parvient pas à déceler des lésions cornéennes. Les lésions épithéliales de la conjonctive ont déjà effectué plus tôt leur disparition. Cette kératite due à la rougeole, du fait de la seule atteinte de l'épithélium, ne laisse pas de cicatrice sur la cornée.

On a pu observer la kératite due à la rougeole chez 76% des 148 enfants atteints. Les petites lésions confluent en macro-lésions chez 4% des enfants atteints de kératite.

Ces érosions guérissent sans cicatrice cornéenne grâce à un traitement habituel.

Chez quelques autres enfants pour la plupart gravement atteints, on va constater l'apparition d'un ulcère cornéen par dissiccation, résultant de la fermeture insuffisante des paupières.

L'application routinière d'une pommade à tous les enfants atteints paraît être suffisante pour prévenir les érosions cornéennes aussi bien que les ulcères à base de dissiccation. Le tableau clinique de la kératite fait penser que ces lésions épithéliales dans la conjonctive et la cornée sont provoquées par le virus de la rougeole. On peut confirmer ces dires par un examen immuno-fluorescent et microscopique électronique de 10 biopsies conjonctivales et un examen microscopique électronique d'une cornée.

La kératite à rougeole et la nutrition

Une sous-alimentation est en général la cause des complications de la rougeole. Il est tout aussi possible que la gravité des symptômes de la rougeole est liée aux habitudes nutritives.

Pour affirmer cette possibilité pour la kératite à rougeole (comme symptôme de la rougeole), nous avons créé un système de score quantitatif ayant trait à la kératite, qui par après a été comparé avec des paramètres
biochimiques et antropométriques de nutrition. On n’a pas pu établir une liaison entre la gravité de kératite à rougeole et la nutrition.

Ce n’est pas pour cette seule raison, qu’on peut poser l’hypothèse que cette kératite chez les enfants atteints de rougeole, les frappe sans distinction d’habitat, dans les pays du Tiers-monde ou dans nos pays d’occident.

La genèse de la cécité après la rougeole

Ces données doivent contribuer à la conversion dans chaque théorie, au sujet de la genèse de la cécité cornéenne, bien que cela reste encore naturellement dans le domaine du spéculatif.

On peut démontrer dans cette étude qu’au moins 76% des enfants ont été atteints d’une kératite à rougeole, tandis que 4% de nombre total des enfants ont subi une détérioration de la cornée (soit qu’il s’agisse d’une érosion ou ulcère d’exposition).

Pour cette seule raison, la cornée est beaucoup plus sensibilisée aux infections secondaires et à d’autres complications. Au moyen d’une recherche microscopique-électronique d’une cornée, d’un œil perdu après une infection à rougeole, on a découvert des particules dans les kératocytes, lesquelles sont probablement des particules virales à rougeole.

Cette découverte est tout à fait inattendue et à l’heure qu’il est le sens en est encore confus. Le rôle principal, la rougeole mise à part, est joué par la malnutrition. Une sous-alimentation aigue (kwashiorkor) peut être provoquée par la rougeole.

Celle-ci mène à une perturbation grave des procès biochimiques qui peut conduire à une défense immunologique moindre de l’organisme et une perturbation du collagène, qui rend à son tour la cornée plus vulnérable.

Le sens qu’on veut accorder à l’emploi de médicaments ophtalmologiques africains et traditionnels, est cependant controversé. Certains ne prennent pas ce facteur en considération, alors que d’autres certifient que l’emploi de ceux-ci est la cause primordiale de cécité, après une rougeole. Je suis tout à fait persuadé que ces médicaments y ajoutent souvent un facteur funeste.

Prévention

La cécité provoquée par la rougeole est probablement causée par une association de trois facteurs: la rougeole, la sous-alimentation et le traitement. La prévention devra se concentrer sur la vaccination contre la rougeole, la lutte contre la sous-alimentation et éviter l’usage de médicaments traditionnels funestes. Cela ne veut pas dire qu’il faut totalement en éliminer l’emploi.

La conclusion de cette thèse se résume sans peine: l’existence de la kératite à rougeole peut expliquer la cécité cornéenne de préférence après une rougeole; l’épidémiologie de la sous-alimentation, la raison pour laquelle cela se produit seulement dans les pays du Tiers-monde.
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The Kenyan Government offered indispensable help. Dr. M.L. Odouri, Chief Paediatrician of the Kenyatta National Hospital, offered the possibility to examine his patients and to take biopsies. Dr. P. Muthiga, Paediatrician at the Kakamega Provincial Hospital, accepted, that the clinical trial was done in his measles-ward. The Director of Medical Services of the Ministry of Health, Dr. J.C. Likimani gave consent to perform the Shikusa Survey.

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Elly Berkers-Mecking, my secretary, typed and retyped the manuscript, with great accuracy.

Haaren, 18-2-1981

Legends to

COLOURPLATE 1

Figure 3.3. The differential staining of Rose Bengal 1% and Fluorescein 1% in a case of herpetic keratitis. The Rose Bengal stains devitalized cells, whereas fluorescein is seen to diffuse into the surrounding epithelium. This herpetic keratitis was seen in a 4 years old Luhya girl, 4 weeks after the measles rash. She is — anthropometrically — mildly malnourished, whereas the biochemical estimations are in the well-nourished range (Retinol Binding Protein 2.7, Albumin 3090). Traditional medicines had been applied (§ 4.4) (Pat P172)

Figure 4.4.b. Descemetocoele, with incarceration of the iris, in a 6 years old Elgeyo boy, two weeks after the outbreak of the measles rash. The child was well nourished, no nightblindness, no Bitot's spots. Traditional medicines had been applied. Pat Tamb VA 120

Figure 4.4.c. 2 years 8 months old, severely malnourished Luhya girl with a corneal perforation and secondary (?) infection, 2 weeks after the measles rash. The clinical impression “panophthalmitis” was confirmed on evisceration, when a purulent hyalitis was found. In the corneal stroma possibly measles virus could be detected on electron-microscopical examination (fig. 5.3.h). The serum albumin (2460 mg%) is rather low. The Retinol Binding Protein in the serum is normal (3.2) Pat Muk P 258

Figure 4.4.d. The same patient as fig. 4.4.c. Descemetocoele, with incarceration of iris of the left eye, 2 weeks after the measles rash.

Figure 4.4.e. Hypopyonulcer in a malnourished Elgeyo boy, 16 months of age, 3 weeks after the outbreak of the rash. The hypopyon has an unusual localization: prior to examination the boy had been sleeping for several hours at his left hand side. The ulcer healed with conservative treatment. Traditional medicines had been applied. Pat Tamb P 273

Figure 4.4.f. Descemetocoele in the right eye of a 8 years old girl, 4 weeks after a measles rash. This 8 years old Kalengin girl is overtly malnourished, and is treated for pulmonary tuberculosis. The left eye demonstrates a phlyctenular keratitis (fig. 4.4.g) Pat Tamb VA 28

Figure 4.4.g. The same patient as fig 4.4.f. The cornea of the left eye showed several phlyctenulae

Figure 4.4.h. Corneal perforation in a 7 months old Luo boy, 5 weeks after the measles rash. The child was breastfed. The Weight for Age is 77%. Pat SiaVA 16

Figure 4.4.i. Corneal staphyloma in an 18 months old, overfed girl, 2 months after the measles rash. Pat Bung VA 83
Legends to:
COLOURPLATE II (see page 112)

Figure 4.1.2.a. Conjunctival lesions intensely staining with Lissamine Green occurring in the early exanthematous stage of measles, localized in the interpalpebral fissure. (Pat M 234, day R + 1)

Figure 4.1.2.b. Larger magnification of the conjunctival epithelial lesions of early exanthematous measles, stained with Rose Bengal. Pat M 206, day R + 1

Figure 4.1.2.d. Conjunctival epithelial lesions are present in continuity with the same lesions in the cornea. It is difficult to show a good picture of this transition because of the extreme difference in contrast at the limbus. The conjunctival lesions are stained with Rose Bengal, the corneal lesions stain with fluorescein. Pat M 206, day R + 1

Figure 4.1.2.e. At this stage the measles keratitis is more or less restricted to the corneal periphery. Pat Cr 63, Day R + 2

Figure 4.1.2.f. The conjunctiva and limbus are now free. The keratitis is localized at the midperipheral and central parts of the cornea. Pat Cr 129, day R + 4

Figure 4.1.2.g. The last manifestation of the epithelial measles keratitis: only the centre of the cornea is involved. Pat M 18, day R + 3
Legends to:
COLOURPLATE III (see page 113)

Figure 4.2.1.a. Corneal erosion in a child with measles, 9 days after the outbreak of the rash, 3 days after the administration of 200,000 IU Vit. A. Pat K 19

Figure 4.2.1.b. Large central corneal erosion, surrounded by an extensive measles keratitis, 9 days after the outbreak of the rash. Pat M 149 (Electronmicroscopy in Fig 5.3.b and c)

Figure 4.2.2.a. (cfr Figure 4.2.2.b) Exposure ulcer in the cornea of a 4 years old boy, 3 weeks after outbreak of the rash. Pat M 117

Figure 5.1.2.b. Positive immunofluorescence test for measles antigen in the subepithelial tissue of a Lissamine Green negative biopsy. The yellow colour indicates the presence of measles antigen

Figure 5.1.2.c. The presence of measles virus antigen in the epithelium of a lissamine Green positive conjunctival biopsy is demonstrated by a positive immunofluorescence test
APPENDIX

In this appendix the numerical data regarding the 148 patients in the group Mukumu I are tabulated. The methods used are described in § 3.3 and 3.5.

The age is given in months.

H = height
H/A = height for age
W = weight
W/A = weight for age
W/H = weight for height
Vacc = history of vaccination against measles
Alb = Serum albumin. Normal value in European adults: 3,500–5,500 mg%.
RBP = Retinol Binding Protein. Normal value in European adults: 3–6 mg%.
Ker = Keratitis score
Adm = Data of admission compared to the day of outbreak of the rash (= R)
Obser = Observation period

For the calculation of H/A, W/A and W/H the Harvard standard as commonly in use in Kenya as the “Road to Health” is used.
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Curriculum vitae

*N.W.H.M. Dekkers*

31-12-1940 geboren te Tilburg

05-07-1958 Eindexamen Gymnasium β aan het St. Odulphuslyceum te Tilburg


01-06-1972 Aanvang opleiding tot oogarts aan het Oogziekenhuis te Rotterdam, onder Prof. Dr. H.E. Henkes en Prof. Dr. A.Th.M. van Balen. Deze opleiding werd van 01-11-1974 tot 15-07-1978 onderbroken voor het gaan leiden van een groot oogheelkundig project van de Prof. Weve Stichting in W. Kenya, waar ook de basis van dit proefschrift werd gelegd.

15-07-1979 Volgde de inschrijving in het Specialisten Register als oogarts.

01-10-1979 Vestiging als oogarts in Tilburg aan het St. Elisabeth Ziekenhuis.