Radiocarbon Dating of the Human Eye Lens Crystallines Reveal Proteins without Carbon Turnover throughout Life

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Background. Lens crystallines are special proteins in the eye lens. Because the epithelial basement membrane (lens capsule) completely encloses the lens, desquamation of aging cells is impossible, and due to the complete absence of blood vessels or transport of metabolites in this area, there is no subsequent remodelling of these fibers, nor removal of degraded lens fibers. Human tissue ultimately derives its 14C content from the atmospheric carbon dioxide. The 14C content of the lens proteins thus reflects the atmospheric content of 14C when the lens crystallines were formed. Precise radiocarbon dating is made possible by comparing the 14C content of the lens crystallines to the so-called bomb pulse, i.e. a plot of the atmospheric 14C content since the Second World War, when there was a significant increase due to nuclear-bomb testing. Since the change in concentration is significant even on a yearly basis this allows very accurate dating. Methodology/Principal Findings. Our results allow us to conclude that the crystalline formation in the lens nucleus almost entirely takes place around the time of birth, with a very small, and decreasing, continuous formation throughout life. The close relationship may be further expressed as a mathematical model, which takes into account the timing of the crystalline formation. Conclusions/Significance. Such a lifelong permanence of human tissue has hitherto only been described for dental enamel. In contrast to dental enamel it must be held in mind that the eye lens is a soft structure, subjected to almost continuous deformation, due to lens accommodation, yet its most important constituent, the lens crystalline, is never subject to turnover or remodelling once formed. The determination of the 14C content of various tissues may be used to assess turnover rates and degree of substitution (for example for brain cell DNA). Potential targets may be nervous tissues in terms of senile or pre-senile degradation, as well as other highly specialized structures of the eyes. The precision with which the year of birth may be calculated points to forensic uses of this technique.

INTRODUCTION

Living tissues undergo a continuous turnover, substitution and remodelling, both on the cytological and tissue level. In the human organism, an exception to this is the dental enamel, which after formation does not undergo any change [1]. Another highly specialized tissue is the eye lens. The transparency and refraction of the normal eye lens is dependent on the so-called crystallines. These proteins fill the cytoplasm of the highly organized eye lens fibre cells [2]. To improve optical quality, these proteins need to exist both as concentrated solutions in the outer cortical region and more glass-like in the core. The high amount of crystallines gives the lens a high refractive index which is necessary for focusing light to the retina. Lens crystallines can be classified as either classical or taxon specific [2]. The classical crystallines include members of the alpha-crystalline family and beta/gamma-crystallin superfamilies [3]. An important function of these proteins is preventing protein aggregation which will make the lens opaque. Cataract occurs when the ageing crystalline proteins are no longer evenly distributed on the scale of light wavelength [4].

The lens fibre cells in the nucleus of an adult lens are thought to be produced during early embryonic life [2]. The lens plate is formed in the 4 mm embryonal stage. The lens fibre cells degrade all membrane-bound organelles and the older lens cells become compressed into the nucleus of the lens by the continuous formation of new fibres at the surface. The lens substance is thus composed of fibres that are constantly formed from elongation of the equatorial epithelium. The human lens grows rapidly in the embryo and during the first postnatal year [2]. The rate of lens growth slows between ages 1 and 10 years, and continues at a much slower, nearly linear, rate throughout life. Because the epithelial basement membrane (lens capsule) completely encloses the lens, desquamation of aging cells is impossible, and due to the complete absence of blood vessels or transport of metabolites in this area, there is no subsequent remodelling of these fibers, nor removal of degraded lens fibers [2].

We have exploited the radical variations of the atmospheric 14C content during the last 50 years to date the formation of the lens crystallines. The concentration of 14C in living tissues reflects the atmospheric 14C content at the time of growth. This is because cosmogenic 14C in the atmosphere reacts with oxygen to form...
carbon dioxide (CO₂), which is incorporated by plants, and then ingested by animals. The plants and animals are ultimately ingested by humans. If there is only minimal turnover in the eye lens, the content of ¹⁴C in the lens crystallines should thus reflect the atmospheric concentration at the time the lens crystallines were formed. The amount of ¹⁴C in the atmosphere (see Fig 1) was almost constant until about 1955, when nuclear-bomb tests caused it to rise dramatically [5]. Subsequent to the Test Ban Treaty in 1963 the concentration has decreased rapidly, mainly because the atmospheric ¹¹C has diffused into the oceans, while radioactive decay (the half-life of ¹⁴C is 5730 years) is of minor importance [6]. Since the change in concentration is significant even on a yearly basis this allows very accurate dating [7].

RESULTS
The content of ¹⁴C in the lens crystallines was compared with the bomb pulse curve, and the intercept indicates the year of formation (Fig 1). Our results show a tight fit of less than three years between the predicted and the actual year of birth (Fig. 2). This is assuming that the lens crystallines have been formed as a single event at the year of birth, which is only a crude approximation. We therefore constructed a mathematical model based on the above assumptions on lens growth. The model was subsequently adjusted to fit the experimental data. In this way we were able to obtain detailed information concerning the timing of the formation process. The key element of our model is the formation function \( F(x) \) which describes the incorporation of carbon into the lens crystallines as a function of the age \( x \) of the individual. It is characterized by a high rate of formation (≈ 1.0 in relative units) between –0.5 to 1 year after birth; an exponential decrease (half-time 0.8 years) in formation rate until the fourth year after birth; a very low linear formation rate (≈ 0.027) until 25 years after birth; and finally an even lower linear formation rate (≈ 0.006). The maximum lifespan is limited to 80 years. We then use the formation function to calculate the content of ¹⁴C in the lens crystallines (see Fig. 1) for a given year of birth \( (b) \) and age at death \( (a) \) on the basis of the atmospheric ¹⁴C concentration \( \text{pmCatm} \):

\[
\text{pmC}_{\text{crystalline}}(b,a) = \int_{-0.5 \text{year}}^{a} F(x) \text{pmCatm}(x+b+R)dx
\]

We apply a “delay” \( R \), as the source of ¹⁴C for the lens crystallines is ¹³C ingested with foodstuffs (either directly crop-derived, or as meat/dairy products from animals, which have fed on crops). These foodstuffs thus “delay” the atmospheric ¹⁴C reaching the eye lens by a growth season (set to 1 year, i.e. \( R = –1 \text{ year} \)). For that reason we also base our ¹⁴C model calibration curve on biomass turnover [8] and not on the atmospheric equilibrium. Although our simple model may not describe all details of the formation process correctly, comparisons with the experimental data show that it is a good approximation and leave absolutely no possibilities for radical adjustments.

The model enables us to calculate the year of birth (Fig. 2). This was done by comparing the content of ¹³C in the lens crystallines with the plotted model curve (red curve in Fig. 1) using the ¹³C calibration program OxCal 4.0 [9,10]. There exists good agreement between the model-predicted years of birth and the actual ones. Thus, in all cases the actual year of birth is within the predicted range. The precision of the method is demonstrated by the average magnitude of the uncertainty, which is ±1.5 years (95% confidence), excluding the case from 1922. The latter case is the oldest included in the study, and the year of birth was 30 years before onset of the bomb pulse rise. Given the extremely low formation rate of crystalline at the point when this individual lived through the bomb-pulse, this means that the ¹⁴C content only shows that the
person was born at least 10 years before the bomb-pulse. The only other pre-pulse case, an individual born in 1940 shows increased ¹⁴C content, indicating that this individual, approximately 6 years old at the start of the pulse, was still forming lens crystallines.

**DISCUSSION**

In our analysis we have made the assumption that the diet of the subjects consisted mainly of terrestrial food and in particular that marine food did not contribute with a significant fraction. Consumption of fish would have a significant effect on the ¹⁴C content of the lenses, because the ¹⁴C concentration in the sea is smaller that in the atmosphere (marine ¹⁴C reservoir effect). However, marine food generally constitutes only a small fraction of the diet in modern Denmark, and reservoir effects are not expected to have influenced the analysis. Consumption of marine food would also have been visible in the stable-isotope ratio ¹³C/¹²C, which was also measured, but no indications of this were observed.

Our results allow us to conclude that the crystalline formation in the lens nucleus almost entirely takes place around the time of birth, with a very small, and decreasing, continuous formation throughout life. Such a life-long permanence of human tissue has hitherto only been described for dental enamel [1,11]. In contrast to dental enamel it must be held in mind that the eye lens is a soft tissue, subjected to almost continuous deformation, due to lens accommodation, yet its most important constituent, the lens crystalline, is never subject to turnover or remodelling once formed. The determination of the ¹⁴C content of various tissues may be used to assess turnover rates and degree of substitution (for example for brain cell DNA [12]). Potential targets may be nervous tissues in terms of senile or pre-senile degradation, as well as other highly specialised structures of the eyes.

The precision with which the year of birth may be calculated points to another use of this technique. In forensics, the identification of an unidentified corpse relies on matching of data, e.g., fingerprints or DNA from the deceased with registered information. Consequently, sometimes there is a need to gather introductory data on the deceased in terms of probable sex and age. The eye lens is easily extracted, removal does not disfigure the corpse, and a result regarding the age at death of the victim may be obtained in as little as 24–48 hours.

**MATERIALS AND METHODS**

We obtained 13 eye lenses from 13 deceased persons of varying age. Ethical consent was obtained from the Copenhagen Board of Medical Ethics. The lenses were extracted by a minimally invasive procedure as part of autopsies performed at the Institute of Forensic Medicine, University of Copenhagen. The lens of young people is rather elastic, while the lenses of older people harden due to the accumulation of degraded by-products. It was our experience that the lens could be removed up to three days post-mortem. After this period, post-mortem degradation and putrefaction made it impossible to extract the lens, as it became more and more fluid and not intact in its capsule. After extraction, the nucleus of the lens was cut out. The nucleus of the human eye lens contains 10–15 mg tissue, corresponding to approximately 3–5 mg carbon, which is sufficient to perform a high precision accelerator mass spectrometry (AMS).

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**Author Contributions**

Conceived and designed the experiments: NL HK SH. Performed the experiments: HK CJ SH. Analyzed the data: NL HK JH. Contributed reagents/materials/analysis tools: CJ SH JH. Wrote the paper: NL HK.

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