The human eye lens is a unique organ; it grows continuously throughout life by the accretion of new cells which are produced in the equatorial zone of the lens and then elongate until they cover existing fiber cells inside the surrounding elastic capsule. The old cells are not shed, but instead, are densely packed into the center of the organ and their contents do not undergo any turnover. Knowledge of how this continuous growth generates the properties of the lens is important for understanding the age-related development of pathologies such as presbyopia and cataract. A lot has been learnt from observations on animals, but significant differences exist between human and non-primate lenses, making it necessary to undertake studies on the human lens.

There have been many studies on the growth of the human lens, both in vivo and in vitro. Most previous in vivo studies have been limited to the measurement of lens thickness using ultrasound or the slit lamp. However, in vivo measurements suffer from the need to correct for sonic or optical distortions which vary with age and with the internal structure of the eye. In addition, the accommodative state of the eye markedly influences lens dimensions. Consequently, there has been little agreement between different studies. The recent application of magnetic resonance imaging (MRI) and optical coherence tomography (OCT) methodologies to ocular measurements has offered solutions to some of these problems, but the current MRI resolution (200 µm) is too poor for accurate measurements and OCT depends on mathematical manipulations to compute the whole lens shape as it cannot image the lens periphery; however, advances in these two technologies should, eventually, provide more uniform data on in vivo lens dimensions.

Lens weights and dimensions can be more conveniently studied in vitro, where the lens is free from the influence of other ocular tissues and measurements do not require corrections. In particular, lens weights, both fresh and fixed/dried, can easily be obtained from human donor eyes. Since the first major study by Smith in 1883, a large body of data on the changes in lens wet weight with age and limited data on lens dimensions have been accumulated from measurements in numerous laboratories. Unfortunately, there are huge variations in the data, most of which can be attributed to the effects of prolonged eye bank storage, generally beyond the control of the investigator, or differences in tissue handling. These have led to various contradictory conclusions regarding lens growth, as well as lens properties.

Analysis of published and unpublished data has revealed that growth in human lens wet weight is biphasic, asymptotic before birth, and linear postnatally. The transition in growth pattern appears to occur at or very shortly after birth, with the completion of the clinically defined adult nucleus, but the available data are too variable to pinpoint the time or to accurately describe further postnatal growth. There is a great need for more reliable data collected under carefully controlled conditions to define growth more precisely.

Although the simplest way to assess lens growth is by determination of changes in wet weight, the measured wet weight may not be indicative of the amount of tissue laid down. Human lenses are generally obtained from eyes that have been...
stored for variable times in eye banks. During such storage, water may be taken up by the lens, increasing its weight. Since there is no loss of cells or their contents (other than water), a more reliable assessment of lens growth would be to measure the accumulation of protein/dry matter. Such an approach, in which the lens is fixed and then dried, is commonly used in methods for determining the ages of animals culled in the wild. Dry weight determination is generally not done with human lenses since it would spoil the precious lens for any further studies. Consequently, very little is known about the increase in dry weight with age. Such data would be valuable in assessing the extent of compaction in the aging lens and its relationship to the development of presbyopia.

The purpose of the current study is to undertake a long-term investigation, which attempts to overcome some of the difficulties of the past by measuring both wet and dry weights and dimensions of lenses collected at a single site from a racially uniform Indian population, using a rigorously followed protocol, which reduces the risk of postmortem changes. Of particular interest are the changes in lens dry weight with age. The data from this study are compared with data obtained from western populations. It is expected that access to a large number of lenses, improved handling methods, and the measurement of several parameters on each lens will provide valuable and reliable information on the growth of the human lens.

Materials and Methods

The donor tissues used in this study were received from the eye bank belonging to a tertiary eye care center in South India. The age, gender, time of death, time of enucleation, cause of death, postmortem time, and time of use were noted for each eyeball used.

Lenses were obtained from donor eyeballs after their corneas were removed under sterile conditions for the purpose of transplantation. Eyes obtained more than 4 days postmortem were rejected. The iris was excised first and then the lens was extracted with a lens spoon after freeing it from the zonular attachments with Vannas scissors. The lens was then carefully blotted dry and its dimensions (diameter and thickness) were measured with a digital caliper (INOX IP54 Caliper, Micro Precision Calibration Inc., Grass Valley, CA, USA). Lenses with visible capsular damage or cataract were excluded from the study.

The lens was weighed using an analytical balance (KERN ABS 120-4, Kern & Sohn GmbH, Balingen-Frommern, Germany) before being placed in 5 ml of 5% buffered formalin in small glass containers. After 1 week fixation, the lenses were dried at 80°C in a hot air oven (Sisco, Thane, India) until constant weight was achieved. This complete drying of lenses resulted in seven wet weight and five dry weight points being obtained by the eye bank in accordance with its practices and procedures. The study protocol was approved by the Institutional Ethics Committee attached to the tertiary eye care center (approval number LEC 08025 dated 23 December 2008) and adhered to the tenets of Declaration of Helsinki.

Results

The dimensions (diameter and thickness) of the lenses are shown in Fig. 1. Diameter increases continuously from around 9 mm at age 16 years to near 10 mm at age 90 years (diameter = 7.40 + 0.51 × ln(age), R² = 0.43, P < 0.0001). The thickness also increases from age 16 years (thickness = 1.38 + 0.76 × ln(age), R² = 0.60, P < 0.0001). The aspect ratio (thickness: diameter) reflects these changes [Fig. 2]. The data suggest that the ratio may increase with age at around 0.001/year (aspect ratio = 0.40 + 0.001 × age, R² = 0.40, P < 0.0001), consistent with the difference in the rates at which thickness and diameter increase.

![Figure 1: In vitro changes in human lens diameter (blue) and thickness (orange) with increasing age. The dimensions were measured with a digital caliper](image-url)
As can be seen in Fig. 3, lens wet weight increases linearly from around 160 mg at 16 years of age to over 250 mg by 90 years of age \[\text{wet weight} = 146.04 (\pm 3.50) + 1.24 (\pm 0.06) \times \text{age}, R^2 = 0.67, P < 0.0001\]. This is very similar to the trend reported earlier by Augusteyn.\(^7\) However, the scatter in the data is substantially less, probably reflecting the more controlled lens isolation conditions in the present study compared with the previous one.

Fig. 4 presents the lens dry weights as a function of age. The dry weight also increases linearly throughout life from around 50 mg at 16 years of age to around 85 mg by 90 years of age. Regression analysis shows the rate of increase to be 0.44 mg/year \[\text{dry weight} = 47.05 (\pm 1.15) + 0.44 (\pm 0.02) \times \text{age}, R^2 = 0.69, P < 0.0001\]. The average lens dry weight increases from ~31% to ~34% of the lens weight over the age range studied. This corresponds to an increase in the concentration from 330 to 370 mg/ml.

The availability of both wet and dry weights makes it possible to calculate lens volumes. Assuming that dry weight represents essentially only protein, the partial specific volume of lens proteins (0.75) can be used to derive the formula: volume = wet weight – (0.25 × dry weight). The change in the calculated volume with age \[\text{Figure 5}\] can be described with the logarithmic expression: volume = 52.43 × ln(age) – 9.05 \(R^2=0.66, P<0.0001\). The volume of the lens, which is around 150 mm\(^3\) in the late teens, reaches around 230 mm\(^3\) by the age of 90 years.

The postmortem time was included as an independent variable and multiple regression analysis was done to study its possible effect on the dependent variable. The results showed that the postmortem time did not have a significant effect on the dependent variables diameter \(P=0.20\), thickness \(P=0.97\), aspect ratio \(P=0.52\), wet weight \(P=0.82\), dry weight \(P=0.11\), and volume \(P=0.89\).

**Discussion**

Among the most debilitating of ocular dysfunctions are the development of presbyopia and senile cataract.\(^{14,15}\) Both can be attributed to age-related alterations in the lens. Detailed knowledge of the growth and biophysical properties of the lens, and their variations with age, is essential for understanding...
The variation was much less in the
provides a reliable measure of lens growth and can be used
controlled conditions and at shorter postmortem times.
lens diameter.

Augusteyn.
the lifespan comparable to the 1.38 mg/year reported earlier by
the Indian lenses increases at a rate of 1.24 mg/year.
the growth takes place in two distinct modes: asymptotic during
as has been noted previously. This is because human lens
described with many of the self-limiting growth equations, such
slowing gradually throughout life. This can be adequately
this preliminary analysis on adult lens growth.

Previous studies have shown that water uptake takes place rapidly in the lenses left in the eye for extended periods (beyond 30 h postmortem). This might be responsible for the huge variations in wet weights of human lenses observed in earlier studies. The variation was much less in the present study and only seven lenses had to be eliminated in the present analysis because they were more than three SDs from the mean. This low attrition can probably be attributed to the following reasons: the data were collected from fresh human donor lenses with shorter postmortem times (median of 22 h), the use of a single collection protocol and, possibly, the racially more uniform Indian population from which the lenses were collected.

Our results indicate that beyond 16 years of age, the in vitro dimensions increase continuously, but gradually slowing, throughout life. However, there are differences in the rate at which diameter and thickness increase. This is reflected in the variation of aspect ratio (T/D) with age. The ratio increases at a rate of 0.001/year, which is very similar to the previous observation of Rosen et al.

The growth in lens weight in most species is asymptotic, slowing gradually throughout life. This can be adequately described with many of the self-limiting growth equations, such as the logistic and Gompertz functions. However, attempts to fit the adult human data to such functions were not successful, as has been noted previously. This is because human lens growth takes place in two distinct modes: asymptotic during prenatal life and for a very short period postnatally, followed by a linear mode thereafter for the remainder of life. The growth in adult population observed in the present study is similar to that reported for the Western population. The wet weight of the Indian lenses increases at a rate of 1.24 mg/year throughout the lifespan comparable to the 1.38 mg/year reported earlier by Augusteyn. This is consistent with continued growth of the lens diameter. The slight difference in the rates could be due to the fact that the lenses were collected under more carefully controlled conditions and at shorter postmortem times.

As there is no turnover of cells or proteins inside the lens capsule, the accumulated protein content or dry weight provides a reliable measure of lens growth and can be used for determining age. This approach has been applied to many species in wildlife control programs. In order to obtain further information on human lens changes with age, we have also measured dry weights. To the best of our knowledge, this is the first study to report dry weights on a large number of human lenses from adult donors. The dry weight of the lenses from the Indian population increased steadily at a rate of 0.44 mg/year. From around 50-55 mg at age 17 years, the dry weight reaches a value of 85-90 mg by 90 years. This linear model of growth would permit simple determination of adult human age from lens dry weight. Such information could find application in forensic medicine.

It had been expected that the variability in dry weights would be significantly less than that in the wet weights. This did not appear to be the case, possibly because the variability in wet weights had been reduced by the shorter postmortem times. However, it was noted that for many lenses, all parameters measured indicate that the age may have been greater than that written in the medical records, i.e. the apparent scatter was not due to random errors or variations in the measured parameters, but rather could be attributed to incorrect ages. Given the nature of the population from which the lenses were obtained, this is highly likely.

Some of the variability in dimensions, especially in the small values of thickness, may also be due to the measuring technique. Although the resolution of the digital calipers used in this study is 0.01 mm, the lack of consistency between operators and day-to-day variations in measurements may be of concern. In future, the data on dimensions will be obtained by using the better technique of shadow photography.

As the lens fiber cells are packed into the center of the lens, they gradually lose water. This involves hysteresis, the transfer of water from the bound (to protein) form to the free form, and subsequent movement out of the cell. The ratio of dry weight to wet weight can provide a measure of the extent of compaction. Our results indicate that the average dry weight concentration increases from 32 to 34% over the age range observed. Most of the increase is due to the lens nucleus which reaches a maximum of 38%, whereas the cortex has a gradient of concentration which remains relatively constant. The compaction is responsible for the development of a refractive index plateau in the nucleus and a gradient in the cortex with increasing age.

Based on the partial specific volume of proteins (0.74–0.76 ml/g), the calculated volume of lens increases from around 150 mm³ in teens and reaches a maximum of 240 mm³ in the elderly. This is very similar to the observations estimated by computing the solid of revolution of the cross-sectional plane of postmortem lenses around the optical axis. Our measurements are much more accurate than those from the ellipsoid calculation frequently used and simpler to perform than the complex algorithms sometimes used. Since the calculation of volume from weights would destroy the lens, it is necessary to complete all other studies on the lens before fixation and drying and, as long as there is no damage to lens, this will yield accurate volumes.

To conclude, the preliminary observations show that the growth of the human lens in the Indian adult population follows the linear portion of the biphasic pattern described
earlier. Over the age range examined, lens growth in the Indian adult population appears to be very similar to that in Western populations. However, insufficient data are available for early ages to determine if the prenatal growth is also the same. It is intended to continue with the collection of data in this project until this shortfall is overcome.

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