Rapid Method for the Detection of Cystic Fibrosis of the Pancreas in Children

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One of the main criteria for the diagnosis of cystic fibrosis of the pancreas is the presence of a raised sweat sodium level in the patient (Di Sant’Agnese, Darling, Perera, and Shea, 1953). The standard method of using pilocarpine iontophoresis to produce sweat (Gibson and Cooke, 1959), and estimating the sodium present by conductivity or flame photometry, is a workable technique but has some drawbacks. Considerable discomfort is experienced by some patients and the process is time-consuming, as it takes at least 30 minutes to obtain sufficient sweat for a reliable sodium determination. Pilocarpine iontophoresis is therefore not suitable for routine use as a screening test on large numbers of patients.

The use of a sodium responsive electrode for direct readings of the sodium concentration of sweat has not been successful in our hands (Goldbloom and Sekelj, 1964). We found that iontophoresis was necessary before reliable readings could be obtained. Difficulties also occurred due to sweat evaporation and variations in skin temperature.

McGrady and Bessman (1955) and Johnston (1956) have found that cystic fibrosis patients have a raised parotid saliva sodium level compared with that of normal subjects. We have devised a method for measuring the sodium concentration of unstimulated parotid saliva using a micro-dual sodium electrode, which is simple and reliable in children and adults.

The results are presented of a trial with this electrode on normal subjects, cystic fibrosis heterozygotes, and cystic fibrosis homozygotes.

Material and Methods

The age ranges of the subjects investigated were as follows: normals (who could be expected to include about 4% of unknown heterozygotes), 1-48 years (mean 24·9 years); heterozygotes, 25-49 years (mean 35·6 years); and homozygotes, 1-19 years (mean 6·24 years).

It is hoped in future to develop this technique for the rapid screening of the newborn.

The electrode. The sodium-responsive electrode used in this survey was developed in the laboratories of Electronic Instruments Ltd. as a modification of a Jena micro-dual-pH electrode (Fig. 1). When not in use the electrode was stored with the bulb immersed in 0·1M NaCl.

The electrode system was used with an E.I.L. Vibron Blood pH Meter, model 48B; readings of pNa were taken on the pH 6·6-8·0 scale.

![Fig. 1.—A micro-dual sodium responsive electrode for the determination of parotid saliva pNa values.](image)

Standardization of the electrode. The electrode system and the pH meter were standardized, using sodium chloride solutions of 100, 75, 25, and 10 mEq Na⁺/litre. The readings obtained from the standard solutions were marked on the pH scale and a standard curve of log mEq Na⁺/litre against pH reading was plotted.

The pH meter when used with the sodium electrode did not give a fixed check point on the metre scale. The calibration of the system was therefore checked daily with a standard sodium chloride solution of 50 mEq Na⁺/litre.

Disinfection of the electrode. As the electrode was to be used in the mouth to measure the sodium concentration of parotid saliva, it was necessary to find a quick and effective means of sterilization that would not affect its working. The effect of wiping the electrode with

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various sterilizing agents after infection with overnight broth cultures of *Aerobacter aerogenes*, a coliform, *Proteus sp.*, *Pseudomonas aerugonosa*, *Staphylococcus aureus*, and a haemolytic streptococcus, was determined. The most reliable agent tested was 70% isopropanol: this did not affect the working of the electrode provided that after wiping with isopropanol the electrode was immediately immersed in 0.1M NaCl. The sterilization of the electrode in this survey was therefore done by wiping the electrode with a pad soaked in 70% isopropanol (Medi-Swab, Medi-Pack Ltd.).

**Measurement of sodium content of parotid saliva.** The pH meter was first adjusted with the temperature compensating dial to read sodium concentrations at 37°C, and the saliva was tested as follows: the electrode was removed from the 0·1M NaCl and the excess saline was wiped from the electrode with a tissue. The electrode was then placed in the upper region of the cheek next to the second molar tooth at the opening of Stensen’s duct. Care was taken to see that the ceramic plug in the reference electrode was uppermost and also that the electrode was tilted at an angle of 45° to avoid any airlocks that might have formed beneath the ceramic plug. The meter was switched to the read position, and the sodium content of the parotid saliva was read after equilibrium had been reached, usually after 30 to 60 seconds. The meter was switched to the check position, the electrode removed from the mouth, wiped with a ‘medi-swab’, and replaced in 0·1M NaCl.

**Results**

**Parotid saliva sodium levels.** The histogram (Fig. 2) summarizes the unstimulated parotid saliva sodium levels of 31 normal subjects, 10 cystic fibrosis heterozygotes, and 15 cystic fibrosis homozygotes.

The normal subjects gave parotid saliva sodium levels from 3-15 mEq Na+/l., with a mean value of 5·9 ± 3·6 mEq Na+/l. The cystic fibrosis heterozygote range of parotid saliva sodium content was between 4·5-12 mEq Na+/l., with a mean value of 5·69 ± 2·76 mEq Na+/l. All the heterozygotes tested were parents of known homozygotes. The parotid saliva from the cystic fibrosis homozygotes gave sodium levels from 12-52 mEq Na+/l., with a mean value of 27·2-14·2 mEq Na+/l.

No correlation between parotid saliva sodium levels and ages of the test subjects within the groups was found in this survey.

**Discussion**

**Comments on the electrode.** The sodium responsive electrode used in this survey was well tolerated by the children tested and well suited for measuring the sodium content of parotid saliva. However, the following precautions have to be taken to avoid errors.

(1) The electrode has to be placed close to Stensen’s duct, as different pNa readings are obtained at different points along the jaw. pNa readings taken from both sides of the jaw were found to be in satisfactory agreement.

(2) Fluctuations in the pNa readings were observed when the electrode was moved in the mouth, presumably due to pressure changes on the electrode surface. These fluctuations were reduced when the electrode was held steady, but fluctuations due to natural movements of the mouth by the subjects could not be avoided.

(3) Cooling of the electrode due to the subject breathing through the mouth was another cause of
fluctuating pNa readings. This was avoided by encouraging normal nasal breathing.

(4) The electrode was also sensitive to violent pH changes, as complete electrode insensitivity to hydrogen ions cannot technically be obtained. Any food, drink, or oral medicine which could cause marked pH changes in the mouth was avoided before measuring the parotid saliva pNa. Changes in the potassium or protein concentrations that might normally be expected in the mouth did not affect the pNa readings obtained.

**Stability of the electrode.** Readings of the parotid saliva sodium levels were taken at 1-minute intervals for 5 minutes in the same subject. Serial readings on a single person never varied by more than 3 mEq Na+/l. (Fig. 3). Fluctuations occurred in all pNa readings but these were never more than 0·05 of a pH unit on the scale on either side of the equilibrium point.

**Conclusions**

The results of this survey showed that there was a highly significant difference between the sodium levels of the parotid saliva in normal subjects and cystic fibrosis homozygotes. Approximately 4% of the values fell in an overlap range of 12-15 mEq Na+/l. This overlap between the groups was similar to that found by McKendrick (1962) in his survey on cystic fibrosis sweat sodium levels, but it could be increased by stimulating parotid saliva flow which raises the sodium levels in normals, heterozygotes, and homozygotes. Chauncey, Levine, Kass, Schwachman, Henriques, and Kulczycki (1962) have noted that there was only a 'borderline' significance between the sodium levels found in stimulated normal and cystic fibrosis parotid saliva.

Cystic fibrosis heterozygotes did not show any significant difference in pNa values from normals.

The pNa values of parotid saliva were not tested in neonatal children as the electrode was too large for this purpose. We hope to reduce the size of the electrode to enable a survey on neonatal children to be done, and this will be reported in due course.

Our results show that the method of detecting cystic fibrosis of the pancreas by measuring the pNa of unstimulated parotid saliva, using a dual sodium electrode, has possibilities as a diagnostic technique. There was a range of pNa values between 12 and 15 mEq Na+/l. which occurred in normals, heterozygotes, and homozygotes.

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**References**


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