LETTERS TO THE EDITOR

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The editors will decide, as before, whether to also publish it in a future paper issue.

Sweat chloride and conductivity 1

**Editor,**—As a principal author of the sweat testing document published by National Committee for Clinical Laboratory Standards (NCCLS) and consultant to the Cystic Fibrosis Foundation (CFP) (USA), I write to address an inaccuracy in the article by Heeley *et al.* The authors misrepresent the NCCLS document on the role of conductivity analysis. Nowhere does the NCCLS document address an inaccuracy in the article by Heeley *et al.*

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**VICKY A LEGRYS**
Professor, Division of Clinical Laboratory Science, School of Medicine, University of North Carolina at Chapel Hill, Carolina, USA
vlegrys@med.unc.edu


**Dr Heeley et al respond**

**Editor,**—As the principal author of the NCCLS guideline on sweat testing methodology, Dr LeGrys should be better informed of its content. It includes the clear statement that when sweat test results are obtained by conductivity measurement “the patient should be referred for quantitative sweat electrolyte testing”. In our paper we refer to this statement as implying that sweat conductivity measurement should be regarded as “unreliable for diagnostic purposes”. This surely cannot be conceived as misrepresenting the NCCLS position, as claimed by Dr LeGrys. Although the NCCLS does, by reference, attribute this advice to Cystic Fibrosis Foundation (CFP) (USA) policy, by including it in their guideline without comment or qualification, the NCCLS authors are actively promoting the advice.

The medical politics of the USA do not concern us, but rather the question as to whether there is any scientific evidence underpinning this advice which the NCCLS uphold. The result of our study suggests there is none.

Dr LeGrys quotes research findings which support the conclusion that sweat conductivity measurement is appropriate only for initial screening purposes. We contend that there is no data presented in this otherwise excellent paper which provide scientific justification for that conclusion.

Dr LeGrys is of the opinion that the conclusion we draw from our own study should have been supported by appropriate linear regression and bias plots of the data. The *Archives*’ professional statistical adviser reviewing our manuscript, which included such data analysis, thought otherwise and requested us to remove it.

Dr LeGrys suggests that his current conductivity methods are subject to evaporation error. The NCCLS document describes in the paper as unreliable; it does refer to the current conductivity methods as with chloride (albeit a very limited sample size).

Sweat chloride and conductivity 2

**Editor,**—As I understand the Scientific Method, a statement purporting to be factual, either in a scientific article or in a discussion with peers, must be supported by cited evidence that may be publicly examined for its scientific veracity.

The paper by Heeley *et al.* provides data to illustrate the equivalence of conductivity and chloride in cystic fibrosis (CF) diagnosis, and therefore corroborates the findings of an earlier clinical trial by Hammond *et al.* Further, a statistical comparison of the extensive published sweat chloride data of Shwachman *et al.* with the conductivity data of Hammond shows that the two are of equal discriminant power in CF diagnosis.

Despite this evidence, Dr LeGrys has authored a document that contains a number of assertions on this subject and on other aspects of sweat testing, that are not supported by any published results of original work of which I am aware. No clinical trial data exist which show that conductivity should only be used as a screen, that it is in any way inferior to chloride as a reliable diagnostic discriminator, or that conductivity readings of 50 mmol/l are positive for CF. Dr LeGrys’ call for more studies on this matter may be seen as an evasion of the true issue. I suggest that the time has come, albeit belatedly, for her to substantiate her case, not with opinions, but by providing proper citations for relevant experimentally obtained data to support her contentions in the said document.

In a separate article Dr LeGrys refers to conductivity as a “qualitative” assay, appearing to infer that it is less reliable than chloride analysis. The term “quantitative”, used in the pad-absorption method merely indicates that...
Letters, Book review

Dipstick examination for urinary tract infection

Editor,—We read with interest the letter by Thayyil-Sudhan and Gupta reporting their study on the role of dipsticks in the detection of urinary tract infection. We therefore, there is no information to indicate whether children who were being treated with antibiotics at or immediately before admission were included in the study. If this is the case, the possibility of false negative culture results cannot be excluded and this will add further bias to the results. No data are provided for the number of infants included in the study. It has been reported that negative dipstick tests have a higher false negative rate in rates of urinary culture frequency because decreased bladder incubation time diminishes in vivo bacterial multiplication. We are not told about the percentage of the samples, which were collected by pads, as compared with catheter specimens as this may further add to the inaccuracy of the culture results.

In our prospective study of 325 children in whom urinary tract infection was a clinical possibility, all urine was sent for laboratory examination. The laboratory was unaware of the results of the dipstick tests until the end of the study. Analysis of our data showed that the combination of negative dipstick tests for nitrite and/or leukocyte esterase gave a negative predictive value for urinary tract infection of 96.9%, with a specificity of 98.7%. The figures for infants were 96.7% and 99.2%, respectively. A positive nitrite and/or leukocyte esterase had a positive predictive value of 60% and a sensitivity of 54.6%, compared with 50% and 20.0% respectively in infants. In our series we found that there were four false negative and six false positive nitrite tests.

The dipstick tests are most likely to be useful as a screening test to exclude urinary tract infection in children but may be less suitable for infants. They should not be used to diagnose urinary tract infection. We therefore disagree with Thayyil-Sudan and Gupta in their view that if nitrites are positive, starting empirical treatment for urinary tract infection seems to be reasonable until cultures are reported.

H L WEBSTER
Senior Research Scientist, Wescor, Inc., 459 South Main Street, Logan, Utah 84321, USA
lewis@wescor.com

6 South Main Street, Logan, Utah 84321, USA
lewis@wescor.com


Dipstick examination for urinary tract infection

Editor,—We read with interest the letter by Thayyil-Sudhan and Gupta reporting their study on the role of dipsticks in the detection of urinary tract infection in children. We believe that this is a very important subject and believe that this is a very important subject and would like to comment on our results in this study.

Our study involved a selected group of children who were at an increased risk of having urinary tract infection. The inclusion criteria were the presence of any of the following: firstly, clinical suspicion of urinary tract infection; secondly, history of previous urinary tract infections or renal anomalies; thirdly, children needing antibiotics (urine culture was sent before starting antibiotics); and finally, any of the dipstick tests (nitrites, protein, leukocyte esterase, or blood) being abnormal.

Out of the 500 children admitted to the hospital during the study period, only 312 met the above criteria and were included in the study. Urine culture was done for all these children, which reflects the local practice at our hospital of sending urine for culture. We wanted to see if there was a change in practice to urine culture being done only if nitrites or leukocyte esterase were positive would be effective in reducing the number of urine cultures.

The inclusion criteria for Sharief and colleagues' study were a clinical suspicion of urinary tract infection, when urine cultures were sent and dipstick testing was done. We found that urinary tract infection could easily be missed if urine culture was used. If nitrites or leukocyte esterase are positive. Surprisingly, the results of both our study and theirs are similar: sensitivity was 34.4% and 90.0%, and specificity was 99.7% and 99.2% in our study and Sharief’s study respectively. Negative predictive value was 92.4% in our study and 96.7% in Sharief’s study. Only the interpretation of the results is different.

A test with such a low sensitivity cannot be recommended as a screening test to exclude urinary tract infection. Urinary tract infection may result in irreversible renal damage in infants and therefore most care should be given to the detection of this infection in this age group. Unfortunately, there is no group where sensitivity of dipstick testing is the lowest, although it is imperative to be aware that there is a group where sensitivity of dipstick testing is the lowest, because of its high negative predictive value, dipstick testing may have some role as a screening test for urinary tract infection in situations where the incidence is very low. Positive nitrites have a high specificity for urinary tract infections, which was the basis of our suggestion that if nitrites are positive, especially in a febrile infant, empirical treatment with antibiotics may be considered until the result of urine culture is obtained. However, it should not be the whole criterion for diagnosis of this infection.

S THAYYIL-SUDHAN
S GUPTA


Should repeat lumbar punctures be routinely done in neonates with bacterial meningitis? Results of a survey into clinical practice

Editor,—Neonatal meningitis remains a very important cause of morbidity and mortality, with 30% death or handicap rate reported in a recent study. In common with other clinical situations, the evidence base for some of the management recommendations for good clinical practice is hard to find. One particular aspect of the management of neonatal bacterial meningitis is whether or not a repeat lumbar puncture should be undertaken routinely. Several standard textbooks of neonatology recommend repeating the lumbar puncture routinely in the course of neonatal bacterial meningitis to ensure that “meningitis” continues to be proved. This recommendation is based on past practice, and current evidence in favour or against repeating the lumbar puncture is not overwhelming.

However, we have observed that day to day clinical practice appears to have changed and fewer repeat lumbar punctures are being done. To investigate this we performed a simple questionnaire survey across the north...
EDITOR,—Dr Elliman is noted for his careful analysis of evidence to support a change, yet in some instances, especially when there is little new published evidence, there is a widely held and practised view that recommends a particular management practice. It will not help in deciding the duration of therapy—63 (70%) do not routinely repeat LP and will only repeat if clinically indicated, because: It is recommended in textbooks—9 (45%); It is a good mix of experience—58% consultants and 42% trainees in paediatrics/neonatology; 47% had more than 10 years neonatal experience (table 2). We agree that corticosteroids do not inhibit, except at very high concentrations, degradation of the eosinophils induced by incubation with opsonised particles, such as Sepharose beads in vitro. However, there is overwhelming evidence that cytokines such as IL-5 prime eosinophils for increased release of granule proteins in this situation, and that they inhibit cytokine-mediated prolongation of eosinophil survival.

Two randomised trials of non-abusive smacking were significantly better than six alternative methods, such as reasoning or time out. Causal evidence of detrimental effects of customary physical punishment was less conclusive and limited to overly frequent smacking—for example, three times weekly for 6–9 year olds. In head-to-head comparisons, the effects of non-abusive or customary smacking rarely compared unfavourably with any disciplinary alternative, whereas its effects were significantly better than six alternative disciplinary tactics, such as reasoning or time out. Causal evidence of detrimental effects of customary physical punishment was less conclusive and limited to overly frequent smacking—for example, three times weekly for 6–9 year olds. In head-to-head comparisons, the effects of non-abusive or customary smacking rarely compared unfavourably with any disciplinary alternative, whereas its effects were significantly better than six alternative disciplinary tactics, such as reasoning or time out.

The American Academy of Pediatrics’s co-sponsored scientific consensus conference on corporal punishment used a more scientific approach than the Elliman-Lynch summary. First, it carefully defined spanking as a subset of corporal punishment. Second, it incorporated a range of scientifically validated perspectives into summary statements, that were more balanced than the Elliman-Lynch perspective. Third, it solicited the first systematic review of child outcomes of non-abusive or customary physical punishment by parents, which was recently updated.

Both reviews concluded that non-abusive smacking had consistently beneficial child outcomes in the most causally conclusive studies—for example, randomised trials. Both non-compliance and fighting decreased in 2–6 year olds after non-abusive smacking was used to back up milder disciplinary tactics, such as reasoning or time out. Both reviews concluded that non-abusive smacking had consistently beneficial child outcomes in the most causally conclusive studies—for example, randomised trials. Both non-compliance and fighting decreased in 2–6 year olds after non-abusive smacking was used to back up milder disciplinary tactics, such as reasoning or time out. Both reviews concluded that non-abusive smacking had consistently beneficial child outcomes in the most causally conclusive studies—for example, randomised trials. Both non-compliance and fighting decreased in 2–6 year olds after non-abusive smacking was used to back up milder disciplinary tactics, such as reasoning or time out.

We need the full picture on both aetiology and management. We agree that the asthma attacks may have resolved spontaneously in some cases, which was precisely why we stated that the markers fell in association with steroid therapy, and not as a consequence of the inflammatory process. Nevertheless, the statistical analysis suggests that the chances this occurred at random are extremely low. We agree that corticosteroids do not inhibit, except at very high concentrations, degradation of the eosinophils induced by incubation with opsonised particles, such as Sepharose beads in vitro. However, there is overwhelming evidence that cytokines such as IL-5 prime eosinophils for increased release of granule proteins in this situation, and that they inhibit cytokine-mediated prolongation of eosinophil survival.

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Progress in the management of disease in the newborn has carried with it a recognition of the substantial risk of injury to the immature nervous system. The aspiration to localise the substantial risk of injury to the immature newborn has carried with it a recognition of the problem of signs available in the early newborn period is easily understood. The problem is that the signs available are in themselves usually insufficient to allow precision. In addition, the signs of the lesion(s) alters, or may become silent, often to reappear later as a different but nevertheless highly significant impairment.

The evaluation of the newborn nervous system was originally based upon concepts learnt from adult neurology. The baby was seen as demonstrating little or no cortical or cerebellar activity and the study of primary reflexes predominated. The approach of adult neurology, with emphasis on localisation of the lesion, becomes less applicable in the younger child. In the newborn period, focal insults to the brain will often give rise to generalised disturbances and, contrarily, generalised disturbances may show focal deviations. Recognition of these phenomena has led to a progression from the concept of a localisation based neurology to one which sees the infant displaying a neurological/behavioural repertoire. Over the past several decades Saint Anne Dargassies, Prechtl, Amiel Tison, Brazelton, Dubowitz, and others have, through meticulous study, done much to illuminate this area. Through these studies, awareness of the importance of the behavioural state of the baby, as well as the more detailed neurological items has evolved.

A second problem in this area, particularly in relation to research studies, has been the development of a systematic newborn neurological examination which is reliable and repeatable. This has been the subject of the two editions of this work. The first, published in 1981, gave a detailed, easily understood and applied system for the neonatal neurological examination. The current edition brings that work up to date. New material is presented, refinement of the scheme has occurred, and the examination is described. Items which were less discriminatory of pathology from the 1981 version have been withdrawn and, following the work of Prechtl, more emphasis is placed on the analysis of general movements. There is a further post neonatal to two year old infant neurological examination proforma presented briefly at the end of the text.

The text is essentially a manual on the application of this neurological examination scheme. It is easy to follow and the segments of the examination are presented clearly with excellent photographs and line drawings of each manoeuvre. There is also a useful addendum ("cautionary tales") to each section of the examination, giving guidance on possible pitfalls and sources of error. There is a lot of very useful information on the variations in findings in term and preterm infants, and particularly the changes in the neurological features of preterm infants as they grow towards term. There follows a section on the development of an optimality score from the observed items of the assessment. This section deals with the results of a survey of 224 normal term infants. In this study each item of the scheme was plotted, and centile values (and thereby optimality scores) were computed. This provides quantification of the assessment, a sense of the range of findings to be expected, and can be useful in correlating lesions observed on neuro imaging with clinical findings. Chapter six deals with the scheme in relation to findings in infants with recognised brain lesions.

The book is not designed to be a text of neonatal neurology and readers looking for discussion of neurological disease states will be disappointed. As a description of a comprehensive and easily applied system of neonatal neurological examination the new edition succeeds admirably.

MICHAEL F SMITH
Neonatal Intensive Care Unit, Jessop Hospital for Women, Leaegreave Road, Sheffield S7 1RE, UK