Background: There is a need to prognosticate the severity of cystic fibrosis (CF) detected by newborn screening (NBS) by early assessment of CF Transmembrane Conductance Regulator protein function (CFTR). We introduce novel instrumentation and protocol for evaluating CFTR activity as reflected by β-adrenergically stimulated sweat secretion.

Methods: A pixilated Image-Sensor detects sweat rates. Compounds necessary for maximum sweat gland stimulation are applied by Iontophoresis, replacing intradermal injections. Results are compared to a validated β-adrenergic assay that measures sweat secretion by evaporation (Evaporimetry).

Abstract:

IMAGE-BASED β-ADRENERGIC SWEAT RATE ASSAY CAPTURES MINIMAL CYSTIC FIBROSIS TRANSMEMBRANE CONDUCTANCE REGULATOR (CFTR) FUNCTION

Danieli Barino Salinas* [Assistant Professor of Pediatrics],
Children’s Hospital Los Angeles, University of Southern California (USC), Los Angeles, CA

Yu-Hao Peng,
Viterbi School of Engineering, University of Southern California, Los Angeles, CA

Brian Horwich,
Keck School of Medicine, University of Southern California, Los Angeles, CA

Choo Phei Wee,
Children’s Hospital Los Angeles, University of Southern California, Los Angeles, CA

Eric Frisbee,
Cystic Fibrosis Research Laboratory, Stanford University, Palo Alto, CA

Jean-Michel Maarek
Viterbi School of Engineering, University of Southern California, Los Angeles, CA

Disclosure: Authors have no conflicts of interest referable to this research.

Category of the study: clinical translational research
**Results:** Ten healthy controls (HC), 6 Heterozygous (Carriers), 5 with cystic fibrosis screen positive, inconclusive diagnosis (CFSPID), and 12 cystic fibrosis (CF) individuals completed testing. All individuals with minimal and residual function CFTR mutations had low ratios of β-adrenergically stimulated sweat rate to cholinergically stimulated sweat rate (β/chol) as measured by either assay.

**Conclusions:** β-adrenergic assays quantitate CFTR dysfunction in the secretory pathway of sweat glands in CF and CRMS/CFSPID populations. This novel Image-Sensor and Iontophoresis protocol detect CFTR function with minimal and residual function and is a feasible test for young children because it is insensible to movement and it decreases the number of injections. It may also assist to distinguish between CF and CRMS/CFSPID diagnosis.

1. Introduction

Cystic Fibrosis (CF) is a genetically transmitted disease caused by a defective anion channel, Cystic Fibrosis Transmembrane Conductance Regulator (CFTR). Defective CFTR results in abnormal mucus secretions causing obstruction in the lungs and pancreas leading to life-threatening lung infections and malnutrition. Thousands of infants in the US are identified annually through newborn screening (NBS) to be at risk for developing CF and are referred to CF Foundation accredited centers for confirmation of diagnosis (www.cff.org). The standard sweat chloride test (SCT) uses quantitative pilocarpine iontophoresis (QPIT) to diagnose CF. (1) This test is based on the pathophysiology of defective chloride reabsorption in sweat ducts that causes elevated chloride concentrations in sweat from patients with CF. (2) (Supplemental figure S1) The SCT is accepted worldwide as the gold standard diagnostic test for CF; however, it was established with patients with severe disease vs. healthy subjects. (3) It assesses CFTR protein function through analysis of the level of chloride content in the sweat collected after 30 minutes of electrical-chemical stimulation of the skin. (1,7) Sweat chloride ≥60 mmol/L confirms the diagnosis; ≤29 mmol/L is considered negative or “CF unlikely”; and 30 mmol/L to 59 mmol/L is considered intermediate. (4) Intermediate sweat chloride results (and even negative results with 2 identified CFTR mutations detected by NBS) represent a significant challenge to physicians when determining prognosis and the appropriate level of management, because those individuals can potentially develop moderate to severe disease and conversely may not develop CF symptoms at all. Treatment is only initiated upon confirmed diagnosis or presentation of symptoms. Currently, positive NBS-children with intermediate range will be followed and standard SCT repeated multiple times along the child’s life. Signs and symptoms and confirmed diagnosis have been reported in individuals with negative and intermediate SCT in specific genotypes: D1152H and 3849+10kbC>T. (5–8)

Since the CF gene was discovered in 1989, > 2,000 different CFTR variants have been identified (http://genet.sickkids.on.ca), and the disease is now better understood as a spectrum expressing mild to severe phenotypes. A broader use of genetic testing in NBS revealed a subset of subjects genetically screen-positive, who are asymptomatic and have negative or intermediate sweat chloride test results. These individuals are given a diagnosis of CFTR-Related Metabolic Syndrome (CRMS) or CF screen positive, inconclusive diagnosis (CFSPID) and they are at risk to develop disease associated with CFTR protein
dysfunction in the future. Genotyping is now part of most NBS programs and an important component of confirmation of diagnosis in the US and in other parts of the world. The focus on genotyping is justified by the fact that there are three recently approved drugs capable of partially improving function of the defective protein. With this shift in paradigm, the standard SCT has become limited in defining diagnosis. Thus, there is a clear need to develop an assay that can better assess CFTR function in vivo, not only to define diagnosis and prognosis in patients with mild to severe disease, but also to define potential responses to CFTR targeted drugs.

The rate of adrenergically stimulated sweat secretion test provides a unique, essentially non-invasive, physiological approach to determine CFTR function in vivo. Human sweat glands secrete sweat via stimulation of two apparently independent neural pathways: cholinergic and adrenergic. The cholinergic pathway controls thermoregulation and the β-adrenergic pathway is thought to be related to a “fight or flight” response. The cholinergic pathway appears to be independent of CFTR, while the β-adrenergic pathway is highly CFTR-dependent. Consequently, β-adrenergic sweat rates can be used as a physiological marker of CFTR activity. The initial sweat secretion test was developed in adults by measuring the ratio of the rate of evaporation of sweat with an Evaporimeter (cyberDERM inc) after sequentially stimulating the two pathways independently. Figure 1 shows how the standard SCT differs from the β-adrenergic sweat secretion test. We applied the Evaporimeter technology to test 30 children who were screen-positive for CF. Although safe and well tolerated, measurements in infants (n=5) were inaccurate due to movement artifacts; and in the preschool age group the sweat rate response to β-adrenergic stimulation (CFTR function) was not statistically different among those with symptoms and deleterious mutations (CF n=16) versus those who were asymptomatic and with benign variants of the gene (CRMS/CFSPID n=10). Thus, sweat secretion assays as measured by the Evaporimeter did not adequately discriminate between verified CF subjects and the CRMS/CFSPID group.

The present work presents new technology and protocol that determine in vivo CFTR function as reflected by β-adrenergically stimulated eccrine sweat secretion. We hypothesize that this new technology and protocol are minimally invasive, suitable to be applied to young children, and sensitive to detect minimal and residual CFTR function. The technology may be used to develop a clinical test to confirm the diagnosis of CF after a positive NBS, to define prognosis of CF screen-positive infants to personalize care, and/or to quantify responses to CFTR targeted therapy in vivo.

1. Methods

2.1. Subjects

The Institutional Review Boards from Children’s Hospital Los Angeles (CHLA) and the University of Southern California (USC) approved the study. Written informed consent was obtained from all subjects ≥13 years old, and from the subjects’ parents for all minors (<18 years old). Assent was obtained from subjects 7–12 years old. Only non-CF health controls (HC) ≥18 years old were recruited during protocol development. HC were recruited based on self-reporting of no chronic respiratory symptoms and no family history of CF; they did not.
not have SCT or genotyping. CF and CRMS/CFSPID patients were diagnosed based on accepted Cystic Fibrosis Foundation (CFF) criteria and recruited from CHLA CF center.(4) The parents of patients who were previously genotyped were also recruited for testing.

2.2. Instrumentation

The measurement is based on the change in local electrical capacitance that results from the generation of sweat by eccrine sweat glands stimulated by cholinergic or adrenergic agonists. The capacitive fingerprint module (HF-EM401, HuiFan Technology Co., China) senses the capacitance on each pixel of the detector to produce an image of the fingerprint ridges and valleys, the air gaps in the valleys having a higher capacitance than the ridges. Sweat production decreases the local capacitance and modifies the image. A thin piece of filter paper placed on the sensor masks the texture of the skin surface. The resulting image is generated as sweat wets the paper non-uniformly as a function of local sweat secretion from individual glands.

To conduct the sweat test, the capacitive sensor module is positioned on the ventral surface of the forearm with a 3D-printed holder that maintains the module in tight contact with the skin with a Velcro strap wrapped around the forearm. Since the image intensity varies with the pressure with which the module is held on the skin, a resistive pressure sensor (FSR 400, Interlink Electronics, Westlake Village, CA) was installed behind the fingerprint module. A cushion of stiff plastic foam was used to normalize the contact pressure between the skin and the sensor module and standardize the contact pressure of the sensor module on all subjects (Figure 2).

Two types of filter paper (GE Healthcare, Pittsburgh, Pa) were used: 1. Whatman AE 99 (120 μm thickness, 8 μm pore size) was used to measure sweat production after cholinergic stimulation; 2. Whatman Cyclopore polycarbonate membrane (7 μm thickness, 0.1 μm pore size) was used after adrenergic stimulation. Two different filter papers were used because the thinner Cyclopore paper allowed detection of small amounts of sweat observed in CF subjects after adrenergic stimulation. However, the thinner paper saturated quickly with the larger amounts of sweat observed in HC after cholinergic stimulation and therefore required thicker paper.

2.3. Image acquisition and analysis

Custom software written in LabVIEW (National Instruments, Austin, TX) was used to transfer the sensor image every 5 seconds from the fingerprint module controller to the computer using serial communication. The grey-scale images (200 X 153 pixels) coded on 8 bits were displayed on the computer monitor to visualize the production and accumulation of sweat on the sensor surface. Analysis of the image sequence during a sweat stimulation protocol and evaluation of the sweat rate was carried out as shown in Figure 3.

2.4. Study protocol

2.4.1 Protocol development

2.4.1.a) Iontophoresis of Pilocarpine 1%: Iontophoresis was performed to decrease the number of intradermal injections. Initial measurements were performed comparing the
standard Pilegal 0.5% (ELITechnology) and Pilocarpine 1% (USC Compounding pharmacy). Maximum sweat rate was measured by Evaporimeter (Model RG1, CyberDERM inc., Broomall, PA). The subject was seated comfortably with both arms resting on a table. A small area of skin on the ventral surface of the forearms (about 4x4 cm$^2$) was gently cleaned with distilled water, dried, and covered by a thin layer of mineral oil to minimize transepidermic water loss. Evaporimeter probes were placed on these cleaned areas and secured with Velcro bands. Baseline activity was measured for 5 minutes. Sweat Secretion was then stimulated by iontophoresis of Pilegal® on one arm and subsequently by Pilocarpine 1% on the opposite arm. Circular-shaped gels (Pilegal®) or circular-shaped felt patches soaked in Pilocarpine 1% solution were mounted on the sweat-inducer iontophoresis disc (Model 3700, Wescor Inc, Logan, Utah), which applies 1.5 mA current for 5 min between electrodes to iontophorese drugs transcutaneously into the intradermal space (Figure 2).

2.4.1.b) Intradermal injections compared to iontophoresis: Quinton et al validated the Evaporimeter protocol using subsequent intradermal (ID) injections of Carbachol, Atropine, and a β-adrenergic cocktail.[14] Serial injections were well tolerated by adults in the validation study, but would be less manageable in a pediatric population. We then compared ID Carbachol 0.1 ml = 0.01 μg (Miostat intraocular, Alcon®) and Iontophoresed Pilegal 1% measured by Evaporimeter in similar fashion as described for Pilegal 0.5% and Pilocarpine 1% experiments. Similarly, ID Atropine 0.2 ml = 8.8 μg (Bausch & Lomb) and Atropine 5% (USC Compounding pharmacy) were compared. Several attempts were made to Iontophorese the β-adrenergic cocktail; however, sweat rate results were not reproducible when Atropine, Aminophylline, and Isoproterenol were iontophoresed concurrently or sequentially.

2.4.1.c) Sweat gland potentiation: We evaluated the use of the ID β-adrenergic cocktail in HC (n=2) with and without prior “sweat gland potentiation” with ID Carbachol followed by ID Atropine measured by both methods.[17]

2.4.1.d) Reproducibility of Image-Sensor: Reproducibility of the sweat response measured by the Image-Sensor was assessed in 3 HC who received 3 ID Carbachol at neighboring sites on the ventral surface of the arm. We also map the darkened spots visualized in response to Pilocarpine and β-adrenergic cocktail to demonstrate that the same sweat glands were stimulated.

2.4.2 Final protocol—in the final protocol, sweat rate was measured on both arms concurrently (Figure 2). The subject was seated and skin preparation of the forearms was as described above. Evaporimeter was applied to the right arm and the Image-Sensor to the left arm. Baseline secretion was measured for 5 minutes. Sweat secretion was then stimulated simultaneously on both arms by iontophoresing Pilocarpine 1%, followed by Atropine 5%. The skin was marked with a pen to reposition the probes in the same site. The effect of blocking cholinergic response with atropine was confirmed by the Evaporimeter only on the right arm. Once the Evaporimeter signal stabilized (usually 2 to 5 Evaporimeter units above baseline), an intradermal injection of 0.2 ml β adrenergic cocktail was administered.
containing: 8.8 μg of Atropine, 940 μg of Aminophylline (Regent Laboratories Inc.), and 8 μg of Isoproterenol (Hospira Inc.). After injecting both sides, the Evaporimeter and the Image-Sensor probes were repositioned over the sites of intradermal injections on the right and left arms, respectively.

### 2.5. Statistical analysis

Patient characteristics and distribution of maximum sweat rate were described by summary statistics. Mean and standard deviation (SD) were used to describe the variables with a normal distribution, while median and interquartile range was used for the variables with non-normally distributed. Frequency and percentages were used to summarize categorical variables. Maximum sweat rate measurements were natural log-transformed to appropriately adjusting for the non-normal distribution and analysis of variance was used to examine the differences in these measurements for both Evaporimeter and Image-Sensor assays across HC, Carrier, CRMS/CFSPID, and CF groups. Then, multiple comparisons with Tukey adjustment was used to further assess which of the pairwise comparisons differ in maximum sweat rate measurements. Taking into account that the two methods have different units, z-scores were calculated based on total sample mean for each method in order to assess β/chol ratio difference for all groups. The difference in the z-scores between two methods was assessed by repeated-measure ANOVA. Statistical significance was set at 5% level with 2-sided throughout the analyses. All statistical computations were done in Stata/SE 15.1 (StataCorp, College Station, TX).

### 3. Results

#### 3.1. Initial results during protocol development

**3.1.a) Iontophoresis of Pilocarpine 1% and Pilogel 0.5% were equivalent—**Iontophoresis of Pilogel 0.5% and Pilocarpine 1% on the same HC individuals were equivalent (104 ± 7.4 and 104 ± 13.4 mgH2O/m2/hr respectively, n=5; p=0.986).

**3.1.b) Intradermal injections compared to iontophoresis—**Intradermal Carbachol 0.1 ml = 0.01 μg and Iontophoresed Pilocarpine 1% measured by Evaporimeter had similar results: 84 ± 11.5 and 82 ± 12.7 mgH2O/m2/hr (p=0.49, HC n=33, CF n=13). Atropine injected versus iontophoresed blocked 95% and 83% respectively of the cholinergic sweat rate (n=11, p = 0.345).

**3.1.c) Sweat gland potentiation with cholinergic stimulation is necessary for maximum β-adrenergic stimulated rate—**When evaluating the use of the ID β-adrenergic cocktail in HC (n=2) with and without prior “sweat gland potentiation,” maximum sweat rate in response to adrenergic stimulation without potentiation was about half the rate with potentiation, when measured by both Evaporimeter and Image-Sensor: Evaporimeter - no potentiation: 40 (36–44) median (95%CI) and with potentiation: 77 (66–83) (p=0.004). Image-Sensor - no potentiation: 36 (33–38) and with potentiation: 65 (49–112) (p=0.001) confirming previous findings. (17, 18)
3.1.d) **Sweat Rates measured by the Image-Sensor were reproducible**—Sweat rates after injection of Carbachol in 3 different locations on the volar aspect of the arm of 3 individual subjects showed: subject 1 = $102.57\pm1.89$, subject 2 = $69.66\pm16.60$, and subject 3 = $184.03\pm19.03$ intensity/time/mm$^2$. The sequence of sensor images gathered after stimulation of the sweat glands demonstrated the production and gradual accumulation of sweat at the stimulated site. Mapping the darkened spots visualized in response to Pilocarpine and β-adrenergic cocktail indicated that the same sweat glands were measured with each repeated stimulation (Figure 4).

3.2. **Evaporimeter and Image-Sensor measurements were comparable**

A total of 10 HC, 6 Heterozygous (Carriers), 5 CRMS/CFSPID, and 12 CF individuals completed testing using the final protocol (Table 1). Group characteristics were as follows: Age in years (SD): HC = 32 (13), Carrier = 38 (8), CRMS/CFSPID = 8 (1), and CF = 16 (4); Race/ethnicity: HC = 100% White/10% Hispanic, Carrier = 100% White/67% Hispanic and 33% Caucasian, CRMS/CFSPID = 100% White/60% Hispanic and 40% Caucasian, and CF = 83% White and 17% Black/33% Hispanic, 50% Caucasian, and 17% African descent. 

*CFTR* mutations were identified by newborn screening and parent DNA testing. Evaporimeter and Image-Sensor measurements had a linear relationship when quantifying maximum sweat rates in response to β-adrenergic stimulation in HC and CF (spearman correlation = 0.8379, supplemental figure S2). There is no significant difference in the sweat rate measurements between Evaporimetry and Image-Sensor according to z-scores analyses. (Table 2)

3.3. **Minimal and residual function *CFTR* mutations yield low β-adrenergic/cholinergic sweat rate ratios**

Table 1 shows individual β/chol ratios measured by both assays. Figure 5 shows individual Image-Sensor-β/chol ratio plotting and genotype with corresponding *CFTR I-VI* classes and functional classification. All CRMS/CFSPID subjects had one or both *CFTR* mutations classified as residual function as did 2 of the 12 CF subjects (siblings with the same genotype: 3849+10KbC>T/935delA). All minimal and residual function *CFTR* mutation combinations had low β/chol ratios when measured by both assays.

3.4. **β-adrenergic/cholinergic ratio in natural log from Evaporimetry and Image-Sensor measurements were able to differentiate between CFSPID and CF.**

β/chol ratio (after natural log transformed) from Evaporimetry and Image-Sensor measurements were equally successful to show significant differences between HC and CFSPID, HC and CF, Carrier and CFSPID, Carrier and CF, and CFSPID and CF. (Figure 6)

**Discussion**

The Image-Sensor and Iontophoresis protocol presented here illustrate initial development of a novel application of technology to detect minimal and residual CFTR function *in vivo*. The Image-Sensor assay detects sweat rates in response to β-adrenergic stimulation in individuals diagnosed with CF and CRMS/CFSPID that can be performed on pediatric subjects. As seen previously with the Evaporimeter, both groups had low β-adrenergic sweat
rate, (16) which suggests CFTR dysfunction in CFTR dependent sweat secretion in glands of both CF and CRMS/CFSPID children. The advantages of this technology and proposed Iontophoresis protocol over other proposed assays are that: a) it can be completed in less than thirty minutes, b) it requires only one ID injection, as Iontophoresis of Pilocarpine and Atropine are applied prior to the β-cocktail injection; c) the apparatus is insensitive to subject movement, making it more feasible for use in a younger population. and d) it detects β-induced sweat secretion over the full range of CFTR function, which may assist in differentiating between CF and CRMS/CFSPID diagnosis.

Since the mid-80s when Sato and Sato discovered that CF glands do not secrete sweat in response to β-adrenergic stimuli, but secrete normally after exposure to cholinergic secretagogues, there have been several attempts to develop an assay with detection limits that are low enough to capture differences in β-adrenergic sweat rate responses among CF subjects with different CFTR mutations.(15, 17, 18) A ratiometric assay comparing β-adrenergic to cholinergic sweat rates using the Evaporimeter was validated several years ago for differentiating heterozygotes from CF genotypes.(14) However, Evaporimetry was not sufficiently sensitive to differentiate among CF subjects with different levels of disease severity such as pancreatic sufficient versus insufficient subjects. As there was a significant difference between CFTR-related disorder (which is a single organ disease caused by two CFTR mutations) and CF, our group attempted to apply Evaporimetry to determine differences between CF and CRMS/CFSPID.(16) To our surprise, the CFTR function in CRMS/CFSPID assayed by this method appeared just as low as in CF individuals. The current Image-Sensor protocol shows no significant difference with the Evaporimeter measurements, but with a slight advantage of a lower detection limit for the CFTR-related response. The Image-Sensor detected β-adrenergic sweat rate responses in 2 individuals who had zero response (consequently zero ratios) when measured by the Evaporimeter (genotypes: G330X/G480C and F508del/W1282X). Since these genotypes are known to express very little protein, these results may be an artifact and not the actual β-adrenergic sweat secretion. However, in view of the cautious phases of protocol development including reproducibility and mapping of the sweat glands, this possibility seems unlikely. The β-adrenergic induced sweat secretion was low regardless of genotype, CFTR class, or category of residual/minimal function (Figure 5). Yet, the Image-Sensor detected some level of β-adrenergic sweat rate responses in 100% of CF subjects while the Evaporimeter detected sweat rate in 80% of subjects with residual/minimal CFTR function (Table 1).

An optical ratiometric technology assay also comparing β-adrenergic to cholinergic sweat rates has been developed to measure CFTR function in persons with CF-pancreatic sufficiency/insufficiency and with CFTR-related disorder.(17, 19) This assay was used to measure the effect of a CFTR modulator (Ivacaftor) on single, individual sweat glands in subjects carrying G551D and R117H-5T.(17, 20) β-adrenergic response was seen in 16–21% of sweat glands of individuals treated with the drug and in no glands in subjects on placebo. This finding suggested that the optical ratiometric method is more sensitive than the Evaporimeter to measure sweat rate response to CFTR-targeted therapy. (21) Although highly sensitive, the optical ratiometric assay does not detect β-adrenergic induced sweat secretion in individuals carrying CFTR mutations classified as minimal function and in others with CFTR-related disorder measurements were highly variable (0 to 0.6). (17, 19)
Additionally it requires maintaining the same position without any movement for a prolonged period of time along with several ID injections, making it less suitable for children.

The Image-Sensor and the Evaporimeter were equally successful to differentiate between groups (HC, Carrier, CRMS/CFSPID, and CF). Interestingly, the natural log of β/chol ratios showed significant difference between CRMS/CFSPID and CF, suggesting that both methods and this Iontophoresis protocol may complement standard SCT in defining the diagnosis of screen-positive children carrying CFTR mutations/variants of unknown clinical significance. Another possible application in this NBS population would be to use this assay to monitor CRMS/CFSPID to CF reclassification over time. The current findings are different than our previous work, which showed no difference between CF and CRMS/CFSPID pre-school age children when sweat rate was measured by Evaporimetry after Carbachol and β-cocktail injections given in different days. The difference may be explained by several factors: 1. Different protocol (sweat glands were not potentiated); 2. Different age groups (sweat glands may not be fully developed in 4 to 6-year old children); and 3. Small samples in both studies. Also, in our experiments the Evaporimeter (or the Image-Sensor) did not detect a significant difference between HC and Heterozygous (Carriers) as previously seen by other groups.(14, 17, 19) Thus, it is important to plan for a larger study using this new technology and Iontophoresis protocol, including all four groups.

The present protocol is better tolerated because it is less invasive, as it uses 1 versus 3 ID injections. The Iontophoresis of Pilocarpine and Atropine was developed after our group repeated and confirmed that pre-stimulation of the sweat gland with cholinergic drugs potentiates β-adrenergic induced sweat secretion.(17) Ideally, we would eliminate all injections, but Iontophoresis of β cocktail were not reproducible when testing the same individual on the same or different days. A possible explanation is the fact that the 3 drugs in the cocktail (Atropine, Aminophylline, and Isoproterenol) are not pharmacologically compatible. These compounds have different charges, pH, and Aminophylline accelerates degradation of Isoproterenol. As in previous protocols β cocktail drugs were mixed immediately before injecting or iontophoresing so the drug effect would not be compromised. For Iontophoresis more promising results were achieved when drugs were iontophoresed sequentially. However, three sequential Iontophoresis prolonged the protocol to about an hour and caused skin irritation, which created measurements artifacts and was judged to be more uncomfortable than injections.

Limitations to this work include: 1. Small sample size that completed the final protocol, which can be justified by the fact that this is an introduction to this new technology and protocol; validation studies will require bigger samples (precisely n=15 in each of the 4 groups for 80% power). On the other hand, within the CRMS/CFSPID group we recruited a broad representation of genotypes: from individuals carrying benign variants (F508del and (TG)11–5T) to others carrying variants of varying clinical consequence (F508del and D1152H).(22, 23) 2. The Image-Sensor does not yield results expressed in physical units. To accommodate the different units, we used z-scores to compare the two methods. 3. There was no direct comparison to the standard SCT. We considered this step irrelevant to the
proof of concept of this protocol as these two assays measure different functions of CFTR in the sweat gland (secretory versus absorptive).

In conclusion, the β-adrenergic assays remain as a valid method to complement the standard SCT, since CFTR secretory rather than absorptive function is assayed. The advantages of the Image-Sensor and this Iontophoresis protocol over previously proposed assays are the detection of CFTR function with minimal and residual function and its practical utility for measurement in young children. The fact that both Image-Sensor and the Evaporimeter assays detect β-adrenergic sweat in CRMS/CFSPID almost as low as in CF suggests CFTR dysfunction in the secretory pathway of sweat glands in both populations. The β/chol ratio (after natural log transform) successfully differentiate between CRMS/CFSPID and CF groups, when measured by both Evaporimetry and Image-Sensor. Once these findings are validated in a larger trial and by other groups, the Image-Sensor assay and Iontophoresis protocol can assist in defining CF and CRMS/CFSPID diagnosis after a positive NBS and monitoring for CRMS/CFSPID to CF reclassification over time. Future directions include testing in a larger sample of young children from NBS, and also before and after use of CFTR targeted drugs.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgements:**

Paul Quinton, Ph.D. (University of California, San Diego), for outstanding mentorship, conceptual support, and critical revision of the article for important intellectual content; Dennis Briscoe (Ellitech) for conceptual and technical support; and Lilit Grigoryan, Nadine Afari, Michael Bauschard, Alexandra Franquez, Alex Katz, and Dipayan Roy for their support in recruiting, coordinating, and executing the experiments.

**Funding:** This project received funding from NIH/NCRR/NCATS: Grant KL2TR000131; SC-CTSI Pilot Grant 8UL1TR000130; and Cystic Fibrosis Research Incorporated (CFRI Grant# Salinas15-SC01).

**References**

1. Wainwright CE, Elborn JS, Ramsey BW. Lumacaftor-Ivacaftor in Patients with Cystic Fibrosis Homozygous for Phe508del CFTR. N Engl J Med. 2015;373(18):1783–4.
2. Quinton PM. Cystic fibrosis: lessons from the sweat gland. Physiology (Bethesda). 2007;22:212–25. [PubMed: 17557942]
3. Shwachman H, Mahmoodian A. Pilocarpine iontophoresis sweat testing results of seven years’ experience. Bibl Paediatr. 1967;86:158–82. [PubMed: 6054626]
4. Farrell PM, White TB, Ren CL, et al. Diagnosis of Cystic Fibrosis: Consensus Guidelines from the Cystic Fibrosis Foundation. J Pediatr. 2017;181S:S4–S15 e1. [PubMed: 28129811]
5. Terlizzi V, Carnovale V, Castaldo G, et al. Clinical expression of patients with the D1152H CFTR mutation. J Cyst Fibros. 2015;14(4):447–52. [PubMed: 25583415]
6. Dugueperoux I, De Braekeleer M. The CFTR 3849+10kbC->T and 2789+5G->A alleles are associated with a mild CF phenotype. Eur Respir J. 2005;25(3):468–73. [PubMed: 15738290]
7. Burgel PR, Fajac I, Hubert D, et al. Non-classic cystic fibrosis associated with D1152H CFTR mutation. Clinical genetics. 2010;77(4):355–64. [PubMed: 19843100]
8. Peleg L, Karpati M, Bronstein S, et al. The D1152H cystic fibrosis mutation in prenatal carrier screening, patients and prenatal diagnosis. J Med Screen. 2011;18(4):169–72. [PubMed: 22156145]
9. Ren CL, Borowitz DS, Gonska T, et al. Cystic Fibrosis Transmembrane Conductance Regulator-Related Metabolic Syndrome and Cystic Fibrosis Screen Positive, Inconclusive Diagnosis. J Pediatr. 2017;181S:S45–S51 e1. [PubMed: 28129812]

10. Farrell PM, White TB, Howenstine MS, et al. Diagnosis of Cystic Fibrosis in Screened Populations. J Pediatr. 2017;181S:S33–S44 e2. [PubMed: 28129810]

11. Taylor-Cousar JL, Munck A, McKone EF, et al. Tezacaftor-Ivacaftor in Patients with Cystic Fibrosis Homozygous for Phe508del. N Engl J Med. 2017;377(21):2013–23. [PubMed: 29099344]

12. Ramsey BW, Davies J, McElvaney NG, et al. A CFTR potentiator in patients with cystic fibrosis and the G551D mutation. N Engl J Med. 2011;365(18):1663–72. [PubMed: 22047557]

13. Behm JK, Hagiwara G, Lewiston NJ, Quinton PM, Wine JJ. Hyposweating of beta-adrenergically induced sweating in cystic fibrosis heterozygotes. Pediatric research. 1987;22(3):271–6. [PubMed: 2889182]

14. Quinton P, Molyneux L, Ip W, et al. beta-adrenergic sweat secretion as a diagnostic test for cystic fibrosis. Am J Respir Crit Care Med. 2012;186(8):732–9. [PubMed: 22859523]

15. Sato K, Sato F. Defective beta adrenergic response of cystic fibrosis sweat glands in vivo and in vitro. The Journal of clinical investigation. 1984;73(6):1763–71. [PubMed: 6327771]

16. Salinas DB, Kang L, Azem C, Quinton P. Low Beta-Adrenergic Sweat Responses in Cystic Fibrosis and Cystic Fibrosis Transmembrane Conductance Regulator-Related Metabolic Syndrome Children. Pediatr Allergy Immunol Pulmonol. 2017;30(1):2–6. [PubMed: 28465863]

17. Wine JJ, Char JE, Chen J, et al. In vivo readout of CFTR function: ratiometric measurement of CFTR-dependent secretion by individual, identifiable human sweat glands. PLoS One. 2013;8(10):e77114. [PubMed: 24204751]

18. Sato K, Sato F. Cholinergic potentiation of isoproterenol-induced cAMP level in sweat gland. Am J Physiol. 1983;245(3):C189–95. [PubMed: 6311023]

19. Bergamini G, Tridello G, Calcaterra E, et al. Ratiometric sweat secretion optical test in cystic fibrosis, carriers and healthy subjects. J Cyst Fibros. 2018;17(2):186–9. [PubMed: 29292091]

20. Char JE, Wolfe MH, Cho HJ, et al. A little CFTR goes a long way: CFTR-dependent sweat secretion from G551D and R117H-5T cystic fibrosis subjects taking ivacaftor. PloS one. 2014;9(2):e88564. [PubMed: 24520399]

21. Kim J, Farahmand M, Dunn C, et al. Evaporimeter and Bubble-Imaging Measures of Sweat Gland Secretion Rates. PLoS One. 2016;11(10):e0165254. [PubMed: 27768743]

22. Salinas DB, Azem C, Young S, Keens TG, Kharrazi M, Parad RB. Phenotypes of California CF Newborn Screen-Positive Children with CFTR 5T Allele by TG Repeat Length. Genet Test Mol Biomarkers. 2016;20(9):496–503. [PubMed: 27447098]

23. Salinas DB, Sosnay PR, Azem C, et al. Benign and Deleterious Cystic Fibrosis Transmembrane Conductance Regulator Mutations Identified by Sequencing in Positive Cystic Fibrosis Newborn Screen Children from California. PLoS One. 2016;11(5):e0155624. [PubMed: 27214204]
Figure 1: The two images on the left depict the traditional sweat chloride test that distinguishes between normal and CF phenotypes on the basis of the differences in concentration of chloride in the final sweat. Sweating is stimulated by the iontophoresis of a cholinergic stimulus (typically pilocarpine), which triggers both normal and CF glands equally. The normal duct readily reabsorbs NaCl, resulting in the output of hypotonic sweat. In contrast, the CF duct poorly reabsors NaCl, yielding sweat with abnormally high NaCl concentrations. The images on the right show the ratiometric sweat rate test; it distinguishes CF from normal based on the defective secretory functions of the CFTR-mediated adrenergic pathway. That is, when an adrenergic stimulus is applied independent of cholinergic stimulus in a normal gland (i.e. cholinergic stimulus is blocked by atropine), sweat is secreted via CFTR-dependent pathway. Because CFTR function is defective in the CF gland, the volume of sweat secreted is significantly decreased or absent. The ratiometric sweat rate test uses the adrenergic sweat rate normalized by the individual’s cholinergic sweat rate (hence, “ratiometric”) to account for variability in sweat gland density and individual sensitivity to sweat gland stimulation. A higher adrenergic-to-cholinergic ratio indicates better in vivo CFTR functional activity while a lower ratio indicates impaired CFTR function.
Drugs were administered identically on both arms: Pilocarpine 1% by Iontophoresis → Atropine 5% by Iontophoresis → β-adrenergic cocktail (Atropine 8.8 μg, Aminophylline 940 μg, and Isoproterenol 8.0 μg) by intradermal injection. 

**Figure 2: Diagram of the study protocol and equipment.**

Drugs were administered identically on both arms: Pilocarpine 1% by Iontophoresis → Atropine 5% by Iontophoresis → β-adrenergic cocktail (Atropine 8.8 μg, Aminophylline 940 μg, and Isoproterenol 8.0 μg) by intradermal injection. **A** = drug soaked felt patches for iontophoresis; **B** = Sweat Inducer (A and B were applied to both sides); **C**, **D**, and **E** = Image-Sensor; **F** = Pressure sensor; and **G** = cyber-Derm Evaporimeter and Evaporimeter probes as applied.
Figure 3: Analysis of the image sequence during a sweat stimulation protocol and evaluation of the sweat rate with the image sensor.

Panel A shows the sequence of images acquired every 5 seconds during the sweat stimulation protocol. Images collected during baseline conditions are sampled first and are followed by images collected after cholinergic stimulation. The sensor is briefly removed from the measurement site to allow for pilocarpine iontophoresis between the two sequences. The baseline image acquired 50 seconds after the start of the sequence is examined visually after the data collection is completed to select the largest region of interest (ROI) free of artifacts and edge effects (B). The mean and standard deviation of the grey levels of all pixels in the ROI are computed to estimate a grey level threshold above which a change in the image associated with detection of sweat is measured above the background grey level. In this example, the grey level threshold is 30 on a scale 0–255 where 0 corresponds to white and 255 to black. On panel C, a typical image collected during cholinergic stimulation displays spotty dark regions in the ROI that correspond to the production of sweat. Panel D shows the number of pixels on this image (ordinate) for each grey level between 0 and 255 (abscissa). As more dark spots appear on the image collected during the sweat stimulation sequence, the image shows more and more pixels with elevated grey levels above the threshold. Thus, the right side of the curve gradually rises above 0 and the area under the curve (hashed) increases. The threshold level (30 in the example) is
subtracted from each image in the sequence to correct for the background grey level of the image. The sum of all pixel intensities above the threshold (hashed area under the curve) is computed as a measure of the darkness of the image in the ROI. Panel E shows a plot of the area under the curve in each image (hashed line) as a function of time. The steepest slope of the curve corresponds to the fastest change in image darkness and is used to represent the maximum rate of sweat production measured with the image sensor. Because different image collection sequences had ROIs of different sizes, the hashed area under the curve in panels D and E is scaled by the size of the ROI expressed in mm² (right vertical scale). Panel E also shows the evaporimeter readings (solid line, left vertical scale) collected on the contralateral site. The maximum rate of sweat evaporation measured by the evaporimeter is observed when the evaporimeter output is maximum and near flat.
Figure 4:
Sweat glands identified by imaging sensor. Mapping shows the same glands were stimulated after cholinergic stimulation (A) and after β-adrenergic stimulation (B).
**Figure 5:**
Individual ratios between maximum sweat rate in response to β-adrenergic and cholinergic stimulation, measured by the image sensor in heterozygotes for one CFTR mutation (Carrier), cystic fibrosis screen positive, inconclusive diagnosis (CFSPID), and CF subjects. The abscissa shows subjects’ genotype with corresponding CFTR functional classification below.

*Siblings: the brother was detected by newborn screening and the older sister was diagnosed during sibling screening. Please refer to Table 1 for phenotype details. These are the only pancreatic sufficient CF patients in this group.*
Figure 6:
β-adrenergic and cholinergic ratio in natural log from Evaporimeter and Image-Sensor measurements in all 4 groups non-CF healthy controls (HC), heterozygotes for one CFTR mutation (Carrier), cystic fibrosis screen positive, inconclusive diagnosis (CFSPID), and CF subjects. Letters denote significant comparisons.
Evaporimeter: a = HC vs CFSPID (p<0.0001), b = HC vs CF (p<0.0001), c = Carrier vs CFSPID (p=0.001), d = Carrier vs CF (p<0.0001), and e = CFSPID vs CF (p=0.026).
Image-Sensor: f = HC vs CFSPID (p<0.0001), g = HC vs CF (p<0.0001), h = Carrier vs CFSPID (p<0.0001), i = Carrier vs CF (p<0.0001), and j = CFSPID vs CF (p=0.022).
Table 1: Groups were defined as follows: non-CF healthy controls (HC), heterozygous for one CFTR mutation (Carriers), cystic fibrosis screen positive, inconclusive diagnosis (CFSPID), and cystic fibrosis (CF). Pancreatic insufficiency (PI) was defined based on fecal elastase results. SCT (sweat chloride test). FEV\(_1\) % predicted = most recent measure. Evap \(\beta/chol\) (ratio of maximal sweat rate with \(\beta\)-adrenergic stimulation to maximal rate with cholinergic stimulation measured by Evaporimetry). Sensor \(\beta/chol\) (maximal sweat rate ratio in response to \(\beta\)-adrenergic to cholinergic stimulation measured by Image Sensor).
Footnote:

^ not available or not assessed

* Evap and Sensor tests were repeated in this CF subject and results were reproducible.
Table 2: Difference between Evaporimetry and Image-Sensor; mean (SD).

|                      | Evaporimetry | Image Sensor | p-value |
|----------------------|--------------|--------------|---------|
| Baseline z-score     | 0.001 (1.00) | 0.03 (1.02)  | 0.7940  |
| Cholinergic z-score  | 0.00001 (1.00)| 0.00002 (1.00) | 0.9530 |
| Adrenergic z-score   | -0.0001 (1.00) | 0.00002 (1.00) | 0.8762 |
| β/chol ratio z-score | -0.003 (1.00)  | 0.01 (1.01)   | 0.7464  |