

Skin electrical properties and physical aspects of hydration of keratinized tissues

by

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List of abbreviations

SC - stratum corneum

RH - relative humidity

EDR - electrodermal response

EP - electrode polarization

DC - direct current

AC - alternating current

LF - low frequency

HF - high frequency

TEWL - trans-epidermal water-loss

CPE - constant phase element

DVS - dynamic vapor sorption

SSCD - skin surface conductance density

TS - test subject

IQR - interquartile range

EEA - effective electrode area

Preface

The first four chapters in this thesis are written with the purpose of introducing the reader to the field of bioimpedance, skin sorption properties as well as the thrilling memristor circuit element that all have been key topics in my research. The topics covered are naturally biased, as well as limited, by my own overview of these fields, but nevertheless the introduction is meant to serve as guide and motivation to the papers that are the final outcome of my research during the last years.

The anatomy and physiology of the human skin are treated in chapter 1. Chapter 2 introduces skin electrical properties, whereas its corresponding water holding capacities are briefly discussed in chapter 3. Memristor theory, suitable for biological systems, is introduced in chapter 4. Finally, in chapter 5 I summarize the main findings in my thesis, sharing also some thoughts about natural follow-up studies.

Electrical bioimpedance is a field of research that has evolved mainly in the last 30 years and this research has provided a detailed knowledge about the passive (i.e. no internal current source) electrical properties of human as well as other tissues. One possible application that has been quite thoroughly studied, is the ability of giving a description of the hydration state of the stratum corneum (SC), the uppermost part of the human skin, by means of electrical measurements. As the general hydration state of the SC is connected to its passive electrical properties as well as to the overall function of the SC, serving as a protective shield against the environments, the role of water in the skin has been studied with interest within dermatology as well as by researchers concerned with bioimpedance and bioelectrical phenomena. In general, information gathered from impedance measurements is a fingerprint of the tissue properties as well as of tissue alterations, and so as an example skin hydration can be linked to the physical parameters of skin barrier functions or alterations can be linked to specific skin related diseases.

Thus, being able to estimate the actual content of water within the SC can be expected to be a strong clinical tool, enabling better diagnostics, and a good precision of the estimates is thus desirable. As one of the foremost

functions of the SC is to provide a diffusion barrier towards the environments, its water hydration characteristics have been and will be important in a structure function understanding of the SC. This argument also applies for keratinized tissue, that is a more general grouping of tissue in the human integument such as hair, nail and SC.

Modeling the human SC as well as the rest of the skin is important in order to actually understand the underlying physics and hence possibly also enable improved diagnostics in some applications. It is desirable that the modeling, often based on a more or less intricate combination of elementary circuit elements is not only precise in its description (i.e. mimicking the observations to a good precision), but also conceptually correct, giving an improved understanding of the physical phenomena at their fundamental level. This is where the memristor comes in.

It is important to keep in mind that electrical measurements on or within the human body involve electrodes, i.e. the site where charge carriers shift from electrons to ions, and electrodes will in general influence the signals that are measured. Hence, electrodes should always be chosen carefully in order to be most suitable for the intended use.

Six papers are presented in this thesis and are referred to by Roman numbers I-VI throughout the text. These papers represent, at least in part, answers to some of the key questions that during the last four years have served as a motivation for my daily work. The two first papers describe methods for estimating the stratum corneum water content, both based on electrical measurements (Paper I) as well as on skin water evaporation measurements (Paper II) and are steps towards an objective measure of the hydration state of the stratum corneum. These are methods that, if sufficiently precise, would be useful in evaluating the protective properties of the stratum corneum as well as within understanding of skin diseases. Paper III and IV deal with the water sorption properties, intimately related to skin electrical properties, of keratinized tissues. In Paper V the memristor, which is the fourth and final passive circuit element, complementing resistors, capacitors and coils, is introduced to the world of bioimpedance, and human sweat duct conductivity in particular. The memristor shows promising abilities in explaining some puzzling non-linear phenomena in circuit theory, and hence also in bioimpedance measurements. Finally, Paper VI deals with the same sweat ducts, but now with focus on how different types of electrodes will influence the final signals that are recorded, and we will see that an electrode is not necessarily just an electrode.

List of papers

- Paper I:** GK. Johnsen, ØG. Martinsen, and S. Grimnes,
Estimation of in vivo water content of the stratum corneum from electrical measurements,
Open Biomed Eng J, **3**, 8-12, 2009.
- Paper II:** GK. Johnsen, AB. Haugsnes, ØG. Martinsen, and S. Grimnes,
A new approach for an estimation of the equilibrium stratum corneum water content,
Skin Res Technol, **16**, 142-145, 2010.
- Paper III:** GK. Johnsen, L. Norlen, ØG. Martinsen, and S. Grimnes,
Sorption properties of the human stratum corneum,
Skin Pharmacol Physiol (submitted), 2010.
- Paper IV:** GK. Johnsen, ØG. Martinsen, and S. Grimnes,
Sorption studies of human keratinized tissues,
J Phys Conf Ser, **224**, 012094, 2010.
- Paper V:** GK. Johnsen, CA. Lütken, ØG. Martinsen, and S. Grimnes,
A memristive model of electro-osmosis in skin,
J Phys D (submitted), 2010.
- Paper VI:** C. Tronstad, GK. Johnsen, S. Grimnes, and ØG. Martinsen,
A study on electrode gels for skin conductance measurements,
Physiol Meas, **31**, 1395-1410, 2010.

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Part I
Introduction

Chapter 1

Anatomy and functionality of the skin

The skin is the largest organ of the human body with a size of approximately 1.8 m^2 and a thickness ranging from $5\text{ }\mu\text{m}$ to as much as 1 mm or more at the sole of the foot. The skin is, provided being properly hydrated, an elastic barrier controlling the evaporational loss of body water as well as protecting vital organs against chemical and physiological substances of the environments.

Despite consisting of merely dead cells, the stratum corneum (SC), which is the uppermost layer, holds many remarkable and interesting properties, and is interesting for many reasons: Trans-dermal drug delivery has been studied and developed since the famous attempt by Munk [1]. The SC barrier function itself is of central importance within the pharmaceutical industry as well as in medicine, especially in the branches focusing on penetration and the understanding of skin diseases characterized by a deficient barrier function such as atopic dermatitis [2, 3]. The construction of a barrier between the body and environment is perhaps the foremost function of the skin.

The skin consists of three primary layers: Epidermis, dermis, and subcutis. The epidermis is the outermost layer and provides a chemical and diffusional barrier towards the environment. The dermis is located beneath the epidermis and is constructed from connective tissue, containing proteins, collagen, elastin and reticulum in a ground substance which is more or less structureless [4]. The dermis is much thicker than the epidermis and serves as a shield against any external mechanical influence. Beneath the dermis is the subcutis, containing mostly fat tissue serving as a heat isolating shield. The subcutis is a basement layer that connects the skin to underlying structures such as bone and muscles.

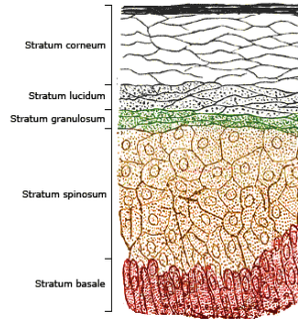


Figure 1.1: Different layers of the epidermis. Figure: Wikimedia commons.

1.1 The epidermis

The epidermis consists of three main cell types: Keratinocytes, constituting roughly 95% of the tissue volume; melanocytes responsible for melanin production, and Langerhans cells that are believed to be immunologically active [4, 5]. Also, Merkel cells are present, responsible for sensory reception [4]. The keratinocytes contain keratin, which is a protein, and are vital for the maintenance of the water diffusion barrier present in the stratum corneum which is the uppermost layer of the epidermis [4, 6]. Furthermore, the keratinized cells constitute a stiff and viscoelastic scaffold for the lipid content that, depending on its structure, determines the function of the diffusion barrier.

The epidermis can be divided into [4] the stratum corneum providing a diffusion barrier to the environments, the stratum granulosum that serves the SC with dead and keratinized cells, the stratum spinosum, and the basal cell layer stratum basale at the border, the basala lamina, towards the dermis, see figure 1.1. Those areas of the skin that experience substantial friction (i.e. plantar and palmar skin) have an extra epidermal layer, the stratum lucidum, between the SC and the stratum granulosum. The epidermis is avascular, meaning there is no blood supply through the capillary network. The viable part, the stratum Malpighii, is supplied with nutrition through a long diffusion process starting from the dermis [4, 6].

When the epidermis cells move outwards from the growth layer at the stratum basale they change both shape and size. The initial vertical cylinders transform into horizontal and hexagonal-shaped disks less than $0.5 \mu m$ thick and with a diameter of approximately $30 \mu m$ [5]. The cell content is also altered in this process usually denoted as the epidermis differentiation [4, 7].

The cells are keratinized, i.e. filled with keratin, which is a protein, and therefore called keratinocytes.

Due to the anaerobic metabolic activity above the basal lamina, nutrition needed for the cell processes is taken from the phospholipids which are the main constituent in cell membranes. Thus, the further up in the epidermis, the more decomposed cell membranes are expected. There is also a synthesis of new lipids, and those lipids that eventually are going to constitute the stratum corneum lipid phase are gathered in a special structure called the lamellar bodies or the Odland bodies [8, 9]. These bodies are membrane coated granules, and thus giving name to the stratum granulosum cell layer.

As the epidermal cells enter the stratum granulosum from beneath, their cell nuclei start decomposing and the cell membrane is replaced by a new and much more solid membrane, the corneocyte envelope [4] that is very important in the water diffusion barrier located in the stratum corneum [10, 11, 12, 13, 14]. This last stage of the epidermal cell differentiation is a sort of a programmed cell death where the final product is the stratum corneum cells and their lipid rich surroundings.

1.1.1 The stratum corneum and its barrier function

In this section I will shortly go through some of the main principles of the SC barrier function, as this is also related to the skin hydration state that is one of the main topics in this thesis. The stratum corneum is considered to be the barrier to diffusion, being 10 to 30 cell layers thick [4, 15, 16, 17, 18], and its water hindrance is depending on the content and structure of extracellular lipids. There are two main components: a hydrophobic part (extracellular lipids), and a hydrophilic part (the keratin of the corneocytes), where the lipid part is particularly vital in the regulation of water transport through the skin [19]. The SC cells, the corneocytes, are “bricks” that are surrounded by extracellular lipids, that constitute approximately 10 – 20% of the total cell volume [3, 19, 20]. Apart from the secretion of water from the sweat glands, an amount, depending on the environmental temperature, of about 380 – 700 ml in total is lost through the skin every day [21]. This is the so-called *perspiratio insensibilis*, that is an unnoticed perspiration of water from the skin. This provides the water needed as a plasticizer for the keratin in the corneocytes to remain their function and the total rate of water loss from the skin will be a measure of the barrier function. As the epidermal top layer is constantly abraded due to mechanical and chemical environmental conditions, a continuous renewal of the diffusion barrier is needed. In order for the SC to remain a constant thickness in time (not location), the adding of new SC material must be correlated with the pace of the shed of the surface

layer, which is often referred to as the stratum disjunctum.

The flat, protein rich hexagonal corneocyte cells are interconnected through desmosomes and filled with keratin fibrils. The fibrils span the cell interior horizontally, and this robust net of threads hinders any expansion in the horizontal plane when the cells take up water and swell, which then only is possible vertically. According to Norlen et al. [22] the SC cells can swell 20% vertically due to the elastic corneocyte envelope, but only a few percent horizontally. This ensures area preserving, preventing skin ruptures and is necessary for a functional barrier. The keratin in the corneocytes is characterized by a high sulfur content, and its high affinity for water [23].

The stratum corneum lipids provide the main barrier to diffusion of water soluble substances across the epidermis and variations in the barrier function is often due to alterations here [19, 24]. The lipid synthesis deeper in the epidermis provides the SC with the proper lipids for constituting this barrier. Those lipids that are meant to constitute the SC lipid phase are gathered in the Odland bodies in the stratum granulosum, and the lipids form a continuous compartment in contradiction to the more open spaces of the Malpighian layers of the epidermis [16, 25]. These membrane coated granules are smooth structures surrounded by a double membrane not very different from the cell membrane [6]. The lipids therein are in a crystalline arrangement, and on the SC side of the stratum granulosum these bodies extrude their lipid content into the extracellular space of the SC, that is, the lipids surround the corneocytes, making the SC a dense layer.

1.1.2 The Barrier properties

The lipids capable of forming biological membranes typically have one hydrophilic and one hydrophobic part. The lipids will seek the most stable state by placing the hydrophobic parts, that consist of long hydrocarbones, towards one another so that the hydrophilic parts face the extracellular water in the SC. Hence, there are several water tight layers arranged on top of each other [13], containing both non-polar and polar domains.

How well the stratum corneum is hindering diffusion of water depends on

- The length of the hydrophobic chains.
- The amount of saturation of the lipids.
- Temperature.

The chains are held tightly together by attractive van der Waals forces, and the longer the chains, the less amount of water is able to penetrate. As the

attractive van der Waals forces are only effective on short, atomic diameter scales, the amount of unsaturation (double bonds) of the lipids will influence the barrier density [19]. Chains with a large amount of double bonds will not be able to be packed densely enough since the double bonds will induce “bends” on the chains. This means that the van der Waals forces can no longer contribute to tighten the barrier, and thus increasing saturation of the lipids will induce a more water tight envelope of the body.

The temperature is very vital in this sense. Lipids in the stratum corneum can exist either in a crystalline state or a liquid crystalline state [19]. If temperature is sufficiently low, i.e. below a certain transition limit, the lipids turn into a crystalline form [19, 26] which is non-permeable to water contrary to the more transparent liquid form. The transition temperature depends on the composition of the lipids as well as their chain lengths. The longer the lipid chains, the higher the transition temperature. Also, if the chains are saturated, the transition temperature increase. For human cells, the transition temperature for lipids is in the range from about 0 – 40°C, which corresponds to chain lengths of roughly 18 carbon atoms [19]. However, as these lengths are found to increase substantially from the basal cell layer to the SC, the transition temperature can be expected to be higher than normal skin surface temperature [19]. In the SC the lipids are of a length of 30 C-atoms or more [27], and thus SC is in the water tight phase at normal skin and environmental temperatures.

1.2 Nail and hair

Both nail and hair, along with SC, are parts of the family called keratinized tissues [28]. Nail tissue is composed of two main constituents which are the dorsal (upper) and ventral (lower) nail plate [29, 30]. The dorsal plate consists of flat, keratinized and dead cells that are oriented in such a way that the shortest typical diameter is perpendicular to the nail surface. Studies of nail by electron microscopy have shown that there are very tight-junction-like contacts between the cells that essentially are glued together [6].

The ventral part lies beneath the much more stiff dorsal top, and is a less tightly bound structure allowing flexibility of the nail. The orientation of the fibrils are more ordered than in the dorsal top and is perpendicular to the growth direction [29], preventing the nail plate from being split all the way down to its tunnel from where it originates. Mechanical strength of the nail plate, which is a very convenient property for nails, is ensured by the two main curvatures of the plate, one fairly large in the direction of growth, the other, much shorter, horizontally but perpendicular to the direction of

growth. A curved sheet of paper (i.e. cylinder shaped) has a much greater mechanical stability than a correspondingly flat, and pays as an illustration of how nail composition ensures a stiffness suitable for the intended use.

Hair fibers consist of cortex cells that are protected by about 10 cuticle cell layers, concentrically oriented around them. The building blocks are keratinized cells that are cemented tightly together and filled with protein filaments [3]. The structure is relatively dry and compact, and the hair fibers are the final result of a differentiation process, just as in nail and SC [4, 6]. The cells in the hair follicle are pushed outwards, through a tight tunnel, and eventually run through a programmed cell death. The cells of hair fibers are about 120 μm long and elongated in shape in the direction of growth. The elongated shape is due to the squeezing of the cells as they are pushed out of a very rigid and tight hair follicle-tunnel. The cortical cells are held tightly together in a substance that is present between the cortex membranes. Within each of the cortical cells, there is a micro filament matrix structure composed by filaments. These filaments are constructed from three α - helices that are coiled into a larger coil, similar to the composition of an old fashion rope. This provides a robust structure, especially in the longitudinal direction [3].

1.3 Sweat ducts

The human skin has two types of sweat ducts. The by far common most is the eccrine. These ducts are distributed mainly all over the skin, and their main function is to spread water onto the skin surface, contributing to the temperature regulation of the human body as the water evaporates.

The highest concentration of these ducts are found in palmar and plantar skin sites where hair is absent [4], and one way to think of this is that this increased density is to ensure a proper and necessary hydration of the hands and the sole of foot to give a decent grip needed for running and climbing and so on. The eccrine ducts are long, thin tubes, originating in dermis or even in subcutis. In the SC the ducts are spiral shaped, a shape that ensures the ducts being squeezed to close when SC swells, preventing too much water entering within the integument barrier or that essential nutrients being washed out of the skin. Contrary to sweat activity at other skin sites, the palms and soles are thought of to produce sweat only due to sensory or mental stimulation, and not due to thermal activity [31]. This kind of sweat activity is often called exosomatic electrodermal response (EDR) [32] and will be treated in Paper VI considering electrode gels used for sweat measurements based on skin conductance recordings.

The other type of sweat ducts are the apocrine that segregate fluids from the upper part of the hair follicle where the duct opening is located. Bacterial decomposition of this secreted material results in a more or less personal odor. The apocrine ducts are mainly found around genitals and in the axilla [4].

Chapter 2

Electrical properties of human tissue

In this chapter I will try to give a short overview of passive electrical properties of tissue, that essentially is the key focus within the field of bioimpedance. In particular, the electrical properties of keratinized tissue will be paid attention since I in this thesis am not much interested in things being alive, but actually only care about the thin and dead envelope of the human body.

Electrical admittivity of human tissue may be predominantly conductive or capacitive, or a combination of these, depending on tissue type, availability of charge carriers, frequency of the applied electric field, and is carried out by ions, dipoles or electrons, as well as holes (semiconductor) [33]. Electrolytes, both intra and extracellular, are responsible for a conductivity of the order of 1 S/m, and are considered to be more or less frequency independent up to roughly 10 MHz [34]. The dominant factor regarding charge carriers in biological materials are ions, but also purely electronic currents have been reported to take place, for instance in DNA molecules [33]. In keratinized tissues, as will be the main topic of interest in this study, admittivity is very much dependent on the amount of water therein and to what extent it is bound to the tissue structure [35, 36, 37]. As water is highly polar it can be bound to other polar groups, for instance in proteins, with a strength that varies from very bound (essentially a solid phase) to more or less free water, and water can be regarded as the most relevant parameter concerning the electrical properties of such protein-rich tissue.

The high relative permittivity of water is perhaps the main reason of its dissociative properties due to a weakening of the electrostatic forces between polar biological molecules. Since binding of water in biomaterials varies from very firm, with water practically immobile to more or less free water, it contributes to both ionic and dielectric admittivity. Protonic conductivity is

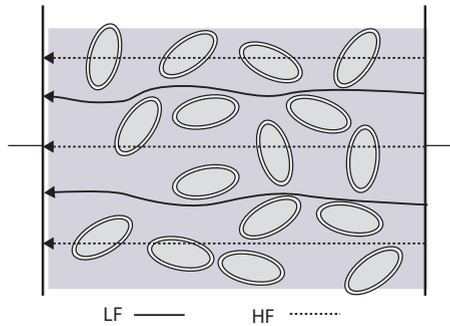


Figure 2.1: Principle of high and low frequency current pathways in tissue. Figure redrawn from [32].

purely DC and is carried out by the H^+ and H_3O^+ cations where the protons are hopping from neighboring water molecules by a tunneling effect that is much faster than ordinary ion mobility in an electric field, since the latter is slowed by viscous forces.

2.1 Cell membranes and tissue

The polar lipids (i.e. with a net dipole moment) are the most abundant constituents of the capacitive cell membrane, which is a necessary condition for living cells by keeping the interior constituents in a controlled environment. Cell membranes are thin films, 7 nm thick, consisting of phospholipids, and have a very low conductivity, but on the other hand a high capacitance, that is assumed to be frequency independent [32]. However, admittance across a cell or a suspension of cells is highly dependent on frequency, and figure 2.1 reflects the capacitive versus the resistive nature of cells and tissue. At low frequencies, current must pass extracellularly as the cells will have a shadow effect because the poorly conducting cell membrane forces the current to go around the cells. This may induce very narrow current paths where the current density can be much higher than the average in the volume. As frequency is increased, current passes also through the cells due to the capacitive shunting of the cell membranes.

Tissue homogeneity, or more likely the lack of it, is important regarding current paths in human tissues, and from an electrical point of view, tissue is as good as always inhomogeneous. Such an anisotropy is present all the way from lipids and cell membranes at a micro level up to organs and long muscle

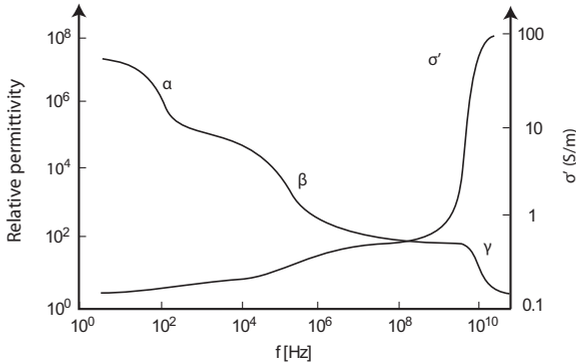


Figure 2.2: Dispersions in human tissue. Figure taken from [32].

fibres, and would highly influence the recorded signals at low frequencies, but the effect of it is more or less absent as the frequency is very high.

Polarization mechanisms

In human tissue, there are a high number of frequency-dependent admittance mechanisms, and these were sorted out and classified in dispersions by Schwan in the 1950's [38] and are illustrated in figure 2.2. The dispersions are drops in relative permittivity over well-defined and sometimes very narrow frequency ranges. Permittivity of biological materials typically decrease as the frequency is increased as charges (i.e. dipoles) are too slow to follow the changes in the applied electric field. The most distinct dispersions are those of cell suspensions and other simple biological systems. In tissue and skin, however, complexity is much higher, yielding a high variety of polarization mechanisms, and dispersions are much broader in the frequency range and overlap, sometimes giving a continuous fall with increasing frequency.

Literature reports on at least four different dispersions or dominating polarization mechanisms in human skin, and some of the mechanisms behind are due to [32]: the counter-ion effect at the cell surface (α), Maxwell-Wagner polarization at interfaces between different dielectrics and capacitive charging of cell membranes (β, δ), large and small polar molecules (α, β, δ), and due to the dipolar nature of the water molecule (γ).

In a Wessel plot, the imaginary part of the electrical recordings are plotted as a function of the real part, and in 1940 K.S. Cole proposed an empirical

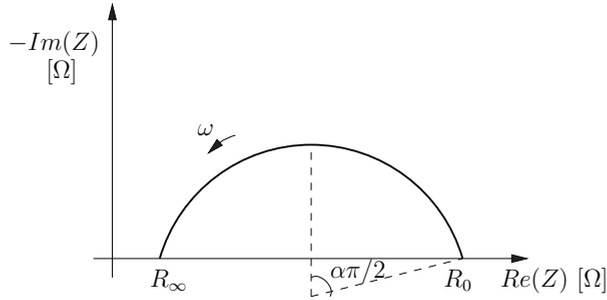


Figure 2.3: Wessel plot of the Cole equation.

equation to describe tissue impedance

$$Z = R_\infty + \frac{\Delta R}{1 + (j\omega\tau)^\alpha}, \quad \Delta R = R_0 - R_\infty, \quad (2.1)$$

where R_∞ is the impedance in the high frequency limit, often called the series impedance, R_0 the corresponding impedance at DC currents, ω is the signal frequency and τ is the characteristic time constant of the system of interest. The entry of this simple equation was strongly motivated by the fact that many biological systems produce circular arcs in impedance space, however with a locus located beneath the real axis. A typical illustration of such a depressed arc is shown in figure 2.3.

If $\alpha = 1$, the Cole equation gives a full semicircle in the complex Wessel plane with its center on the real axis, and there is now also a unique correspondence to physical elements in equivalent models of the biological material, in this particular case consisting of two (ideal) resistors and one (frequency independent) capacitor that often may be given physical interpretations such as current pathways through extra -and intracellular tissue and capacitive current pathways of lipid cell membranes.

However, in more complex tissue, $\alpha < 1$, and there is no longer a direct relation to pure physical elements, but the Cole equation now includes what is often called a constant phase element (CPE) that is not actually a physical component and can not be realized in terms of a finite number of traditional passive circuit elements (RLC), but is related to the complex theory of dielectric dispersions and often thought of as a distribution of time constants.

2.2 Keratinized tissues

Keratinized tissue is composed of membrane-coated cells that are cemented tightly together, and the cells are normally arranged in layers. Electrical immittance across such a layer is dependent on how the cells are connected, the amount of intercellular space, the hydration state as well as any shunting mechanism through channels such as sweat ducts perforating the entire SC.

The most important parameter governing the admittivity of keratinized tissue is the amount of water therein, and admittivity typically shows an exponential dependence with relative uptake of water [35, 36, 37]. A possible explanation to the behavior of conductance upon the amount of water is suggested in the early work of Algie [39], and is connected to “proton hopping” between water molecules. As water content increases, water is eventually more loosely bound, and the binding energy to hydrophilic groups is reduced. Along with shorter average distance between the water molecules, this yields a rapid and exponential increase in conductance.

The SC, hair and nail are relatively dry compared to other bodily tissues. For the SC this is particularly true in the uppermost parts, and so it will dominate impedance with an increased dominance as frequency is lowered, since stratum corneum in general is a poorly conducting layer on top of a well conducting viable skin. It has been shown, performing finite element analysis, that at frequencies of the order of 10 Hz, the SC would totally dominate the measured tissue information, but at frequencies above 100 kHz, less than 10% of the received information stems from the top epidermal layer, and as a rule of thumb skin impedance is dominated by SC at frequencies below 10 kHz [40, 41]. Hence, an increase of frequency then generally leads to measurements on deeper tissue layers. However, this will very much depend on the hydration state of the skin, and electrode configuration such as geometry and size. Skin immittance also varies between different sites on the skin as well as between individuals [32]. In particular, sweat activity and relative humidity in the ambient air will play a key role due to increased duct filling and hydration of the SC. Increased RH will also result in increase of water bound in nail and hair, and in general the electric properties of nail and hair resemble those of SC, although there are reported on some differences in nail concerning low frequency susceptance values [37]. Different sorption properties, as seen in Paper IV, may also influence on how these tissues respond to an applied electric field.

Data on skin impedance have been retrieved by Yamamoto and Yamamoto [42] by means of tape stripping, and relevant parameters are shown in figure 2.4. These data again illustrate the following important point: As the frequency increases the contributions of the SC on the recorded signals

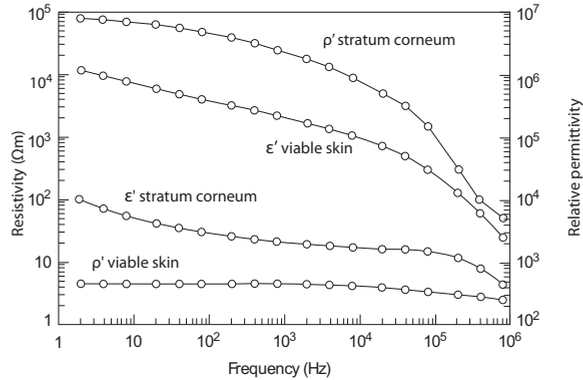


Figure 2.4: Resistivity and relative permittivity of human SC and skin. Figure taken from [32].

are no longer dominant as the impedivities of SC and viable skin converge.

Keratinized tissue, such as the SC, may be modeled by an electrical equivalent as shown in figure 2.5 when doing impedance measurements. The sensed signals from measurements can thus contain contributions from electrode polarization (EP), SC, sweat ducts and viable tissue and include dispersions from each Cole element. The SC has been shown to have a broad dispersion in the α -domain [42, 43], the EP has one or two as such [44], whereas viable and deeper tissue, being more complex, will have multiple dispersions, sometimes overlapping completely in the frequency domain.

Such a model may be simplified based on detailed knowledge of each of the constituents in the circuit. For instance, as the electrical properties of the sweat ducts are dominated by conductance, and since the ducts show negligible signs of polarization admittance [45], the Cole element of the ducts will practically be reduced to a general resistor.

2.3 Electro-osmosis

As impedivity of the SC is high, although depending on its hydration state and applied signal frequency, the sweat ducts form ionic shunts that have large influence on the admittance of the skin layer. SC admittance is roughly doubled within a few seconds when the sweat ducts are filled [46]. Since sweat is an electrolyte, the filling of the ducts result mainly in a conductive contribution to electrical admittance. The amount of current passing through

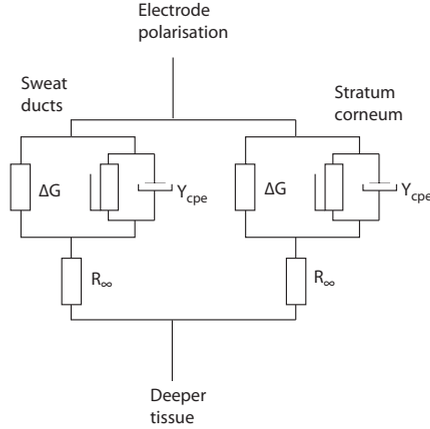


Figure 2.5: Equivalent electrical modeling of human skin.

the human skin is in some cases found to be markedly different in absolute value depending on whether an anodic or cathodic signal is being applied. This difference has been explained as an effect of electro-osmosis, first by Munk [1] and then later refined by Grimnes [47]. Water is dragged along in the sweat duct capillaries by viscous forces as the mobile parts of the charged thin double layer in the ducts are influenced by an external electric field, leading to increased or decreased current flow depending on the polarity of the applied signal. A negative surface potential will normally attract water [47], resulting in increased conductivity, whereas a positive potential will repel the capillary water, and yields a decrease in conductivity. A schematic model of the duct capillaries and their corresponding charged wall films and baseline water content is shown in figure 2.6. The flux F of sweat duct liquid through a capillary is generally proportional to current and resistivity of the fluid, and so inversely related to the concentration of the electrolyte. It can be expressed as [48]

$$F = \frac{\zeta \epsilon I \rho}{4\pi \eta}, \quad (2.2)$$

where ζ is the electro-kinetic zeta potential, ϵ , ρ , and η are permittivity, resistivity and viscosity of the liquid, respectively, and I is the current. Such an electro-osmotic transport of water comes in addition to the ongoing diffusional transport of water across the skin, that in general also will lead to a build-up of fluid beneath any measuring electrode due to occlusive effects.

Sweat ducts of human skin are electrical shunts for DC current and are

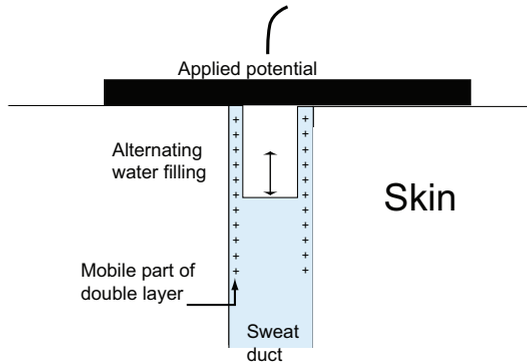


Figure 2.6: Schematic model of sweat ducts in human skin.

not responsible for the capacitance of the SC, due to the lack of counterion relaxation phenomena [45]. In the light of the discussion in Paper V, the sweat ducts can in some cases be memristive rather than conductive (resistive), due to the memory effect of the resistance upon the amount of charge having passed the duct. The modeling and investigation of sweat duct conductivity has been one of the main subjects in this thesis.

2.4 Influence from electrodes

Electrodes used for electrical immittance measurements will practically always influence the signals that are measured, and so a proper choice of type is very vital [32, 49, 50]. By onset of electrodes on the skin surface, water will immediately start building up underneath it due to the water concentration profile present in the SC. This process will generally lead to increased admittance with time, and an important question in this manner is when a stable state is reached, or more likely: To what degree does one eventually have stable conditions?

Not only the amount of water building up beneath the electrode, but also the characteristic time for this process will in general be important for the result. The rate of build-up will typically be largest just after onset, and then decreases with time, reflecting the nature of many physical processes where the rate of change is proportional to the concentration difference between the present and equilibrium value of the actual physical quantity. Thus, one does not measure on the skin “as it was“ prior to onset of the electrode for monitoring. Still, it is the pre-occlusional value that is most representative

for the actual hydration state of the skin, and this is a challenge in most set-ups for skin hydration recordings.

Also, the initial hydration state of the skin will influence the immittance, and thus environmental conditions such as RH and temperature will govern the content of water prior to any electrode onset. This in particular means that it is not possible to construct an electrode that just stabilizes the hydration level beneath the electrode.

Electrode gels are constructed with the aim of establishing an electrical contact between the electronic conduction in any metal wire and the relatively dry SC layer. Depending on the relation between the water vapor pressure of SC and the gel, water will flow either to or from the SC just after electrode onset. Furthermore, electrolytic concentration of the electrode gel used as the contact medium plays a significant role. Depending on this concentration the skin will be wetted or dried out due to osmotic effects. A high electrolytic concentration in the gel will likely induce an increased water transport from deeper skin, containing much more water than the dry top layer, towards the surface and so results in increased skin admittance. For a dry metal plate electrode, admittance increases instantly after onset due to a water contact film build-up between the metal and the SC surface. Wet and dry gels can result in both increased or decreased admittance, depending on which way there is a net transport of water. However, wet gels will usually give a markedly jump in admittance due to their rapid wetting of the SC. Using wet hydrogels with low viscosity opens for the possibility that the gel penetrates the sweat ducts that normally are empty or partly filled [47]. Since such gels may have much higher conductivity than the sweat electrolyte and also open for an increased duct filling, the result is an increased admittivity of the skin. And as the ducts are believed to have negligible polarization, the factor influenced is the conductance. Especially in skin conductance measurements, where among others sweat activity can be monitored using a low frequency set-up [51], such gel behavior would for instance yield a decrease in skin conductance during a sweat response, due to the replacement of high conductive gel with the lower, but still highly conductive sweat as seen in Paper VI. Yet, as gels usually have a high conductivity at low frequencies compared to the SC, individual differences in gel conductivity will therefore not be expected to influence low frequency conductance unless the gels are able to penetrate down in the ducts, giving an effective change in filling.

Occlusion over time may not necessarily only influence the SC hydration state (i.e. a low-frequency phenomenon), but may also induce changes in immittance in the high-frequency area, normally ascribed to effects found in deeper tissue. Irritation and altered water content are possible explanations, but still only at a speculative level, as discussed briefly in Paper VI.

Electrode size and shape, as well as the distance between them will also affect the final outcome, and I refer to Grimnes *et al.* (p. 190-191) for a more rigorous treatment of this subject [32].

In Paper VI the role of different electrode choice on the final result for skin conductance measurements and sweat activity has been studied, comparing low frequency skin conductance recordings with the water sorption properties of the gels of the electrodes.

Chapter 3

Skin hydration properties

As the hydration state of the epidermal stratum corneum is vital for a proper function as well as visual aspects of the skin, a large variety of techniques and approaches have been carried out with the purpose of giving better estimates of skin water content. The methods include, among others, microwave, spectroscopic, gravimetry or by means of nuclear magnetic resonance [32]. Hence, a direct, simple, low-cost, non-invasive and reliable method for estimating human SC hydration will be of great interest and has for over 30 years been developed within the field of bioimpedance [52, 53, 54, 55, 56].

Although it has been challenging to reach consensus on how to get the best estimate of skin hydration, a low and single frequency susceptance method will be suitable if the aim is to measure superficial hydration such as in the human SC [40, 56]. Since the sweat ducts are predominantly conductive, the proper electric parameter to be used for SC hydration assessments is low frequency susceptance or AC conductance [32]. However, if one wants to find absolute water content (as gram water per gram dry skin per unit area) a calibration of such a susceptance based hydration method against the weight of SC water content must be carried out, as is attempted gravimetrically *in vitro* in Martinsen *et al.* [36]. *In vivo* the water concentration profile across the SC, that is present due to the ongoing transport of water, complicates things further. The possibility of estimating absolute water content *in vivo* from low frequency electrical measurements is a challenging task, at least to a good precision, and Paper I represents a step in that direction. A different approach is tested out in Paper II where measurements of *perspiratio insensibilis*, often called trans-epidermal water-loss (TEWL), are combined with water desorption properties of SC *in vitro* to yield an estimate of the average and stable state *in vivo* SC water content.

3.1 Sorption of water in keratinized tissue

Keratins have, as we have seen, an important function forming the interface between the bodily interior and the external environment. However, their overall functions may be influenced by ambient factors that may alter their nature. In particular water is important in this manner due to its high dipole nature and the ability of forming bonds with side-chain-endings in keratin, and it is known that SC needs a content of roughly 10% water in order to remain its primary functions [57]. Water has a very strong influence on the physical properties of hair, nail and SC, and for instance the elasticity modulus changes with a factor of about 100 from dry to humid conditions [29, 58]. The electrical dependence upon water content is already described, and further details can be found in [32].

3.1.1 The water sorption isotherm

A common way to present sorption data of water into keratinized tissue is by means of sorption isotherms, which give the amount of water that can be contained at a given relative humidity when the temperature is fixed.

A typical example is shown in figure 3.1 for keratinized tissue, others are found in [29, 57, 59, 60], where the SC curve shows a sigmoid shape that is more profound compared to those of hair and nail. Such isotherms are typical for tissue rich on proteins [3]. As the relative humidity is at its high end, SC takes up considerably more water than the other two types of tissue, an effect that probably is ascribed to the restrictions upon swelling for hair and nail as they have stronger internal constructions and filament networks [6, 29, 62]. The isotherms provide knowledge about the interactions that take place between water and different chemical groups in the tissue under investigation, and Paper III is concerned with some of these questions.

The increase or decrease in water uptake will in general be diffusion controlled as long as the temperature is above a certain limit, T_g , that defines the transition between more complex diffusion-relaxation sorption processes and processes governed by Fick's laws of diffusion. For keratinized tissues, T_g is below 20° C [63, 64], and so the amount of water sorbed, $M(t)$, as a function of time, is a solution of Fick's 2. law of diffusion [64]

$$\frac{M(t)}{M_\infty} = 1 - \frac{8}{\pi^2} \sum_{m=0}^{\infty} \frac{1}{(2m+1)^2} \exp \left[- \frac{D(2m+1)^2 \pi^2 t}{d^2} \right], \quad (3.1)$$

where M_∞ is the mass when the sample is in a stable state with its environments, D is the diffusion coefficient, assumed to be concentration independent (which is not always a good approximation as relative humidity is much

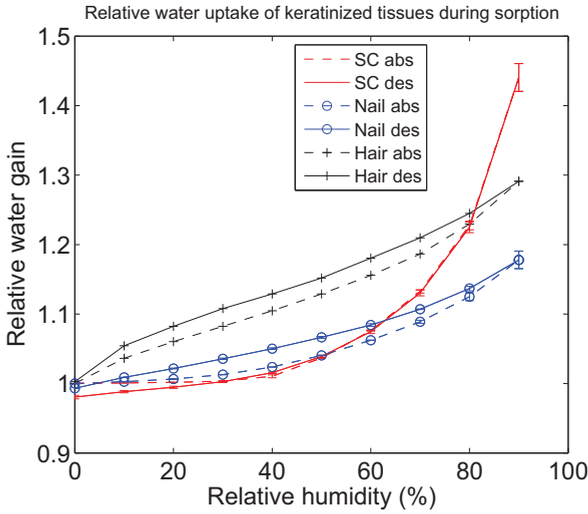


Figure 3.1: Typical water sorption isotherms for keratinized tissue. Figure taken from Johnsen *et al.* [61].

altered, cf. Paper III), and d the thickness of the sample. In contrast to the isotherms described earlier, the time course given in equation (3.1) will provide information about the rapidity of water sorption into the specimen of interest, such as SC, hair and nail. This will for example be relevant when ambient RH changes, inducing an altered hydration level of the SC, or when electrodes are attached, with the inevitable effect of hydration or depletion of the underlying SC.

It is worth noticing that isotherms in general are *in vitro* results. *In vivo* there is a different situation as there is a water concentration profile across the SC thickness that is responsible for the never ending flow of water through the human skin. Also segregation of sweat will influence *in vivo* sorption of the SC.

When water is absorbed in SC, it is typically bound as three different phases: strongly bound, bound, and bulk water, although their respective amounts vary in the literature [3, 59, 60, 65]. The characteristic time for SC to reach its equilibrium state will be different depending on which of the three types of water that is being bound. This is further discussed in Paper III, where the general sorption characteristics of SC were investigated.

3.1.2 Our findings and the link to bioimpedance

In our sorption measurements, using a DVS Intrinsic setup (SMS Ltd, London, UK) we found no hysteresis of significance for the SC samples from heel and breast. This means that the sample contained the same amount of water, having reached its equilibrium state during absorption and desorption as long as the relative humidity of the ambient air was fixed. This result is in contradiction to a previous study where the same setup was used, however with a lower resolution as discussed in Paper III. If present, a hysteresis could have been an indication of structural changes of the tissue during a sorption cycle, i.e. giving altered water holding capacities. However, in Paper IV, where the sorption properties of nail, hair and SC, all belonging to the keratinized tissues, were investigated, a hysteresis was visible for hair and nail. Seemingly, keratinized tissues responded differently to changes in relative humidity and showed different abilities to bind water, a property that most likely can be expected to influence the output in i.e. immittance measurements.

The observed differences may, at least in part, be ascribed to differences in structure as hair and nail cells are formed by restrictions imposed by the shape of their anatomical site, whereas SC cells do not have such restrictions, and hence are able to swell, at least vertically [3, 22, 29].

The triphasic behavior of the absorption constant of SC is perhaps the most interesting finding in our study in Paper III. In short, this means that the time needed for SC to reach equilibrium with the environments in the DVS measurement chamber was highly dependent on the relative humidity of the surrounding air. We propose as a potential explanation to this phenomenon the possibility of the SC keratin filament network within the corneocytes to exist in various “phases” [66]. The observed increase in absorption time constant τ with water content is somewhat contra intuitive as the diffusion constant satisfies $D \propto \tau^{-1}$ and will normally increase in a medium as its water content rises. We believe that this can be explained by the non-hydrophilic behavior of the keratin filaments in the SC as is discussed in further detail in Paper III. This pattern in absorption time constants is present both for heel and breast samples, although with somewhat different curve forms. It should be commented, however, that since these two sample types had been stored very differently prior to the measurements, a comparison is of reduced value.

As water is highly influencing tissue admittivity, and water bound in different ways and configurations (water strongly bound to polar groups give dielectric properties, bulk water contribute to ionic conductivity) this is also one of the key questions within bioimpedance, as more detailed knowledge may contribute to more detailed explanatory models without losing the high

precision that usually is present for more overall descriptive models based on equivalent circuits.

3.2 Trans-epidermal water-loss

Water can be lost from the skin surface both by sensible and insensible perspiration, where sweating is the main contributor to the former, whereas passive diffusion processes are dominant in the latter. Trans-epidermal water-loss (TEWL) includes both as it is the quantity of water passing through the epidermis, starting from the highly hydrated viable tissue of the human body. TEWL is a useful measure of the insensible water loss, and hence the SC barrier function as long as sweating is not present. One then measures purely diffusional water loss. A reduced SC barrier function will generally lead to increase in TEWL-values as water then diffuses more rapidly from deeper skin towards the skin surface from which it evaporates. However, TEWL is much sensitive to the relative humidity in the surrounding air as well as a series of other factors such as temperature, anatomical sites, skin diseases and so on [3].

3.3 Sweat measurements

Sweat activity on palmar and plantar skin sites are very sensitive to psychological stimuli [32, 51], and is in general not related to thermal sweating due to i.e. physical activity, but rather psychological factors such as consciousness or stress-related situations [31]. Such an activity is called electrodermal activity [32] and has been studied since the late 19th. century. Later, the origin of psychogalvanic phenomenon was suggested to be in relation with the sweat ducts [67]. In the recent decades, electrodermal measurements have been studied also in connection with the field of psychophysiology [68].

By means of TEWL-measurements, the amount (given in liter pr min pr area) of sweat evaporating from the skin can be estimated, and so yield a quantitative estimate of the sweat production. This amount is not the same as the sweat filling of the capillaries that is more related to electrodermal measurements. Skin conductance levels are hence more related to sweat gland activity and capillary filling than sweat evaporation from the skin. Conductance is in general a better measure of the sweat activity than resistance since conductance is related to susceptance and a parallel setup, whereas resistance correlates to reactance and a corresponding series circuit. In the human skin the ionic conductivity in the sweat ducts is in parallel with

the remaining parts of the SC where the capacitive parts are found [45, 56].

Tronstad *et al.* has reported on a portable instrument capable of quantifying sweat activity by means of low frequency monopolar conductance measurements [51]. Capacitive contributions will not be of interest since they in general are related to the SC hydration and not duct filling [45]. Clinically, these types of sweat recordings can be used as a diagnostic tool as well as in treatments of diseases such as hyperhidrosis, where the sweat level of the patient is elevated. The sweat pattern of hyperhidrosis is partly unknown [51, 69], and so objective methods for studying these patterns are important. A part of such a set-up involves also the understanding of the influence from different electrodes on the final measured electrical signals. This was the main motivation for the studies performed in Paper VI.

3.4 Ethical aspects of our study

In studies where measurements are performed on human test subjects as well as with animals, it is, depending on how much the test subjects can be expected to be exposed to hazards as well as the sensitive information needed, normal practice to apply the local ethics committee for approval prior to the initiation of the study. In the measurements that have been performed in our study, including both the preparation of SC samples with the dermatome “cheese knife” as well as in the conductance measurements with the Sudologger [51] and the Solartron 1260+1294 equipment, we have come to the conclusion that an application to the regional ethics committee was not required prior to the onset of these studies. This can be argued for based on the following knowledge on which aspects our studied contained as well as the actual equipment that was used.

- The dermatome (Braun, Tuttlingen, Germany), that was used to prepare the SC samples used in Paper II, Paper III, and Paper IV, is CE marked, meaning that it fulfills the EU regulations with respect to security, health and environmental control.
- In Paper VI, where electrode properties were monitored, both instruments used (the Sudologger and Solartron setup) are CE marked, enabling secure measurements on the group of volunteer test persons. Furthermore the test persons were well informed prior to the initiation of the measurements, and were allowed to abort the study at any time without any reason for this.
- All the methods that have been carried out in this thesis have been non-invasive. The dermatome used to harvest SC from the test subjects

had a high precision, ensuring a non-invasive procedure where only the outer layers of the heel SC were removed.

- No measurements were performed on test subjects that can be regarded as particularly vulnerable. The test subjects involved in our studies were mainly gathered from our scientific milieu.
- No sensitive personal information was gathered or stored during our different studies.

Chapter 4

Memristance of human skin

In this chapter I will introduce memristors as a potential new building block in circuit modeling of electrical properties that are found in biological systems. The emphasis will be on possible applications within the field of bioimpedance. The memristor is probably equally fundamental as it is uncommon in electric circuit theory. Predicted to exist from purely theoretical symmetry arguments [70], the memristor, as a passive physical component remained unrealized until quite recently when scientists at the Hewlett-Packard lab constructed it as a nano-device that was in accordance with conventional semi-conductor theory [71].

Based on the four fundamental circuit variables; current i , voltage v , charge q , and flux ϕ , six distinct relations can be found. Two of them are the definition of electric current $q(t) = \int_{-\infty}^t i(t')dt'$ and Faraday's law of induction $\phi = \int_{-\infty}^t v(t')dt'$, and in addition the definitions of the four axiomatic circuit elements, which are the resistor $R = \frac{dv}{di}$, the capacitor $C = \frac{dq}{dv}$, the inductor $L = \frac{d\phi}{di}$, and finally the memristor $M = \frac{d\phi}{dq}$. The complete set of combinations is illustrated in figure 4.1. As we see from the diagram, memristors are fully different from the other and more traditional circuit elements. This means that no combination of passive RLC circuit components can replace the memristor, no matter how non-linear these resistors, capacitors and inductors might be. An extra circuit component may not sound that spectacular, but consider you have been living for 40 years with shoes, socks and trousers, but no sweater available, and then suddenly get one. Initially, it may be hard to use it properly (mixing it up with the other three) and to see its final potential, but who knows how nice it eventually may be if the sweater also turns out to be of a convenient size.

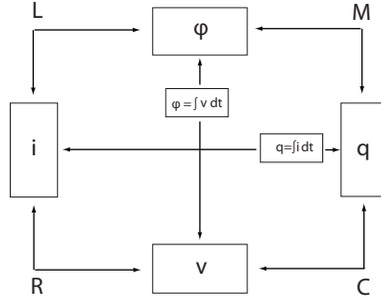


Figure 4.1: Symmetry diagram that shows the need of introducing memristors.

4.1 Properties of memristors

Although defined as such, memristors do not necessarily involve explicit magnetic fields. This is easily seen by realizing that memristance, in terms of Faraday’s law of induction, can be rewritten as

$$M(q) = \frac{d\phi}{dq} = \frac{v(t)}{i(t)}, \quad [\Omega], \quad (4.1)$$

and so looks very much like ordinary resistance, which it also is under certain circumstances. When Strukov *et al.* discovered the memristor (often called Maxwell’s hidden solution), or more precisely they found a memristive system, they took advantage of the fact that memristors require a nonlinear relationship between the integrals of current and voltage, and not magnetic fields explicitly. Earlier searches for memristors probably involved magnetic systems, which was a natural starting point considering how memristors are defined. This search in “wrong places” is likely to be one of the main reasons why memristors, although equally fundamental as the RLCs, were undiscovered for an additional century after Maxwell came with his famous equations.

Memristors can be generalized into memristive systems, and so was done in 1976 by Chua [72]. These systems can be defined as

$$v = R(x)i \quad (4.2)$$

$$\frac{dx}{dt} = f(x, i), \quad (4.3)$$

which actually is a generalization of Ohm’s law where v is the voltage, the resistance, R depends on the state variable x whose time derivative is in turn

a function of itself and the current i . The system above equals Strukov's [71], but can also be further generalized with additional state variables (not just x).

Letting charge be proportional to the state variable, x , resistance will be a function of the amount of charge having passed the element and in general not a constant, which furthermore results in a non-linear iv -characteristic parametric curve. In other words, a non-linear version of Ohm's law. The resistor is therefore now said to have some memory since its resistance depends on past events, and from that the name memristor arose quite naturally. As memristors are rather uncommon in traditionally circuit theory, it may be of some pedagogical value to illustrate their most important properties by an example following the procedure that actually led to the discovery of the memristor as a physical component [71]. This memristor was realized as a device that possessed a net resistance R that switched between a more conductive state R_{on} and a less conductive state R_{off} depending on the current passing the component

$$R(x) = x(t)R_{on} + (1 - x(t))R_{off}, \quad (4.4)$$

where the state variable $x \in [0, 1]$ and was set to be proportional to electric charge. This principle is also the basis for the memristive description of skin electrical conductance in Paper V where $\frac{dq}{dt} \sim i(t)$ and where the charge carriers, through electro-osmosis, will influence the degree of water filling in the sweat duct, and hence also the net resistance.

The memristor produced in the HP lab was made of titanium oxide that had been ion doped and then tightly squeezed in between two metal contacts separated by a distance D as illustrated in figure 4.2 where 17 memristors are shown in parallel. The electrical properties of this nano-device alter when the ions drift in an applied electric field and total resistance can then be given as [71]

$$M(q) = R_{off} \left(1 - \frac{\mu_V R_{on}}{D^2} q(t) \right), \quad (4.5)$$

where μ_V is just the average ion mobility in this material. This expression in particular serves as a good illustration of typical properties of memristors: The scale dependence and the charge dependence on the resistive memory-effect. As the typical scale, D , of the system decreases, the term involving the charge-dependency (i.e. giving the non-linear characteristics of the memristor) rapidly plays a more dominating role in the total resistance, and so memristors are typically to be found in small scale systems. As the system is sufficiently small, memristive effects will totally dominate the resistivity of the given component, but as D is large, the memristor degenerates to

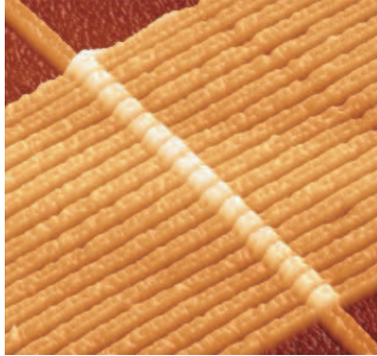


Figure 4.2: The first memristor realized as a physical component. Figure: Wikimedia commons.

an ordinary resistor. The charge-dependence in equation (4.5) ensures the behavior as a non-linear resistor with memory.

Typically memristive devices show double-loop hysteresis in a parametric current-voltage plot. An example is illustrated in figure 4.3 where suitable parameters are chosen based on the model in Paper V. At higher frequencies the memristor degenerates to a simple resistor with a straight iv line. Using the state variables q and ϕ as a basis for a parametric plot, we retrieve a single-valued and non-linear characteristic for memristors as seen in the inset in figure 4.3. Again, if the $q\phi$ curve is simply a straight line, memristance has degenerated to resistance, or to state it otherwisely, M is just a constant.

4.2 Memristors and bioimpedance

So far we have gone through some generic properties of memristors that perhaps were mainly of general interest, but have not yet considered the new possibilities that necessarily arise with the introduction of a new fundamental circuit element. Within the field of bioimpedance and bioelectricity, as well as in biology, a lot of non-linearities are present in biological systems [32], but may sometimes be of an unknown origin. Especially in systems at small scales, where memristive effects are expected to be of increasing significance, one should open the possibility that non-linear phenomena are sometimes better modeled memristively than by the conventional RLC circuit elements, or by the CPE for that matter.

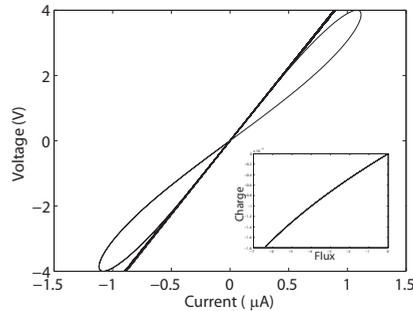


Figure 4.3: Hysteresis loops characteristic for memristive systems. At high frequencies the hysteresis vanishes and the memristor has turned into an ordinary resistor. Inset: Charge-flux characteristics are single-valued for memristors.

The scales where memristors have been reported so far [71] are also of the order of size where biophysics and bioelectricity start to emerge, i.e. from molecular levels towards larger systems, and it is therefore to be expected that memristors are relevant also here. The fact that memristors have been hidden for so long, compared to resistors, inductors and capacitors, increase the possibility that memristive systems have been reported wrongfully, for example as anomalous current-voltage characteristics. Apart from its fundamental interest, the possibilities and excitement of the memristor are that it enables new and conceptually more correct descriptions of some phenomena within bioimpedance. However, it is hard to say something specific, at the present stage, of how widely it can be used.

A new physical component opens for improved equivalent modeling of tissue electrical properties as memristor theory should also be rich enough to cover non-linear phenomena. This would be very promising in our field where circuit modeling plays a central role [32, 73]. As introduced by Chua [72], memristive systems, with all four RLCM, form a complete basis for constructing circuit models for all kinds of electrical phenomena. In Paper V we have introduced memristors to skin electrical properties and showed that this “new” element is capable of mimicking the observed non-linearities in the current response as the resistance of the sweat duct capillaries can be thought of as having some memory depending on the current history.

Apart from describing the skin capillary physics more conceptually correct than traditional RLC-modeling, the memristor is likely to capture also some cell electrical phenomena where the current-voltage behavior of cell po-

tentials are due to the flux of ions. This is a charge controlled system, first described by Hodgkin and Huxley [74] in 1952, that can be described memristively as was later mentioned by Chua [72]. Such cell electrical phenomena, although not passive (the ion pumping needed for a potential build-up is energy consuming), satisfy the two criteria for memristors since they are charge controlled as well as in the nanometer scale where memristive effects are believed to be present.

Although the concept of memristance has not yet been adapted in modeling biophysical electricity, a few things are important in order to include also this building block: Any current responding to an applied sinusoidal voltage should from now on be interpreted with also the memristor element ready in the modeling toolbox. Especially if the iv curve turns out to be double looped with hysteresis pinched in the origin, memristors may provide just the suitable description of the observed phenomenon. This is also the case if one observes a “positive differential resistance” in a iv parametric plot, i.e. that the current continues to increase for some time after the applied voltage started to decrease. Such a positive feedback effect is visible in figure 4.3. It is worth remembering, looking for memristive signs, that the memristor is basically an element that works under AC conditions where the applied voltage depends sinusoidally with time.

Moreover, memristors have been shown to be useful in describing the process of learning for very small and simple organisms [75]. It is believed that this understanding of very primitive intelligence, and the corresponding memristor circuit describing this process, is similar to neurons in the brain and applications are believed to be found in describing neural networks someday in the future.

It is not easy at the present stage to see the full potential of the memristor in bioimpedance. It is possible that it turns out trivial with very few applications far from our everyday measurements. However, as stated above, there are typical memristive fingerprints that should be kept in mind when interpreting the measurements. Also, there are a number of non-linear phenomena that are not memristive. As an example the traditional resistor may also be non-linear, i.e. if the resistance is current dependent. This need not necessarily be due to a memory function, and so current dependence is not the same as charge dependence.

4.3 A few final remarks on our model

In Paper V we have generalized the memristor concept into memristive systems containing two coupled state variables (not just one as in Strukov *et al.*

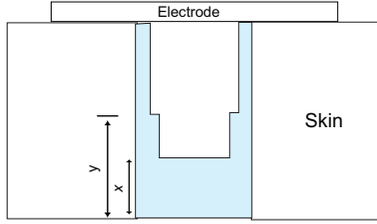


Figure 4.4: A simple illustration of a memristive system containing two state variables.

[71]) so that in a general form the governing set of equations is given by

$$v = R(x, y)i \quad (4.6)$$

$$\frac{dx}{dt} = f_1(x, y, i) \quad (4.7)$$

$$\frac{dy}{dt} = f_2(x, y, i), \quad (4.8)$$

where f_1 and f_2 are general, coupled functions. This is shown in a simple sketch in figure 4.4 where, as in Paper V, y is the relative height of the wetting film, and x is the relative height of the water filling in the duct. I refer to Paper V for further details. Such a memristive system, with the coupled equations, quickly gets intricate. However, as the value of y is constant for all times greater than $t = T/2$, where T is the period of the applied voltage signal, our model simplifies to a memristive system with only one state variable.

An open question left in this thesis is whether a constant phase element (CPE), that frequently is used in circuit modeling of the human skin, can be identified as a memristive system. It is generally known that such an element cannot be realized by a finite number of RLC-components, and it has been thought of as purely unphysical. The CPE is known to have a continuous distribution of time constants [32], and Chua [72] shows that memristive systems, under certain circumstances, may be realized as systems having similar properties. A problem that necessarily arises in this manner is that the definition of memristive systems yields zero current as the potential vanishes, as seen in equation (4.2) and (4.3). This induces a phase angle that is not constant with time. Thus, within memristive systems, the phase angle seems non-constant which is incompatible with a CPE. As this is not entirely investigated, and presently only at a speculative level, it demands a more thorough study before such a very useful connection eventually can be established (or rejected), and is without the scope of this thesis.

Part II
Conclusion

Chapter 5

Summary of my results

The papers that are presented in this thesis do not necessarily form subsequent steps, all in the same direction and within the same branch in bioimpedance. On the contrary, they approach the core topics in my study from different angles such as skin hydration assessments (Paper I and II), water holding properties of keratinized tissues (Paper III and IV), electrical properties, that intimately are linked to SC hydration (Paper V), and skin conductance measurements viewed in the light of electrode gel sorption properties (Paper VI).

Here, I will in short summarize the main findings in my papers, and finally share some thoughts about potential future topics of relevance.

Paper I

In this paper, which purely is a theoretical work, a method for estimation of *in vivo* SC water content is carried out. The method combines the experimentally established dependence of electrical susceptance on water content with the shape of the water concentration profile across the SC.

Paper II

Trans-epidermal water-loss (TEWL) is combined with SC desorption properties to form a non-invasive method for estimating *in vivo* SC steady water content, and so this paper has the same aim as Paper I. Baseline TEWL-data and evaporation rate of water from the SC enable a calculation of the total content of water in the SC area of interest.

Paper III

This paper is concerned with the water sorption properties of the human SC, that is, how water is bound and released depending on time as well as external factors such as relative humidity. SC samples from both heel and breast are investigated. The implications of hysteresis, or the lack of it, during a full sorption cycle are discussed as well as the triphasic patterns seen in the absorption time constants.

Paper IV

In this paper we followed up the previous with a broader study regarding water holding properties of keratinized tissues such as hair, nail and SC. Water uptake was found to be quite different between these three sample types, although being similar in composition and structure. The SC was found to take up considerable more water than hair and nail, and this is related to differences in internal structure, and “armoring” that prevents swelling.

Paper V

In this paper we developed a new model for sweat duct conductivity based on memristor theory. Memristors are the fourth fundamental circuit element, but has remained “unused” for many years, due to their late realization as a physical component. Current responses of human sweat capillaries are shown to behave as memristive systems since the resistance of the capillaries depends on the current history, i.e. behaving memristively.

Paper VI

Electrodes are known to influence the electrical properties of human skin (see i.e. [32]), and hence a proper choice of electrodes for the intended use is vital. In this paper we found a clear correspondence between gel water content free to evaporate and skin electrical recordings after electrode onset. In addition, the viscosity of the gel was found to be important, as a low viscosity gel opens for sweat duct penetration, giving an *inverse sweat response*, increase the probability of mechanical instability as well as reduce the ability of the recordings to return to baseline after any response.

5.1 Look ahead

Below are a few of many topics that could be possible for future studies:

- The knowledge of how water is bound in the SC should be used in order to better understand the observed SC electrical properties.
- The possibilities of a calibration of *in vivo* water content against low frequency susceptance measurements should be investigated further.
- The “new” element - the memristor - opens for improved circuit modeling within bioimpedance, and these possibilities should be followed up, including e.g. a potential CPE-modeling.
- The memristive model of skin conductance should be developed further, including also the “clipping”-effects that are believed to influence the sweat duct conductance.
- The inverse sweat response, present for gels with low viscosity, should be studied further with the aim of revealing the mechanisms that govern this interesting phenomenon.
- How will the electrode induced effects to skin conductance behave if AC is replaced by DC current, that is more frequently used in clinical studies? Electro-osmotic effects are likely to be present in the DC situation.
- The characteristic time constants should be found also for hair and nail with the aim of better understanding water holding properties of keratinized tissues.

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Part III
Papers

Estimation of *In Vivo* Water Content of the Stratum Corneum from Electrical Measurements

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Abstract: *In vivo* water content in the epidermal stratum corneum can be estimated by means of low frequency susceptance measurements. In the *in vitro* calibration necessary to find the *in vivo* water content, the stratum corneum will have a uniform distribution of water across its thickness. However, *in vivo* stratum corneum has an increasing water concentration profile from the outermost towards the innermost parts. This paper will investigate the possibility of estimating the equilibrium water content in the *in vivo* stratum corneum non-invasively from electrical susceptance measurements. Given a known shape of the water concentration profile in the *in vivo* stratum corneum and the dependence of susceptance on the water content, it is possible to calculate the water content *in vivo* based on analytically derived expressions for the water concentration profile. A correspondence between *in vivo* and *in vitro* water content needed for this purpose is also established.

Keywords: Bioimpedance, electrical susceptance, stratum corneum, hydration.

INTRODUCTION

The vital functions of the stratum corneum (S.C.) are strongly dependent on the water content therein [1-3]. Knowledge of the water content of the S.C. *in vivo* has been shown to be an important factor in clinical determinations of non-visible skin diseases such as atopic eczema [4-7]. Electrical methods for measuring skin hydration have been studied for several decades and a low frequency susceptance is found to be the most adequate [8-10]. In this paper the term low frequency means a constant frequency below 100 Hz. In general higher frequencies will measure also on deeper skin layers [11]. Furthermore, the conductance of the S.C. is influenced by sweat duct filling which is an unwanted error to the measurements. The method of Martinsen *et al.* [12] is not affected by sweat duct filling, viable skin or temperature as it is focused on S.C. only [9]. Often a four-electrode system is used for the electrical bioimpedance measurements [13], but the susceptance data used in this paper are found with monopolar recording using a three-electrode setup following the principles of Grimnes [14].

In this theoretical paper we want to investigate the possibility of estimating the equilibrium water content of the *in vivo* S.C. by electrical susceptance measurements. The method we propose in this paper is non-invasive, rapid and simple. The estimation of the *in vivo* S.C. water content is done by calculating the relation between *in vivo* and *in vitro* susceptance, and thereby also the connection between the corresponding water contents. Viable skin contains about 70-80 % water [1], whereas the surface of the S.C. is drier. There will therefore be a net water gradient throughout the

S.C. *In vitro*, however, the water content may be kept uniform, due to a moisture chamber as in the measurements of Martinsen *et al.* [12], where an exponential correspondence is found between measured susceptance and water content of the S.C. given in gram water per gram dry skin. This means that any *in vivo* susceptance corresponds to an *in vitro* water content. As also discussed in Johnsen *et al.* [15], assuming a linear water concentration profile in the *in vivo* S.C. it is possible to analytically find a relation between *in vitro* and *in vivo* measurements, and therefore also possible to predict the water concentration profile which yields the water content of *in vivo* S.C. The linearity of the water concentration profile is indicated in the literature [16-20]. Also from diffusion theory the water concentration profile of the S.C. will be linear as a first approximation. The passive diffusion of water across the S.C. is governed by Fick's laws of diffusion. For the equilibrium transport of water across the thickness of the S.C. Fick's law [21, 22] yields

$$F = D \frac{dC(z)}{dz}, \quad (1)$$

where F is the flux of water, D is the diffusion constant, and $C(z)$ is the concentration of water across the S.C. thickness, z . A constant value of D gives a linear gradient of water in the S.C. whereas a non-constant value, as indicated by [20, 23], will give deviations from linearity. However, a slight variation in the value of D with water content results in only small deviations from linearity with respect to the water concentration profile in the S.C. [20, 22] where the former predicts linear water profiles based on a non-constant value of D with respect to the water content of the S.C. We will show that the assumption that the concentration profile being linear is valid for our calculations even if the profile deviates from linearity, i.e. containing also higher order terms. Thus, as a first approximation we consider the *in vivo* water concentration profile to be linear across the entire S.C. when

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we perform our estimates of *in vivo* water content. Any given analytic expression of the water profile will in turn make it possible to estimate *in vivo* water content.

METHODS

The S.C. is modeled so that the water content increases with relative depth, z , where $z \in [0,1]$. The S.C. consists of dead and deformed keratinised cells containing water in and around them [24]. This results in strong capacitive properties at low frequencies [25, 26], and a section across the S.C. thickness is therefore modeled as a series of capacitors, each with infinitesimal thickness and a dielectric dependency on the water content at the given depth. Thus, the S.C. is viewed as a continuous series of capacitors and resistors as illustrated in Fig. (1). The current density will be equal in all infinitesimal slices of the S.C. that is being considered (i.e. under the electrode). This is ensured by using a three-electrode setup as described first by Grimnes [14]. For sufficiently low frequencies, that is, below 100 Hz, impedivity of the S.C. is much higher than for viable tissue [27]. In addition, the much larger dimension of the measuring electrode compared to the S.C. thickness also justifies a uniform representation of the current density across the S.C. thickness. Yamamoto and Yamamoto [28] report on an upper limit of linearity of skin current density to be about $10 \mu\text{A}/\text{cm}^2$ at 10 Hz and $100 \mu\text{A}/\text{cm}^2$ at 100 Hz. In this paper we will be operating well within the linear range. Electrical admittance is basically measured by applying a constant voltage and measuring the corresponding current. Furthermore, a lock-in amplifier is used to measure only the imaginary part of the admittance, i.e. the susceptance. As a general model of the water concentration profile we will introduce a linear term and also higher order terms, covering a broad class of profile shapes (depending on the values of the parameters)

$$w = a + bz + cz^2, \tag{2}$$

where w is water content given in gram water per gram wet skin (i.e. skin and water), a (dimensionless), b ($[\text{length}]^{-1}$) and c ($[\text{length}]^{-2}$) are constants. Thus, the outermost water content of the S.C. is then given by a and

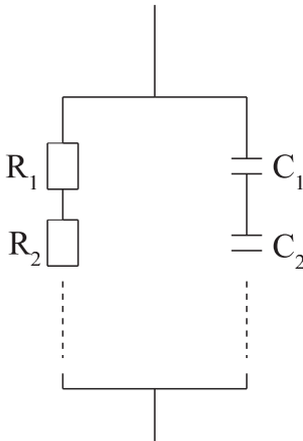


Fig. (1). The electrical model of the stratum corneum.

the innermost by $d = a+b+c$. The parameters of the water concentration profile in our model are unbiased, meaning that the value of b and c can be negative giving a negative slope. Such a shape of the profile may appear if the relative humidity of the ambient air is sufficiently high giving a larger value of a than d .

The exponential behavior of the susceptance with respect to *in vitro* water content, v given in gram water per gram dry skin found in Martinsen *et al.* [12], then yields the general form of the susceptance

$$B = B_0 \exp(\alpha v), \tag{3}$$

where B_0 represents the susceptance measured when the S.C. is only one, uniform dielectric, and α is an experimental constant that can be found in Martinsen *et al.* [12] in the case of *in vitro* S.C. taken from the heel. We want to use the linearity of the water concentration profile in the S.C. to find *in vivo* water content. However, the susceptance depends exponentially on gram water per gram dry skin and must therefore be converted to be a function of gram water per gram wet skin, that is, a function of w rather than v , in order to be combined with the linear water distribution. Converting v into w is done by

$$w = \frac{v\rho_s\rho_w}{v\rho_s + \rho_w}, \tag{4}$$

where ρ_s and ρ_w are the densities of dry S.C. and water, respectively. The density of S.C. is estimated by Scheuplein [29], and the density of water is set to 1.00 g cm^{-3} . Eq. (3) then turns into

$$B = B_0 \exp\left(\frac{\alpha}{\rho_s} \frac{a + bz + cz^2}{1 - a - bz - cz^2}\right) \tag{5}$$

An expression of the infinitesimal *in vivo* impedance, dZ , for each thin capacitor in our model can be found through the relation $X = (\omega C)^{-1}$, where X is reactance, ω is signal frequency, at which the susceptance is being measured, and C is the capacitance depending on the water content of the dielectric. It is worth noticing that the outermost and driest parts of the S.C. in general will contribute more to the impedance than the deeper layers. Using that relative permittivity is proportional to susceptance, we get that

$$\frac{dZ}{dz} = \frac{1}{\omega A \epsilon_0 \epsilon(w)} \tag{6}$$

$$= \frac{1}{\omega A \epsilon_0} \exp\left(-\frac{\alpha}{\rho_s} \frac{a + bz + cz^2}{1 - a - bz - cz^2}\right), \tag{7}$$

where A is the cross-section area of the S.C. region being considered, ϵ_0 is the electrical permittivity of vacuum, and $\epsilon(w)$ is the relative dielectric permittivity which is proportional to the susceptance and therefore also a function of the water content, w . Total *in vivo* impedance of the S.C. is found by integrating up (7)

$$Z = \frac{1}{\omega A \epsilon_0} \int_0^1 \exp\left(-\frac{\alpha}{\rho_s} \frac{a + bz + cz^2}{1 - a - bz - cz^2}\right) dz \quad (8)$$

We would now like to find the *in vitro* water content, w_{in_vitro} that corresponds to the *in vivo* impedance given in (8). This is done by combining the *in vivo* result with the fact that impedance decreases exponentially with the water content, w (gram water per gram dry skin). Thus, from (8) we get that

$$Z = \frac{1}{\omega A \epsilon_0} \exp(-\alpha v_{in_vitro}), \quad (9)$$

which results in

$$v_{in_vitro} = -\frac{1}{\alpha} \ln \left[\int_0^1 \exp\left(-\frac{\alpha}{\rho_s} \frac{a + bz + cz^2}{1 - a - bz - cz^2}\right) dz \right] \quad (10)$$

The *in vitro* water content, w (gram water • gram wet tissue) is then given by

$$w_{in_vitro} = \frac{v_{in_vitro} \rho_s}{v_{in_vitro} \rho_s + 1}, \quad (11)$$

with v_{in_vitro} given as in (10).

RESULTS

The correspondence between *in vivo* and *in vitro* water content is given by the set of equations (10) and (11) where the *in vivo* water content is given by the right-hand-side (RHS) of (10). We will now show that the estimated value of *in vitro* water content, based on any physical *in vivo* concentration profile is more or less the same when $c = 0$ (linear profile) as when $c \neq 0$ (deviations from linearity). For any measured *in vivo* susceptance value, the water concentration profiles must satisfy the condition

$$f(v_{in_vitro}, a, b, c) = 0, \quad (12)$$

where

$$f(v_{in_vitro}, a, b, c) = v_{in_vitro} + \frac{1}{\alpha} \ln \frac{1}{\omega A \epsilon_0} \int_0^1 \exp\left(-\frac{\alpha}{\rho_s} \frac{a + bz + cz^2}{1 - a - bz - cz^2}\right) dz \quad (13)$$

We see that the water concentration profiles are in general not unique for a given value of the *in vitro* water content (i.e. for an *in vivo* susceptance) due to the fact that both a and c are unknowns. This is as expected since two independent parameters can not be obtained from a single measurement. Hence, an *in vivo* susceptance will in general not be sufficient to determine the profile of the water concentration of the S.C. exactly, but rather serve us with a continuum of possible solutions.

The assumption that *in vivo* water content of the S.C. can be estimated from susceptance measurements and a linear concentration profile is reasonable as a first approximation. We show this in Tables 1 and 2, and Fig. (2) where we compare the results of *in vitro* estimated values for a purely linear concentration profile (i.e. $c = 0$) and profiles containing also non-linearities for two different cases. The

Table 1. In Vitro Water Contents Corresponding to a Linear Water Concentration Profile and Non-Linear Profiles when $a=0.212$ and $b=0.288$. The Constants, a , b , and c are as defined in Eq. (2)

c	v_{in_vitro}
-0,4	9
-0,25	16,3
-0,1	19,32
0	20
0,1	20,42
0,25	20,86
0,25	21,39

Table 2. In Vitro Water Contents Corresponding to a Linear Water Concentration Profile and Non-Linear Profiles when $a=0.132$ and $b=0.268$. The Constants, a , b , and c are as defined in Eq. (2)

c	v_{in_vitro}
-0,4	11,4
-0,25	27,12
-0,1	29,53
0	30
0,1	30,3
0,25	30,67
0,25	31,1

deviation is significant only for very large negative (or positive) values of c . Physically, the numeric value of the parameter c , governing the deviation from linearity, may be limited to the interval $[-0.5, 0.5]$. This is due to the fact that one can not have negative values of the S.C. water content. Values exceeding the relative value of 100 % are also unphysical. The most likely value of c is that it is slightly positive due to the fact that there is a gradient from the outermost S.C. (dry) towards the interface at the vial epidermis (wet). We note that the deviations from the linear results are small in this manner. Thus, it seems to be a reasonable approximation to model the concentration profile to be linear when we want to find the correspondence between *in vivo* and *in vitro* water content, and the term cz^2 will therefore be omitted in the following. Water content of the S.C. *in vivo* can possibly be found on the basis of the water concentration profiles. The profiles are in general given by the RHS of (10). We can think of this process of estimating *in vivo* water content as illustrated schematically in Fig. (3). Any susceptance corresponds to an *in vitro* water content which again is equivalent with *in vivo* water content in the S.C. It should be commented that whereas the relation between *in vitro* susceptance and water content in Fig. (3) is one-to-one, the correspondence between *in vitro* water

content and *in vivo* water content is not unique. This is due to the fact that a , b and c are in general not constant.

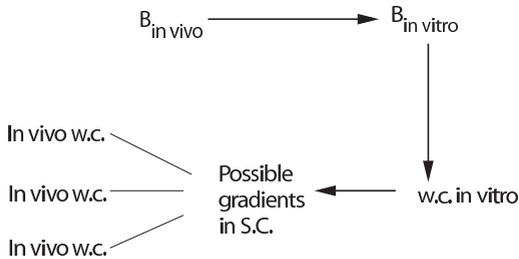


Fig. (2). The derivation of possible *in vivo* water gradients and then again *in vivo* water content with basis in an *in vivo* susceptance measurement.

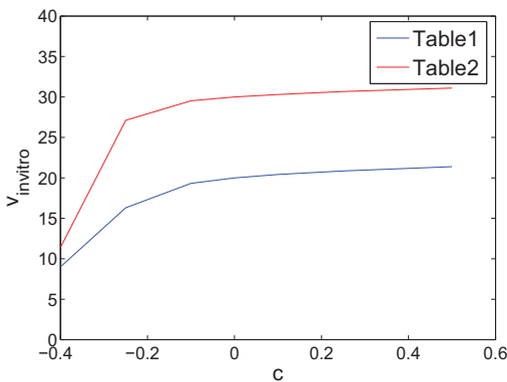


Fig. (3). The data in Table 1 and 2 illustrated graphically. Note the nearly flat curves for positive and slightly negative values of c .

DISCUSSION

We have in this paper studied the possibility of estimating the water content in the S.C. by means of electrical susceptance measurements performed *in vivo*. We have found a correspondence between *in vivo* and *in vitro* water content and shown that the estimated *in vivo* water content from a measured susceptance value is approximately the same when a linear water concentration profile is assumed as when a non-linear profile is used for the basis of our calculations. This is at least correct if the parameters of the profile have physically realistic values. In this paper we have found relative *in vivo* water content (gram water per gram skin) and not absolute content. In order to find absolute water content of the S.C. from electrical susceptance measurements, the exact shape of the profile would be needed as well as the S.C. thickness of the area being considered. This should be a topic for future studies and would include more experimental methods in order to reveal the parameters in our model. If either of the parameters in the gradient are fixed, it would be a step towards an analytical expression of the absolute S.C. water content. The value of a , the water content of the S.C. surface, will depend

on the relative humidity and temperature of the ambient air and will probably vary with time. Thus, the most likely candidate to be fixed is the water content, d , at the boundary of the S.C. towards viable skin. Viable skin contains a more or less stable content of water, independent of ambient variations such as relative hydration and temperature, which opens the possibility of boundary conditions to the innermost parts of the S.C.

An important question in this context is whether the interface towards the epidermis provides natural boundary conditions to the value of the innermost water content in the S.C. This should be a topic for future studies since literature seems ambiguous at this point. Occluding the S.C. while measuring susceptance may result in an asymptotic behavior of the susceptance due to a stable content of water in the S.C. corresponding to the level of hydration of the viable skin. Another possible method to search for the value of d is from trans-epidermal-water-loss (TEWL)- measurements. Adjusting the relative humidity of the ambient air until TEWL vanishes probably means that the S.C. is hydrated at a level equal to d throughout its entire thickness. The S.C. thickness varies due to swelling and that the number of keratin layers vary at different parts of the human body [19, 30]. This is a problem yet to be solved. However, the interpersonal differences with respect to the S.C. thickness at equivalent parts of the body may be expected to be smaller, and therefore possible tabulated values of S.C. thickness with respect to location, age, sex etc. may show a better uniformity and possibly be good enough for our purpose.

It is also worth noticing that the *in vitro* water content is in general not the average of the *in vivo* water concentration profiles. This is as expected due to the capacitive properties of the S.C. where the outermost and driest parts of the S.C. contribute more to the impedance, and hence more to the estimated corresponding *in vitro* water content. This is, however, also a weakness in our model since then the estimated *in vivo* water content may not vary much with the changes in the shape of the innermost part of the water concentration profile.

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Sorption Properties of the Human Stratum Corneum

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Abstract

Water sorption is important for the overall structure and function of keratinized tissues such as the human epidermal stratum corneum (SC). In this study we report on a gravimetric method for studying sorption properties of human SC, both from heel and female breast skin. Changes in mass were measured as the relative humidity was altered in steps under controlled environmental conditions. The possibility of hysteresis is also discussed. Furthermore, we have found that the sorption time constants show triphasic behaviour during absorption, but not during desorption. This behaviour is connected to the three different types of water present in the SC. Water also enters the SC much more rapid compared to its exit at relative humidities lower than 50 %. Finally, the amount of time between sample preparation and onset of measurement seemed to have an effect on the absorption rate, but less on the total amount of water absorbed.

Keywords: Skin, stratum corneum, hydration, time constants, hysteresis.

1. Introduction

The foremost function of skin is to constitute a barrier between the body and the environment. The SC, i.e. the superficial most layer of skin, is directly responsible for skin barrier function. It is composed of dead keratinized cells embedded in a continuous extra cellular lipid structure. The major role of the keratinized cells is to constitute a both stiff and viscoelastic protective scaffold for the lipid structure. The protective mechanical properties of the stratum corneum cells are intimately linked to their water content [1-3]. Skin function is thereby largely dependent on stratum corneum water balance. Also, the stratum corneum hydration properties are important for the understanding of skin diseases characterized by deficient barrier function, such as atopic dermatitis [4-6]. The SC water content will generally vary

with both relative humidity (RH) and temperature as the partial pressure in ambient air and SC tries to equal. Water contents of more than 10 % are necessary in order for the skin to remain healthy and pliable [1].

In this study we report on a gravimetric method for studying the sorption properties of human SC, both from heel and female breast skin obtained from plastic surgery. Knowledge about SC sorption properties is central for our understanding of SC hydration. It will also aid the interpretation of electrical skin hydration measurements [7]. First we investigated the absorption and desorption properties of SC in order to evaluate if hysteresis was detectable, meaning that the amount of water absorbed at a fixed RH during absorption is different from the corresponding water content during desorption. Water uptake as a function of RH was studied as well as time constants, which are the characteristic time needed in order to reach about 63 % of the equilibrium value.

The study was performed with a dynamic vapour sorption (DVS) instrument from Surface Measurement Systems Ltd (London, UK), where the relative humidity is altered in steps under controlled environments. Kilpatrick-Liverman *et al.* previously performed a similar study on porcine skin [8], but to our knowledge no such study has been performed on human skin. However, some results were presented in a preliminary study [9]. Also few, if any, consideration of the sorption time constant have earlier been carried out, although Martinsen *et al.* commented on some of these aspects [7], and literature reports on concentration dependent diffusion of water within the SC [10,11].

2. Materials and methods

The SC heel samples were taken from the heel of 10 volunteer test persons, 5 men and 5 women at the ages ranging from 24 to 29 years, all with no signs of unhealthy skin and with no history of dermatological diseases. The SC was removed by means of an Aesculap dermatome (Braun, Tuttlingen, Germany), and the sample pieces were about 4 by 5 mm in size after preparation. All samples were prepared with as equal size as possible in order to reduce any effect of geometry on the final results. Their thicknesses were 0.20 mm, which was the thinnest possible configuration provided by the dermatome.

Breast skin from 10 healthy females aged between 32 and 68 years was obtained from reconstructive surgery and kept in a freezer at -20° C for 3 months. After dissection of the subcutaneous fat, the samples were heated for 3 min at 60°C with the dermal side in contact with a metal plate. The superficial epidermis was then peeled off with a razorblade. Next, the samples were incubated in 0.5 % trypsin with the epidermal side down for 2 h at ambient temperature to remove remnants of stratum granulosum adhering to the epidermal side of the SC. Finally, the sheets were washed four times in distilled water and dried and stored at ambient temperature under nitrogen for about 2 years.

One SC heel sample from one of the 5 male test subjects was also hydrated in HPLC water for 4 hours and then contained in a freezer at -20° C for 72 hours. This was done in order to investigate the effect of freezing on the sorption properties of the SC. Also, 3 heel samples from one of the 5 male test subjects were contained *ex vivo* for several days prior to experiment onset in order to discover any effects on sorption properties with respect to time between sample preparation and measurement onset.

Furthermore, a DVS was used in order to reveal the SC sorption characteristics by altering the RH in steps from 0 % until 90 %, and then back to 0 %. The steps were all pre-programmed to be of a 10 % increase in RH and temperature was kept constant at 25° C throughout the entire experiment. The DVS apparatus provides a very sensitive microbalance weight with a resolution of 0.1 µg. The SC heel samples were prepared so that their weights

were about 3-5 mg, whereas the breast samples, being much thinner, were about 0.4-0.6 mg, but still well within the resolution of the DVS instrument. The DVS was set to jump to the next stage, i.e. a new RH value, when the rate of sample mass, dM/dt , was less than 20 ppm per minute over a period of at least 10 minutes.

The sorption time constants were found by fitting the data to an exponential time course, $y = a + \exp(-kt)$. A course proportional to $\exp(-kt)$ would then yield time constants equal to $\tau = 1/k$.

3. Results

Figure 1 shows the time course of one of the SC samples, taken from the heel, as the RH in the DVS chamber was altered in steps of 10 %. The increment in mass corresponds to absorption of water caused by an increase in RH, whereas decreasing mass corresponds to desorption, i.e. the sample loses water to the environment. The increase (or decrease) in weight then equals the percentage of the weight of water in the skin sample relative to the dry weight of the sample. Initially, the SC sample was set to be in equilibrium with the environment at 0 % RH, giving the dry weight of the sample. Parts of this calibration is seen to the far left in the time course in figure 1 where the sample has a negative mass gain. The time evolutions of the 10 SC heel samples showed very similar time courses. We notice that the first steps in RH resulted in very small mass changes. These steps are better viewed in figure 2 where the steps corresponding to RH at 10, 20, 30 and 40 % are enlarged. The corresponding desorption time courses at low RH followed similar patterns, only now with a negative mass rate.

The alterations in mass of the samples, initiated by the instantaneous increments in RH, followed courses that were close to exponential at high RH, indicating that the mass rate was proportional to the difference between the actual mass and the equilibrium mass for each step. This is in accordance with Fick's diffusion theory [11], and time courses of the mass for each step was therefore approximated as

$$M(t) \approx \exp(-k_i t)(M_1 - M_2) + M_2, \quad (1)$$

where M_1 and M_2 are the sample masses before and after a step in RH was initiated, respectively, $i = a, d$, so that k_a^{-1} is the time constant for the absorption process, whereas k_d^{-1} is time constant for desorption. The steps at the lower part of the RH interval deviated somewhat from a purely exponential behaviour, especially courses at 30 and 40 % RH which are shown in figure 2. However, all regressions produced fits to the measured data with root mean square error (RMSE) less than 0.2%.

In figure 3 a corresponding course is shown for a full sorption cycle for one of the samples from SC isolated from breast skin. The two different types of SC samples, heel and breast, showed very similar full sorption courses, however, with more distinct steps for the latter sample type. This may be due to the fact that breast SC samples were much thinner than SC taken from the heel, which again resulted in much larger surface to volume ratio, resulting in less time in water diffusing into the samples.

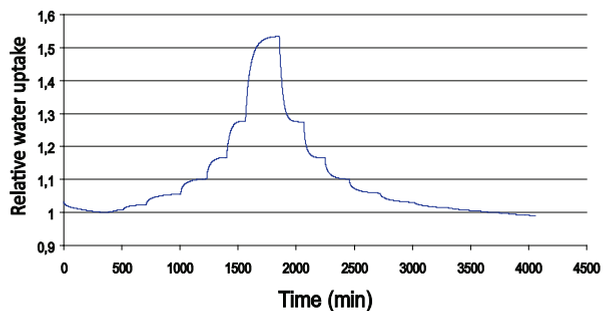


Figure 1: Time course of relative water uptake for SC from the heel during absorption and desorption.

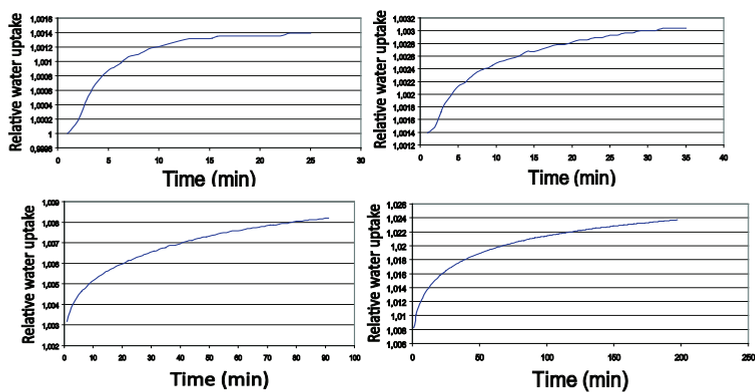


Figure 2: Time courses of relative water uptake at RH corresponding to 10, 20, 30 and 40 % for heel SC during desorption.

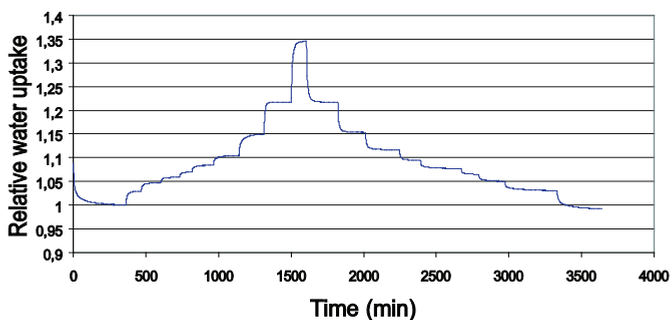


Figure 3: Time course of relative water uptake for SC isolated from breast epidermis during absorption and desorption.

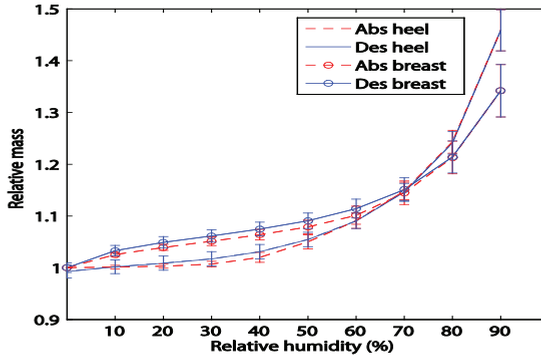


Figure 4: Relative water uptake for heel and breast SC during absorption and desorption.

Figure 4 shows the average water uptake of the two different sample types of human SC at 25° C as a function of ambient RH. The red, dotted graphs correspond to absorption data, and the blue, continuous graphs show desorption data. We observe that the absorption and desorption graphs were very similar, but with the desorption data slightly elevated compared to absorption, indicating that the samples contained nearly equal amounts of water during absorption and desorption at any given value of RH in the DVS closed chamber. The water sorption isotherms for heel SC showed small changes in water uptake for RH below 50 %, but a more rapid and exponential behaviour at the highest RH values. The isotherms for the SC isolated from breast skin were more elevated compared to heel data at RH below 70 %, indicating larger water uptakes. At the highest RH values, breast SC contained substantially less amount of water compared to heel SC. These shapes of isotherms correspond well with others found in previous sorption studies [1,12,13]. There was, however a much less hysteresis than reported in similar studies in porcine skin, and the tendency in our study was that desorption water contents were slightly higher than absorption data. Only at the mid range RH there were small signs of deviation between water content from absorption and desorption for heel SC, however mainly within the uncertainties in our measurements. For breast SC, the small signs of hysteresis were broader, spanning over a wider RH interval.

Figure 5 and 6 compare water uptake of heel SC from male and female test persons. The two courses were as good as identical in both cases, and no differences of significance with respect to the water contents between the genders were visible in this study. Furthermore, the sorption time constant, that is the characteristic time of water entering or leaving the SC, of the heel samples are illustrated in figure 7. The rapidity of water entering or leaving the SC varied as a function of RH. We note the significant difference of the time constants between absorption and desorption at low RH. During absorption, SC rapidly reached its equilibrium state at very low RH and then decreased with an increase of RH until 50 %. Above 50 % the absorption time constant decreased with increasing RH until 80 %, from which it again increased. The desorption time constant followed the pattern of absorption for RH above 50 %, but at lower humidities the desorption time constant was much higher and flattened out as RH approached zero. Thus, the SC seemed to allow water uptake very rapid, but as water left, the time needed was much longer.

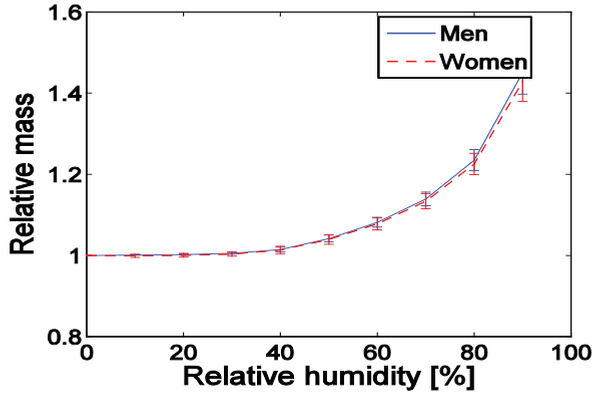


Figure 5: Relative water uptake of male and female heel SC during absorption.

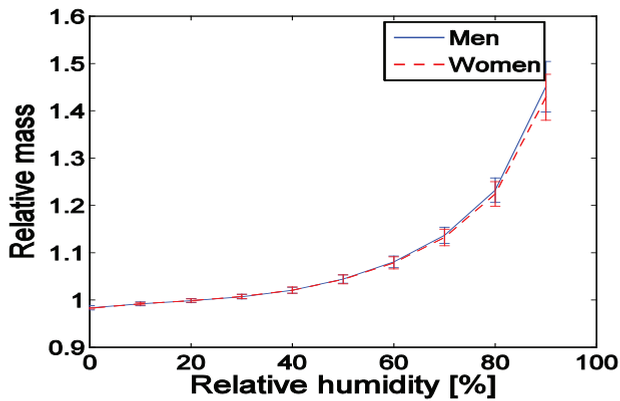


Figure 6: Relative water uptake of male and female heel SC during desorption.

The corresponding sorption time constants, now for the breast samples, are shown in figure 8. Here, the time constants are normalized with respect to the value of the time constant of the first absorption step. This was done in order to be able to compare at least curve forms with the heel data. The breast samples were much thinner than the heel samples, and the thickness of the samples will influence the numeric values of the time constants, so that thinner samples result in smaller values of the time constants [14]. However, as the thicknesses of the breast samples were unknown, and thus potentially of different values, we had to normalize the results of each sample in order to find the shape of the average curve.

We note that there were similar trends in curve form for the time constants of heel and breast SC, although there were also substantial differences. The time constants of both sets of samples experienced a peak followed by a rapid drop during absorption, but with different rates compared to the corresponding absorption trends. In figure 7 the time constant of a heel SC sample that had been hydrated and then frozen for 72 hours is shown in order to see if freezing, which had been applied to all breast samples, would affect the time constants. We note that the absorption data were elevated compared to the 10 other heel samples when RH was lower than 60 %, but that the desorption curve followed more or less the same trend as for the 10 unfrozen heel samples.

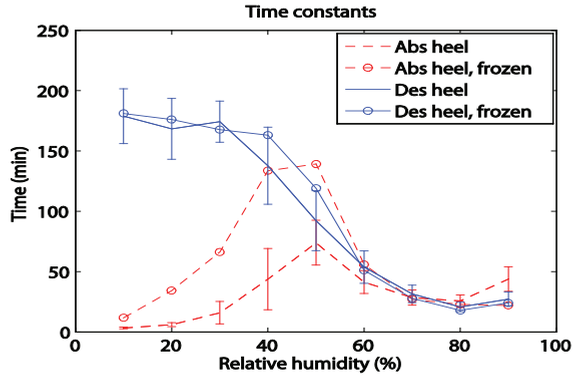


Figure 7: Time constants of heel SC during absorption and desorption.

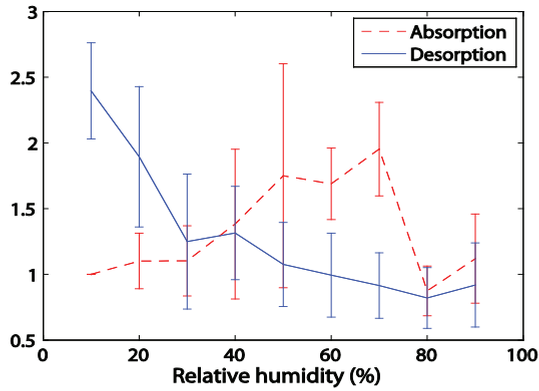


Figure 8: Normalized time constants of SC isolated from breast epidermis.

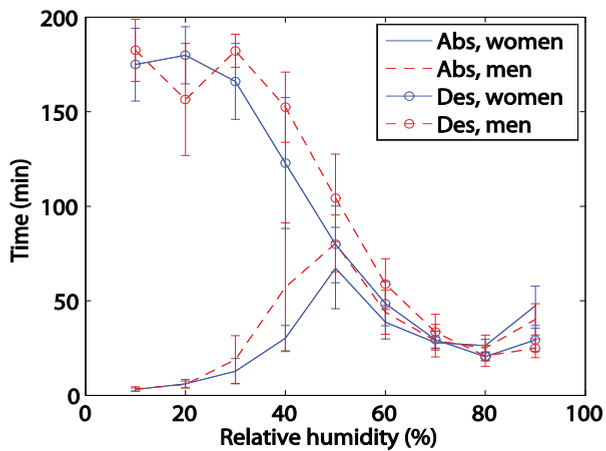


Figure 9: Time constants of heel SC from men and female during absorption and desorption.

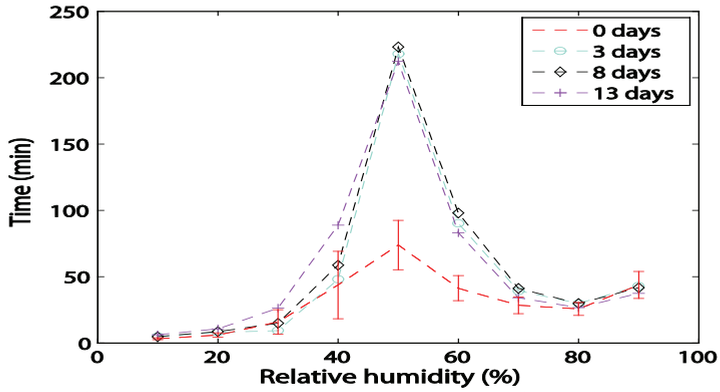


Figure 10: The effect of time between heel sample preparation and onset of measurement on the absorption time constant.

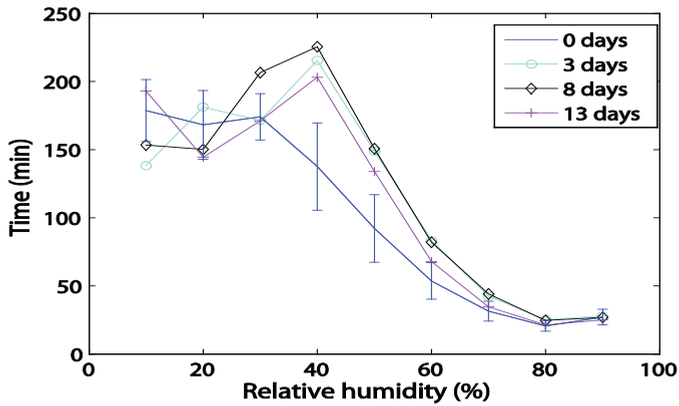


Figure 11: The effect of time between heel sample preparation and onset of measurement on the desorption time constant.

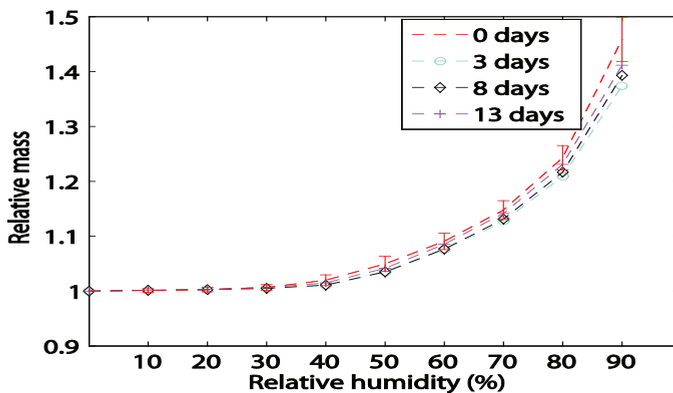


Figure 12: The effect of time between heel sample preparation and onset of measurement on the relative water uptake during absorption.

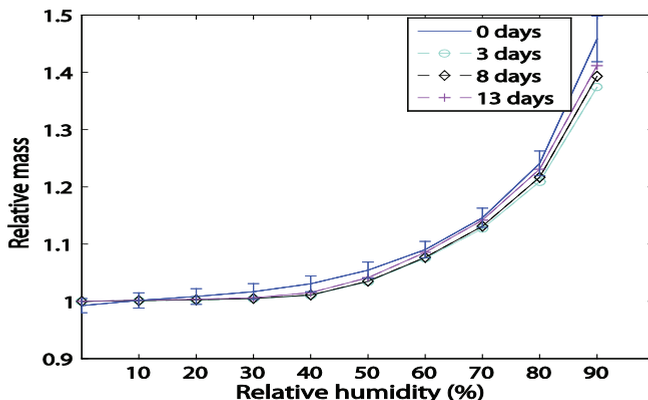


Figure 13: The effect of time between heel sample preparation and onset of measurement on the relative water uptake during desorption.

A comparison of the sorption time constants between men and women are shown in figure 9. The tendency was that female skin reached the equilibrium state more rapid than male skin, and thus seemed to have a higher affinity for water. During desorption this means that male skin held water a little longer before stable state was reached.

It is also interesting to note that the absorption time constant seemed to be dependent on time between heel sample preparation and onset of experiment with the DVS instrument. Figure 10 shows that the absorption time constant increased substantially at the RH interval of 40 to 70 % when the samples were contained in the lab for several days prior to onset of experiment. The same effect was found for desorption data, however with smaller deviations from the average value of the time constants. The number of days greater than three did not seem to influence the data much.

In figure 12 and 13 the corresponding time-effect is viewed through the relative water uptake during absorption and desorption. We note only small differences in water uptake in the RH regions where the time constants altered, but there may be a weak tendency that the samples took up less water after having been stored for several days prior to the onset of measurement. One should, however, notice that in this manner only samples from one male test person was tested against this potential long-term effect of SC from the heel.

4. Discussion

In this paper we have studied sorption properties of human SC and found that the sorption time constants varied greatly with RH of ambient air with triphasic behaviour during absorption. The sorption time courses were best described by single exponential time courses at high RH, whereas at the lowest RH both absorption and desorption showed a behaviour with slightly more pronounced deviations from exponential courses. A single exponential dependence with time may indicate that there is one underlying sorption mechanism that is strongly dominating, and a deviation from such a course could therefore be a combination of two or more mechanisms, both sorbing water, in the SC.

Studies on the diffusion properties of polymers such as keratins have shown that this type of materials show a behaviour governed by Fickian diffusion above a certain, and

material dependent temperature, the glass transition temperature, but being more complex below T_g . This has usually been ascribed to the differences in mobility, above and below T_g , of keratin filaments. Diffusion of water in between the keratin filaments is therefore possible since the filaments are free to move above T_g [15]. A possibility for the behaviour at low RH (i.e. at low sample water uptakes) may therefore be due to the effect of vapour induced glass transition, below which water content diffusion can be expected to deviate from purely exponential. A topic for further studies should therefore be to reveal more of the underlying mechanisms of SC water binding, and to see if SC experiences a vapour induced RH.

From figure 4 we note that our measurements show very small, if any at all, amounts of hysteresis, being of a size comparable to the uncertainties between the different samples, and may therefore be due to the relatively large inter-individual spread with respect to skin sorption properties. Hysteresis is reported, however much more substantial, in porcine skin [8], but not found in human SC from the hip [13]. Some hysteresis has been reported to be found in horn keratin [16], and Hey *et al.* reported on a low pressure SC hysteresis [17]. The hysteresis in our measurements was so small that it in general was within the uncertainties of the measurements. Hysteresis associated with sorption isotherms may indicate that there are mechanisms keeping more of the water within the samples during desorption compared to absorption. Significant hysteresis will in general require structural changes in the SC which again affects the water holding capacity. Such changes are, according to literature expected to take place at temperatures below the glass transition limit. For SC this limit is reported to be below 20° C [18], meaning that large amounts of hysteresis can not, in our study, be expected to occur due to glass transition structural changes.

One can not rule out the possibility that some of the deviation in absorption and desorption water uptakes are due to the necessarily present gap between absorption and desorption equilibrium masses due to the fact that one has to specify a cut-off in mass-rate (we used 20 ppm per minute) as a criterion for RH to step to the next level as well as the fact that SC varies much between individuals. The numeric value of the smallest allowed mass fluctuation in the DVS setup will influence the result, especially if it is chosen too large. A large equilibrium rate will result in short time intervals before RH is programmed to initiate a step of 10 %. This would allow little time for the SC samples to fully reach the equilibrium state, resulting in deviations between absorption and desorption, that is hysteresis. Still, as the mass rate criterion was chosen very small, and much smaller than in the similar study with SC from porcine skin [8], this effect can be assumed to be small, and is not likely to produce large gaps between absorption and desorption water uptake in a sorption cycle.

The difference between SC from heel and breast in curve form, as seen in figure 4, is hard to explain. An effect that may play a role in this manner is freezing artefacts. Ice crystal formation during freezing of skin will probably somehow disrupt the keratin filament network structure which could potentially influence the absorption and desorption properties of the corneocytes. To date little is known about such structural rearrangements upon freezing which is why speculation becomes very difficult. Cryo-electron microscopy of vitreous skin sections is likely the method to obtain such structural information in the future.

We notice that the absorption time constant showed triphasic behaviour. Our results are similar to those found in Martinsen *et al.* [7], but in this study the SC samples were allowed sufficient time to reach the equilibrium states after the initial calibration at 0 % RH, and a full sorption study was performed, including also desorption data. The observed triphasic behaviour may be related to the three different types of water present in the SC (strongly bound, bound and bulk) [13,19-21] and to their corresponding sorption mechanisms. Silva *et al.* reported on three regimes for isolated corneocytes where water contents were below 5 wt %, 5-17 wt %, and above 17 wt %, respectively [22]. This corresponds to our findings of the time constants for absorption in figure 7 and 8 combined with the mass gains

in figure 4. The strongly bound water, potentially bound to polar functional groups of lipid bilayers until monolayer coverage was reached [12,20], entered the SC very rapidly at low RH, and the increase in time constants with RH could be induced by less available sites to attach. Thereafter, as a new type of water was bound in the SC, the time constant decreased, which is an expected behaviour since the diffusion coefficient of water in SC, inversely related to the absorption time constant, usually increases with concentration [14,15]. The final and slight increase in the time constant above RH values of 80 % may be due to bulk water entering the SC combined with the onset of a swelling process that is time consuming, i.e. higher time constants, although bulk water in itself is expected to yield a more rapid diffusion. This part of the RH domain is within the swelling area which has been reported to start at 22-33 % wt [12,13,21,23].

The initial increase in absorption time constant, meaning that it took more time for water to enter the SC as RH was increased, is the opposite of the effect caused by an increasing diffusion coefficient with concentration. Rouse reported on decreasing diffusion constant (i.e. increasing time constant) with increasing concentration for polymers that were not strongly hydrophilic [24]. This effect was proposed to be due to increasing proportion of immobilized water as the water content increases, leading to water entering the polymer more slowly. It is clear that keratin filaments are not very polar as they are quite insoluble in non-denaturing buffers [25]. This relatively low polarity of keratin is further underlined by its close association to lipids *in vivo* [3]. It has also been reported on a rapid increase in water uptake at low and high RH, but with a slower uptake in the intermediate range [21].

The absorption and desorption time constants from heel SC differed substantially at RH lower than 50 % as seen in figure 7. This regime is likely to be the strongly bound water, and thus it seems reasonably clear that this type of water entered the SC much more rapid compared to its corresponding exit. The underlying mechanism for this peculiar difference is not clear other than that bound water prefers to remain bound. However, the increasing time constant with decreasing RH was as expected since the diffusion coefficient usually decreases with decreasing water content [13-15]. Thus, free and more mobile water will evaporate first and eventually bound water, being more strongly attracted will desorb.

Although the sorption time constants of heel and breast SC showed similarities with respect to curve forms, including triphasic absorption, there are clearly also different patterns which are hard to explain. The time constant of heel SC stored under freezing conditions, shown in figure 7, did not follow the pattern of a low valued time constant during desorption, as was seen for SC isolated from breast skin, indicating that freezing of samples, possibly inducing distortions to the keratin matrix, was not solely responsible for the new pattern in time constants. However, a combination of freezing effects and alterations with time during storage may have contributed to this difference, as well as to the relatively large spread in time constants for breast SC. Also, the amount of water replacers such as urea and glycerol varies with age of the subjects and is expected to have an effect on the water holding properties of human SC [26].

The peculiar effect of time between preparation of the SC samples and the onset of the DVS measurements, present mainly for RH between 40 and 70 %, may be due to structural changes in the SC when being stored *ex vivo* over time. With time one can expect the extracellular lipid matrix as well as the keratin filament matrix of the corneocytes suffer from irreversible distortions, such as sorting of lipids in the lipid matrix with crystalline domain building, or irreversible glass transition of the keratin filament matrix as consequences [27]. This effect did not seem to have much influence on the total water uptake, comparing the time constants in figure 10 and 11 with the relative water gains during absorption and desorption in figure 12 and 13, although the water holding capacities seemed to decrease slightly with time.

The statistical material in this manner is very limited with only one test sample investigated, and more effort is needed to reveal a more complete picture.

An improved understanding of the characteristics and location of the three different water phases in the SC will be of importance regarding clinical treatments and knowledge of the human skin and it is therefore to be a topic for future studies.

5. Conclusion

In this study we did not find signs of significant hysteresis for human SC, although there were small spreads in mass from absorption and desorption data. Any hysteresis of SC is important and must be accounted for in calibrations of susceptance based hydration measurements, which are robust methods for determining the hydration state of human SC.

We found the time constants to be triphasic during absorption, possibly reflecting the three different types of water in the SC, and increasing with decreasing RH during desorption. The latter may be interpreted as a protective property, providing the skin with sufficient moisture in order to retain all its protective functions, whereas the former is connected to the weak polarity of SC keratin filaments. Knowledge of these types of human SC water content and their sorption characteristics is vital for a structure-function based knowledge of SC properties.

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A memristive model of electro-osmosis in skin

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Abstract. We show that some of the non-linear conductance properties of electro-osmosis in sweat-duct capillaries may be modeled by a memristive circuit. This includes both the observed phase shift and amplitude modulation of the electrical current response to a simple harmonic driving potential. The memristor, a new type of elementary passive circuit element recently invented by the semi-conductor industry, may therefore be expected to play a role in modeling the electrical properties of skin, and perhaps also in other systems where non-linearities are observed in their bio-impedance.

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1. Introduction

It is well known that non-invasive measurements of the electrical conductance of skin, obtained by placing flat electrodes on top of the epidermis, depends on the current flowing through the skin. This may, at least in part, be due to electro-osmotic transport of water through the sweat-duct capillaries [1]. Our purpose here is to reexamine this explanation by showing that electro-osmosis can be described using a *memristive* model, whose defining property is that it has memory of the current having passed through it, whence the impedance is not constant but some non-linear function of the current history.

The *memristor* is a new elementary passive circuit element, complementing resistors, coils and capacitors, whose *memristance* $M(q) = d\varphi/dq = v/i$ is defined by how the magnetic flux $\varphi = \int v(t)dt$ responds to the charge $dq = idt$ transported by the electrical current $i(t)$. Predicted from symmetry arguments by Chua [2], it remained a theoretical curiosity for almost four decades until a team from Hewlett-Packard invented an electronic nano-memristor [3]. We want to show that the memristor concept may be useful in describing bio-physical systems, and in particular that the electro-osmotic behavior observed in human skin can be thought of as a memristive system. Such

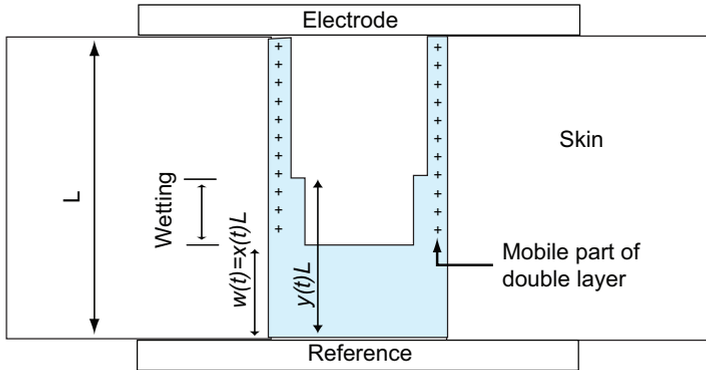


Figure 1. Schematic illustration of a sweat-duct capillary with wetting layers.

memristive properties are now believed to be rather common in very small systems (cmp. Ref. [3]). This includes the extremely tiny electronic components built from semi-conductors, but this is also the scale where the collective degrees of freedom used in bio-physics and biology start to emerge from the underlying molecular chemistry.

Memristive systems are generalizations of the memristor concept where the memristance is controlled by a number of state variables [4]. This study extends the preliminary results reported in an earlier study [5] to include two microscopic state variables, whose physical origin is explained and which explains further properties of electro-osmosis.

In dry skin most pores and capillaries are empty or only partly filled, but conductive films are believed to be present also in the otherwise empty parts of these ducts [6, 7]. With dry electrodes applied on the skin surface on the ventral forearm, an increase in conductance may occur if water is dragged by the applied electric field from deeper layers towards the surface where the electrode is located. The amount of filling in the capillary will be influenced by the electro-osmotic transport of water, as illustrated in figure 1.

When the mobile part of the conductive double layer is subjected to an externally applied electric field, the ions which are set in motion drag along the fluid column in the filled part of the duct because of viscous forces. Consequently the volume V of water in the capillary changes at a rate (flux) dV/dt , since fluid is supplied or extracted at the bottom of the duct from or to the surrounding tissue. This flux is proportional to the cross-section area $A = \pi r^2$ of the capillary, the applied electric field E , the electrokinetic potential ζ at the interface between the duct wall and the fluid and the relative permittivity ϵ , and inversely proportional to the viscosity η of the fluid [8]. Lumping the systemic parameters into one with suitable dimensions for our problem, $\alpha = \zeta\epsilon/(4\pi\eta)$, we obtain Glasstone's fundamental equation:

$$dV = \alpha AE dt \quad . \quad (1)$$

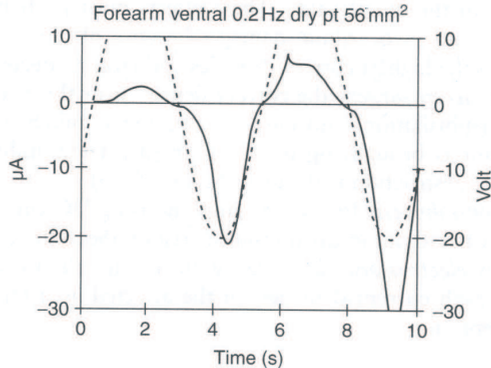


Figure 2. Current response (solid curve) and applied voltage (dashed curve) on ventral forearm using dry disc electrodes (data from Ref. [1]).

Since $E(t) = v(t)/L$, with L the total length of the capillary, and $\alpha < 0$ because $\zeta < 0$ in a sweat duct [9], we see that the volume of the water column increases when the electrode potential is negative. A rising column reduces the total resistance so we expect the current to increase in value.

A typical electrical current response measured on human skin using dry disc electrodes is shown in figure 2. As expected, the value of the current increases as the applied field changes from positive to negative polarity at the skin surface electrode, because of the electro-osmotic transport of the filled part of the ducts. A negative potential attracts the water column, whereas a positive potential depresses it.

Furthermore, the current amplitude was found to increase from one cycle in the applied periodic voltage signal to the next, described as effects of “wetting” in Ref. [1]. This effect is thought of as a build-up of fluid in the duct over a series of current cycles. As the fluid column oscillates, it deposits more water on top of the thin film covering the wall (the duct wall is “wetted”), leading to reduced duct resistivity, and hence increased current. The height $w(t)$ of the water table will be a function of the amount of charge having passed through the duct, whence we recognize this as a memristive system, i.e. a system with some “memory” of its electrical history.

2. Memristive modeling of electro-osmosis

A partial explanation of these data was proposed in Grimnes [1], by relating the observed changes in skin conductance to fluid motion through electro-osmosis. An improved understanding, and a better quantitative analysis, may be obtained by using the concept of memristance. The imittance M defined below in eq. (2) is physically distinct from ordinary bio-electric impedance since it has memory, and is therefore an example of the memristance of a memristive system defined by Chua [2, 4]. More precisely, by deriving

expressions for the charge function $q(t)$ and the corresponding current $i(t)$, we shall see that an electro-osmotic capillary behaves like a charge-controlled memristive system.

From figure 1 it is clear that an electrical current flowing through a capillary of total length L meets a total *instantaneous* (i.e., at any instant in time) resistance $M(t)$ composed of three resistances coupled in series:

$$M(t) = R(x(t), 1) + R(y(t) - x(t), \tau_F) + R(1 - y(t), \tau_f) . \quad (2)$$

Observe that not only are these “resistors” time dependent, their time evolutions are inextricably intertwined with each other.

The electrical resistance of each section of the capillary depends on the length l and thickness d of its wet volume. We normalize so that the (unobservable and microscopic) state variable is the longitudinal filling fraction $z = l/L$, while the transverse filling fraction $\tau(d)$ for a film of thickness d is the fraction of the total cross-sectional area of the capillary that is filled by the film:

$$\tau(d) = \frac{A(d)}{A(r)} = \frac{d}{r} \left(2 - \frac{d}{r}\right) \stackrel{d \ll r}{\approx} \frac{2d}{r} . \quad (3)$$

Thus $\tau(d = r) = 1$ in the full part, while $\tau \ll 1$ for both the thin film (f) of thickness $d_f \ll r$ and the fat film (F) of thickness $d_F \ll r$. We then have

$$R(z, \tau) = \frac{z(t)}{\tau(d)} R_{\bullet} , \quad (4)$$

where $R_{\bullet} = \rho L/A$ is the minimum possible total resistance, which obtains when the capillary is completely filled ($z = \tau = 1$) with our electrolyte (sweat), which has electrical resistivity ρ . The maximum possible total resistance $R_o = R_{\bullet}/\tau_f \gg R_{\bullet}$ is reached when the capillary is completely empty save the thin film coating the capillary wall. The longitudinal fractional filling of water is called $x(t) = w(t)/L$, and $y(t)$ is the fractional height reached by the thick film, cmp. figure 1.

The wetting problem is most naturally parametrized by

$$\delta = \frac{\tau_f}{\tau_F} \approx \frac{d_f}{d_F} \leq 1 , \quad (5)$$

with $\delta = 1$ when there is no wetting, i.e., $d_F = d_f$. We can use δ to express the total resistance (actually, memristance) in terms of a single dimensionless function (measured in the natural unit R_o):

$$M[R_o] = \frac{M}{R_o} = 1 - \delta x(t) + (\delta - 1)y(t) . \quad (6)$$

Note that this memristance depends on time through the two microscopic and unobservable state variables $x(t)$ and $y(t)$, and in order to progress we must now deal with these. We first trade x for a measurable quantity (charge or current) by exploiting Glasstones law.

During a small time interval dt the volume of water in the capillary changes by $dV = Adw = ALdx = V_{\bullet}dx$, where $V_{\bullet} = AL$ is the total volume of the capillary.

Another expression for dV follows from Glasstones law, eq. (1), if we assume that all of the current $i(t) = dq/dt$ passes through the water column. The potential drop

over this part of the capillary is $Ew = \rho wi/A$, which is used to eliminate the electric field strength from Glasstones law, giving $dV = \alpha \rho dq$. Equating the two expressions for dV we find the simple differential equation $dx = -c_1 dq$, where $c_1 = -\alpha \rho / V_\bullet > 0$ is a systemic constant.

This immediately integrates to $x(t) = x_0 - c_1 q(t)$, where x_0 is the initial position of the water table at $t = 0$. The sign of q is chosen so that a negative current gives the water table a positive velocity $dx/dt > 0$, as required by the experimental findings in Ref. [1]. Inserting this expression for $x(t)$ into eq. (6) gives the memristance:

$$M[R_o] = c_0(\delta) + \delta c_1 q(t) \quad , \quad (7)$$

where the new function:

$$c_0(\delta) = 1 - \delta x_0 + (\delta - 1)y > 0 \quad (8)$$

at first sight appears to depend on a time-dependent function $y(t)$. However, this is a general expression for the memristance in the duct, valid with or without wetting, and it will turn out that $y = y_0$ is a constant in the former case.

In the latter case $\delta = 1$ and y drops out. This describes the first half-cycle after a negative driving potential $v(t) = -v_0 \sin(\omega t)$ ($v_0 > 0$) is applied. This ensures that the water table that initially only is filling the lowermost parts of the sweat duct is attracted to the electrode. Electro-osmosis will, as long as the driving potential remains negative, drag the water table in the duct towards the electrode located at the skin surface [1]. When the driving potential after a time $T/2$ changes from negative to positive the water table is at its maximum height $x(T/2) = y_0$, and the memristance is at its lowest value.

With a positive potential the electro-osmotic force flips sign, pushing the water table back into the sweat duct and leaving behind a thickened film. This additional film is the wetting-effect, and now the total memristance will consist of a series of three volume conductors: two films and the decreasing water table. The total memristance is now lower than it would have been without wetting. The current flow is therefore larger, pushing the water table deeper down in the duct, beyond its starting point x_0 .

In other words, after a full period ($t = T$), the first half without wetting and the second half with, we find that $x(T) < x_0$.

If we now assume that further wetting does not take place, i.e. that the film can only have two thicknesses, then the subsequent periods are quite different since the half-cycles are now symmetric. The water table will therefore return to the same maximum height y_0 at times $t_n = nT/2$ for each odd integer n , and to its minimum value $x(T)$ at times t_n for each even positive integer n . This heuristic description will be verified analytically below.

In this model we have ignored the so-called ‘‘clipping’’ effect which is expected to happen when the driving potential is positive and the water table is repelled by the surface electrode. Then the film eventually loses contact with the electrode on top of the duct, yielding no current for a short period until the film diffuses back and remains in contact with the electrode which again repels the film, etc. This repeats as long as the potential is positive and leads to a reduced current during this interval, and can

be seen in the data reproduced in figure 2. There is therefore less current to drive the water table downwards by electro-osmosis and the table does not reach back to its lowest position where it was one period earlier. With $x(t_n) > x_0$ (n odd) the water table will reach higher up in the duct at its next maximum, giving a higher value of the current due to an increase in the wetting-effect for each period of the driving potential. This is also as reported in the original paper by Grimnes [1].

In summary, in our mathematical model, which includes wetting but not clipping, we need a negative initial potential in order to account for the observed increase in current amplitude with time. When clipping is present the current amplitude will increase for each cycle as discussed above, and a positive initial potential could have been applied.

We now look in detail at the two basic intervals of interest, where we have either 1-film or 2-film memristance.

2.1. Memristance without wetting (thin film of constant thickness)

This is the first half-period $0 < t < T/2$ after the driving potential is switched on. The memristance during this time is simply:

$$M(q) = R_o(c_0 + c_1q) \quad , \quad (9)$$

with $c_0 = c_0(1) = 1 - x_0 > 0$.

By integrating Ohms law $v(t)dt = M(q)dq$ we find the charge as a function of magnetic flux $\varphi = \int v(s)ds$:

$$c_1q(t) = -c_0 + \sqrt{c_0^2 + 2c_1\varphi(t)/R_o} \quad . \quad (10)$$

As explained in the introduction we choose a driving potential $v(t) = -v_0 \sin(2\pi t/T)$ that starts out negative ($v_0 > 0$), which gives the flux $\varphi(t) = -(v_0 T/\pi) \sin^2(\pi t/T)$. The flux-charge characteristic is shown in the inset in figure 3.

The current is given by $i = Gv$, where the *memductance* $G = 1/M$ is obtained from eq. (9) by inserting our result eq. (10) for $q(t)$:

$$M = R_o \sqrt{c_0^2 + 2c_1\varphi(t)/R_o} \quad . \quad (11)$$

2.2. Memristance with wetting (film with varying thickness)

As soon as $t > t_1 = T/2$ wetting starts to modify the total memristance of the system:

$$M(q)[R_o] = c_0(\delta) + \delta c_1q \quad , \quad (12)$$

where $\delta = d_f/d_F$ is the ratio of the thickness d_f of the thin film and the thickness d_F of the thick film left behind by wetting the thin film. We have no restrictions on δ in our model, apart from $\delta \leq 1$.

This M governs the current for all $t > t_1$, and the charge and current functions are obtained by the same argument used in the non-wetting case discussed above. We find the charge valid for $t > t_1 = T/2$

$$\delta c_1 q(t) = -c_0(\delta) + \sqrt{c_0(\delta)^2 + 2\delta c_1\varphi(t)/R_o} \quad , \quad (13)$$

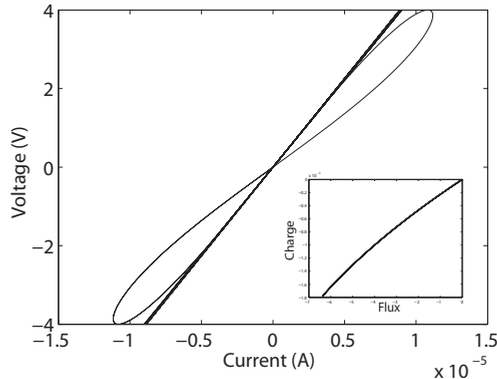


Figure 3. Current-voltage characteristic of the memristive model deduced for skin sweat ducts, shown for $\omega = 2\pi/T = 0.2$ Hz and $\omega = 5$ Hz. Inset: Charge-flux characteristic of the memristor.

where the $c_0(\delta)$ -term has been updated by the fractional height of the thick film is $y = x_{\max} = x(t_1) = x_0 - c_1 q(t_1)$ in order to include also the effects of wetting.

The current is given by $i = Gv$, where the memductance $G = 1/M$ is obtained from eq. (12) by inserting our result eq. (13) for $q(t)$:

$$i(t) = v(t)/R_0 \sqrt{c_0(\delta)^2 + 2\delta c_1 \varphi(t)/R_0} . \quad (14)$$

It is now easy to verify that $q(t_3 = 3T/2) = q(t_1)$, from which it follows that $x(t)$ returns to the same value for all later times $t_k = (2k + 1)T/2$ with $k = 0, 1, 2, \dots$, as claimed above. The expression for $i(t)$ deduced from $M(q)$ is therefore valid for all $t > T/2$.

Figure 3 shows this current plotted against the potential $v = -v_0 \sin(\omega t)$, with $\omega = 2\pi/T$ and $\delta = 0.5$. For a suitable choice of parameters hysteresis in the iv -characteristic, which is the fingerprint of memristance, is in evidence at low frequencies, with curves running through the origin in double loops. At higher frequencies, the iv -characteristic degenerates to a straight line, i.e., the memristor behaves like an ordinary resistor.

The inset in figure 3 shows the charge-flux characteristic with the harmonic driving potential, for which the flux is $\varphi(t) = -(2v_0/\omega) \sin^2(\omega t/2)$. Evidently the $q\varphi$ -characteristic is single valued, as it must be for charge-controlled memristive systems [2].

A smaller δ will result in a reduced memristance M and an increased current $i(t)$. If $\delta = 1$, then there is only one single film and no wetting at all, meaning no increase in the current amplitude. Figure 4 illustrates how the current in the sweat duct given by our memristive model behaves for a suitable choice of system parameters, for some possible thicknesses of the wetting layer. The current response is similar to the behavior

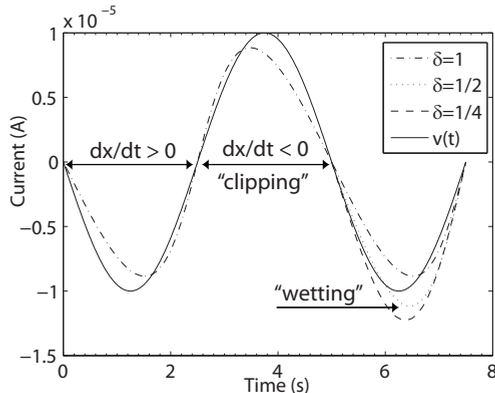


Figure 4. Scaled applied voltage (solid) and current responses for different thicknesses of the wetting layer.

seen in the data displayed in figure 2. The current amplitude increases with wetting when the applied potential is negative, and the phase shift is positive so that the current lags behind the driving potential. The positive current amplitude is larger in our model than seen in figure 2 since the depressing clipping-effect is not accounted for.

3. Discussion

We have demonstrated that an improved understanding of skin conductance can be obtained by using the memristive paradigm defined by Chua, which goes beyond the traditional “resistive (RLC) framework”. This shows that that the memristor may be a suitable circuit component for modeling bio-physical systems whose electrical parameters depend on their (electrical) history, in particular when hysteresis and phase-shifts are present. The expression for the memristance of sweat duct capillaries obtained in eq. (7) clearly exhibits the explicit charge dependence of this “device”, showing that it is memristive rather than resistive.

A closer examination of the memristance M is instructive. Resurrecting the dependence on systemic parameters we can write eq. (7) as:

$$M[R_o] = c_0(\delta) - \frac{\alpha\delta R_\bullet}{L^2} q(t) , \quad (15)$$

which exhibits how M depends on the scale L of the “device”. It is clear that M will only deviate significantly from a resistance when L is small, as the charge-dependence is suppressed by L^2 . Since there are many bio-physical systems, including skin capillaries, that are very small, we suggest that memristance may be useful for explaining a variety of non-linearities seen in bio-physics and bio-electricity, effects that in many cases are not yet fully understood.

The frequency band where memristive properties are seen will in general depend on the “speed” of the system, that is how rapid the water table responds to changes in the driving potential. High frequencies will depress the memristive character of the system, since the liquid column has inertia and therefore needs time to respond to the applied field.

Figure 3 shows the typical feature of memristors when studied in the $q\phi$ plane, giving a single-valued and non-linear characteristic. This agrees with Ref. [3], which also reported a non-linear $q\phi$ -characteristic in the semi-conductor nano-memristor that they invented.

The effect of “clipping” has been left out of our model due to its complexity, but it should probably be taken into account in order to explain the continued increase in current amplitude over several cycles observed in Grimnes [1]. The net effect of clipping is a reduced average current, which is actually a highly jagged current smoothed out at the small frequency ($\omega = 0.2\text{Hz}$) used in the experiment shown in figure 2. At higher frequencies the resolution improves, and at $\omega = 500\text{Hz}$ the current curve appears to be ragged around the peaks, but not elsewhere [1]. While his observation supports the clipping-explanation, including clipping in our model would increase its non-linearity substantially, beyond our ability to analyze the model properly.

Figure 4 shows how the value of the wetting parameter $0 < \delta = d_f/d_F \leq 1$ determines the amount of current passing through the capillary duct. Since the thickness of the wetting-film is unknown it is hard to reach a definite conclusion about the importance of the wetting-mechanism in real systems. A reasonable estimate is perhaps that wetting doubles the film thickness ($\delta \approx 0.5$), in which case it should not be neglected.

Another uncertain aspect of our model is that we assume that the wetting film remains more or less constant in height y during a cycle, i.e., that diffusion processes do not have sufficient time to smear out the film.

Our model does not so far include electrical activity in the epidermal stratum corneum itself, which usually acts in parallel to the sweat ducts [9]. Some of the activity in the stratum corneum is conventionally modeled using a theoretical device called “the constant phase element (CPE)”. This is not a real device which can be built from a finite number of conventional (RLC) passive circuit elements [9, 10]. It would be of great interest to study how the new memristive circuit element can be used in this type of more sophisticated modeling.

4. Conclusion

We have shown that the fourth fundamental circuit element, discovered theoretically by Chua but “missing” in real life until HP invented such a nano-device recently, can be used in modeling electro-osmotic properties of capillaries in skin. The memristive paradigm should be useful in describing other non-linear phenomena that appear at small scales. Such phenomena have frequently been observed in electronics, physics and

biological systems, but have often remained unexplained. Memristors clarify and enable circuit modeling of non-linear systems, and therefore promise to be useful in circuit modeling of bio-physical mechanisms, particularly bio-electrics.

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A study on electrode gels for skin conductance measurements

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Abstract

Low-frequency skin conductance is used within several clinical applications and is mainly sensitive to sweating and the moisture content of the stratum corneum (SC), but also how electrodes introduce changes in the electrical properties. Four electrode-gels were investigated with regard to sorption characteristics and electrical properties. Skin conductance time-series were collected from 18 test subjects during relaxation, exercise and recovery, wearing different pairs of electrodes contralaterally on the hypothenar and the T9 dermatome. Pressure test was applied on the T9 electrodes. Impedance frequency sweeps were taken on the T9 electrodes the same day and the next, parameterized to the Cole model. ANOVA on the initial skin conductance level change, exercise response amplitude, recovery offset and pressure-induced changes revealed significant differences among gel types. The wetter gels caused a higher positive level change, a greater response amplitude, larger recovery offset and greater pressure-induced artifacts compared to the solid gels. Sweating on the T9 site lead to negative skin conductance responses for the wetter gels. Correlations were found between the desorption measurements and the initial skin conductance level change (hypothenar: $R=0.988$ T9: $R=0.901$) RM-ANOVA on the Cole parameters revealed significant decrease in R_s of the most resistive gel. Clinical implications are discussed.

PACS codes:

82.45.Fk	Electrodes
84.37.+q	Measurements in electric variables (including voltage, current, resistance, capacitance, inductance, impedance, and admittance, etc.)
84.37.+q	Electric impedance measurement
87.19.R-	Electrical properties of tissues

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Keywords: Sweat, skin conductance, electrodes, water sorption, stratum corneum, electrodermal activity, sweat glands

1. Introduction

The electrical admittance of the skin is dependent on several factors. The epidermal Stratum Corneum (SC) layer, consisting of mainly dead skin cells, gives a large impedance in series with the viable skin, dominating the measurement at low frequencies (below 10kHz) (Martinsen et al 1999). The low-frequency skin admittance measurement then becomes highly sensitive to mechanisms occurring in this layer, mainly the ionic shunting provided by the sweat ducts as they are filled with sweat and the absorption of water in the SC. The admittance contribution from the sweat duct is predominantly conductive (Martinsen et al 1998), while the SC moisture content also significantly influences the low-frequency capacitive part of the admittance (Martinsen et al 1995, 1998, Martinsen and Grimnes 2001). The high sensitivity of the skin conductance to sweat activity has opened up for several clinical applications.

Due to the link between palmoplantar sweating and the sympathetic nervous system, this parameter has found its use in psychophysiology as a tool for stress assessment (Healey and Picard 2005, Setz et al 2010), assessment of psychiatric disorders (Iacono et al 1999, Nilsson et al 2006) among other uses, often in combination with other

physiological parameters. Within anesthesia, it has been proposed as a tool in pain assessment (Ledowski et al 2007, Hullett et al 2009). Skin conductance at other skin sites has been proposed for the diagnosis and treatment evaluation of hyperhidrosis (Tronstad et al 2008), and may be a promising tool for diabetes (Hoeldtke et al 2001), cystic fibrosis (Quinton 2007) and various neurological disorders.

It is known that the choice of electrodes for electrical measurements will heavily influence the signals that are measured (McAdams et al 1996, Mayotte et al 1994 and Rahal et al 2009), and so it is important to find the best suited electrode for the intended use. Depending on how the measurements are interpreted, artifacts introduced by the electrodes could lead to incorrect diagnostic assessments.

Studies of electrode properties have previously been carried out in different ways (Rahal et al 2009, Egging 1993), but not with focus on monopolar skin conductance measurements combined with an *in vitro* sorption study of the water holding properties of the electrode gels. In addition, the authors are not aware of any studies where parameters from skin conductance time-series have been compared between different electrode gel types recorded simultaneously on the same test subjects.

In this study we investigated four different types of electrode gels for skin conductance measurements. As a low frequency monopolar set-up will focus mainly on the SC, its general hydration state, and sweat filling in particular, will be very dominant for what is being measured. SC hydration may be affected by occlusive effects, but also through a direct and sometimes rapid hydration from the electrode gel. We compared the amount of free water in the electrode gels *in vitro* with the results from electrical *in vivo* measurements with the aim of finding how different electrode types will affect the skin conductance recordings. In particular, we wanted to investigate the stability of the skin conductance level, the response to sweating stimuli, the post-stimulus recovery and the susceptibility to mechanical artifacts. As both SC properties and innervation differ according to skin site, especially on palmoplantar skin compared to non-palmoplantar skin (Kuno 1956), we included the T9 dermatome on the abdomen in addition to the hypothenar site on the palm for this study.

2. Materials and methods

2.1. Electrode types

Four electrode types were investigated, all with similar Ag/AgCl metallic part, but different composition of the gel. The *a priori* gel properties of the different electrodes are listed in table 1.

Table 1. Overview of the electrode gel types examined in this study according to gel material, the intended use, effective electrode area (EEA) and type of adhesion.

Type	Gel material	Intended use	EEA [cm ²]	Adhesion
A	Solid hydrogel	ECG, neonatal	5.05	Adhesive gel
B	Wel gel	ECG, short-term	2.54	Outer adhesive part
C	Isotonic EDA jelly	EDA	1.95	Outer adhesive part
D	Karaya solid gel	ECG, hypoallergenic	6.16	Adhesive gel

The brands of the electrode types were as follows:

Type A: Kendall™ KittyCat™ 1050NPSM

Type B: Ambu® Blue Sensor Q-00-A

Type C: Discount Disposables, model TD-246 Skin Resistance – Skin Conductance Electrode Paste

Type D: Kendall™ Arbo™ H83V

2.2. *In vitro* sorption characteristics

The sorption characteristics of the four different electrode gels were obtained using a gravimetric method where water loss and water gain could be monitored carefully. A dynamic vapor sorption instrument (DVS) from the Surface Measurements Systems (London, UK) was used in this purpose. This instrument enables very sensitive gravimetric measurements that can be performed under controlled environmental conditions, ensuring high-precision and reproducible results. The mass-resolution of the instrument is 0.1 µg, and the gel-samples in our

study had a mass in the range of 4-10 mg at measurement onset. This means that the sample masses during the entire sorption cycle were well above the resolution of the instrument, resulting in smooth time series. The gel-samples were loaded into a metal pan in a closed chamber where temperature and relative humidity (RH) was pre-programmed and carefully monitored. The two solid-gel samples, A and D, were prepared by means of sterile scalpels to be of approximately the same size and geometrical shape (cubics). The wet-gel B was scratched off the attach-surface of the electrode and placed in the metal pan as a small droplet.

Finally, the cream-based gel C was loaded into the pan by gently squeezing the bottle containing the cream. The samples were measured one at the time and all four measurements were initiated immediately after sample preparation was finished. The RH was pre-programmed to increase in steps of 10%, from 0% up to 90% and then back to 0%. The temperature was kept stable during the entire measurement at 25°C. The mass reading found at 0% RH was taken to be the dry mass of the sample. A step in RH was initiated when the mass rate, dm/dt , was less than 20 ppm for a period of at least 10 min, ensuring the samples to reach their equilibrium state at every level of RH.

The characteristic time constants, which yield the time needed for the sample mass to step approximately 63% of the entire increase needed to reach its stable state value, were obtained by fitting the time courses for each increment in RH to $y = a \exp(bt) + c$. The time constants are then given by $\tau = b^{-1}$.

2.3. *In vitro* electrical characteristics

Electrical impedance frequency sweeps were taken on fresh pairs of each electrode type attached front to front using a Solartron 1260+1294 setup with a constant voltage excitation of 30 mV rms and a frequency range from 1Hz to 100 kHz.

2.4. *Skin conductance measurement*

The three-electrode system (Grimnes 1983b and Grimnes et al 2009) was used to obtain unipolar measurements of the skin surface conductance density (SSCD) of the area below the effective electrode area of the measuring electrode. AC measurement was used to avoid polarizing the skin or electrodes and to avoid contribution from endogenous potentials in the skin. For single-frequency SSCD time-series recording in four channels, the Sudologger (Tronstad et al 2008) was used. Impedance frequency sweeps from 1 Hz to 100 kHz were taken using the Solartron 1260+1294 equipment. Constant voltage excitation of 30 mV RMS magnitude was used for both equipments to keep the current density low and the measurement within the linear range.

2.5. *Test subjects and protocol*

18 (14M, 4F) healthy volunteers who gave informed consent were each assigned to two of the four electrode types for contralateral comparison of measured SSCD parameters. Initially, the test subject (TS) was asked to sit down in a chair while measuring electrodes were placed simultaneously on the left and right hypothenar site of the palms and about 4cm to the left and right of a point slightly above the navel on the T9 dermatome. The measuring electrodes on each side of the body were of the same type and the selection was based on a matrix of all six possible pairwise combinations of the four types.

After electrode fixation, the SC was recorded for 10 minutes while the TS was sitting still in the chair. During the last four of these minutes, the TS was asked to remain quiet and focus on relaxing in order to obtain a baseline. At the 10th minute, a loud sound (2.5s sound sample of a glass breaking played by a set of speakers at an intensity measured to be 82dB 1m in front) was played to induce a startle response and the TS was asked to stand up and perform squats for two minutes to induce thermal sweating. This was followed by 15 minutes of sitting relaxation until a test for pressure artifacts on the abdomen electrodes was performed by an operator. The TS was asked to relax the diaphragm while the operator pushed the electrodes equally and simultaneously inwardly on the abdomen for 10 seconds, pushing as far as to the point where the tissue would no longer compress easily. This was followed by two minutes of recording to check whether any SSCD offset would remain or recover to the previous level.

The palmar electrodes were then removed and electrical impedance frequency sweeps were taken on both of the abdomen electrodes. The TS was then wearing these electrodes until returning the next day, approximately 24 hours later, for a control measurement. For the prewired electrodes, the wire was taped to the skin in order to

avoid pull on the electrode. The TS was instructed to pursue normal activities and not to give the electrodes any consideration except for not showering the abdomen area. For all the electrodes which were still attached on the next day, new impedance sweeps were taken both with the electrodes as they were and after a refixation of the electrodes by gentle pressure on the adhesive area. After gentle removal of the electrodes, photographs of the skin areas were taken. The pictures were evaluated by a dermatologist for signs of skin irritation caused by the electrode gel.

2.6. Data analysis

Sudollogger measurements were scaled to a unit of $\mu\text{S}/\text{cm}^2$ (SSCD) for comparison between electrodes of different effective electrode area. For the SSCD time-series, the following parameters were extracted:

- Level change during the initial 10 minutes after electrode-skin contact.
- Amplitude of the SSCD change during the 2 min exercise interval, calculated as the maximum deviation from the prestimulus baseline.
- Offset between the prestimulus baseline and the level reached after the 15 minutes of recovery following the exercise interval. In some cases there were spontaneous responses during the end of the recovery, hence the minimum value within the last two recovery minutes was subtracted by the prestimulus baseline to obtain this parameter.
- Change induced by the pressure test on the electrodes, calculated as the maximum deviation from the prestimulus baseline during an interval of the applied pressure time + 5 s.
- Offset between the level two minutes after the pressure test and the level before the applied pressure.

Separate but equal statistical analyses were done for the hypothenar and abdomen measurements. ANOVA was done on all the parameters. Test for normality was done using the Shapiro-Wilk test and the Kruskal-Wallis ANOVA on ranks was used when the test for normality test or equal variance did not pass. For pairwise multiple comparison, the Holm-Sidak t-test was used when the normality test passed and the Student-Neuman-Keuls test was applied otherwise. For cases of missing values, the Dunn's Test was used instead.

For the Solartron impedance frequency sweeps, the measurements were parameterized using a Cole system electrical equivalent model (Grimnes and Martinsen 2005). Using the curve-fitting tool in the ZView 3.0a software (Scribner Associates Inc.), the Cole-parameters R_p , R_s , Y_{cpe} and alpha were determined with a median accuracy of 1.77% fit error. In a few cases with poor electrode-skin contact on day 2, the impedance measurement was inaccurate leading to a poor fit. The parameters with an error larger than 10% were excluded from the analysis. Repeated measures ANOVA was done on each of the Cole parameters, grouped into the day 1 measurements and both the day 2 measurements. The same tests as for the time-series parameters were applied for pairwise multiple comparison.

For comparison between the wetness of the gels and the skin conductance measurements, Pearson product moment correlation coefficients and p-values were calculated between the initial relative mass change of the gels in the DVS at 0% RH and the median initial level change of the SSCD time-series.

3. Results

3.1. In-vitro results

Figure 1 shows the relative water uptake of the four different gels as the RH was altered in steps from 0% up to 90%. The mass value at 0% is taken as the dry mass of the gel, meaning that the initial values at the far left in the figure represent the relative amount of water in the gel as it will be before any electrode to skin attachment in the electrical measurements. The wet gel in our study, the type B gel, contained the highest amount of water initially, almost doubled compared to the cream based C gel, whereas type A and D were very similar, with little amounts of free water to be depleted into the environments. The types B and C gels did not regain their initial water content, even at 90 % RH, although B took up an amount of water corresponding to several times its own dry weight. The A and D gel, on the other hand, more than doubled their weight during the absorption process.

There were few signs of hysteresis in our measurements, which would correspond to a gap between absorption and desorption steady-state mass values at a given value of RH. Only for the B gel there was hysteresis in a range where the RH value was between 60 % and 80 %, where the gel remained the more of its water content during desorption compared to absorption. The time courses following a step in the RH were found to closely follow an exponential curve which is in accordance with Fick's law of diffusion.

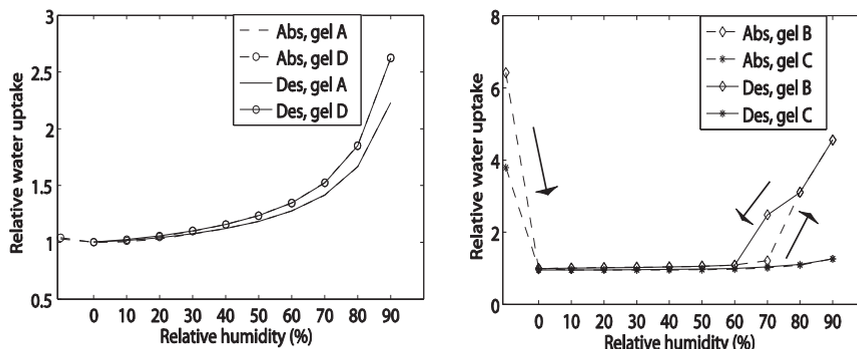


Figure 1. Relative water uptake of the four gels tested in our study. The arrows in the figure to the right indicate the direction of the sorption process, both initially after sample placement in the DVS chamber as well as in the hysteresis.

The characteristic time constants for water entering and leaving the gels are shown in figure 2. The values to the far left in the graphs are the typical values for how rapid the gels, starting from their initial (manufactured) hydration state, will reach equilibrium with the dry air in the DVS chamber at 0% RH. The smaller this value was, the faster the gel deposited water into its environment. Again, the A and D gels showed very similar patterns, both during absorption and desorption. The B gel gained and released water very rapidly for almost all RH values, whereas the cream based C gel gained water with a markedly decreasing rate as the RH was increased. During desorption the C gel needed long time to reach its equilibrium state for each decrement in RH.

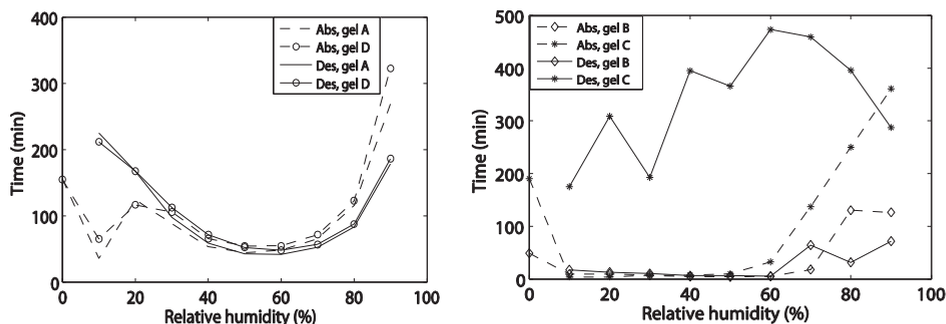


Figure 2. Time constants for the four different gels tested in our study. The points to the far left in the graphs illustrate the characteristic time for the gels to reach equilibrium with the dry chamber, starting from their initial hydration state.

The in vitro electrical measurements of the gels revealed a higher AC resistance of gel type D than the others, represented in table 2 at 1 Hz and 100 kHz for a single electrode by halving the measured resistance. These measurements are sensitive to the dispersions from both the metal-gel polarization impedance and the gel, and would not parameterize with a good fit according to a single Cole system.

Table 2. Front to front AC resistance [Ω] at low and high frequencies for all the gel types.

Frequency	A	B	C	D
1 Hz	92.6	195.9	156.4	4870
100 kHz	48.7	24.2	41.0	177.6

3.2. In-vivo results

The ANOVA revealed significant differences among the electrode types for all the time-series parameters. Figure 3 a) and b) shows that on both the hypothenar and T9 sites, electrode type B had the highest change in SSCD after electrode fixation followed by electrode C. Type A and D had lower and roughly equal medians, but with a slightly lower interquartile range (IQR) for electrode A. Figure 3 c) reveals that on the hypothenar site, electrode type B gave a significantly higher response amplitude than the other types. This is also true for the T9 site, however the type B response is negative in this case. The ability to recover towards baseline after a period of sweating is presented in figure 3 e) and f). Again, we see the highest offset for type B on both sites, while type A has the lowest median/mean offset and IQR/variance. A difference between the sites is seen on the type C offset, with clearly lower ability to recover on the T9 site than on the hypothenar site.

Correlations were found between the initial change in SSCD (Fig 3 a) and b)) and the initial relative mass change in the DVS at 0% RH with an $R=0.988$, $p<0.05$ for the hypothenar and $R=0.901$, $p<0.1$ for the T9 site.

Significant differences were also found in the susceptibility of the electrode types to pressure artifacts. As figure 4 a) and b) shows, types B and C were more affected by the pressure test than types A and D. Again, type A scored the lowest offset and IQR with type D not far behind.

The differences between the SSCD measurements from different electrode types were most clearly seen on the recordings using certain pairs of electrode types simultaneously on the same subject. Typical examples of these findings are presented in figure 5. The tendency of types B and C to initially increase the SSCD was absent for types A and D in the same subjects. Figure 5 b) displays how type B can feature a *negative* response to sweating on the T9 site while type A at the same time expectedly gives a positive response on the same site. In figure 5 a) and c), the most visible difference between the electrode types on the hypothenar is in the long-term changes of the SSCD level where type A seem to be more stable than type B and D. However, there were also some differences in the spontaneous SSCD responses recorded, where a few responses could be seen in type B which were absent or vanishingly small in type A. Figure 5 a) and d) shows the tendency of either type B or C to remain elevated after the sweating compared to type A or D. An intraindividual comparison of type B and C is shown in figure 6, exhibiting a special case on the T9 site where type B and C initially drift apart, converges and overlaps during the exercise interval and proceeds to drift apart again.

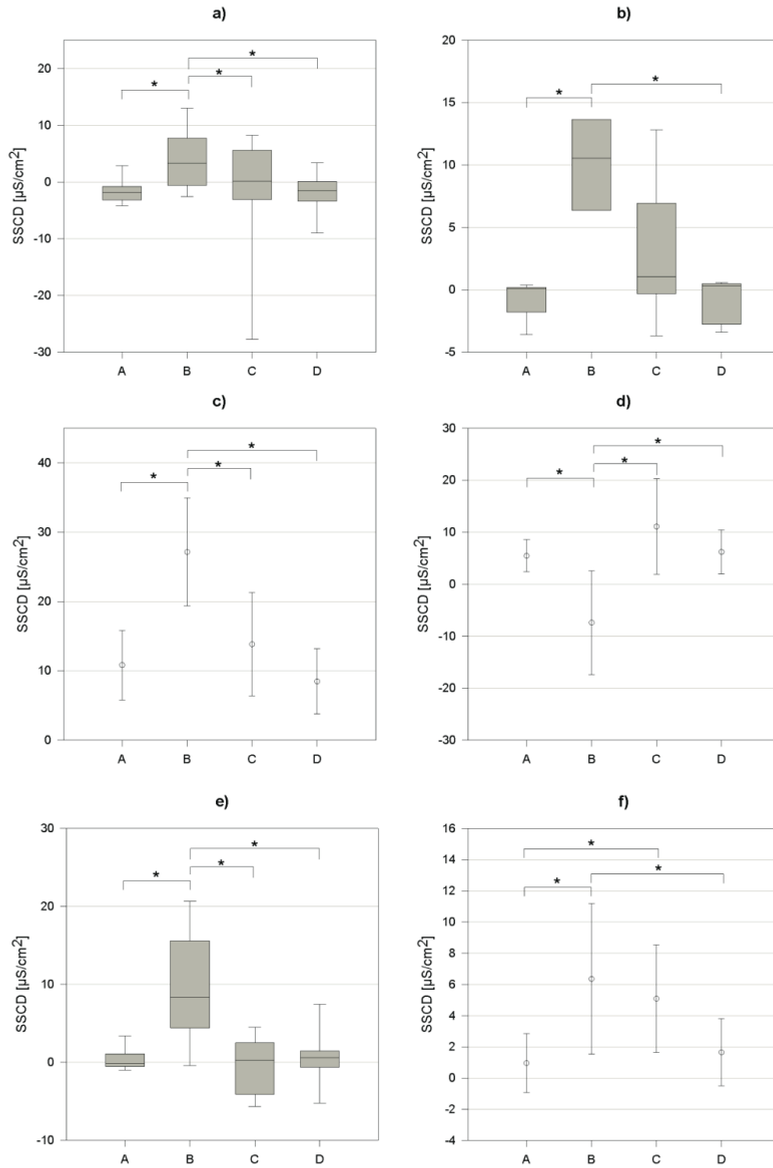


Figure 3. Time-series parameters from the hypothenar (left) and the abdomen (right). a) and b) initial difference in SSCD after 10 minutes of relaxation. c) and d) maximum change from baseline during sweating stimulus. e) and f) offset from baseline at 15 minutes after end of the sweating stimulus. * $p < 0.05$.

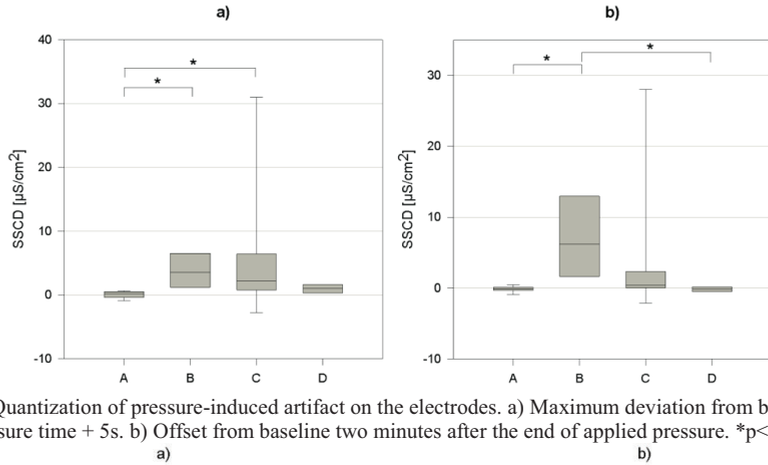


Figure 4. Quantization of pressure-induced artifact on the electrodes. a) Maximum deviation from baseline during pressure time + 5s. b) Offset from baseline two minutes after the end of applied pressure. * $p < 0.05$

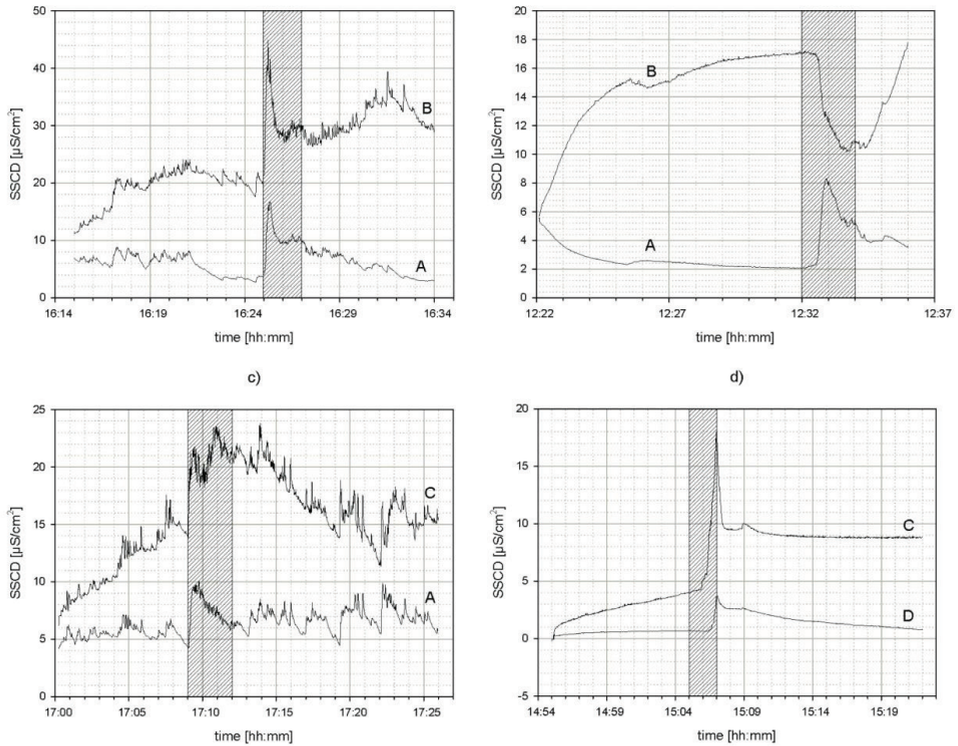


Figure 5. Examples of features found with simultaneous recordings using different electrode types on the same subjects. a) hypothenar recording using types A and B. b) T9 recording using types A and B. c) hypothenar recording using types A and C. d) T9 recording using types A and C.

and C. d) T9 recording using types C and D. The hatched area indicates the 2 min exercise interval.

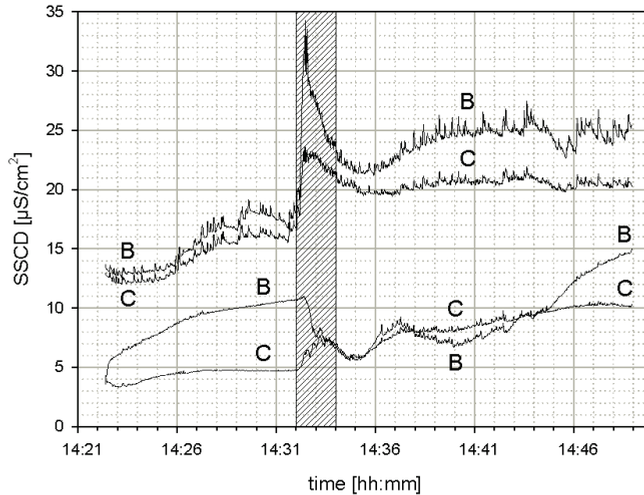


Figure 6. Example measurement from one test subject wearing electrode types B and C on the hypothenar (upper plots) and the T9 site. The hatched area indicates the 2 min exercise interval.

For the Cole-parameters from the impedance frequency sweeps, the ANOVA revealed no significant differences between day 1 and day 2 except for three cases:

- The Y_{cpe} for electrode A was significantly higher on day 1 than on day 2, also after refixation of the electrode with a difference in means of 77 nS.
- The R_s for electrode D was significantly higher on day 1 than on day 2, also after refixation of the electrode with a difference in means of 212 Ω .
- The alpha parameter for electrode D was significantly lower on day 1 than on day 2, also after refixation of the electrode with a difference in means of 0.072.

From day 1 to the control on day 2, one of type A, two of type B, and two of type D electrodes had fallen off the skin of the TS. This should not be interpreted as a result on adhesion properties, as the electrodes are constructed differently in this regard. In three of the type C electrodes, the gel chamber was completely void of gel, evident after removal of the electrode and corresponding to very high impedance values which were not included in the Cole-parameter assessment.

The dermatologist assessment of the pictures revealed no significant signs of skin irritation. There were some signs of very faint erythema, but its degree was always lower than on the areas which had been taped.

4. Discussion

In this paper we have shown differences in both the *in vitro* properties and the *in vivo* electrical measurements of the gels, which are discussed below.

4.1. *In-vitro* evaporation

Figure 1 shows that the gels we tested in our study contained very different amounts of water free to evaporate, and this means that they also likely will hydrate the SC very differently after electrode onset. However, the amount of water released from the gels initially in the *in vitro* sorption measurements is not necessarily the same

as the amount of water entering the SC from the gel after electrode onset. The transport of water to or from the gels is in general determined by the difference in partial pressures of water between the gel and the material it is in contact with. The water in the SC will in general have a different partial pressure than the air initially in the DVS measuring chamber. Also, other transport mechanisms such as skin occlusion and osmotic transport of water to or from the electrode gel, due to differences in electrolytic concentrations between the SC and the gel, can be expected to alter the skin conductance. The type B and C gels contained large amounts of water, whereas A and D barely lost any water to the dry environments at 0% RH in the DVS measuring chamber. This is correlated with the results from the skin conductance measurements, especially in how the SSCD changes initially after the application of the electrode. This indicates that the moisture content and viscosity of the electrode gel has a major impact on the quality of the skin conductance recordings. A too wet gel such as type B will bring moisture to the SC or penetrate sweat ducts if the viscosity is low enough, introducing errors in the skin conductance measurement. For electrode A and D the lack of initial free water to hydrate the SC, opens for the possibility that the SC was depleted of moisture for a short period of time after electrode onset, depending on the ionic concentration of the hydrogel, by means of osmotic transport of water into the gel. It is seen from figure 3 a) and b) that skin conductance actually decreased during the 10 first minutes of relaxation. However, a slight reduction in skin conductance level due to a decrease in sweat activity is to be expected as the test subject is sitting down and relaxing. Also, the relatively cold gel may cause contractions in skin pores, resulting in a small decrement in sensed conductance values (Eggsins 1993). Finally, the sorption results showed that the type A and D gel had very similar in vitro patterns, both regarding the relative water uptake as well as how rapid water entered or left the gel samples, supporting our findings from the skin conductance time-series.

4.2. In-vitro sorption time constants

Both the amount of water free to evaporate from the gel as well as how rapid such a diffusion process takes place can be expected to influence the skin conductance values. From figure 2 we see that the type B gel deposited its free water more rapidly into the dry DVS chamber than the other gels. A corresponding effect was seen in our electrical time-series measurements, e.g. figure 5b). The type C gel, although containing high amounts of free water initially, gave lower initial skin conductance level change (figure 3 a), b)) than gel B. This effect is probably a combination of less free water, but also a larger time constant initially for this water to evaporate from the gel into the dry chamber air. Depending on the initial hydration state of the SC, a similar trend can therefore be expected to be seen also when water diffuses from the gel into the SC. The evaporation of water from a sample, and hence the time constant, is depending on the geometry and size of the sample (Crank and Park 1968). However, as the four samples were of relatively equal size and mass, the patterns in time constants seen in figure 2 can be expected to give a representative measure of the hydration characteristics of the gels. Although gels can be expected to hydrate the SC giving increased conductance levels after onset, the water transport may potentially also go from the SC to the gel, especially during a sweat response when the moisture content locally in and around the sweat ducts increase significantly. In that sense the gel with the lowest absorption time constant (gel B) will pick up this water most rapidly, whereas gel C will not be able to gain much water. Such effects are probably slow, and only significant in long time monitoring.

4.3. Inverse sweat response

From figure 3 c) all electrode types gave increased SSCD values at the hypothenar sites during the sweating stimulus, and electrode type B, the wettest gel, gave the highest response. However, for the T9 site, electrode B gave a significant *decrease* in SSCD during the sweat response initiated by the squats. It is our belief that this inversed sweat response is due to the effect of electrode gel having penetrated down into the sweat duct. As the conductivity of the wet gel of the type B electrode is much higher (roughly 200 $\mu\text{S}/\text{cm}$) than the sweat electrolyte solution (5.56 $\mu\text{S}/\text{cm}$, Licht et al 1957), a penetration of gel down into the ducts, that are empty or partly filled (Grimnes 1983a), will result in an increased conductivity. On the other hand, during a sweat response, the higher conducting gel will be substituted with a lower conducting sweat liquid causing a decrease in the measured skin conductance (figure 5b, figure 6). Figure 6 shows a special case where the type B and C SSCD initially drift apart, but lie on top of each other for a while during and after the exercise interval. Within the overlapping period, there are equal proportions of sweat and gel in the ducts below both electrodes, yielding equal SSCD. The

question remains why the inverse sweat response occurred on the T9 site, but not on the hypothenar site. The dependent factor may be the thickness of the SC, being several times thicker on the palmar skin than the abdominal skin (Montagna and Parakkal 1974, Ya-Xian et al 1999). Comparing figure 3 a) and b), we see that the initial SSCD change on the T9 site was more than three times as large as the change on the hypothenar. This points to the electrical shunting effect caused by gel penetrating the sweat ducts being dependent on the SC thickness and thereby the sweat duct length within the SC. It is likely that more complete shunts will occur in thinner SC than in thicker SC where partly shunting will contribute a lot less to the measurement, due to the main restricting factor to the skin conductance being the SC. Thus, a larger proportion of ducts completely shunted by the gel at the T9 site may explain why sweating induces a negative response on this site contrary to the hypothenar where the thicker SC yields a larger proportion of partly shunted ducts.

The partly shunting of the hypothenar ducts may also explain the higher SSCD amplitude of type B (fig 3 c) by a sensitivity to a greater number of sweat glands where the sweat is not pumped all the way to the orifice of the duct, which also explains why some responses are present for this type but absent for a solid hydrogel type (fig 5 a)). This could not be due to differences in EEA as type B had the smallest EEA of all the gels.

4.4. Post-sweating recovery

After the end of the exercise epoch, the SSCD is expected to return to a level close to baseline within a few minutes as the sweat activity diminishes. As seen in figure 3 e) and f), there is still a large positive offset after 15 minutes of relaxation for electrode type B on both sites, and type C on the T9 site. We believe that this can also be attributed to the penetration of gel into the sweat ducts along with the sweat reabsorption process in the ducts or glands. As sweat is reabsorbed in the glands following the sweating period, gel is again allowed to enter the ducts, perhaps even actively transported through the ductal reabsorption process. This may explain why type B exhibits a positive offset even though the response during the exercise was in the negative direction. Another plausible explanation for the recovery offsets, also attributal to the solid gels, is the absorption of sweat in the SC following the sweat emergence, leading to a change in the total conductance. A third plausibility, concerning only the wetter gels, is that there may have been cases of a constant wetting of the SC during the whole recording period, leading to a somewhat steady increase in level as with type B in figure 5 a) and figure 6.

These hypotheses are also supported by the lower variance of the solid gel types in all the results presented in figure 5, suggesting that the artifacts introduced by the wet-gels depend on intersubjective properties of the SC such as its thickness.

4.5. Pressure artifacts

The pressure artifacts showed the same pairwise pattern as the sorption results with electrode A and D very similar and relatively unaffected, but with large effects on electrode B and C. The two latter electrode types have a gel that is believed to have a much less mechanical stability and are thus sensitive to applied pressures, smearing the gel giving both better contact and possibly also an increased effective electrode area. Due to its low viscosity gel B may actively be pushed into the sweat ducts, which in that case would explain the significantly elevated level in skin conductance also after the final relaxation period.

4.6. Changes in Cole-parameters from day 1 to day 2

Although the repeated measures ANOVA resulted in few significant changes in the Cole parameters from day 1 to day 2, the test may have suffered from unapparent adhesion changes of the effective electrode areas. Even though the electrodes were gently re-fixed for a second measurement on day 2, persistent poor adhesion could alter the impedance measurements. This, along with a lowering of the number of samples due to electrodes having fallen off, may have weakened the ability of the test to find significant changes in skin impedance changes caused by the electrodes. We believe that the significant change found with type D, evident in both the Rs and the alpha parameter, could be attributed to a skin reaction occurring below the SC, but can also be explained by a lowering of the gel series resistance, as type D had a relatively high front to front resistance compared to the other gels. The latter case is also supported by the dermatologist assessment of skin reaction, where no sign of any reaction beyond very faint erythema was found. Thus, it is likely that this gel may cause a drift in long-term skin conductance measurements because of the initial low-frequency resistance of type D which is relatively high

compared to the SC resistance during sweating. This high gel resistance may also explain the slightly lower SSCD response on the hypothenar for this type compared to the other solid gel type A (fig 3 c). Another effect that may have been responsible for the changes in the type D gel is its ability to gain an amount of water corresponding to about 2.5 times its own dry weight under humid conditions, as seen in figure 1. During a 24h electrode placement, water is building up underneath, giving a much increased water content locally beneath the electrode. Some of this water may potentially have been taken up by the gel, which surely would have resulted in a much reduced series resistance. However as such a water uptake typically has a characteristic time constant of a few hours, this effect is not seen initially, but will be present in long-time measurements.

The three (33%) cases of empty gel chambers on day 2 for type C may be due to absorption of all its moisture in the skin corresponding to its low in-vitro ability to reuptake moisture.

The decrease in Y_{cpe} from day 1 to day 2 may have been due to a net reduction in the effective electrode area due to only a partly electrode contact with the SC. Thus, a reduced number of ducts in contact with the gel would result in a net reduced low frequency admittance as a reduction of electrode area would probably also yield less water build-up in the SC due to occlusion.

4.7. Clinical implications

The inverse sweat response (figure 3 d, figure 5 b) can obviously lead to misinterpretation in sweating-related measurements. Even though this effect did not occur on the hypothenar site in this study, we cannot exclude the possibility that it could occur on this site as well. The type B recordings were often seen to have a steady increase in SSCD level throughout our 29 minute recordings (e.g. figure 5 a), and it is not unlikely that inverse sweat responses could occur also on this site after a longer time of electrode-skin contact when the gel has been allowed to penetrate deeper into the ducts.

The implications of these electrode-related phenomena depend on how the measured signal is interpreted or computationally parameterized. Traditionally, both the skin conductance level and the characteristics of the skin conductance response have been used as psychophysiological parameters (Boucsein 1992) and the results in this paper points to potential sources of errors introduced by the electrodes regarding both features. The frequency of the skin conductance responses is a parameter which may not be as susceptible to these errors as the abovementioned. However, the response detection usually depends on an amplitude criterion beyond a certain threshold, which may cause electrode gel dependency in this parameter as well due to differences in amplitudes. In addition, these results indicate that the gel to duct penetration may alter the sensitivity to the sweat glands below the EEA. In other words, a moist gel with low viscosity could detect a greater number of responses than a solid gel or gel-free electrode. For the wetter gels, the results will likely have a significant dependency on the time since electrode attachment and the SC thickness.

The pressure artefacts are more relevant in situations where the patient is moving or physically active during the recordings than for controlled setups in the laboratory. Pressure on the electrodes is also likely to occur during sleep monitoring as the test subject is turning around. Push or pull on the electrodes may give transient artifacts which could be detected as single responses, but the greatest source of error comes from the electrodes which cause changes that remain after the pressure is no longer applied.

The mentioned errors are probably equally relevant for dc and ac methods of skin conductance measurement, as long as these are performed within the linear range.

4.8. Summary of findings

- The electrode gels can introduce large changes in the level of the skin conductance over time, corresponding with the amount of free water in the gel and its viscosity.
- Sweating can lead to negative skin conductance responses for wet gels with low viscosity. This inverse sweating response effect probably depends on the thickness of the SC.
- Wet gels may provide an increased sensitivity to sweat activity in partially filled ducts.

- Solid gels have in general a better ability than the wetter gels to return the skin conductance to baseline during recovery after a period of sweating.
- Wet gels seem to give less repeatable measurements due to the changes introduced by the gel which likely depend on the interindividual SC properties.
- Expectedly, the gels with lower mechanical stability are more susceptible to pressure artifacts.
- The karaya-based solid gel may introduce a drift in the skin conductance measurement due to lowering of its initially high resistivity.
- The Isotonic EDA Jelly (Type C) may be completely absorbed by the skin during long-term wear.

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