OPINION

Calcitox-aging counterbalanced by endogenous farnesol-like sesquiterpenoids: An undervalued evolutionarily ancient key signaling pathway

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ABSTRACT

Cells are powerful miniature electrophoresis chambers, at least during part of their life cycle. They die at the moment the voltage gradient over their plasma membrane, and their ability to drive a self-generated electric current carried by inorganic ions through themselves irreversibly collapses. Senescence is likely due to the progressive, multifactorial damage to the cell’s electrical system. This is the essence of the “Fading electricity theory of aging” (De Loof et al., Aging Res. Rev. 2013;12:58–66). “Biologic electric current” is not carried by electrons, but by inorganic ions. The major ones are H⁺, Na⁺, K⁺, Ca²⁺, Mg²⁺, Cl⁻ and HCO₃⁻. Ca²⁺ and H⁺ in particular are toxic to cells. At rising concentrations, they can alter the 3D-conformation of chromatin and some (e.g., cytoskeletal) proteins: Calcitox and Protonox. This paper only focuses on Calcitox and endogenous sesquiterpenoids. pH-control and Ca²⁺-homeostasis have been shaped to near perfection during billions of years of evolution. The role of Ca²⁺ in some aspects of aging, e.g., as causal to neurodegenerative diseases is still debated. The main anti-Calcitox mechanism is to keep free cytoplasmic Ca²⁺ as low as possible. This can be achieved by restricting the passive influx of Ca²⁺ through channels in the plasma membrane, and by maximizing the active extrusion of excess Ca²⁺, e.g., by means of different types of Ca²⁺-ATPases. Like there are mechanisms that antagonize the toxic effects of Reactive Oxygen Species (ROS), there must also exist endogenous tools to counteract Calcitox. During a re-evaluation of which mechanism(s) exactly initiates the fast aging that accompanies induction of metamorphosis in insects, a causal relationship between an endogenous sesquiterpenoid, namely the farnesol ester named “juvenile hormone,” and disturbed Ca²⁺-homeostasis was suggested. In this paper, this line of thinking is further explored and extended to vertebrate physiology. A novel concept emerges: horseshoe-shaped sesquiterpenoids seem to act as “chemical valve” or “spring” in some types of multi-helix transmembrane proteins (intramolecular prenylation), from bacterial rhodopsins to some types of GPCRs and ion pumps, in particular the SERCA-Ca²⁺-pump. This further underpins the Fading Electricity Theory of Aging.

Introduction

In most animal species the typical morphological and physiologic aging features do not start manifesting themselves right after birth, but after a species-specific delay, usually at reproductive maturity. Yet, instant aging soon after birth is a possibility as illustrated in Hutchinson-Gilford Progeria, a rare fast-aging syndrome in humans. Which mechanism is responsible for the delay? None of the numerous possible driving mechanisms of aging, neither alone nor in combination, adequately explain this phenomenon in full. In insects, the partial answer is: a high titer of an ester of the sesquiterpenoid farnesol, which acts as a “juvenile, anti-aging hormone” (JH). Vertebrates also pass through a juvenile/larval stage, just like their common ancestor with the invertebrates, but they do not use some (known) “juvinilizing hormone.” Is this evolutionary divergence due to the loss in an ancient (in)vertebrate ancestor of some essential gene in the biosynthesis of farnesol or some endogenous farnesol-like substance (FLS)? Or do vertebrates use FLS in some other way to obtain a similar juvenilizing effect? Or do we have to search for a still evolutionarily older juvenilizing/aging mechanism that was already present in (some) prokaryotic ancestors of eukaryotes? How to proceed to answer such questions?

In search for the still unidentified membrane receptor for insect JH, a fortuitous flash of new insight emerged when it was realized that farnesol (and related substances (FLS)), the plant toxin thapsigargin (a potent blocker of the SERCA-calcium pump), as well as retinal/retinoids all belong to the family of sesquiterpenoids. Retinal is the key cofactor in the proton-pump rhodopsin of prokaryotes (Archaea and Bacteria) and eukaryotes. The cis-trans isomerization of retinal under the influence of light
energy is essential for visual perception. It is often overlooked that retinoids not only structurally resemble the isoprenoid farnesol (both are isoprenoids), but that some retinoids, e.g., retinoic acid (a metabolite of vitamin A (retinol)) also have "juvenile hormone" activity when tested in appropriate insect bioassays. Apparently, animals lost the capacity to synthesize retinoids by themselves, but they continued to need them in some signaling pathways. This is well documented from studies in developmental biology. Animals solved this problem by ingesting vitamin A with their diet followed by its metabolization into retinal/retinoids. In insects, the onset of metamorphosis when aging phenomena start manifesting themselves is not caused by (absence of) some retinoid, but by the sudden drop of the titre of the endogenous sesquiterpenoid "juvenile hormone." Despite 50 y of research, the mode of action of JHs, in particular at the level of cell membranes, is not yet fully understood. In contrast, substantial progress has been made in the search for the protein signaling cascade and nuclear receptors/transcription factors (Methoprene-tolerant and its paralog Gce in *Drosophila*). In animals endogenous sesquiterpenoids are synthesized in the mevalonate pathway. In vertebrates, this pathway is best known for its role in the biosynthesis of cholesterol and steroids. In insects it is best known for its role in the biosynthesis of juvenile hormone. Because some ancestor of the arthropods seems to have lost the gene coding for the enzyme squalene synthase, the direct precursor of cholesterol in vertebrates (see later), insects cannot synthesize cholesterol by themselves. They have to ingest it as a vitamin. Hitherto, a key role of the mevalonate pathway in Ca\(^{2+}\)-homeostasis has not been emphasized in either vertebrates or invertebrates, but exactly this may, in my opinion, be the key functional role of this pathway (this paper).

The elucidation in 2011 of both the structure of the binding pocket and the mode of action of thapsigargin, the sesquiterpenoid that is the best documented blocker of the SERCA-Ca\(^{2+}\) pump in all eukaryotes, was, in my opinion, a landmark in thinking about control of Ca\(^{2+}\)-homeostasis. It suggested an overlooked causal link between *endogenous sesquiterpenoids* and control of some aspects of Ca\(^{2+}\)-homeostasis. This mechanism seems to have been well conserved in evolution in all eukaryotes. It triggered the formulation of the "Calcitoxaging" concept as outlined in this paper. This (novel) term denotes (accumulating) damage to cells due to rises ("explosions," "puffs," "sparks") in intracellular Ca\(^{2+}\) ([Ca\(^{2+}\)]i), in particular long-lasting ones. A serendipitous side aspect of the Calcitox concept is that 50 y after the chemical identification of insect juvenile hormone, the (fast) mode of action at the membrane level of this and other endogenous sesquiterpenoids seems to get a plausible explanation.

Because some readers may not be familiar with the basic principles of the divergent topics that are needed to construct the argument, a few concise reminders (probably superfluous for specialists) highlighting the important issues will be given first.

### Some introductory principles about Ca\(^{2+}\)-homeostasis

To understand why the Ca\(^{2+}\) concentration in cells can drastically and abruptly change, one has to have an idea about the magnitude of the concentration gradient outside-inside cells, as well about the ways that Ca\(^{2+}\) can be compartmentalized in cells.

#### How steep is the trans-plasma membrane Ca\(^{2+}\)-gradient? Where can Ca\(^{2+}\) be stored in the eukaryotic cell?

[Ca\(^{2+}\)]i can quickly rise from very low to very high because: 1. there is a huge trans-plasma membrane calcium concentration gradient; 2. Ca\(^{2+}\)-channels with various gating mechanisms are present; 3. a cell can store Ca\(^{2+}\) in high concentrations in some intracellular membrane compartments (in particular the endoplasmic reticulum and the mitochondria) from where part of the Ca\(^{2+}\) can be (temporarily) released upon proper stimulation. [Ca\(^{2+}\)]i in unstimulated (= resting) cells amounts to about 100 nM (nM). In such low concentrations Ca\(^{2+}\) is not toxic. [Ca\(^{2+}\)] in the extracellular environment (blood e.g.) is orders of magnitude higher than in the cytoplasm, from around 2 mM in blood to even 50 mM in milk (Fig. 1). [Ca\(^{2+}\)]i = 100 nM versus [Ca\(^{2+}\)] = 1 mM represents a gradient of 10,000 times.

The steepness of this gradient can be illustrated as follows. Imagine that you are 1.7 meters tall and that you are standing at the bottom of a canyon of which the walls on both sides are about twice as high as the Himalaya mountains, thus about 17,500 m high. Debris and dust are continuously falling down from the walls. If you do not shovel them away, you will be buried by the debris in a short time. This example (1.7 vs. 17,500 m) corresponds to an altitude gradient of 10,000 times. Such a concentration gradient would not pose a problem if the plasma membrane would be impermeable to Ca\(^{2+}\). However, due to the presence in all cell types of Ca\(^{2+}\)-channels with different gating mechanisms that is not the case. As a result, Ca\(^{2+}\) enters the cell downhill the Ca\(^{2+}\)-gradient (= passive transport), and the resulting excess Ca\(^{2+}\) has to be dealt with one way or another.

Ca\(^{2+}\) can become toxic when its concentration rises above [Ca\(^{2+}\)]i of resting cells for longer time intervals.
Cells can cope with increasing [Ca\textsuperscript{2+}]i in various ways. For the transport of Ca\textsuperscript{2+} against the concentration gradient (also called uphill transport), the major proteins involved are the plasma membrane Ca\textsuperscript{2+}-ATPases or PMCAs and the Ca\textsuperscript{2+}-ATPases present in the 2 types of endoplasmic reticulum ATPases (smooth (SER) and Rough (RER)), the (SERCAs). For the passive transport of Ca\textsuperscript{2+}, thus down the Ca\textsuperscript{2+}-gradient (also called the downhill transport) the various types of Ca\textsuperscript{2+}-channels are responsible (Fig. 2). As will be dealt with in some more detail later, there are 4 gross mechanisms for keeping [Ca\textsuperscript{2+}]i low.

**Ca\textsuperscript{2+}-explosions, puffs, sparks, and waves: A lifelong phenomenon. Life begins and ends with a Ca\textsuperscript{2+}-explosion.**

Ridgway et al.\textsuperscript{19} were the first to report that upon fertilization free calcium increases explosively in activating medaka eggs, hence the name “Ca\textsuperscript{2+}-explosion.” Later it became documented that Ca\textsuperscript{2+} is the universal signal for egg activation in all sexually reproducing species. It is a finely tuned system. In Xenopus oocytes, the rise in [Ca\textsuperscript{2+}]i lasts several minutes,\textsuperscript{20} in some other species it even lasts longer. In Xenopus, PMCAs, which are attached to elements of the cytoskeleton in the egg cortex, are internalized during egg maturation because the cytoskeleton contracts due to the rise in [Ca\textsuperscript{2+}]i.\textsuperscript{21} Therefore they are unable to extrude Ca\textsuperscript{2+} out of the ooplasm. One could think of this Ca\textsuperscript{2+}-rise as a near death experience for the egg. Sensitivity of the cytoskeleton to Ca\textsuperscript{2+} is a ubiquitous phenomenon in cell physiology. The most obvious example is muscle contraction. The cytoskeleton may also be of crucial importance for storing “cognitive information.”\textsuperscript{22} At death, the Ca\textsuperscript{2+} gradient over the different membrane systems of cells collapses. This is the final Ca\textsuperscript{2+}-explosion cells experience. In between the 2 key explosions, there are numerous smaller rises in [Ca\textsuperscript{2+}]i.

According to\textsuperscript{23,24} Ca\textsuperscript{2+}-puffs are localized Ca\textsuperscript{2+} signals mediated by Ca\textsuperscript{2+} release from the endoplasmic reticulum (ER) through clusters of inositol triphosphate receptor (IP\textsubscript{3}R) channels. Ca\textsuperscript{2+}-sparks result from the opening of a single, or the coordinated opening of many, tightly clustered ryanodine receptor (RyR) channels in the
sarcoplasmic reticulum. They can be frequent. Normal sparks usually last 10 to 100 milliseconds, but long lasting sparks lasting several hundred milliseconds are also widely observed. The recruitment of IP3R channels during puffs depends on Ca\(^{2+}\)-induced Ca\(^{2+}\) release, a regenerative process that must be terminated to maintain control of cell signaling and prevent Ca\(^{2+}\) cytotoxicity. Physiological activation is triggered by binding of agonists to G-protein-coupled receptors (GPCRs) on the cell surface, leading to cleavage of phosphatidylinositol biphosphate and release of IP3 into the cytosol. Many different GPCRs have already been described or predicted. The human genome harbours about 800 different GPCR-coding genes. In the worm C. elegans the prediction is that there are about 128 GPCRs for neuropeptides alone. Because many GPCRs use Ca\(^{2+}\)-puffs in...
their signaling system, such high figures indicate that Ca\(^{2+}\)-puffing occurs very frequently in all cell types.

Lionel Jaffe\(^{30}\) measured the speed of Ca\(^{2+}\) waves, which are usually Ca\(^{2+}\)-induced but sometimes also H\(^+\)-induced in a variety of cell types.\(^{31}\) Fast calcium waves (10–30 microns/s) can be propagated through all nucleated eukaryotic cells.

Ca\(^{2+}\) blinks (the corresponding depletion of Ca\(^{2+}\) in the sarcoplasmic reticulum during a spark, and the small Ca\(^{2+}\) release events called quacky SR Ca\(^{2+}\) release (QCR) are also terms used in the literature.\(^{32}\)

**Why and when can Ca\(^{2+}\) be toxic, and even deadly?**

H\(^+\) and Ca\(^{2+}\) probably were primordial regulators of the conformation of some macromolecules that were/are instrumental to control of gene expression in the very first cell on earth. Since then they continued performing key functions in cell physiology. Making the conformation of chromatin and of selected proteins in key signaling pathways change substantially, can be considered as a toxic effect. Undoubtedly, the statement that a Ca\(^{2+}\) puff can act as a secondary messenger, a beneficial effect at the (sub)cellular level, *because it is toxic* is counterintuitive. Whether Ca\(^{2+}\)-puffs cause damage mainly depends upon the duration of the puffs (= the speed of clearance of excess free Ca\(^{2+}\) from the cytoplasm), as well as upon the properties of various cell types with their specific Ca\(^{2+}\)-homeostasis systems in which the puffs occur. The frequency of the heartbeat gives an estimate of what is “short” at the organismal level, namely a few seconds maximally. This is the time needed to restore a low [Ca\(^{2+}\)]i. In cardiac myocytes the Ca\(^{2+}\)-sparks themselves have a typical duration of 20–40 ms.\(^{33}\) The opposite extreme is the long lasting Ca\(^{2+}\)-sparks/puffs which induce apoptosis, an extreme form of cell damage and very rapid aging.\(^{18,34-36}\)

**Control of gene expression by inorganic ions**

In addition to changes in the conformation of macromolecules in the cytoplasm, some inorganic ions also have an effect on control of gene expression in both prokaryotes and eukaryotes as known since the 1960s.\(^{37-40}\) In particular the experiments of Kroeger and of Lezzi\(^{41-46}\) pointed to the importance of the ionic environment for the coarse regulation of gene expression while transcription factors are needed for fine tuning. These authors used induction of puffs (here puff has another meaning than in Ca\(^{2+}\)-puffs) in the polytene chromosomes of some dipteran insects as a model. Their experiments urged for caution when interpreting results on control of gene expression by e.g., hormones, in particular when these experiments are executed with cell-tissue homogenates. Indeed, homogenization results in the release of Ca\(^{2+}\) from its subcellular storage sites. This influences the activity of some enzyme systems. The dilemma: is an observed effect of a given treatment on gene expression due to the altered inorganic ion environment itself, or to the interaction of e.g., the hormone under study, with a nuclear receptor? As an example: it has been reported that the change in concentration in nuclear Ca\(^{2+}\) itself controls the T cell fate decision between a proliferative immune response and tolerance.\(^{47}\)

Furthermore, one should also keep in mind that hydrophobic signaling molecules (e.g., some steroid and farnesol-like hormones) diffuse through all intracellular membranes (Fig. 3) and nestle themselves in suitable binding pockets (see later). This means that such molecules not only act through nuclear receptors/transcription factors (e.g., Methoprene-tolerant and Gce for juvenile hormone), but as well through membrane receptors with a role in Ca\(^{2+}\)-homeostasis, some of which can be located in intracellular membranes. To date, the controversy whether lipophilic signaling molecules like JH

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**Figure 3.** Schematic representation of the 2 major models on the mode of action of some lipophilic signaling substances, e.g., Juvenile hormone/FLS and (some) steroids. The classical “Karlson hypothesis” says that such molecules can pass the plasma membrane unhindered, and somehow end up in the nucleus where they act through nuclear receptors/transcription factors. The inbrome model\(^{15}\) says that lipophilic signaling molecules first enter the whole membrane system and, if they encounter a binding pocket in some of the membrane proteins, they may exert a function. In addition, a second mode of action through nuclear receptors is also a possibility.
and some steroids only act through nuclear receptors/transcription factors (= the classical Karlson hypothesis) or also through membrane receptors as already suggested by Wigglesworth in 1969, has not yet been resolved.

**Proteins responsible for trans-membrane transport of \( \text{Ca}^{2+} \)**

**Structure and main function of \( \text{Ca}^{2+} \)-ATPases: PMCA and SERCA**

The molecular structure of \( \text{Ca}^{2+} \) pumps (Fig. 4 and 5) and \( \text{Ca}^{2+} \) channels is complex. A variety of agents can interfere with their activity. It is the concerted action of all controlling agents that results in keeping \( [\text{Ca}^{2+}]_i \) below the toxic limit by eliminating excess \( \text{Ca}^{2+} \) in various ways. The aa-sequence of PMCAs and SERCAs is very similar, suggesting a common evolutionary origin.

**Plasma Membrane \( \text{Ca}^{2+} \) ATPase (PMCA) resides in the plasma membrane and extrudes \( \text{Ca}^{2+} \) from the cytoplasm of all eukaryotic cells.** PMCAs reside in the plasma membrane of cells and function to remove \( \text{Ca}^{2+} \) from the cytosol of cells. In mammals they are encoded by 4 separate genes, and additional isoform variants are generated via alternative RNA splicing of the primary gene transcripts. They have 10 transmembrane hydrophobic domains (Fig. 4), similar to SERCA2a (Fig. 5). The pump is powered by the hydrolysis of ATP. Another important player for keeping \( [\text{Ca}^{2+}]_i \) very low (at about 100 nmol) is the \( \text{Na}^{+} - \text{Ca}^{2+} \) exchanger (NCX). Various isoforms of PMCA are expressed in different tissues. PMCAs are regulated by several agents, the most important being calmodulin.

**Sarcoplasmic Reticulum \( \text{Ca}^{2+} \) ATPases (SERCAs) pump \( \text{Ca}^{2+} \) from the cytoplasm into the sarcoplasmic/endo-plasmic reticulum.** In higher vertebrates there are 3 SERCA- and 2 SPCA-pump genes which encode P-type ATPases of the P(2A)-group which are respectively found in membranes of the endoplasmic reticulum and the secretory pathway. The SERCA-pump paralogs are named SERCA1, SERCA2, and SERCA3. They are expressed at various levels in different cell types. Various splice variants are known, and the number of proteins reported to interact with SERCA is rapidly growing.

**\( \text{Ca}^{2+} \)-homeostasis: some control mechanisms of the activity of SERCAs and \( \text{Ca}^{2+} \)-channels**

**Binding sites on SERCA for inhibitors/activators, e.g., of thapsigargin.** A variety of inhibitors of the SERCA \( \text{Ca}^{2+} \)-pump are known. We only elaborate on the plant toxin thapsigargin because it is a sesquiterpenoid that, in my opinion, likely acts by displacing a natural endogenous sesquiterpenoid ligand(s) from its binding site.

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**Figure 4.** Cartoon depicting the main structural features of the PMCA at resting \( [\text{Ca}^{2+}]_i \) (A) and elevated \( [\text{Ca}^{2+}]_i \) (B). The PMCA consists of an N-terminal domain, 10 transmembrane domains, 4 cytosolic loops and a regulatory C-terminal domain. When \( [\text{Ca}^{2+}]_i \) is elevated, the Ca/CaM binds to the autoinhibitory CaM-binding domain causing a conformational change which exposes the catalytic site which consists the ATP-binding sites and the aspartate residue that is phosphorylated during the reaction cycle. This increases the \( \text{Ca}^{2+} \)-transporting activity of the PMCA. Reprinted under the Creative Commons Attribution-NonCommercial License from: Bruce J (2013) with thanks.
site on the SERCA pump (Fig. 6). That ligand could be **farnesol itself or a farnesol-like sesquiterpenoid (FLS)**. The importance of FLS will be outlined later.

Thapsigargin is a sesquiterpene lactone originally extracted for the first time from the plant *Thapsia garganica*. Upon binding to its receptor on the SERCA-pump, it causes the release of $\text{Ca}^{2+}$ from an internal store by a mechanism independent of the hydrolysis of phosphoinositides and concomitant activation of protein kinase C. It acts in the subnanomolar range. It binds with high affinity. Inhibition leads to elevated $[\text{Ca}^{2+}]_i$, which, in its turn, induces apoptosis. It binds to SERCA in a hydrophobic funnel-like cavity open to the cytosol formed by TM3, TM5 and TM7 and prevents the movement of the helices relative to each other. Nine out of the 10 transmembrane loops of SERCA undergo rearrangements during a pumping cycle, enabling the release of $\text{Ca}^{2+}$ into the lumen of the SR, and, on the cytoplasmic side, to create a pathway for entry of new $\text{Ca}^{2+}$-ions (Toyoshima and Nomura 2002). This is thought to explain the inhibitory effect of the pump (Takahashi et al. 2007). The amino acids lining the binding pocket for thapsigargin in SERCA1a are fully conserved in SERCA2b (Fig. 5, Fig. 6). Because thapsigargin acts as a SERCA blocker in various eukaryotes, and because SERCAs are very well conserved in evolution, it can be assumed that the binding is well conserved as well. However, this awaits experimental validation.

**Does thapsigargin displace an endogenous ligand? Its nature? Its mode of action in inbrome signaling?** To date, it remains unknown whether thapsigargin binds at a pocket that is normally occupied by an endogenous ligand, or that it just binds haphazardly to that site in the SERCA-pump. It has been suggested that the binding pocket may be sensitive to cholesterol. An alternative hypothesis was advanced namely that thapsigargin, being a sesquiterpenoid, displaces an ubiquitous (in eukaryotic cells) sesquiterpenoid that is formed in the mevalonate biosynthetic pathway, namely farnesol or farnesol-like sesquiterpenoids (FLS). Thapsigargin can displace an endogenous ligand from its receptor if it binds with a greater affinity.

The possible link with aging becomes apparent if one realizes that in insects, the hormone which keeps larval insects in their juvenile/larval stage, namely the well documented juvenile hormone, is nothing other than an ester of farnesol. Furthermore, farnesol that occurs in all eukaryotic organisms, tissues of mammals and **Homo inclusive**, is also active in bioassays that visualize antiaging (juvenilizing) effects in insects. For figure see ref. 15. This raises the question whether, perhaps, farnesol-like endogenous sesquiterpenoids might also be causal to realize and maintain the juvenile state in vertebrates, and why such compounds are not known as hormone(s) in vertebrates? The answer is: they were not sought for. There are several reasons. One reason is that in vertebrates, unlike in insects, one cannot realize a

**Figure 5.** Schematic representation of the structure of the SERCA2a (10 transmembrane helices) and SERCA2b (11 transmembrane helices) calcium pumps. Slightly modified (by adding a horseshoe-shaped string (in yellow) representing the binding of (endo- or exogenous) sesquiterpenoids linking the TM loops 3, 5 and 7): after Dhitavit et al. in Br J Derm 2004, Vol 150, page 823. With permission from the publisher.
viable situation in which farnesol/FLS are totally absent. Another reason is that no bioassays have been established in vertebrates allowing the induction or prolongation of typical juvenilizing effects in a relatively short time. The third reason is that physiologists did not consider the possibility that farnesol/FLS, in addition to serving as precursor for the synthesis of squalene and cholesterol, might have another function on their own, namely as a key adjuvant of the SERCA-pump (see later). Textbooks of biochemistry, physiology and endocrinology do not list this possibility.

**Ca\(^{2+}\)-channels: Various families**
Such channels represent the passive aspect of the Ca\(^{2+}\)-homeostasis system. They facilitate the influx of Ca\(^{2+}\) from the external environment into the cytoplasm. Sometimes Ca\(^{2+}\)-channel is synonymous for voltage-dependent calcium channel. There are also receptor-gated calcium channels and store-operated ones (Fig. 2).

In recent years a relatively large family of TRPs, Transient Receptor Potential channels has been described. Some of these ion channels are non-selectively permeable to cations, including Na\(^+\), Ca\(^{2+}\) and Mg\(^{2+}\). Some chemicals act as channel openers, others as blockers. Several synthetic blockers are of important medicinal use. In some tissues, a particular class of Ca\(^{2+}\)-channels is sensitive to farnesol. Free oxygen radicals (ROS) open some types of plasma membrane Ca\(^{2+}\) and K\(^+\)-permeable channels in plant root cells. The role of ROS in Ca\(^{2+}\)-signaling and disease is well documented.

**Gross mechanisms for coping with excess Ca\(^{2+}\) in the cytoplasm**

**Four key mechanisms**

The well-known fact that different cell types age at a different pace likely depends, in part, on differences in their...
$Ca^{2+}$-homeostasis systems. Such differences are inherent to the principles of differentiation during development. Cells have an array of strategies for coping with rising $[Ca^{2+}]_i$.

The 4 key mechanisms are:
1. If little $Ca^{2+}$ enters the cell, thus if the plasma membrane is only weakly permeable to $Ca^{2+}$, the $Ca^{2+}$-ATPases present in the plasma membrane (PMCAs and the sodium-calcium exchanger (NCX)) will pump the excess $Ca^{2+}$ out of the cell. Such cells do not have abundant internal membrane systems (RER and SER) (red dot with number 1 in Fig. 1).
2. If more $Ca^{2+}$ enters, some excess can be stored in intracellular storage sites, in particular the sarcoplasmic endoplasmic reticulum (SR) and mitochondria. Muscle cells are typical examples (red dot with number 2 in Fig. 1).
3. If both mechanisms do not suffice for extruding $Ca^{2+}$, a third mechanism can be activated, namely the synthesis of proteins for secretion in the rough endoplasmic reticulum (RER)-Golgi system. This system removes excess $Ca^{2+}$ along with the secreted proteins. A typical example is removal of $Ca^{2+}$ by the secretion of milk proteins in mammary gland cells (red dot with number 3 in Fig. 1).
4. If even this is insufficient, the $Ca^{2+}$-dependent apoptosis system is activated, and the cell dies. (Fig. 1B).

**Indirect effects of $[Ca^{2+}]$ on activity of some enzyme systems**

The $Ca^{2+}$ concentration inside the SER is about 10,000x higher than in the cytoplasm. This explains why in cells at rest, some enzymes that are anchored in the SR-membranes are inhibited and why, upon release of $Ca^{2+}$, these enzymes can be temporarily activated. This is e.g., the case for some enzymes with a role in lipid- or steroid biosynthesis. When a lipid- or steroid producing cell is in rest, the SER is fully uploaded with $Ca^{2+}$. The activity of the cited enzymes is very low. Apparently, the high

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**Figure 7.** Comparison of the mevalonate biosynthetic pathways in insects and vertebrates. Modified after. Insects do not have the gene coding for squalene synthase. As a result, they cannot synthesize cholesterol by themselves. Some chemical structures of endogenous sesquiterpenoids, retinoic acid and the SERCA-pump blocker thapsigargin (which is also a sesquiterpenoid) are also represented.
Ca\(^{2+}\)-concentration in the SER inhibits their activity. When the cells are stimulated by e.g., a releasing hormone from the brain, a Ca\(^{2+}\) puff will induce a Ca\(^{2+}\)-induced release of stored Ca\(^{2+}\) from the SER. As a result of the decreasing Ca\(^{2+}\) concentration in the SER-membranes, some enzyme systems are no longer inhibited. For short while, lipids or steroids will be synthesized. When the released Ca\(^{2+}\) is taken up again in the SER, the enzyme activity will be halted. This system may explain why in vertebrates the release of e.g., pituitary hormones happens in pulses. A similar system has also been observed in the mosquito *Aedes aegypti*. These authors reported that Ca\(^{2+}\)-influx enhances neuropeptide activation of ecdysteroids hormone production by mosquito ovaries.

**The mevalonate biosynthetic pathway for farnesyl-PP, farnesol, juvenile hormone, cholesterol and steroid hormones: Similarities and differences in invertebrates-vertebrates**

It may not be superfluous to refresh one’s insight into the nature of the mevalonate biosynthetic pathway with an emphasis on a key difference between the pathway in vertebrates vs. in insects. Vertebrates and arthropods share the same early steps in the biosynthetic pathway of the endogenous isoprenoid farnesyl pyrophosphate (Fig. 7). In vertebrates, farnesyl-PP is converted into squalene by squalene synthase. Squalene in its turn is converted into cholesterol that, in addition to other roles, can act as the precursor of steroid hormones. Arthropods and plants cannot synthesize squalene and cholesterol. The widely accepted explanation says that, in insects, this is due to a silencing mutation of the gene coding for squalene synthase in an ancient ancestor of arthropods. Another possibility is that the ancient ancestors of arthropods never had a gene coding for squalene synthase, and that such gene was a specific acquisition in the evolution of chordates/vertebrates. Whatever the right explanation is, the result is that for insects cholesterol is a vitamin that has to be ingested with the food, or that has to be metabolized from particular steroid precursors present in the food (e.g., plant sterols). This food-borne cholesterol can be enzymatically converted into steroids, e.g., ecdysteroids. If the loss of the squalene synthase gene is the correct explanation, the inability to convert farnesyl-PP into squalene must have prompted the mutant ancestor and its progeny to come up with a solution for the problem as to how to cope with the excess farnesyl-PP. The solution seems to have been to transform this compound into farnesol that was next esterified. The addition of an epoxy group made it into a compound with juvenile hormone activity (in bioassays) that is much more potent than farnesol in the same bioassays. That explains why, in insects, the *corpora allata* (CA), very tiny glands connected to the brain, which are the only site of JH synthesis in larvae, can produce enough hormone activity to serve the physiological role of farnesol in vertebrates where e.g., the liver, a large organ, is the major site of synthesis of FLS and cholesterol.

Recently, Schenk et al. reported that the brain of the annelid worm *Nereis virens* produces methylfarnesolate of which the major function is the suppression of reproduction. This is achieved by inhibiting the release of stored reserves for the production of yolk and eggs in females committed to reproduction. This is probably a Ca\(^{2+}\)-mediated process. This strengthens the hypothesis that sesquiterpenoids are ancient animal hormones that were already active in the marine ancestors of arthropods.

**Farnesol, prenylation, retinoic acid**

**Farnesol: A noble unknown in vertebrate physiology**

*Farnesol* is derived from the name of the Farnese acacia tree *Vachellia farnesiana*, from which the compound was isolated for the first time. *Vachellia* is one of the plant species that is used for extraction of compounds of interest to the perfume industry. Farnesol-related sesquiterpenoids are hydrophobic, thus immiscible with water. Upon contact with membranes, which are invariably very rich in lipids, they insert themselves into such membranes and diffuse rapidly all over the interconnected membrane systems of cells: plasma membrane, SER, RER and nuclear envelope, to name the major ones (Fig. 1). Although farnesol is a ubiquitous isoprenoid compound in animals, plants and fungi, little is known about its possible functions except its role as a precursor of squalene or juvenile hormone.

**Retinal: An ancient functional precursor of farnesol/FLS? Vitamin A**

To better understand the mode of action of farnesol/FLS, some data from the physiological archeology of farnesol may be helpful. The structural- and functional similarities between retinoids (Pubchem: 11-cis-retinal) and FLS/sesquiterpenoids (Pubchem: Farnesol, Juvenile Hormone) are striking.

Retinoic acid is a metabolite of vitamin A. Structurally it is related to insect juvenile hormone. (Fig. 6). It has functions in control of growth, regeneration and development (Hox-genes). It affects calcium signaling, e.g., in adult molluscan neurons and in mouse ovarian granulosa cells. Various retinoids are active in increasing adhesive properties of some cell types.
Retinal (Fig. 8), is a cofactor in bacterial and eukaryotic rhodopsins. Bacterial protorhodopsins were probably $H^+$-pumps. Some evolved into $Na^+$-rhodopsins and $Cl^-$-pumping rhodopsins. The ur-proton pump is also thought to be ancestral to the various inorganic ion transporting ATPases present in eukaryotes. To animal physiologists rhodopsin is best known as a light-sensitive receptor protein involved in visual phototransduction. It is a biological pigment found in the rods of the retina. It is a G-protein coupled receptor (GPCR). When rhodopsin is exposed to light, it immediately photobleaches. In animals this is due to the isomerization of the covalently bound chromophore 11-$cis$-retinal into all-$trans$-retinal by light which induces a conformational change (bleaching) in opsin. The way retinal is positioned inside rhodopsin and the way $H^+$ is transported is well documented.

Animal rhodopsins are thought to be derived from ancestral prokaryotic rhodopsins which functioned as proton pumps. In microbial rhodopsins, it is all-$trans$ retinal that photo-isomerizes into the 13-$cis$ form, thus the opposite from the situation in animals. In evolution, other types of ion-pumping microbial rhodopsins appeared: chloride pump, sodium pump, cation channel, sensor (membrane transducer) and sensor (soluble transducer). Many animal GPCRs have a rhodopsin-like structure and are therefore thought to be possibly evolutionarily related to prokaryotic rhodopsins.

**Rhodopsins and GPCRs**

G protein-coupled receptors (GPCRs), also known as 7TM receptors represent a large protein family of receptors, to which molecules outside the cell selectively bind. They activate internal signal transduction pathways and, ultimately, cellular responses. They are only found in eukaryotes. One of the classification systems group the numerous GPCRs in 6 classes. The largest class that codes for about 85% of all the GPRC genes comprises the Rhodopsin-like GPCRs. Such GPCRs still harbour some of the properties of the ancient rhodopsins, the involvement of isoprenoids inclusive.

The GPCR arranges itself into a tertiary structure resembling a barrel, with 7 transmembrane helices forming a cavity within the plasma membrane that serves a ligand domain. The eventual effect of the binding of an agonist is a change in the relative orientation of the TM helices (likened to a twisting motion). It causes activation of a G protein. G proteins cannot diffuse far away due to the palmitoylation of $Go$ and the presence of an isoprenoid moiety that has been covalently added to the C-
A. DE LOOF

The primary effectors of Gβγ are various ion channels, some types of (voltage-gated) Ca^{2+}-channels inclusive. Inverse agonists and antagonists may also bind to several different sites, but the eventual effect must be prevention of this TM helix reorientation.

Mode of action of farnesol/FLS? Rotatable bond count. FLS: Important inbrones in control of Ca^{2+}-homeostasis?

One of the chemical properties of farnesol that is seldom focused on in studies of its role in endocrinology is its “rotatable bond count.” Rotatable bond is defined as any single non-ring bond, bounded to terminal heavy (i.e., non-hydrogen) atom. Amide C-N bonds are not considered because of their high rotational energy barrier (from Molinspiration). The number of rotatable bonds (nrotb) is a measure of molecular flexibility, an important issue in the functioning of molecules that are cofactors in e.g., multi-helix transmembrane proteins. With its alternation of single and double bonds and a nrotb-value of 5, farnesol is indeed moderately flexible, as are retinal (value 5), retinoic acid (value 5) and JH III (value 8). Thus all these molecules have, in principle at least, the right flexibility to function as “molecular glues” to keep several (usually 3) helices of complex transmembrane proteins well aligned, and bring them back into their “relaxed position” after an activity cycle, e.g., pumping an ion across the membrane. Exactly this property may explain its key function as an inbrome. Inbrome stands for intramembrane signaling molecule, a term introduced for the first time by De Loof et al. This mode of action of some lipophilic signaling molecules, hormones inclusive, is undervalued in physiology and endocrinology.

With respect to the mode of action of FLS at the level of the membrane, antagonism to the mode of action of thapsigargin is, in my opinion, a plausible explanation. As cited before, thapsigargin binds to SERCA in a hydrophobic funnel-like cavity open to the cytosol formed by the transmembrane loops TM3, TM5 and TM7, and prevents the movement of the helices relative to each other. This is thought to explain the inhibitory effect of the pump.

The likely opposite effect brought about by the natural ligand which, in my opinion, is an endogenous farnesol-like sesquiterpenoid, is gluing together the 3 TMs in a moderately flexible way. This is visualized by the yellow string in Fig. 4. Thus, the basic function of FLS is, in my opinion, to allow temporary movements of transmembrane helices of the SERCA pump during its pumping cycle: stretching followed by narrowing. In other words, farnesol may act as a spring inbrome with the function of a “chemical valve” that makes that the TM3, TM5, TM7 transmembrane loops of SERCAs are brought back in their original position after each Ca^{2+}-pumping cycle. To function this way, the sesquiterpenoid must be firmly anchored (covalently like retinal?) to at least one of the helices. It should be noted that the sidechain of thapsigargin does not have double bonds, making it more flexible than farnesol. Neither does it have methyl-groups. This part of the molecule may not be of prime importance for binding of thapsigargin to SERCA and blocking it.

Farnesol has also been reported to inhibit L-type Ca^{2+} channels in vascular smooth muscle cells without affecting membrane fluidity or Ca^{2+}-ATPase activities (in vitro).

Prenylation

Because of the importance of prenylation in the fast-aging Hutchinson-Gilford Progeria syndrome, a short explanation may not be superfluous. Prenylation, a post-translational modification of proteins, involves the enzymatic attachment of a farnesyl-PP or geranylgeranyl-PP moiety to a CAAX motif at the C terminus of a protein (Fig. 9) (Wikipedia: Prenylation). Such moieties are lipophilic. The enzymes involved are farnesyl transferase or geranylgeranyl transferases.

The best known prenylated proteins include the oncoprotein p21ras and other low molecular weight proteins GTP-binding proteins, the nuclear lamins, and the γ-subunit of the heterotrimeric G proteins many of which play a role in GPCR-functioning.

Protein prenylation facilitates the anchoring of proteins into the cell membrane and mediates protein-protein interactions. The alternation of single and double bonds suggests a moderate degree of flexibility in such adhesion system.

Isoprenylated proteins have also key roles in cytoskeletal function, vesicle trafficking and protein functionality, protein maturation, cell growth and in

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Figure 9. The 2 major molecules instrumental to Prenylation. See also Fig. 6 for the horseshoe-shaped configuration of farnesol.
signal transduction via some GPCRs, this list not being exhaustive. In a medicinal context, they seem to have a central role in some types of cancer, and they are likely involved in the pathogenesis of and progression of atherosclerosis, Alzheimer disease, and in the Hutchinson-Gilford Progeria syndrome. In the Discussion, the idea is forwarded to include the gluing of several helices of a multihelix transmembrane protein in prenylation, under the name “intramolecular prenylation,” thereby complementing the classical “intermolecular prenylation.”

The Hutchinson-Gilford Progeria fast aging syndrome in humans: a role for prenylation

Classical analysis of the syndrome

As cited in the introduction, Progeria is an extremely rare (estimated at 1 case in 4 million births) disorder in which symptoms resembling aspects of aging are manifested at a very early age (Fig. 10). The first symptoms develop during the patient’s first few months, indicating that such mutants do not have a normal juvenile stage in their development. Apparently, the juvenilizing factor or mechanism, whatever its nature, is missing.

The cause of the syndrome resides in a mutation in a lamin gene, more specifically at position 1824 of the LMNA gene, thus in a protein that contributes in the nucleus to the formation of the nuclear lamina. Lamins not only give support to the nucleus (Fig. 11), they also help to organize RNA and DNA synthesis.

A-type lamins also have a key role in the maintenance of telomere structure, length and function and they also play a role in the DNA damage response pathway. In normal, non-mutated individuals, the LMNA gene codes for a structural protein called prelamin A. Like all other proteins it is made in the cytoplasm. Here it undergoes a series of processing steps before ending up in its final form, called lamin A, inside the nucleus. One of the steps involves prenylation of prelamin A. This means that the protein undergoes a modification by which a farnesyl- or a geranylgeranyl group is attached. Such lipophilic group helps to associate/attach the protein to membranes. It makes it possible, in proper conditions, to influence other proteins that already reside in the membrane. In the case of prelamin A, the prenylation allows prelamin to temporarily attach to the nuclear rim. Once the protein is attached, it is cleaved by a protease thereby removing the farnesyl group along with a few adjacent amino acids. The truncated protein is now named Lamin A. Lamin A, along with 2 other members of this protein family namely lamin B and lamin C, make up the nuclear rim.

In progeria a mutation generates a cryptic splice site within exon 11. This results in the formation of an abnormal variant of prelamin A from which the farnesyl group cannot be removed. It has been named progerin. It remains permanently attached to the nuclear rim and does not become part of the nuclear lamina. That is thought to be the cause of the abnormal shape of the

Figure 10. Extremes in longevity: insects vs. patients with the Hutchinson-Gilford Progeria syndrome (modified after ref. 15).

Figure 11. Nuclear lamina (from Wikipedia: Nuclear lamina). The localization of lamins just underneath the nuclear envelope is shown in an electron-micrograph and in a schematic representation. For full details, see Wikipedia.
nuclei (Fig. 10). Progerin production is activated in typical human senescent cells.

When the nuclear lamina is deficient, normal chromatin organization and mitosis are likely compromised. To date, it is not known whether or not lamins play a role in enabling the nucleus to control its own ionic environment.

**Something overlooked in the lamin A mutation-explanation in Progeria? A thapsigargin-like effect on $Ca^{2+}$-homeostasis?**

Because farnesyl-groups are lipophilic, prenylation helps prelamin A to bind to the nuclear membrane. Like all membranes of the endoplasmic reticulum, the nuclear membrane also contains $Ca^{2+}$-ATPases. It is thus theoretically possible that the prenyl-group comes in contact with the binding site for thapsigargin on the SERCA-pump, and who knows, also inhibits or overstimulates to some extent the SERCA pumps present in the nuclear envelope.

In my opinion, the misshapen abnormal form of the nucleus may by itself be insufficient for accounting for the numerous symptoms in Progeria. The fact that in Progeria patients mitosis goes on for some years indicates that other mechanisms are playing a role. Furthermore, lamin A does not seem to be necessary for life: it is rather the build-up of prelamin A in the wrong place that matters.

Maybe one of the prenylated forms of prelamin causes thapsigargin-like effects. There may be other as yet unknown endogenous metabolites that might compete with endogenous sesquiterpenoids for binding to their receptor site. If that indeed occurs, treatment with highly active farnesol-esters, like juvenile hormone and its thousands of synthetic analogs, might be tried in rescuing experiments. Treatments with drugs that target some aspects of prenylation, e.g., statins, have been tried, some with positive results. However, no treatment has as yet been proven fully effective.

**The opposite of Progeria: Extra-long larval/ juvenile life in insects**

With respect to aging, insects have 2 features that make them into better models than mammals. First, insects usually have a long lasting juvenile life (as larvae) as compared with the duration of their adult-reproductive life. Passing through a series of larval stages with intermittent moults, enables the larvae to stockpile substantial amounts of reserves. This is beneficial for producing either many eggs, or one or a few very big eggs or a larva (e.g., in tsetse flies). By contrast, the juvenile state of mammals, thus the period before gametes start being shed, is short as compared with their adult-reproductive life. In humans, larval life ends around the age of 12 y (puberty).

The extraordinarily long larval-juvenile life of insects is enabled by the ongoing production of juvenile hormone (JH). The rule is that as long as the JH titre remains high, a larva will moult into another larval instar. It has been argued before that the key role of high titres of farnesol/FLS in maintaining the juvenile state is to keep $[Ca^{2+}]_i$ low enough to prevent the induction of $Ca^{2+}$-induced programmed cell death/apoptosis. At a given moment in their larval development, they stop producing farnesol and juvenile hormone, they degrade the remaining JH that circulates in the haemolymph, thereby creating a situation of complete absence of farnesol and JH, thus of endogenous sesquiterpenoids. That drop to zero initiates metamorphosis, the onset of fast aging in insects. This is a major argument in favor of a crucial role of farnesol/FLS in promoting some aging effects. Such a drop to zero of farnesol/FLS is unknown in vertebrates. This means that in vertebrates FLS are thought to be always present and to be devoid of any hormonal function. However, that does not exclude that they act as “inbromes.”

**Chemical nature of arthropod Juvenile hormone**

The structure of the first JH was elucidated by Röller and Dahm (Figs 5 and 6). Later it was discovered that, depending upon the species, JH occurs in 6 different isoforms. All natural JHs are esters of the endogenous sesquiterpenoid farnesol. Like farnesol, they are horseshoe-shaped, and they are moderately flexible (rotatable bond count of JH III is 8). Wigglesworth assayed 42 compounds for juvenile hormone activity in a typical insect bioassay. He found that the natural hormone of cecropia (= Juvenile hormone I or the DL-, trans, trans, cis isomer) was the most active one. He also emphasized the importance of the trans, trans configuration at C-2 and C-6 in all the compounds, as well as the importance of a suitable balance between apolar and polar properties. This resembles the situation of retinal in some rhodopsins as cited before. In addition to the natural JHs, many JH-analogues have been synthesized during the past decades. The main purpose was to find novel insecticides (acting as growth regulators). This effort yielded a few commercial insecticides.

**JH-receptors: A still ongoing search**

Since 1968, numerous studies aiming at unravelling the mode of action of JH have been undertaken, with
moderate success. Interesting results have been obtained with respect to some proteinaceous (nuclear) receptors (Methoprene-tolerant, Gce).\textsuperscript{9–14} However, it is known from electrophysiological studies that JH can exert very fast effects (in seconds).\textsuperscript{108,109} Such effects can only be generated via membrane receptors. Despite numerous efforts, the isolation of the membrane receptor(s) for juvenile hormone (JH) of insects) and retinoic acid by chromatographic techniques has as yet not been successful.

\textbf{The mode of application, injection or topical application, matters}

One of the reasons of the failure is that JH “sticks to everything,” even to glass, a common saying in the 1960–80ties. But, sticking to everything thereby gluing together transmembrane protein loops may be its overlooked key function. In my opinion, Wigglesworth\textsuperscript{48} was very forethoughtful when he wrote that the juvenile hormone influences the gene-controlled enzyme system within the cells by regulating permeability reactions in cellular membranes. More specifically, he hypothesized that “lipoprotein membranes could be the basic structure concerned, and that the effectiveness of substances with JH activity may rest upon their capacity for gaining access to membranes of oriented lipid.” In an evolutionary perspective, if farnesol substitutes for retinal of prokaryotes as hypothesized in this paper, the binding of FLS in a pocket in a complex multi-helix transmembrane protein may, perhaps, be of the same type as that of retinal in rhodopsins.

When testing JH- and juvenoids for their physiological effects, researchers have to keep in mind that to be physiologically active, the way of application of farnesol/FLS is very important. In insects, injection of farnesol and some other JH-analogs (emulsions prepared with a variety of wetting agents) often gave negative results, while topical application (\(=\) to the surface of the (abraded) cuticle) was very effective.\textsuperscript{48} The best results were obtained if FLS are administered as a slow release formula, e.g., by mixing them with some oil, e.g., arachis oil. Apparently, a relatively long exposure to the active compound is needed for a physiological response, but it is unclear what the underlying principle might be. My hypothesis is that to be active, JH must be able to operate in a well-defined cyto-architecture, e.g., in an epithelium with internal membranes containing the JH-receptor(s). It may take time (days) to realize such structure. To the contrary, the effects of total absence of JH as is typical at the onset of metamorphosis (e.g., on the protein composition of the haemolymph) usually become visible in less than a day.

\textbf{Is the thapsigargin receptor also the membrane receptor for farnesol/FLS/juvenile hormone? The missing link with \(\text{Ca}^{2+}\)-homeostasis?}

A positive answer to these questions has already been suggested before.\textsuperscript{15} In short: as already cited before thapsigargin is, like JH, also a sesquiterpenoid. In my opinion, the most logical explanation why thapsigargin binds to the SERCA pump is that it probably displaces an endogenous farnesol-like sesquiterpenoid from its natural binding site on the SERCA-pump. Both absence of JH and presence of the SERCA \(\text{Ca}^{2+}\)-blocker thapsigargin induce apoptosis, probably according to the principle of \(\text{Ca}^{2+}\)-induced apoptosis (in selected tissues).\textsuperscript{18} However, experimental evidence that FLS bind to the thapsigargin pocket still needs to be forwarded. Kenneth Davey,\textsuperscript{109,110} using a completely different approach, also came to the conclusion that the membrane receptor for JH (and triiodothyronine as well), may be a \(\text{Ca}^{2+}\)-ATPase. The membrane receptor for JH is probably a promiscuous receptor, given the fact that about 4,000 synthetic bio analogues of JH are active in various JH-bioassays.\textsuperscript{15,103}

\textbf{Discussion}

As so often in science, serendipity may open novel ways of thinking. My interest in the role of \(\text{Ca}^{2+}\) in cellular aging was triggered by the study of how, during insect metamorphosis, the absence of juvenile hormone, an endogenous sesquiterpenoid, that is an ester of farnesol, induces apoptosis in some tissues. Such effect can also be induced by the SERCA pump blocker thapsigargin, which is also a sesquiterpene. Because Primordial Signaling Pathways in cell physiology tend to be very well conserved in evolution, comparing data from prokaryotes, invertebrates and vertebrates in an evolutionary perspective may yield novel insights.

\textbf{What characterizes the “juvenile state”?}

The juvenile state precedes the reproductive state at which aging is usually thought to intensify. What changes at the transition? It is difficult to answer this question univocally in organisms in which this transition is smooth and slow, like in invertebrates. In insects, in particular in species which undergo a complete metamorphosis, this transition is abrupt. It starts early in the last larval instar. In many species it is completed in a few days or weeks. In insects, the juvenile state is characterized by a low \([\text{Ca}^{2+}]\).\textsuperscript{15} There is no reason to assume that this should not be the very essence of being a juvenile in vertebrates as well.
Why vertebrates do not need a “juvenilizing hormone”

From an evolutionary perspective, it is intriguing that in contrast to annelids, arthropods/insects and probably other less well studied invertebrates as well, vertebrates do not seem to need a specific hormone for realizing a juvenile stage in their development. This is indeed intriguing because they both descend from common prokaryotic and eukaryotic ancestors. As already outlined in the Introduction, in my opinion the sequence of events was: first some prokaryocytes started to use retinal as a necessary cofactor in proto-rhodopsin, not as a means to perceive light as such, but as means for harvesting solar energy for their metabolism. Later, some early eukaryocytes lost the capacity for synthesizing retinoids, but they could overcome this loss by synthesizing sesquiterpenoids in the mevalonate biosynthetic pathway. Archaeal microbial rhodopsins were probably proton pumps, and had a 7 transmembrane helix configuration. Some of such rhodopsins mutated into bacterial rhodopsins that transport other inorganic ions (e.g., Na⁺, Cl⁻), into the common ion pumps of contemporary eukaryotes (Ca²⁺-ATPase, Na⁺+K⁺-ATPases etc.), and, last but not least into a huge variety of GPCRs. Thus, many different proteins share a common origin and common structural and physiological features. Hence the possibility exists that a given controlling agent concurrently acts on several functionally related proteins, e.g., actors in Ca²⁺ homeostasis.

Farnesol extracted from mammalian tissues has juvenile hormone activity. The reason why vertebrates do not have to use farnesol/FLS as a hormone is that they can synthesize farnesol/FLS in all their tissues, not necessarily in similar (relative) amounts. Thus, farnesol/FLS can end up in the membranes of the cells in which they are synthesized: no need for transport as hormone. This does not exclude that farnesol that is present in blood can be taken up in some cell types. Experimental data are missing. In larval insects, only the corpora allata, which are tiny glands produce JH. It is transported via the haemolymph to other tissues. Thus, the difference between insects and vertebrates with respect to the role of farnesol/FLS in realizing the juvenile state does not seem to reside in major differences in the chemical nature of the family(ies) of controlling agents, but in the way these compounds are delivered to the membrane system of target cells.

Calcitox: A counterintuitive idea

The emphasis that in teaching and in vulgarizing advertising is given to the beneficial effects of calcium (calcareous skeleton, egg shell, milk, sexual differentiation, Calcigender, as secondary messenger etc.) strengthens the perception that it is unlikely that Ca²⁺ might be toxic and/or causal to major diseases and to aging. To non-specialists in Ca²⁺-homeostasis, it may require a drastic switch in thinking to accept that Ca²⁺ is the most ubiquitous toxic pollutant on earth, and that it can serve as a second messenger in cell signaling, undoubtedly a beneficial activity, because it is toxic. In fact, its beneficial effects (at the organismal level) are means to get rid of excess toxic Ca²⁺. It is only at very low cytoplasmic concentrations, up to about 100 nM, that Ca²⁺ is not toxic. Toxicity starts playing a role when [Ca²⁺]i rises too frequently too high.

It is difficult to unequivocally answer the question from which Ca²⁺-concentration on damage occurs. It depends upon the situation: which cell/tissue types in which environment and in which stage of development? Indeed, the Ca²⁺-homeostasis system of any cell is subject to change lifelong. The toxic effects likely occur when there is a Ca²⁺-explosion or a serious of puffs. Because puffs do not last long, it will be rather the frequency of the puffs that is more important for the accumulation of damage, e.g., to the cytoskeleton. For example, the fall to zero of the FLS titer at the onset of metamorphosis of holometabolous insects induces (Ca²⁺-induced) programmed cell death in all tissues that actively secrete proteins. Meanwhile, tissues that are less active in the production of proteins for secretion such as nervous tissue are not affected. The nervous system continues to further develop normally during metamorphosis. Thus the question to which degree Ca²⁺ is causal to aging and more specifically to neurodegenerative diseases, needs to be answered with caution.

The cytoskeleton: A key target of Ca²⁺

Changing concentrations of Ca²⁺ influence a lot of cellular processes, ranging from the synthesis of proteins, lipids, and steroids, over contributing to controlling chromatin structure and the shape of the cytoskeleton, to induction of Ca²⁺-induced apoptosis, this list being far from exhaustive. The most obvious effect of rising Ca²⁺ concentrations is muscle contraction. However, not only the cytoskeleton of muscle cells but of all cell types reacts to changes in [Ca²⁺]i. Many transmembrane proteins are somehow anchored to elements of the cytoskeleton. If an activity cycle of such a protein results in the local release of Ca²⁺ from an internal storage site, local contraction of the peripheral cytoskeleton may result, with possible displacement of attached proteins as a result. This is well documented in egg activation. In internalization of membrane receptors, a similar mechanism may operate. In a recent paper, it has been pointed
out that there are good arguments to attribute a major role to the plasma membrane-cytoskeletal complex, and in particular to the electric phenomena that continuously take place in that system, for storing and retrieving cognitive information in the as yet enigmatic cognitive memory system. Problems with the cognitive memory system are one of the aspects of aging.

**Prenylation: inter- and intramolecular prenylation?**

*A widespread mechanism in multi-helix solute-transporting membrane proteins?*

In endocrine studies on the mode of action of juvenile hormone, farnesol and some GPCRs, little attention is given to the importance of prenylation. Yet, it may be an important issue not only for connecting different proteins, e.g., a γ-type G-protein to an ion channel. Indeed, such a connecting/gluing function may also be at work in selected multi-helix transmembrane proteins, in particular to keep the transmembrane helices that form the core of an ion-pumping unit (e.g., TM3, TM5, TM7 in the SERCA pump) or the helices involved in H+-pumping in rhodopsins back into the original relaxed position after each pumping cycle. In such cycle the helices of the active core move away from each other for a short while and distance. They have to be brought back into their “relaxed positions” to prevent uncontrolled flow of ions through the membrane. It is striking that the physiologically active stereoisomers of farnesol and juvenile hormone all have a horseshoe-shape and a moderate flexibility (rotatable count number around 5). It is also striking that *trans, trans, cis* farnesol is more active in specific JH-bioassays than other stereo-configurations. Acting as a hydrophobic molecular spring in a multi-helix transmembrane protein is a new idea in Ca2+-homeostasis and in endocrinology. Perhaps, the term **intramolecular prenylation** may apply to such activity.

**Are FLS with their link to Ca2+-homeostasis causally relevant, directly or indirectly, for some neurological and other disorders? Alzheimer, Progeria, Parkinson, Obesity**

Progeria (Hutchinson-Gilford) and Alzheimer are diseases for which there is not yet a (pharmacological) cure. The situation is less dramatic for Parkinson (at least temporarily) and obesity. For diseases in which dysfunctional Ca2+-homeostasis is thought to play a role, e.g., Alzheimer disease (AD), or Parkinson, farnesol/FLS attracted only minor attention. The reason may be that up to a few years ago the hypothesis that endogenous Farnesol/FLS sesquiterpenoids likely act as necessary adjuvants of the SERCA-pump that help to keep the cytoplasmic Ca2+ concentration low, these compounds were not considered as key players in Ca2+-homeostasis.

With respect to AD, Liang et al. reported that: “Dysregulation of intracellular Ca2+-signaling has been observed as an early event before the presence of clinical symptoms of AD and is believed to be a crucial factor contributing to its pathogenesis. The progressive and sustaining increase in the resting level of cytosolic Ca2+ will affect downstream activities and neural functions. In AD patients there is an increasing Ca2+ release from the endoplasmic reticulum. Hitherto the cause is sought in mutations in the presenilin genes and amyloid-β oligomers.” I suggest that one should consider the possibility that the early cause of AD may, perhaps, be due to a decrease in the production of endogenous farnesol/FLS sesquiterpenoids, or in the displacement of farnesol/FLS from their natural binding pockets on the SERCA pump by lipoic competitors that can bind to the same pocket with higher affinity. Why not give a chance to think about AD as an FLS-deficiency syndrome or as a sort of thapsigargin-like poisoning by some as yet unknown metabolite?

Excessive lipid deposition in some tissues with obesity as a result, is a normal physiological process in insects and other animals that prepare for hibernation, diapause and metamorphosis. In some insects, a major causal agent is the *disappearance of juvenile hormone from the blood*. The explanation may be that obesity is a normal result of a diminishing Ca2+-gradient over the membranes of the SER. As long as the SER is fully loaded with Ca2+, the enzymes needed for lipid and steroid biosynthesis are inhibited. When Ca2+-release is triggered by e.g., some hormones, the inhibition is temporarily lifted and synthesis of some lipids and steroids can proceed.

**Perspectives for pharmacology**

Drugs for curing the cited aging-related diseases are still missing. It is unclear how to come up with novel approaches. In the sesquiterpenoid domain of pharmacology, it is probably not widely known that during recent decades, over 4,000 synthetic juvenoids (= analogues of insect juvenile hormone) have been synthesized. The purpose was to find novel insecticides that act as insect growth regulators. Many of these analogues are not toxic to vertebrates. Only a few compounds made it into a commercial insecticide. The other compounds are by now largely forgotten. Perhaps, some may fit in the thapsigargin pocket and act as activators of the SERCA pump to keep [Ca2+]i low. To my knowledge, these compounds were never tested for possible beneficial medicinal effects. Given the fact that the pioneers in
this domain, if they are still alive, have passed the retirement age, not much time is left to validate some of their knowledge and recover some of the synthesized compounds.

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No potential conflicts of interest were disclosed.

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