RESUMEN:

Se revisan las evidencias que apoyan la "Hipótesis de Contaminación de Membrana" (HCM) propuesta por Ulrich Altrup (1943-2007).

Altrup centró sus estudios en los mecanismos celulares de la principal manifestación electrofisiológica de la actividad epiléptica-las Despolarizaciones Paroxísticas (DPs). Sus estudios mostraron que: i) aun cuando la epilepsia se manifiesta en una intrincada red de neuronas y glías, una DP puede inducirse en una neurona aislada; ii) las DPs se acompañan de un aumento en la resistencia de membrana; iii) las DPs no son potenciales post sinápticos excitatorios gigantes, sino potenciales marcapasos gigantes; iv) la maquinaria sináptica no necesariamente está afectada durante la epileptogénesis, pero las entradas sinápticas a las neuronas marcapasos epileptizadas pueden modular el desarrollo de las DPs.

Respecto a los mecanismos básicos, la HCM asume que: i) la actividad epiléptica resulta de la inserción descontrolada de sustancias anfífilicas en la membrana celular, conduciendo a un incremento en la presión lateral que daña mecánicamente diferentes canales iónicos, conduciendo a DPs; ii) las células gliales desempeñan una función de "descontaminación";
iii) las neuronas epilépticamente activas liberan sustancias intracelulares desde sus dendritas activando inespecíficamente las neuronas circundantes; iv) la sincronización neuronal aparece después de la despolarización epiléptica y pudiera ser la base de un foco epiléptico.

La HCM contraviene algunos puntos de vista generalmente aceptados que conciben la epilepsia como un fenómeno a nivel de red neuronal con la DP como un potencial postsináptico gigante. Al mismo tiempo, es capaz de explicar la mayoría de los hechos conocidos acerca del desarrollo de la epilepsia y concuerda con las opiniones recientemente expresadas respecto a las insuficiencias de la concepción generalmente aceptada de la epilepsia como un desbalance excitación/inhibición. Lamentablemente, esta hipótesis es poco conocida entre los investigadores.

PALABRAS CLAVE:

SUMMARY:

Evidences supporting the "Membrane Pollution Hypothesis" (MPH) for epileptogenesis - proposed by Ulrich Altrup (1943-2007)- are reviewed.

Altrup centered his research on cellular mechanisms for the electrophysiological hallmark of epileptic activity: Paroxysmal Depolarization Shifts (PDS). His research led to the following observations: i) Even when epilepsy takes part in an intricate neuronal-glial network, a PDS can be elicited in an isolated neuron; ii) PDS is accompanied by membrane resistance increase; iii) PDSs are not gigantic excitatory postsynaptic potentials, instead, they are gigantic pacemaker potentials; iv) Synaptic machinery not necessarily is modified during epileptogenesis, but synaptic inputs to epileptized pacemaker neurons might modulate PDS development.

Regarding basic mechanisms, the MPH assumes that: i) Epileptic activity results from uncontrolled insertion of amphiphilic substances into the cell membrane leading to an increased lateral pressure in the membrane and thus to mechanical damage to different ionic channels, which leads to PDSs; ii) Glial cells play a "housekeeping" role in this process; iii) The pacemaker neuron when epileptically active releases intracellular substances from its dendritic surface on to neighboring neurons which are unspecifically activated; iv) Neuronal synchronization follows the epileptic depolarization and this might be the basis for an epileptic focus.

MPH counters commonly held views about epilepsy as a network phenomenon with the PDS as a gigantic postsynaptic potential. At the same time, it can explain most known facts about epilepsy development and it is congruent with recently expressed views about drawbacks in the usually accepted hypothesis of epilepsy as an excitation/inhibition misbalance. Unfortunately this hypothesis is rather unknown among researchers.

KEY WORDS:
INTRODUCCIÓN

About 0.5% of humankind suffers from epilepsy. Fortunately, available treatments are capable of keeping most of persons with epilepsy free from seizures, but still little is known about the basic mechanisms involved in epileptogenesis.

Theories about epilepsy are as old as philosophical thought. Worth of notice is the treatise about Epilepsy by Hippocrates (c. 460 BC–c. 370 BC) who denied any supernatural origin to the disease:

"It appears to me to be nowise more divine nor more sacred than other diseases, but has a natural cause from where it originates, like other affections."..."The brain is the cause of this affection"..."In this disease as in all others, [the doctor] must strive not to feed the disease, but endeavor to wear it out by administering whatever is most opposed to each disease, and not that which favors and is allied to it...without minding purifications, spells, and all other illiberal practices of a like kind."

It seems obvious that for understanding epilepsy, the basic functioning of the brain needs to be apprehended, and the way for it was paved only after the advent of the scientific method during the so called Scientific Revolution of the 17th Century. Still, the main heroes of that revolution were quite "naive" in their ideas about brain function. Thus Descartes, the genial creator of the orthogonal coordinate system, did not consider it necessary to make any observation or experimentation to discover how does the brain work, he simply knew how the brain must work. This passage from "Passions of the Soul" (1649), illustrates how the imagination of a genius can distort reality:

"...and that this new blood that has just been formed from the alimentary juice, being driven to the heart with greater force than the blood from other parts of the body, enters the heart in greater abundance and produces a stronger heat there because it is coarser than the blood that has been rarefied many times in passing repeatedly through the heart. This makes it send to the brain spirits with unusually coarse and agitated parts; and these spirits, by strengthening the impression formed by the first thought of the loved object, compel the soul to dwell on that thought. This is what the passion of love consists in."

Newton, "the greatest genius of all times" developed a speculative scenario for nerve conduction. In "Opticks" (1704), Newton stated:

"...Vibrations, being propagated along the solid Fibres of the optick Nerves into the Brain, cause the Sense of seeing. For because dense Bodies conserve their Heat a long time, and the densest Bodies conserve their Heat the longest, the Vibrations of their parts are of a lasting nature, and therefore may be propagated along solid Fibres of uniform dense Matter to a great distance, for conveying into the Brain the impressions made upon all the Organs of Sense...I suppose that the Capillamenta of the Nerves are each of them solid and uniform, that the vibrating Motion of the Æthereal Medium may be propagated along them from one end to the other uniformly, and without interruption."

On the other hand, Isaac Newton suggested that the speed of propagation of nerve activity is very high (like that of light).
These views, expressed by the greatest knights of the scientific realm, influenced, and perhaps hindered, the advance toward a better understanding of nervous function.

Thus, when Hermann Helmholtz tried in 1850 to measure the speed of nervous impulse, he found himself discouraged by those who claimed that a process moving at the speed of light and manifested at the centimeter range, lays in the nano-second time scale and it would be practically impossible to detect using the technology available in middle of 19th century.

Helmholtz decided to challenge the infallible Newton, and found experimentally that the velocity of a nerve impulse is 27 m/s, (surprisingly agreeing with modern estimations). This value is even one order of magnitude lower than the speed of sound. This implied, that nerve impulse conduction was based on an unknown for that moment mechanism5.

Second part of 19th century marked important advances in clinical description and cataloguing of epilepsy modalities, but little was clarified regarding putative neurophysiological bases, besides Huggins Jackson’s view that epilepsy "represents a sudden, excessive, disordered and recurrent discharge of cortical neurons"

Advances in understanding the bio-physical bases of nerve conduction allowed the proposal of physiologically sound hypotheses about epileptogenesis6.

In particular, thanks to a better understanding of synaptic mechanisms as well as the important role of inhibitory processes in brain function, the hypothesis of epileptic activity as a misbalance between excitation and inhibition had loomed. Nowadays, among a great diversity of proposed mechanisms, the idea of excitation/inhibition misbalance appears as the first-choice hypothesis for epileptogenesis.

What follows is a brief summary of prevailing ideas about epilepsy mechanisms as suggested by Maya Entenza (2010) in his excellent treatise6:

\[ \text{Epileptogenesis is the process whereby a normal structure becomes hyper-excitabile up to spontaneously elicit an epileptic seizure.} \]

\[ \text{The normal functioning of the brain can be viewed as the result of the interaction of a high number of neuronal circuits where millions of neurons are interwoven. Neuronal networks, both in cortex, in hippocampus and the thalamus arise from the interaction of excitatory and inhibitory neurons, acting in a shrewdly established arrangement. Such balance is kept because the circuits are organized in such a way that excitatory and inhibitory inputs develop collateral connections with local interneurons, whereby inhibitory circuits -both anterograde and retrograde-are established that locally limit the excitation level. This way of functioning warrants that networks are self-regulated, since an increment or a reduction in excitation is accompanied by an increase or a decrease in inhibition. At the cellular level a fine balance is also kept between excitation and inhibition, and each individual neuron is endowed with calcium and sodium-dependent membrane currents, especially at the level of dendrites, whose goal is to potentiate low amplitude synaptic inputs. These depolarizing currents are countered by voltage-dependent potassium currents activated by calcium, thus controlling the membrane potential, which prevents an exaggerated depolarization of the nerve cell.} \]

\[ \text{According to the prevailing conceptions, the basic problem of epilepsy consists in understanding how and why a sudden and transient impairment of brain activity is} \]
produced, triggered by the rhythmic and synchronous discharge of a neuronal population, the pathogenic substrate of the epileptic event. This abnormal and paroxysmal discharge is clinically expressed in different ways, originating a wide repertory of clinical manifestations of the epileptic seizures. It is important to determine in each case, what caused such misbalance.

Most of the hypotheses and models of mechanisms for epileptogenesis arise from animal experimental models, where the mechanisms of paroxysmal discharges appear at the hippocampus after the injection of GABA-antagonists.

Since 1960’s epileptologists are trying to understand cellular mechanisms for epilepsy. High amplitude intracellular depolarizations, known as paroxystic depolarization shifts (PDSs), are followed by a hyperpolarization whose EEG counterpart is the wave following a peak discharge. In most of the studied neuronal groups, the PDS corresponds to a huge excitatory post synaptic potential mediated by glutamate and/or aspartate. To the depolarization resulting from the excitatory post synaptic potential the activation of voltage dependent ionic currents is added, mainly carried by sodium and calcium. The hyperpolarization following afterwards is originated by GABA-dependent currents as well as by intrinsic currents from the neuron, mainly potassium and potassium-calcium dependent.

Epilepsies are characterized by the capability of certain neurons to produce (PDSs). PDSs are initiated with a depolarization of the neuron that reacts with a high-frequency burst of action potentials, accompanied by a sustained depolarization, usually followed by a hyperpolarization of the neuron, finally returning to the resting potential. The initial phase of this burst is attributed to the activation of sodium channels associated to Kainate- or AMPA-type glutamatergic receptors, this leads to a fast inflow of sodium ions, depolarizing the cell. The sustained depolarization and the high-frequency burst are attributed to the activation of NMDA receptors, which produce a slow influx of calcium ions, as well as voltage-dependent calcium currents. The following sustained hyperpolarization has a fast component associated to the activation of GABA-A receptor associated chloride channels, and a slow component due to the activation of GABA-B receptor associated potassium channels; under normal conditions, these prevent the spread of the discharges into other brain areas.

Neuronal synchronization during inter-ictal spikes as well as during the seizure are due to the existence of recurrent excitatory connections. However, other mechanisms may also be involved, among them: i) Electric coupling between neurons; ii) Extracellular ions concentration shifts; iii) Diffuse release of neuro-modulators; iv) Ephaptic interactions. Among the inhibitory mechanisms that might be impaired during epileptogenesis, there are: i) Blocking by Mg++ of NMDA-receptor-associated channels; ii) Na,K-ATPase activation with subsequent calcium outflow and after-discharge hyperpolarization; iii) Clearance of excess potassium and neurotransmitters by glial cells.

Hyperactivity of the excitatory system might lead to calcium inflow, dysfunction of the mitochondrial chain, lowered ATP production, lipases- and proteases activation, fatty acids release (which are substrates to synthesis of prostaglandins, leucotrienes, free radicals, and others). The activation of fast response genes might lead to apoptosis.

The above described picture is the result of intense research in brain mechanisms, observations made on persons with epilepsy as well as animal model research by many
generations of epilepsy researchers.

Nevertheless, almost every piece of this proposed scenario is a matter of deep debate among specialists. Thus, a long standing discussion is related to whether epileptic activity takes place at the level of one neuron or is it manifested at single neuron level. Thirty years ago, one of us, also defended the idea that the minimal physiological entity capable of producing epileptic activity is the cortical column\textsuperscript{7}. Today, this position is still held by most authors, for example, as Goldberg and Coulter\textsuperscript{8} expressed in 2013, "Epilepsy is inextricably a circuit-level phenomenon and cannot be understood outside this context". As it will be shown later, this view might be not as solid as it appears at first glance.

Today, new hypotheses, involving newly described molecular protagonists are being proposed, however, most of them rely on the classic idea of epilepsy being caused by a misbalance between excitation and inhibition. Thus for La Sarge and Danzer\textsuperscript{9} "Altered excitatory/inhibitory balance in the brain, in this case favoring increased excitability via reduced inhibition, might be an important mechanism by which changes in mTOR signaling promote epileptogenesis".

Similarly, Lason et al. stated\textsuperscript{10}: "Neurochemical approach strongly supports the electrophysiological findings that seizures can be generated from excessively enhanced excitatory processes …or from hypoactivity of neuronal inhibition, hyperactivity of glutamatergic transmission functional disturbances of the ligand- or voltage-gated sodium and calcium channels, insufficient GABAA receptor-mediated neurotransmission and extracellular potassium currents".

Huberfeld et al. (2013) addressed the crucial question of why epileptic fits are so rare and unpredictable, and their approach to find an explanation is through a combination of synaptic excitatory/inhibitory balance and ion homeostasis mechanisms\textsuperscript{11}.

At the cellular level, epilepsy studies were benefitted when the theoretical and methodological/instrumental approach introduced by the Cambridge School was applied to find what the neuronal electrophysiological hallmark of epileptic activity is, and which the underlying mechanisms are.

In particular, it was possible to simultaneously obtain an electrocorticogram and an intracellular recording of a pyramidal cell during an epileptic episode in a rat (Figure 1).

As it can be observed from Figure 1, epileptic activity in brain’s cortex is accompanied by high-amplitude and long-duration depolarizations. This is the event coined as "paroxysmal depolarization shift" (PDS).
It seems that understanding the nature of the PDS can be an excellent way to unravel epilepsy mechanisms, in a manner similar to the great boost basic neuroscience experienced after understanding the bases of the action potential. As noticed from the mainstream scenario shown above, most researchers view the PDS as a huge excitatory postsynaptic potential arising from misbalanced synchronic activity of neuronal masses.

The question about the nature of PDSs is also linked to the "epileptic neuron vs. epileptic neuronal network" controversy.

The prevailing opinion, till our days is that the PDS is merely a huge postsynaptic potential taking place in a reverberating pathological network of neurons. This conception fits very well into the widely accepted idea of epilepsy arising from a misbalance between excitation and inhibition.

**Ulrich Altrup and Basic Epilepsy Mechanisms**

Among the alternative hypotheses for epileptogenesis, the name of Ulrich Altrup (1943-2007), deserves a special place. Altrup was a full professor of physiology at the University of Münster, and worked during 35 years at the Institute of Physiology and the Institute for Experimental Epilepsy Research at the University of Münster, Germany. References 13-36 conform a sample of the papers published by him between 1979 and 2006.

Professor Altrup devoted most of his intellectual life to the study of epileptogenesis in the simplest known model for epilepsy: paroxysmal depolarizations shifts elicited in the neuron B3 from the buccal ganglia of the land snail *Helix pomatia* (Linnaeus, 1758), known under the common names of "orchards' snail", "vineyard snail", "Roman snail" or "Burgundy snail"- and probably one of the most known species of terrestrial mollusks.

Molluscan neurons have been a traditional model in neuroscience. In particular, studies by Erick Kandel of memory mechanisms on *Aplysia californica* were awarded with the Nobel Prize in 2000. More widely, invertebrate neurons have often proved useful in the study of basic mechanisms in nervous systems since they offer several decisive technical advantages when compared to vertebrate preparations. Among others are the existence of visually identifiable giant neurons which are well-known neuronal individuals. A further advantage is the structural and functional intactness of the ganglia kept under in-vitro conditions for many hours and even days.

The buccal neural ganglia of Helix contain four easily identifiable giant neurons, called B1, B2, B3, and B4. Among them, only B3 is capable of developing epileptic activity upon the application of proconvulsant substances (e. g. Pentylenetetrazol, Etomidate; Figure 2).
After systemic application of Etomidate (ETO) and Pentylentetrazol (PTZ), mammalian as well as molluscan neurons generate epileptic paroxysmal depolarization shifts.

Both ETO and PTZ elicits PDs in a dose-dependent manner. Full PDSs appear at a ETO- and PTZ- concentrations of 0.8 mM and 40 mM respectively.

Altrup adhered to the working hypothesis that essential parts of the mechanisms underlying epileptic activity are constant irrespective of the considered type of human seizure and irrespective of the studied nervous system. Several observations demonstrate that the epileptiform activities of neurons in the buccal ganglia of Helix pomatia correspond to epileptiform activities recorded in vertebrate preparations and in the human nervous system. As an example, the buccal ganglia of Helix pomatia have been used to develop new antiepileptic substances and it was subsequently shown that the substances which proved antiepileptic in the invertebrate system were also antiepileptic in vertebrate nervous systems.

As expected, PDSs were found to be associated to a long-lasting inward current. However, unlike most of the classically described inward currents, PDS-associated inward current was accompanied by a paradoxical increase in membrane resistance (Figure 3). In particular, this counters to the mainstream idea that the depolarization during a PDS is due to a "fast inflow of sodium ions"
Indeed, the ionic mechanisms of this inward current were not clear\textsuperscript{37}, even when it was initially postulated as being due to "a decreased potassium conductance."

At the same time, PDSs are paralleled by increases in the intracellular concentration of calcium (Figure 4).

The possible synaptic nature of PDSs was ruled out through different means, in particular, PTZ is capable of eliciting PD in isolated \textit{Helix} neurons\textsuperscript{38}.
It was obtained that the effect of both PTZ and ETO upon postsynaptic potentials is completely uncorrelated with respect to their PDS-eliciting actions: ETO does not change the amplitude of excitatory post synaptic potentials whereas PTZ reduces them in a dose-dependent manner (Figure 6). Therefore, an explanation for epileptic actions of these two substances based on their effects upon postsynaptic potentials does not seem to be uncontroverted.

Among neurons B1-B4, only neuron B3 is capable to develop pacemaker potentials. Reports from other labs showed that PTZ is capable of inducing phase-shifting in circadian pacemaker neurons. Altrup explored the effect of PTZ and ETO upon action potential parameters. It was found that PTZ increased the duration of action potentials in a dose-dependent manner (Figure 6).

Further, Altrup studied the effect of convulsant substances upon pacemaker potentials. It was obtained that with the application of an epileptogenic drug, endogenous pacemaker potentials develop into PDSs. With increasing concentration of the drug, both the amplitude of pacemaker-depolarizations and delay of pacemaker-repolarization increased progressively finally resulting in PDSs. Only neurons which generated pacemaker potentials under control conditions could generate PDSs under epileptic conditions (Figure 7).
Synaptic input into neuron B3 could enhance or abolish the development of PDs. Specifically, synaptically induced potentials (induced via the stimulation of a nerve or from intra-ganglionic central pattern generators) did not participate in the generation of epileptiform activity but they can trigger both pacemaker potentials and PDSs.

Thus, the wealth of data collected by Altrup and his colleagues significantly modified prevailing ideas about epilepsy.

1. Even when epilepsy takes part in the intricate neuronal-glial network of the brain, the hallmark of epileptic activity -the PDS-can be elicited in an isolated neuron.

2. Unlike the common situation for excitatory processes, PDS are accompanied by an increase in membrane resistance.

3. PDSs are not gigantic excitatory postsynaptic potentials, as usually assumed, instead, they are gigantic pacemaker potentials.

4. Synaptic machinery not necessarily is modified during epileptogenesis. However,
synaptic inputs to epileptized pacemaker neurons might exacerbate or diminish PDS development.

These facts, taken from the detailed study of epileptogenesis in a simple nervous system, on one hand, radically change some of the prevailing views about epilepsy mechanisms. At the same time, it can give a rational explanation to many known facts about epilepsy in humans.

The membrane pollution hypothesis

As it has been shown, processes—such as membrane resistance, action potential duration, pacemaker potential amplitude and duration—are altered by the tested epileptogenic drugs. Some processes showed a threshold concentration of 1 mM for PTZ and 12.5 µM for ETO. A 40-fold increase in concentration of both drugs induced PDSs, i.e., 40 mM of PTZ and 500 µM of ETO. The similarity in the dose dependency for several different neuronal mechanisms (leakage channels, voltage dependent potassium channels, pacemaker potentials to develop into PDS, membrane resistance) led Altrup to the idea that they resulted from an unspecified effect.

A possible explanation for such unspecified action could be through unspecified incorporation of epileptogenic substances into the cell membrane. To test this hypothesis, experiments were carried out on the air/water interface of a Wilhelmy film balance. After forming the phospholipid monolayer on the saline subphase, the surface pressure in the layer was adjusted to 10 mN/m. At this pressure, all phospholipids molecules are in the liquid-condensed state and packed in a way that reasonably mimics the physical state of a biological membrane and still allows a certain rate of lateral diffusion.

Using a push-pull pump, the epileptogenic drugs were added to the saline beneath the phospholipid layer. During injection of the respective drug, pressure in the membrane rose exponentially. With an epileptogenic concentration of PTZ, the pressure increase was 6.9 ± 0.9 mN/m and the epileptogenic concentration of ETO increased pressure by 6.3 ± 0.9 mN/m. Threshold effects of both drugs were 1/40th of their respective epileptogenic concentration. Dose-effect characteristics of incorporation in a phospholipid membrane thus corresponded to dose-effect characteristics of several different neuronal properties (Figure 8).

![Figure 8. Increase in lateral pressure in an artificial membrane upon the application of both PTZ (left), and ETO (right). Abscissas: PTZ concentration in mM, and ETO concentration in µM; ordinates: logarithm of percent change in measured lateral pressure. Values were computed from data provided in 36.](http://biomed.uninet.edu/2018/n1/valdes.html)
For Altrup, all these results together, did not directly show that epileptogenic drugs enter the neuronal membrane and that their incorporation increased membrane pressure and induced PDSs in neurons, which have expressed the membrane mechanisms of pacemaker potentials. At the same time, taking into account the bulk of observations here summarized strongly suggests that the key mechanism of epileptiform activity evoked by PTZ and Etomidate is their incorporation into the neuronal lipid bilayer, increasing membrane pressure and affecting conformational movements of membrane proteins. The amphiphilic drugs PTZ and Etomidate could attach to the external side of the membrane and then incorporate their lipophilic part into the lipid layer. This should increase membrane pressure and reduce membrane fluidity. Correspondingly, there were effects on membrane mechanisms in all studied neurons (B1, B2, B3, and B4). Epileptiform activity, however, was found in neuron B3 which can generate pacemaker potentials (Figure 9).

Figure 9. Summary of Altrup´s Membrane Pollution Hypothesis for epileptogenesis. Amphiphilic drugs like PTZ and ETO (A) -or byproducts of abnormal brain cells metabolism- are postulated to adhere to the membrane surface (1) and incorporate the lipophilic part thus increasing membrane pressure (2, red arrows). Mechanical effects on the membrane pacemaker machinery are suggested to induce PDS development, and subsequently epileptic activity.

Summarizing, the key-points of Altrup´s membrane pollution Hypothesis are:

- i) Epileptic activity is the result of uncontrolled insertion of substances into the cell membrane leading to an increased lateral pressure in the membrane and by this a mechanical damage to different channels, which leads to PDS.
- ii) Glial cells play a role in this context since they make "housekeeping" and when they cannot do this (e. g. destroyed by Pronase) epileptogenicity increases\(^{28}\).
- iii) The pacemaker neuron when epileptically active liberates intracellular substances from its dendritic surface on to neighboring neurons which are unspecifically activated.
- iv) Neuronal synchronization follows the epileptic depolarization and not (as is generally believed) neurons synchronize and from this generate PDs. This might be the basis for an epileptic focus.

**Aftermath**

Professor Ulrich Altrup passed away in 2007, at the age of 63. It seemed that with him faded the idea of using mollusk neurons to study epilepsy mechanisms, and the membrane pollution hypothesis has not been yet quoted in scientific literature.

On one hand, ideas about epilepsy mechanisms continue to be centered on the assumption of network and synaptically based processes.
On the other hand, Altrup hypothesis has been capable of explaining the bulk of known facts about epilepsy, both clinical and in experimental, and -unlike mainstream views- can provide a reasonable explanation to some well-known facts as the paradox of the favorable effects of ketogenic diet

In recent years mounting evidence is accumulating that reveals the unsustainability of the traditional scenario and claim for the need of alternative hypotheses.

Thus Kaila et al concluded that "Changes in excitation-inhibition (E/I) balance are often used to explain epileptogenesis and seizure generation but, as should be obvious from the work reviewed above, the explanatory value of the E/I balance in the context of epilepsy is limited."

With the advent of Genomic Era, a group of genes associated to epilepsy has been identified. Paradoxically, "The bulk of the mutations associated with epilepsy are not in inhibitory or excitatory ion channels, so the mechanisms by which mutations lead to seizures and epilepsy need to be broadened. It is possible that entirely new mechanisms of epilepsy will be discovered as a consequence of investigations into the cell biology of these newly-discovered gene defects."

Recently, the study of epilepsy mechanisms based on isolated snail neurons has been retaken.

According to Erick Kandel, a good hypothesis is the one that is capable of setting new, experimentally testable questions. In that sense, the membrane pollution hypothesis can pave the way for several questions, such as:

- How can the duration of individual action potentials be increased on the background of an increase of membrane resistance?
- Given a parsimonious mathematical model for pacemaker potential, is it possible to increase its amplitude via manipulating membrane input resistance?
- Which mechanisms is the cell membrane equipped with, in order to fight membrane pollution, and which consequences are expected from weakening or invigorating such mechanisms?
- How does the ketogenic diet influence the physicochemical properties of the lipid bilayer? Is there an alternative antiepileptic diet that can be proposed on the basis of the membrane-pollution hypothesis?
- Are electroshock-induced seizures related to the release of membrane-polluting amphiphilic substances? What about insulin shock-induced seizures?
- Can the membrane pollution hypothesis provide a rational explanation to the wealth of epilepsy-associated genes that do not fit into the mainstream excitation-inhibition misbalance idea?
- Is it possible to find an isolate vertebrate neuron capable of producing PDS?

For now, the authors would like the scientific community to be aware of this nearly forgotten hypothesis that was promoted by a humble researcher who regarded that "We increasingly know more about smaller items even sub-quark particles or conditions, but this appears not to help with the fundamental problems of how the brain is working."
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Este artículo describe principalmente la hípotesis de contaminación de membrana de Ulrich Altrup como mecanismo epileptogénico y la confronta con lo que hoy está más aceptado.

En la actualidad se piensa que la epilepsia se origina por un desbalance a nivel de los circuitos neuronales entre la actividad excitatoria e inhibitoria. En condiciones fisiológicas, un incremento en el nivel de la excitación de las redes neuronales desencadena mecanismos de autorregulación aumentando la actividad inhibitoria de otras redes con las que se interconecta. Las investigaciones a nivel celular se han focalizado en determinar las neuronas capaces de generar paroxismos de despolarización, que no son otra cosa que grandes ondas de despolarización seguidas de una hiperpolarización reactiva. Aunque la aceptación más universal es que el mecanismo epileptogénico es a nivel de una red neuronal y no intracelular.

Altrup mediante sus investigaciones desarrolla su "hipótesis de contaminación de membrana", en la cual postula que ciertas sustancias actuarían a nivel interno de la membrana celular provocando una modificación en su resistencia y generando cambios a nivel de la presión intracelular lateral, lo que finalmente llevaría a una afectación de los canales iónicos ubicados en ella y generarían así una onda despolarizante.

En mi opinión personal ambas teorías pueden complementarse y no son necesariamente excluyentes, es probable que en los próximos años y con las actuales líneas de investigación encontremos una nueva teoría más general y unificadora.

La epileptogénesis es un tema de fundamental interés para los neurólogos. Las convulsiones son uno de los trastornos más impactantes en nuestra práctica y en el imaginario social. Desde la antigüedad y la edad media, en donde a las crisis se le atribuía un carácter divino o demoníaco, estos eventos paroxísticos resultan fascinantes para quienes estudian el campo de la salud y su origen es territorio de grandes controversias.

Como bien revelan los autores, existen múltiples explicaciones, que se basan en la premisa
del desequilibrio entre influencias excitatorias e inhibitorias, ya sea por exceso de las primeras o por falla de las segundas. La "hipótesis de la contaminación de membrana", de Ulrich Altrup, explica varios mecanismos celulares que podrían estar involucrados en el inicio de la actividad epiléptica a nivel neuronal.