

P74 4.0 BESA-Detailed Project Spike-proteins & virus-fragments DNA & Cell Protector Card



Detailed- Project P74 4.0 to BESA expert opinion

Bioenergy informative system analysis within the framework of the BESA seal of approval about the effectiveness of the product "Leela Quantum DNA & Cell Protector Card on Spike-proteins & virus-fragments in the project also called "test object"





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Client:

Leela Quantum Tech, LLC Attn: Eleonora Goldenberg 1421 LUISA STREET, STE G SANTA FEE, NM 87505 USA

Project participants:

Project leader: Wolfgang Hans Albrecht, president and scientific director of the

IFVBESA

Testing person: Eva Krankl, Vice president and deputy scientific director of the

IVFBESA

Respondent: 7 anonymous test persons with different health conditions and

age or gender.

Detail project P74 1.0 Respondent 1: woman aged 58 years, corona unvaccinated,

high vitality level, was tested with digitized mRNA vaccine in

correlation with specific EMSF.

Detail project P74 2.0 Respondent 2+3: The two subjects, female aged 48 years and

male aged 65 years were injected with a so-called Corona

mRNA vaccine.

Detail project P74 3.0 Respondent 4,5 + 6: Woman aged 64 years, man aged 61 years

and woman aged 36 years, all 3 Corona unvaccinated, were

tested for spike proteins and viral fragments.

Detail project P73 4.0 Respondent 7: Subject aged 80 years, 2x Corona mRNA

vaccinated, was tested for the effect of the test object against

spike proteins, virus fragments and graphene oxide.

other participants: none

Project location: location of IFVBESA (international professional association for

bio energie information system analysis), Hauptstraße 1, A-

4861 Kammer/Schörfling am Attersee

Date: 27.10.2021 until 16.11.2021

Project duration: 21 Days

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BESA- legend for interpretation of BESA-measurement result

The measured value of 50 on the tested meridian represents an optimal energetic state in this organ or its subordinate and superordinate levels. Also measured values in the range of 50 to max. 70 still count as a neutral and balanced energy status. The organism is able to regulate irritations of the system (wrong environmental signals) very well.

Readings above 70 to 100 represent the inflammatory range or a so-called energy surplus in response to the irritation of the system by environmental signals corresponding to it. After reaching the maximum values, the energy state tilts into the degenerative (blue) range.

Readings from below 50 to near 0 represent the so-called degenerative measurement range or a lack of energy in response to the stimulation of the system by environmental signals corresponding to it.

Measured values represented by a so-called pointer drop of more than 3 scale lines indicate a total deregulation. The influence of certain environmental signals then leads to such strong system overloads, which can only be harmonized by corresponding new signals.

BESA Indicators:

up to 0,79	very deep energetic regulation disorder (SSD) energy deficiency
0,8 to 1,19	severe energy regulation disorder (SD) degeneration/energy
	deficiency
1,2 to 1,59	energetic regulation disorder (D) degeneration/energy deficiency
1,6 to 1,99	degenerative transition area (DÜ)
2,0 to 2,39	optimal regulation (OR)
2,4 to 2,79	in regulation (R)
	partial inflammation = regional energy excess (PE) total inflammation = strong general energy surplus (TE)



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Basics of the research project creation P74 4.0

The international association for bioenergetic system analysis was commissioned by the company Leela Quantum Tech LLC to test or prove the effect of the test object "Leela Quantum DNA & Cell Protector Card" by means of bioenergetic system analysis (BESA). The testing took place independently of the subjective feeling of the test person.

Description of the "Leela Quantum DNA & Cell Protector Card" by the client:

Our DNA and cells are at the heart of our physical existence, and strengthening and protecting them from potentially harmful substances, influences and external forces is key to our overall well-being. The big questions these days are: Have you been vaccinated or are you being vaccinated? Or do you want to forgo a Corona vaccination, but are concerned that you may ingest or become infected by so-called spike proteins through people who have been vaccinated with an mRNA or vector-based vaccine? In either case, you are concerned about keeping your DNA and cells pure, strong and in the best possible shape. This high power frequency set is a masterpiece that has been developed and tested by the best healers in our network together, for which we are very grateful. It is designed to energize and support body systems and performance. All subjects report that it increases vitality, clarity and well-being. While each person's response to external fields, forces and substances is unique, this set of frequencies has been shown to reduce stress and its influence.

Further, the developer believes that the "Leela Quantum DNA & Cell Protector Card" as follows:

- strengthens the cells
- strengthens the DNA
- creates the awareness in the cells to be able to optimize their function in a natural way, thus building the greatest possible protection against harmful foreign substances and information

Furthermore, the company "Leela Quantum Tech, LLC" claims about tests that were carried out, that side effects caused by the mRNA vaccination on the body or in the brain could be reversed by the "Leela Quantum DNA & Cell Protector Card". Furthermore, it is assumed that these effects should also occur with other similar vaccinations or vector-like vaccinations.

protects unvaccinated people from becoming infected by shedding

Due to its mode of operation, it has a wide range of applications. In the case of this project, the "Leela Quantum DNA & Cell Protector Card" is used for various modern, so-called mRNA or vector-based vaccines and their effect on the human body.

Research Support Services of the IFVBESA - BESA Reference Tests

The project P74 4.0 deals specifically with the proof of concept of the "Leela Quantum DNA & Cell Protector Card" in a subject who was vaccinated with 2x Corona mRNA. The subject suffers from advanced Alzheimer's dementia and hypertension. He takes appropriate medication



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such as donepezil (cholinesterase inhibitor for dementia) as well as anticoagulants (blood thinners) and ACE inhibitors against the increased blood pressure. In addition, the test person is almost permanently in the environment of Corona vaccinated people. Thus, it can be assumed that at least the transmission of spike proteins can occur within these people or groups of people. Special attention is paid in this detailed project to spike proteins as well as virus fragments and graphene oxide, which according to scientific studies (see list of sources on page 59) suppress the repair mechanism of DNA and thus irritate the immune system or cause autoimmune reactions. Spike proteins, virus fragments as well as graphene oxide are digitized and their effect simulated accordingly in the BESA test.

The "Leela Quantum DNA & Cell Protector Card" is tested according to the client's request within the framework of the applicable conditions of the IFVBESA for the award of quality seals. In principle, seals of approval are awarded in three categories depending on the significance of the test results, taking into account all tests of a project. For the "Leela Quantum DNA & Cell Protector Card" it is to be determined whether by its application stresses over already mentioned stress factors like the administered mRNA vaccine to the test person as well as possible correlations of these with certain activated EMSF and in consequence of it in the energy system of the test person (biological system) developing or existing disturbances, problems, blockades, disharmonies can be harmonized, neutralized and thus negative pathological conditions by positive conditions be replaced. This will be questioned in the following commissioned tests of this project.

Research project description

The reason for the test is to prove the functionality of the "Leela Quantum DNA & Cell Protector Card" by test results obtained by exposing the test person to the digitized spike proteins, the virus fragments as well as the graphene oxide in order to significantly prove and compare the reactions without the "test object" and with the "test object". In a BEFORE measurement, the subject is exposed to the listed stress factors and tested. In the AFTER measurement, the subject will be associated and tested with the test object, the "Leela Quantum DNA & Cell Protector Card", in addition to the stated loading factors.

- The BEFORE measurements are performed without the "Leela Quantum DNA & Cell Protector Card"
- The AFTER measurements are performed with the "Leela Quantum DNA & Cell Protector Card"

The question for each AFTER measurement is: "Is the "test object" suitable and able to harmonize or neutralize the so perceived negative effects of the mentioned stress factors on the organism"?

The correspondingly designed tests should provide information about this by comparing the pre-measurement without the "Leela Quantum DNA & Cell Protector Card" with the test



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results of the post-measurements to be carried out using the "Leela Quantum DNA & Cell Protector Card".

The purpose of "Leela Quantum Tech, LLC" is to determine whether the test object, the "Leela Quantum DNA & Cell Protector Card", as noted in the product description, is suitable for harmonizing and/or neutralizing the disturbances, problems, blockages, disharmonies in the meridian system of the test persons (biological object) resulting from the above-mentioned stress factors.

General information transmission of the test object

The information transmission takes place from the hyperspace of the test object to the hyperspace of biolo-gical objects (humans, animals, plants). From there the information reaches the reference space or the energy space via so-called interaction channels. This is a union of, among other things, all organs and energy forms in the biological object. There, the information of the program can dynamically materialize and thus change current states. The changes can show themselves in the form of neutralizations or harmonizations of disturbances, the dissolution of problems, blockages and disharmo-nies.

Conditions:

BESA tests are performed in the premises of IFVBESA under laboratory conditions, at room temperature 20°Celsius, on natural wooden floor. The test persons are deswitcht (made testable) before the BESA tests or the test possibility is questioned with the test persons.

- **Pos.1** BESA 1 Test Basic (bioenergetic status) on the test person.
- **Pos.2** BESA 2 Test Basic-BREVIS (extended bioenergetic status) on the test person.
- **Pos.3** BESA 3 Test when the test person is confronted with the digitized spike proteins and virus fragments.
- **Pos.4** BESA Test when the subject is confronted with the digitized spike proteins, the virus fragments and the graphene oxide.
- **Pos.5** BESA Test when the test person is confronted with the already mentioned stress factors and the test object, the "Leela Quantum DNA & Cell Protector Card".
- **Pos.6** Evaluation of the results in the project as well as summary in a corresponding expert opinion according to the sample.

Procedure and specifications for the implementation

- 1. **BESA basic measurement of the test person** at all previously determined measurement points (TING points) serves to determine the actual condition. The results are determined exactly according to the BESA specifications and documented via the BESA graphs.
- 2. Basic BESA measurement BREVIS of the test person at all previously determined



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measurement points (BREVIS points) serve to determine the actual state. BREVIS refers to an extended matrix of acupuncture points to be measured. These points are applied or tested in order to get a deeper insight into the energy-informative events of the functional circuits of the test person. The results are again determined exactly according to the BESA specifications and documented via the BESA graphs.

- 3. The test person is brought into contact with the listed stress factors depending on the project, whereby the sequence discussed with the developer is considered as a specification and is adhered to accordingly. In order to be able to determine the current energy state, the measuring points mentioned under point 1 are measured in the same sequence and for the same duration for each test of the load factors. The results are determined exactly according to the BESA specifications and documented via the BESA graphs.
- 4. Activation of the "Leela Quantum DNA & Cell Protector Card" test object
 - 4.1. When activating the "Leela Quantum DNA & Cell Protector Card", it is brought into the measurement area according to the instructions of the client..
 - 4.2. The test person is brought into contact with the stress factors. The measuring points mentioned under point 1 are measured in the same order and for the same duration in order to determine the current energy state. The results are determined exactly according to the BESA specifications and documented via the BESA graphs.

Test procedure

BESA 1 Testing BASIC BEFORE as status

In the first step, a basic bioenergetic test (bioenergetic status) is performed on the meridian end points (TING points) of the test person.

BESA 2 Testing BASIC BREVIS BEFORE as satus

In the next step, a basic bioenergetic BREVIS testing (bioenergetic status) is performed on the meridian end points (TING points) of the subject.

BESA 3 Testing BEFORE with the mentioned stress factors like spike proteins and virus fragments.

In the further BESA testing process, the subject is contacted and tested with listed digitized spike proteins and the virus fragments. The questions are: How does the meridian system react? How does the energetic status change when confronted with the stress factors? What are the differences between BESA 1 and 2 BASIC testing BEFORE?

BESA 4 Testing BEFORE with the mentioned stress factors from BESA 3 testing BEFORE and the graphene oxide.



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In the further course of BESA testing, the test person is brought into contact with the digitalized stress factors and tested. The questions are: How does the meridian system react? How does the energetic status change when confronted with the stress factors? What are the differences compared to the BESA 1,2 and 3 tests BEFORE?

BESA 5 Testing AFTER with the mentioned stress factors and the "Leela Quantum DNA & Cell Protector Card".

In the last BESA test, all the stress factors already tested above and the "Leela Quantum DNA & Cell Protector Card" are introduced into the measurement circuit. Now the question is: How does the meridian system of the test person react within the area of effect of the "Leela Quantum DNA & Cell Protector Card" when all the stress factors already tested are activated at the same time?

BESA 1 Testing BASIC BEFORE

BESA 1 Testing BASIC BEFORE as status

Eva Krankl performed a basic BESA measurement on the subject. All BESA tests were performed at the TING points (40 nail fold points on the fingers and toes).

Objective: The creation of a basic test (status) for the representation of the energetic initial situation for all further BESA tests.

BESA-Test evaluation P74 4.0

from 12-11-2021 at 10:35 to 10:39 (4 minutes) page 12 and 13

Result: The measurement result indicated partly strong energetic stresses at the meridian end points and subsequently on the subordinate metabolic situation of the test person.

97 % in the blue area

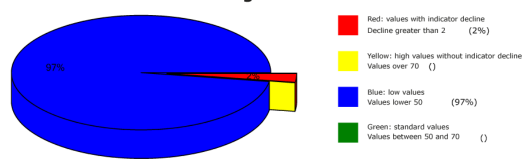
2 % in the red area

1 % in the yellow transition area

Conclution: As the graphs show, all measuring points are in the degenerative blue range (energy deficiency) although 2% of them are even in the red measuring range. The red measured values interpret a current total deregulation in this control circuit (organ large intestine). The comparisons of the BESA graphs confirm the stressful influences on the energy-informative events in the meridian system of the test person.

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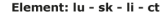
Overview of BESA measuring



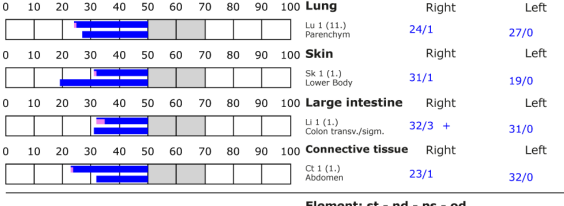
BESA basic test

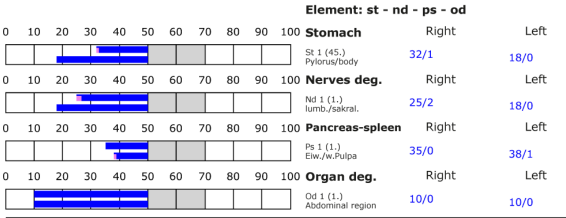
- +++: Indicator decline > 15 Skt.
- T: Total inflammation (>89 Skt.) P: Partial inflammation (70-89 Skt.) ++: Indicator decline 6-15 Skt.
- +: Indicator decline 3-5 Skt.

Standard values: (50-70 Skt.)



D: Degeneration (< 50 Skt.)





						Element: bl	- ly - ki - al	
0	10					100 Bladder	Right	Left
						BL 1 (67.) body	30/1	29/0



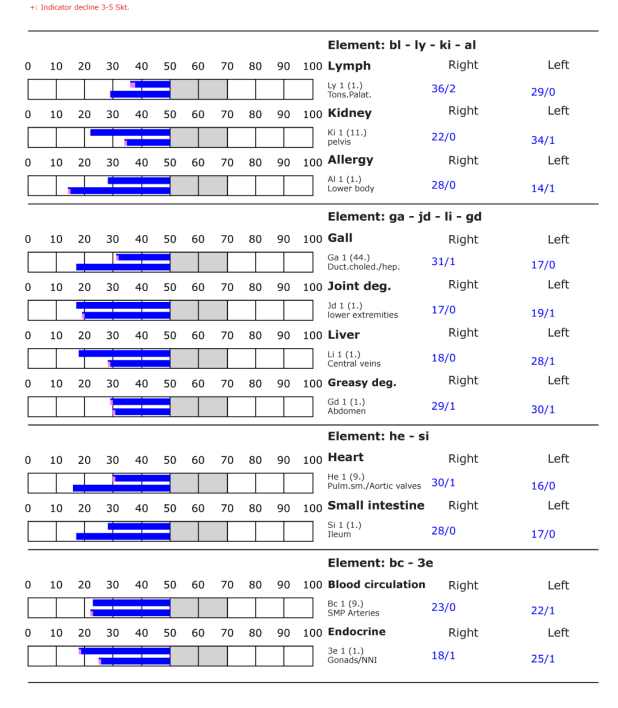
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BESA basic test

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- T: Total inflammation (>89 Skt.) P: Partial inflammation (70-89 Skt.)

D: Degeneration (< 50 Skt.)

Standard values: (50-70 Skt.)



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BESA 2 Testing BEFORE

BESA 2 Testing BASIC BREVIS BEVOR as extended status

Eva Krankl performed a basic BESA measurement on the subject. All BESA tests were performed on the BREVIS points (78 measuring points on the fingers and toes including the base joints).

Objective: To create an extended basic test (status) to represent the energy-informative initial situation for all further BESA tests.

BESA-Test evaluation P74 4.0

from **12-11-2021** at **11:04** to **11:15** (11 minutes) page 15 to 17

Result: The measurement result also indicated again strong energetic stress at the tested meridian points and in further consequence on the subordinate metabolic situation of the test person.

100 % in the blue area

Conclusion: As the graphics show, all measuring points are in the degenerative blue area (energy deficiency). This test variant was chosen in order to get a deeper insight into the energetic events at the meridian system of the test person. As can be seen in the BESA graphs, this matrix allows a larger spectrum of organ details and thus also shows the chronified events in the energy-informative area in contrast to the TING points (acute events). The BESA graphic shows the resolution of the red measured values in the representation of the chronic course in contrast to the acute event, although here the peritoneum (peritoneum) as part of the large intestine with 22/2 also just scrapes a red measured value. The comparisons of the BESA graphs confirm the stressful influences on the energy-informative events in the meridian system of the test person.



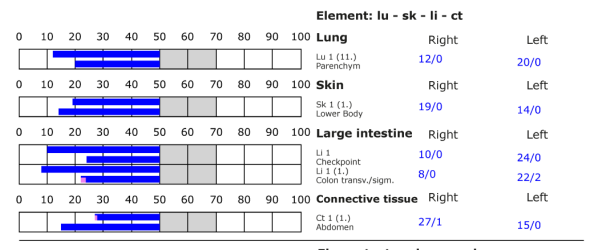
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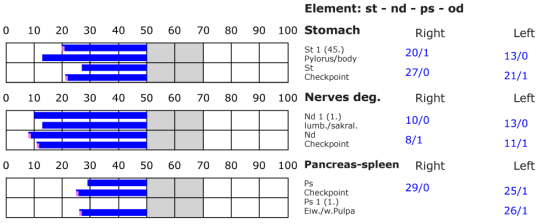
BESA basic test

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- +: Indicator decline 3-5 Skt.
- T: Total inflammation (>89 Skt.)
 P: Partial inflammation (70-89 Skt.)

D: Degeneration (< 50 Skt.)

Standard values: (50-70 Skt.)





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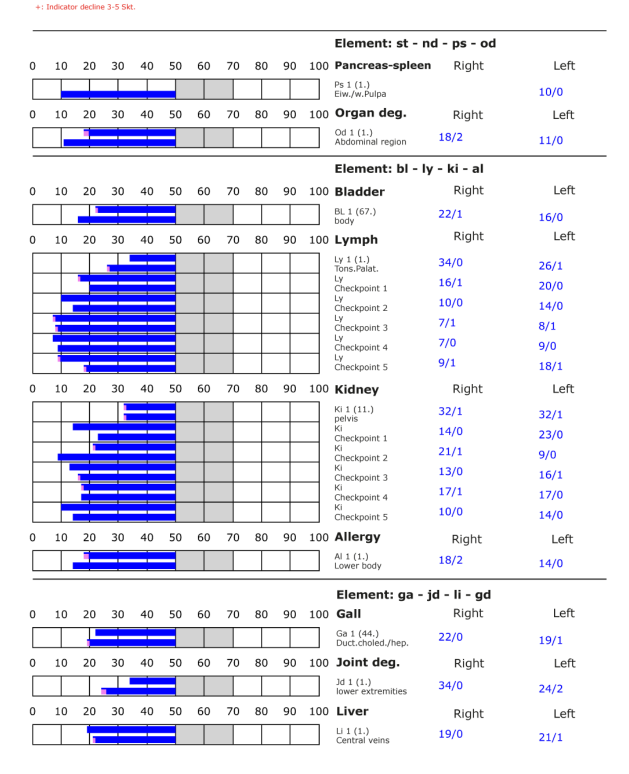
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Standard values: (50-70 Skt.)





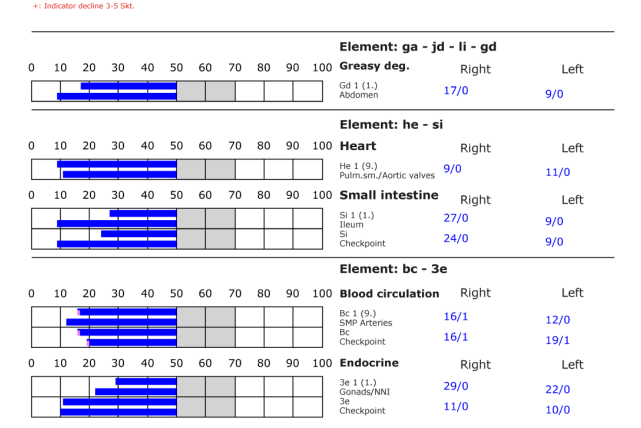
Internationaler Fachverband für BESA I ZVR Nr. 975047937 Hauptstraße 1, A-4861 Kammer-Schörfling am Attersee / Austria Tel.: +43-664-73152899 I E-Mail: <u>info@ifvbesa.at</u>

BESA basic test

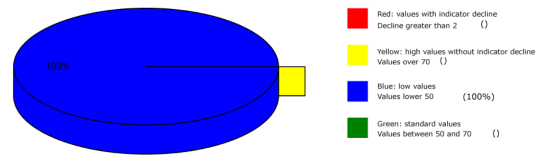
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Overview of BESA measuring



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BESA 3 Testing BEFORE

BESA 3 testing BEFORE; testing for the presence of spike proteins and virus fragments - as a result of the modified spike proteins of the mRNA vaccine.

In the further BESA testing procedure, the subject was tested for the presence of digitized spike proteins and viral fragments in the organism. For this purpose, the digitized loading factors were introduced into the measurement circuit and tested at the specific measurement points. All BESA tests were again performed at the TING points (40 nail fold points on the fingers and toes) as in the BESA 1 test BASIC BEFORE.

Objective: To determine the response in the subject's meridian system to the question in which regulatory circuit spike proteins and virus fragments can be tested. It is important to understand that in the case of the presence of spike proteins, the measuring device reacts with a "measuring response in the green zone", unlike in the classical measuring process.

BESA-Test evaluation P74 4.0

from **12-11-2021** at **10:40** to **10:45** (5 minutes) page 19 to 22

Result: The bioenergetic measurement result shows the load of spike proteins and virus fragments within these tested organs via the display of the green measurement values. Only the green measured values are decisive. The blue measured values play only a subordinate role in this case.

22 % in the blue area

77 % in the green area

1 % in the yellow transition area

Conclusion: As the graphs show, after the digitized spike proteins and the virus fragments have been introduced into the measuring circuit of the test person, at least 77% of the measured values are in the green range (response to the presence of spike proteins and virus fragments within these organs). It is interesting to see in this test person how quickly the ingredients have spread to almost the entire body of the test person, or it can be assumed that they will continue to spread. According to the bioenergy-informative measurement results, these listed and tested ingredients are located in the following organs:

the lung- left, skin- right, large intestine- right and left, connective tissue- left, stomach, nerve degeneration, pancreas/spleen as well as organ degeneration right and left, bladder, lymph, as well as kidney right and left, veins (allergy point) left, heart, small intestine right and left as well as circulation and endocrinum right and left. The presentation of the BESA graphs confirms the existing influences on the subject's organs. Please note the correspondingly marked measurement results of the organs in the green area.



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It is difficult to establish a relation to the symptoms of the test person (especially Alzheimer's dementia) because the subject matter is too complex. However, a closer look at these measured values, also in comparison to other test persons, raises the question of what influence the medication taken by the test person has in correlation with the ingredients of the vaccine.

For a simple explanation: What are spike proteins (S-proteins)?

An S-protein (S-glycoprotein) is an outwardly protruding viral envelope that is studded with so-called spikes. This S-protein consists of viral membrane proteins, which is anchored in a lipid membrane. The outwardly projecting spikes or spines are glycosylated by sugar residues (saccharification of e.g. proteins). These spines enable the S protein to bind to the surface receptors of the target cell, thus enabling the so-called virus to enter the cell.

About the virus: Scientifically, a so-called virus could never be identified as what it is assumed to be, as dangerous for humans! Let us leave for the time being the designations in such a way, as they are quoted currently pseudo-scientifically.

Über die sogenannten "Spikes" tritt das Virus und seine Funktion oder Eigenschaft (für uns eine bestimmte Information) bei der Bindung an die Wirtszelle mit den Antikörpern des Wirtes in eine Interaktion.

Via the so-called "spikes", the virus and its function or property (for us, a specific piece of information) interact with the host's antibodies when it binds to the host cell.

Never has anything attracted so much attention in the past months/years since Corona as the "spike protein". It is important to understand that in addition to the typical S protein found in influenza disease, there is another. This is the one that is produced by genetically engineered vaccines such as the mRNA vaccine or the vector vaccines. In principle, the natural S protein is already a challenge for the human immune system, but the modified S protein that the body produces in response to the vaccine represents a real challenge for the human organism.

In an article by Stephanie Seneff and Greg Nigh titled "Worse Than The Disease: Reviewing Some Possible Unintended Consequences of mRNA Vaccines Against COVID-19: Reviewing Some Possible Unintended Consequences of mRNA Vaccines Against COVID-19), published in the International Journal of Vaccine Theory, Practice and Research is explained that a major part of the problem is that while the natural spike protein is bad, the spike protein that the body produces in response to the vaccine is even worse.

What are modified spike proteins: It is necessary to first return to modified RNA (mRNA). RNA stands for ribonucleic acid. This means that RNA or RNA is a chain of so-called nucleotides. They are the basic building blocks of DNA and RNA and also have regulatory functions in cells. This synthetic RNA (mRNA) has been manipulated to produce an "artificial spike protein". The difference to the natural S-protein is that it does not collapse as soon as it binds to an ACE2 receptor, but it remains open and attached to the ACE2 receptor. This overrides the receptor



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and allows for appropriate immunological reactions, leading to challenges such as cardiopulmonary and autoimmune reactions.

ACE2 (Angiotensin Converting Enzyme 2) is a protein compound that is mainly produced at the vascular endothelial cells of the heart, kidneys as well as respiratory epithelia and the gastrointestinal tract. It thus also plays an important role in the regulation of blood pressure as well as in anti-inflammatory and lung-protective effects.

Exactly these or similar symptoms could be detected in this and the following subjects with BESA.

What are viral splinters:

Here is the further explanation of what happens after the ACE2 receptors are inhibited by the spike proteins. These enter the cell and nucleus through the injected mRNA vaccine, suppressing the human body's DNA repair mechanism and triggering an explosion of immune deficiency, autoimmune, or other severe complications.

New research published in Viruses, part of MDPI's SARS-CoV-2 Host Cell Interactions Edition (open access journals), shows that vaccine spike proteins enter cell nuclei and destroy the cells' DNA repair mechanism by suppressing DNA repair by up to 90%.

The research paper is titled "SARS-CoV-2 Spike Impairs DNA Damage Repair and Inhibits V(D)J Recombination In Vitro" and was authored by Hui Jiang and Ya-Fang Mei at the Department of Molecular Biosciences, The Wenner-Gren Institute, Stockholm University, SE-10691 Stockholm, Sweden, and the Department of Clinical Microbiology, Virology, Umeå University, SE-90185 Umeå, Sweden, respectively.

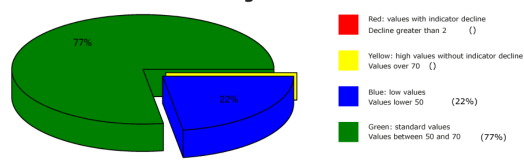
Mechanistically, the researchers found that the spike protein is localized in the nucleus and inhibits DNA damage repair by interfering with the recruitment of the important DNA repair proteins BRCA1 and 53BP1 to the damage site.

This means that the spike protein, which is formed in the cell's ribosomes after cells are hijacked by mRNA vaccines, does not always leave the cell and enter the bloodstream, as proponents of mRNA vaccines tell us.

In some cases, the spike protein enters the nucleus. There it interferes with the DNA repair mechanism, as described in this article. It confirms that such vaccines do indeed destroy genetic integrity and have side effects that were not predicted or described by proponents of mRNA vaccines. These SARS-CoV-2 viral fragments are referred to as "Nsp1, Nsp5, Nsp13, and Nsp14." Overexpression of these viral fragments and spike proteins reduces DNA repair efficiency (NHEJ repair) according to this study).

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Overview of BESA measuring

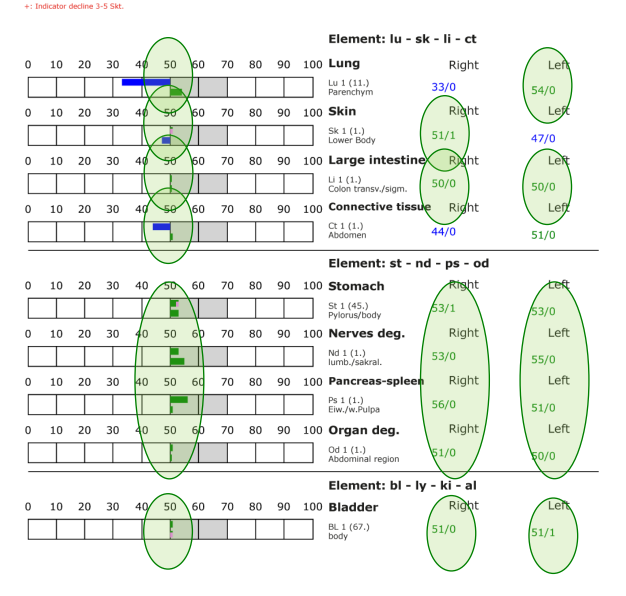


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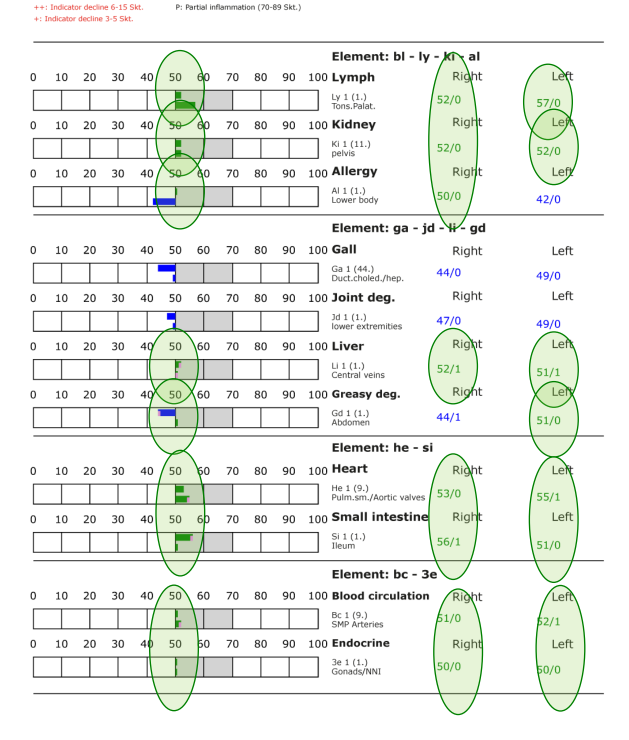
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BESA basic test

+++: Indicator decline > 15 Skt.

T: Total inflammation (>89 Skt.) P: Partial inflammation (70-89 Skt.) D: Degeneration (< 50 Skt.)

Standard values: (50-70 Skt.)



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BESA 4 Testing BEFORE

BESA 4 testing BEFORE; testing for the presence of stress factors from BESA 3 testing BEFORE as well as graphene oxide.

In the further course of the BESA test, the test person was tested for the presence of digitized graphene oxide in connection with the digitized stress factors from the BESA 3 test BEFORE in the respective control circuits of the organism. For this purpose, all the mentioned digitized loading factors were introduced together into the measurement circuit and tested at the specific measurement points. All BESA tests were again performed as in BESA 1, and 3 testing BEFORE at the TING points (40 nail fold points on the fingers and toes).

Objective: To determine the reaction in the meridian system of the test person to the question in which control loop the mentioned stress factors can be tested energy-informatively. It is important to understand that in case of the presence of the stress factors the measuring device reacts with a "measuring response in the green range". See green markings on the BESA graphs.

BESA-Test evaluation P74 4.0

from **12-11-2021** at **10:47** to **10:52** (5 minutes) page 24 to 26

Result: The bioenergetic measurement result shows the load of the indicated load factors within these tested organs via the representation of the green measurement values. The absence of the blue measured values shows that all organs of the test person are stressed!

100 % in the green area

Conclusion: As the graphics show, after the introduction of the mentioned digitized stress factors such as spike proteins, virus fragments and the digitized graphene oxide into the measuring circuit of the test person, all measured values are in the green area (answer to the presence of the stress factors within all organs)! Following the measurement result, these are located in the organs of the BESA graph shown. Therefore, we do not show the green circles separately (delimitation). The representation of the BESA graphics in the green area confirms the existence of influences by the indicated stress factors on the energy-informative control circuits of the test person.

For a detailed description of the spike proteins and viral fragments, please refer to pages 19 to 20.

What is graphene oxide? Graphene oxide is obtained from graphite under the action of strong oxidants. Among other things, graphene oxide is potentially used in the biomedical field as a drug carrier to target organs and cells with the help of the bloodstream. Graphene oxide and



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cationic lipids (cationic lipids are used as lipid shells for drug delivery to the mRNA particles of COVID vaccines) have toxic effects.

Graphene is 200 times stronger than steel, 1 million times lighter than paper and transparent, thus information about laser light or similar beams can be transported optimally and mega fast. Graphene conducts electricity and heat better than a diamond. It conducts 250 times better than silicon, so it is the new superchip.

Under an electron microscope, saliva can be used to see exactly how tiny white or silvery spots or particles move and crosslink with each other after the mRNA injection. Subsequently, they crystallize and branch into rectangular antenna-like structures.

Several scientific works show that graphene oxide is used in gene therapy as a platform for delivering biomolecules such as mRNA into cells. The background lies in its high electrical conductivity and ability to penetrate cell membranes.

The crystalline and rectangular networks that form in the body fluid after mRNA vaccination and in the vaccine itself look like high-voltage antennas.

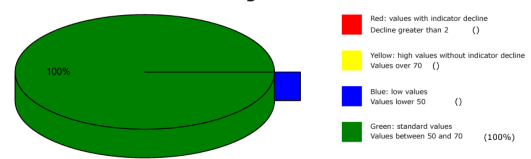
The "National Graphene Institute" is located in Manchester, where extensive research is being conducted. It was at this university that the existence of graphene antennas was first proven. It has been shown that graphene liquid crystals form spontaneously in the presence of an external magnetic field. Further scientific results showed that electric fields can change the crystal structure of graphene.

Graphene is considered a foundational technology for 5G by graphene manufacturers such as Grolltex, who are working on the future of graphene and 5G. Graphene has also been successfully connected to neurons and graphene-based neurotechnologies have been and continue to be the subject of intensive research.

More details or scientific background on graphene can be found at the end of this detailed project.

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Overview of BESA measuring



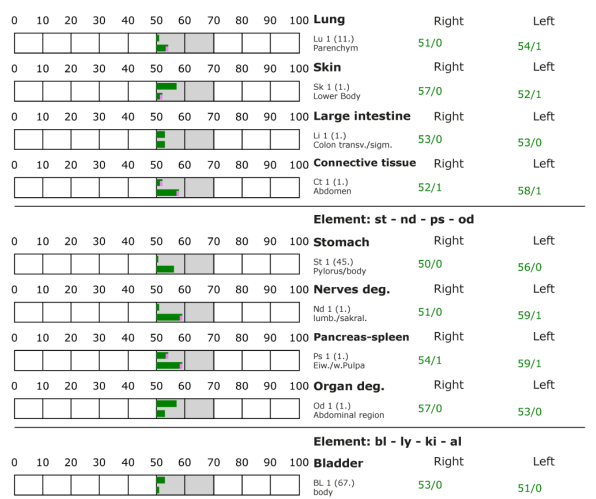
BESA basic test

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- +: Indicator decline 3-5 Skt.
- T: Total inflammation (>89 Skt.)
 P: Partial inflammation (70-89 Skt.)

D: Degeneration (< 50 Skt.)

Standard values: (50-70 Skt.)

Element: lu - sk - li - ct





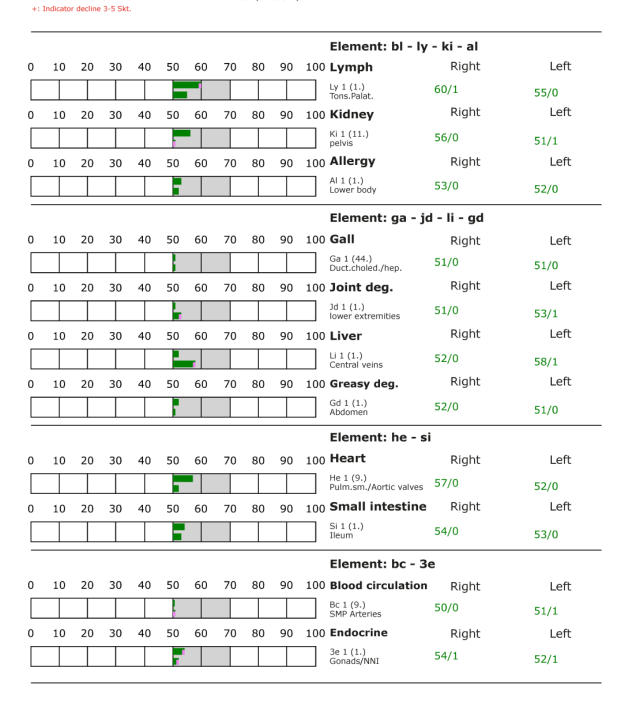
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BESA 5 Testing AFTER

BESA 5 testing AFTER; modified spike proteins, viral fragments and graphene oxide in combination with the "Leela Quantum DNA & Cell Protector Card".

In the further BESA test, the subject was exposed to the modified spike proteins, the viral fragments, the graphene oxide and the test object, the "Leela Quantum DNA & Cell Protector Card". All BESA tests were again performed at the TING points (40 nail fold points on the fingers and toes) as in the BESA 1, 3 and 4 tests.

Objective: to determine the response of the subject's meridian system within the strong influence of the stress factors tested in conjunction with the test object. To determine the differences of BESA 5 testing AFTER compared to BESA 1, 2, 3 and 4 testing BEFORE.

BESA-Test evaluation P74 4.0 from **12-11-2021** at **10:53** to **10:58** (5 minutes) page 28 to 29

Result: The measurement result shows significant improvements in the meridian end points or in the energetic state of the test person.

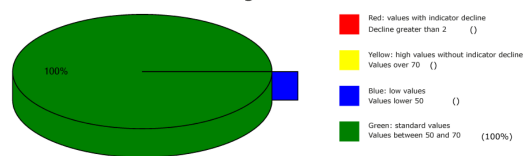
100 % in the green area

Conclusion: As the graphs show, 2 minutes after inserting the "Leela Quantum DNA & Cell Protector Card" into the measuring circuit, all measuring points are in the green, optimal and harmonized range (balanced energy system). The BESA test results in a significant improvement of the energy situation in the meridian system of the test person compared to the BESA 1, 2 and 3 tests BEFORE. All readings were at or just above 50 sct. It can be seen that the "Leela Quantum DNA & Cell Protector Card" is able in a very short time to give the necessary impulse for harmonization (neutralization) into the life-promoting range to the heavy loads of the tested spike proteins, the virus fragments and the graphene oxide (see the green measured values as an expression of the positive measurement response) determined in the BESA 1, 2, 3 and 4 tests BEFORE. The comparisons of the BESA graphs confirm the change and harmonization of the stress factors on the meridian system of the test person.

These first BESA tests show that the test object, the "Leela Quantum DNA & Cell Protector Card" is basically able and suitable to produce a harmonization of the stressful information through the modified spike protein, the virus fragments and the graphene oxide into the deeper structures of the cell or into the DNA.

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Overview of BESA measuring

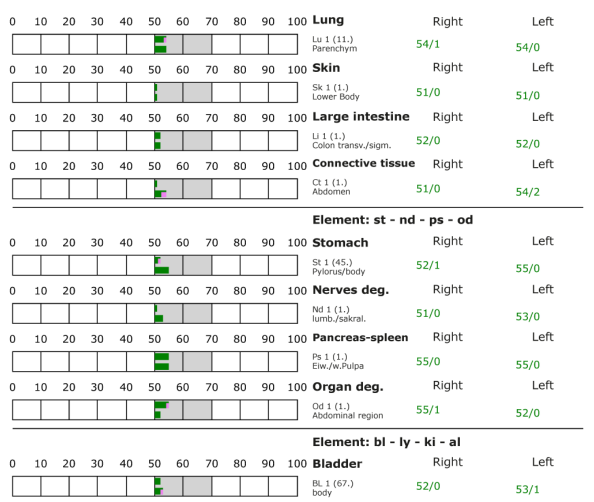


BESA basic test

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Standard values: (50-70 Skt.)

Element: lu - sk - li - ct





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BESA basic test

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- T: Total inflammation (>89 Skt.) P: Partial inflammation (70-89 Skt.)

D: Degeneration (< 50 Skt.) Standard values: (50-70 Skt.)

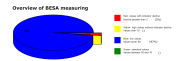
											Element: bl - ly	- ki - al	
	10	20	30	40	50	60	70	80	90	100	Lymph	Right	Left
					-						Ly 1 (1.) Tons.Palat.	52/1	58/1
	10	20	30	40	50	60	70	80	90	100	Kidney	Right	Left
					-						Ki 1 (11.) pelvis	52/0	54/1
)	10	20	30	40	50	60	70	80	90	100	Allergy	Right	Left
											Al 1 (1.) Lower body	51/1	51/0
											Element: ga - jo	d - li - gd	
)	10	20	30	40	50	60	70	80	90	100	Gall	Right	Left
											Ga 1 (44.) Duct.choled./hep.	51/0	52/1
)	10	20	30	40	50	60	70	80	90	100	Joint deg.	Right	Left
					=						Jd 1 (1.) lower extremities	53/0	53/1
)	10	20	30	40	50	60	70	80	90	100	Liver	Right	Left
					-						Li 1 (1.) Central veins	52/0	53/0
)	10	20	30	40	50	60	70	80	90	100	Greasy deg.	Right	Left
					-						Gd 1 (1.) Abdomen	53/1	53/0
											Element: he - s	i	
)	10	20	30	40	50	60	70	80	90	100	Heart	Right	Left
					-						He 1 (9.) Pulm.sm./Aortic valves	52/0	53/0
)	10	20	30	40	50	60	70	80	90	100	Small intestine	Right	Left
					7						Si 1 (1.) Ileum	57/0	52/0
_											Element: bc - 3	e	
)	10	20	30	40	50	60	70	80	90	100	Blood circulation	Right	Left
											Bc 1 (9.) SMP Arteries	50/0	52/0
)	10	20	30	40	50	60	70	80	90	100	Endocrine	Right	Left
_			\top								3e 1 (1.) Gonads/NNI	50/0	55/1

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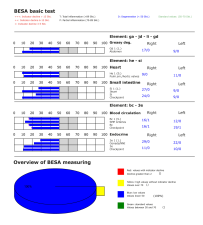
The results of the BESA tests at a glance

Respondent 1

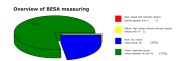
BESA 1 Testing BASIC BEFORE as energetic status



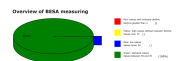
BESA 2 Testing BASIC BREVIS BEFORE as 2. energetic status



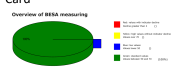
BESA 3 Testing BEFORE with spike-proteins and virus-fragments



BESA 4 Testing BASIC BEFORE with spike-protein, virus-fragments and graphene oxid



BESA 5 Testing BASIC BEFORE with spike-protein, virus-fragments, graphene oxid and "Leela Quantum DNA & Cell Protector Card"



Please note the respective explanations of the BESA graphs following the BESA tests BEFORE and AFTER to avoid misunderstandings in the interpretation of the measurements.

General information about the test result

Man represents a kind of receiving antenna for environmental information. This is because human life depends fundamentally and exclusively on environmental information. Our organism is biologically very sensitive where natural information (fields) are located or where this natural information is subject to interactions and fluctuations. The situation is all the more dangerous when such field-building structures are introduced (injected) into the body as nanoparticles via a so-called mRNA vaccine. For this reason, detected informative electromagnetic interference fields are biologically highly relevant. Any reduction or transformation of these interference fields (ideally to 100 percent) is biologically very important, in the case of these tested vaccines possibly even life-critical. These information loads from the vaccine as well as from our artificial environment are only compatible with life



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if they can be readjusted to a natural fluctuation tolerance. Disturbances, problems, blockades, disharmonies in the biological control circuit of man find their causes in such disturbing electromagnetic information influences.

Neutralizing or harmonizing effects could be proven in this project P74 4.0 to determine the effect of the test object, the "Leela Quantum DNA & Cell Protector Card". The "Leela Quantum DNA & Cell Protector Card" was able to neutralize the biologically detrimental effects and effects of the stress factors tested on the test subject.

The modulation of many ingredients in the Corona mRNA and vector-based vaccines has created a completely new situation and challenge for the human immune system. It will occupy the prevailing science for decades to come to find solutions to the resulting questions. Many solutions and answers already exist. They offer an important approach in understanding what is taking place on the human level.

But the biggest challenge is facing those people who already have pre-existing health conditions and are taking medications. This is because no one can imagine what effects these drugs will produce when combined with mRNA and/or vector-based vaccines. Many question marks have already arisen within this detailed project. It will take countless more projects to resolve them.

But critical to this detail project is the definitive ability of the test object, the "Leela Quantum DNA & Cell Protector Card" to neutralize and harmonize the severe strain factors tested in this P74 4.0 project. The transformation of the tested stress factors into bioenergy information with biological and life-promoting goodness is proven with this project.

Authorized summary

The BESA tests carried out by IFVBESA on the energetic and physical effectiveness of the test object, the "Leela Quantum DNA & Cell Protector Card" have clearly shown that this test object is able to neutralize or harmonize biologically significant stress factors such as spike proteins, virus fragments and graphene oxide at the acupuncture points of the test person. Via the bioenergy-informative system analysis, the effect of the above-mentioned stress factors on the test person, his meridian system and his energetic-biological control circuits was questioned and systemically tested on the energy-informative level. The BESA tests BEFORE - AFTER show significant changes at the tested acupuncture points on the meridian system of the test person. The measurement data as well as their key figures impressively confirm on the one hand the stresses that the tested factors cause on the human organism, and on the other hand clarify how the deregulating energies are transformed into body-immanent and biocompatible energies after the application of the test object, the "Leela Quantum DNA & Cell Protector Card".

From a holistic point of view, it can be assumed that the positive effect on the test subjects also occurs in other people. That the positive influence by the "Leela Quantum DNA & Cell



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Protector Card" is actually possible with high precision is clearly shown by this test through the BESA-PRE-AFTER comparison. All measured values improved significantly from the mostly 100% blue measuring range to the green, mostly 50 Skt. range (Skt = scale value), i.e. the range of optimal measured values. This means that an optimal regulation dynamic has taken place. Here, in the sense of the IFVBESA, one can clearly speak of an optimal, significant improvement of the body's own energy situation.

Result

The test person was brought into contact with heavily loaded, digitized ingredients of an mRNA vaccine during each BESA AFTER test. In contrast to the BESA BEFORE tests, in which the test object, the "Leela Quantum DNA & Cell Protector Card" was not used, consistently positive measurement results were found, indicating that neutralization or harmonization had taken place. The regulation dynamics developed into an optimal effective range.

By demonstrating the energy informative effectiveness of the test object "Leela Quantum DNA & Cell Protector Card" in this project P74 4.0, the requirements for obtaining a BESA seal of approval by the International Professional Association for BESA were met.

Sources directory

to spike-proteins and virus-fragments:

Monster, Zombies and Mutants. HORRIFYING new research reveals how vaccines suppress DNA repair mechanism in your cells

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https://www.naturalnews.com/2021-11-02-science-horror-vaccine-spike-protein-enters-cell-nuclei-suppresses-dna-repair-engine-of-the-human-body-cancer-aging.html#

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Non-Homologous End Joining

NHEJ is an error-prone repair pathway that can occur throughout all cell cycle phases, whereas HR is an error-free pathway that predominantly occurs in late S and G2 phases.

https://www.sciencedirect.com/topics/biochemistry-genetics-and-molecular-biology/non-homologous-end-joining

to graphene oxid:

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DARPA-funded microchip technology optimizes convalescent plasma therapy for COVID-19 patients | School of Medicine | University of California, Irvine https://www.som.uci.edu/news_releases/Convalescent-plasma-emerging-pathogens.asp

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Graphene

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https://www.darpa.mil/work-with-us/covid-19



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PIL genocide - by PCR tests and FFP2 masks (using Darpa hydrogel) - PIL genocide study confirms the - StuDocu

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