

## **Criminal and Scientific Misconduct Involving Neural Prosthesis Research Funded by the NIH/NINDS/NPP and The Alfred E. Mann Foundation**

David A. Larson  
lars1212@verizon.net

---

### **Articles of Evidence:**

The photos contained in this document are of microstimulator devices that were surgically recovered and analyzed by using microscopy and by materials characterization techniques. The recovered devices are an exact match for the devices that were developed by research scientists Joseph Schulman, Gerald Loeb and Philip Troyk under contract from the National Institutes of Health. There are about 4 contracts involved, but the initial contract was #N01-NS5-2325 and funded by the NIH/NINDS/NPP. More information regarding these contracts can be found at the NIH Neural Prosthesis Project website. The following list briefly outlines some of the evidence presented in this document:

1. Microstimulator devices have been surgically recovered by a Physician in Los Angeles CA.
2. Devices have been analyzed using Energy Dispersive Spectroscopy (EDS) at Photometrics (materials characterization lab, Huntington Beach, CA) and confirm the presence of Silicon, Chlorine and other materials used in semiconductor processing.
3. The recovered devices show fabrication and design methods which uniquely belong to research scientists Joseph Schulman, Gerald Loeb and Philip Troyk, and are detailed in their U.S. Patent office filings. The recovered devices precisely match these patents.
4. Additionally, Schulman and Loeb filed US Patent applications for a system of monitoring a patient and methods allowing bi-directional telemetry over greater distances that involves using one frequency typically used for communication between the number of small implants and other near-by devices (between 570Hz and 2Mhz), and then an additional frequency (such as 224.840 MHz, or higher) which carries the data over the necessary long distance to the remote location.
5. Timing 1: My assault occurred during the same quarter in which Schulman and Loeb filed patent applications for implantable devices and patient monitoring systems which employ above mentioned methods of bi-directional telemetry. U.S. Provisional Application No. 60/039,164 was filed on Feb. 26, 1997 and U.S. Provisional Application Ser. No. 60/042,447 on Mar. 27, 1997
6. Timing 2: Schulman and Loeb report the first functional microstimulators in a QPR just prior to my assault and first animal implantations are planned. They subsequently report "tightening up" the accountability and traceability of prototype devices in a QPR following the assault
7. Schulman has an FCC license and broadcasts at 224.840 MHz (K6BWA) and has repeaters operating to the North, South and East of my residence, all within a 22-mile range. Additionally, another repeater in Altadena CA (but with a call sign that begins with a "W"), operating at the same frequency is actually licensed to an individual who lives in Oak Lawn, IL., (14 miles from IIT Chicago).
8. Professor Wise from Univ. of Michigan reports in a QPR that because of bandwidth and impedance issues, 200MHz should be chosen for remote, wireless stimulation using a neural prosthesis. The FCC database shows that Joseph Schulman operates radio repeaters in this frequency as call sign K6BWA.
9. Schulman reported acquiring an ISDN "Connectix" phone as part of contract #N01-NS5-2325 claiming it would be used for file sharing and enhanced communication. Later, a company he founded called "NeuroDyne" markets a new system which used such an ISDN video-phone to enable multiple physicians to tele-conference while analyzing a patients biofeedback signals such as EEG and EMG in real-time as the physiologic data is displayed on the screens and which can be saved for further analysis
10. Schulman reports that a "faulty" diode and resistor required that they be removed in-house (using two weeks worth of labor as well as equipment) and that replacement components would then have to be re-attached, however the patents of Schulman, Loeb and Troyk describe how additional, minimal circuitry can be integrated onto existing circuitry allowing diodes and resistors to be eliminated rather

than replaced and beneficially resulting in a smaller overall device with increased range of telemetry operation.

11. In a table of data listing the specs for components of the "BION" microstimulator, Schulman indicates that the diode used in replacement of the "faulty" diode is a "THD9064", however there doesn't appear to be any such diode in existence, and in fact, THD9064 only references low-power, high frequency microcontrollers and frequency doublers from manufacturers such as Siemens, ICS and New Focus
12. Throughout contract #N01-NS5-2325, Joseph Schulman fabricates an absurd amount of "vendor error" and "faulty product" which was done to conceal the real reason for the excessively low yield of microstimulators which can be attributed to the error prone stacking process of the smaller unethical implants as well as his need to secure a number of chips without accounting for them so that they could be used unethically.
13. Much further evidence exists in Joseph Schulman and Gerald Loeb's U.S. Patent Office filings and the Quarterly Progress Reports from contract #N01-NS5-2325. This evidence shows that implantable device invention and related technology discovery occurred during the contract period, and additionally were not reported or claimed to be made with government support. More importantly, many of the inventions involve integrating or stacking additional circuitry that enables additional functionality including long-distance bi-directional telemetry, and the accompanying ability to use this unethically
14. A vendor of Schulman' contract, Tom Wolf at Slicex reported that he sent hundreds of integrated circuits to Schulman and was never paid for the work. Schulman reported to the NIH that they were faulty, but the CEO of vendor, Tom Wolf, maintains that Schulman was happy with the devices and that they indeed worked fine. Schulman had to cook the books and not pay for the devices so they would be off the books, that way he could keep them and use them illegally without having to account to the NIH for them.

---

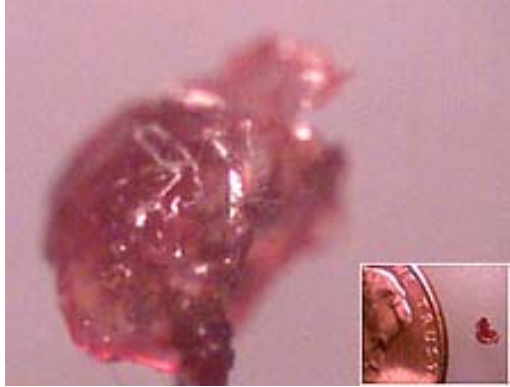
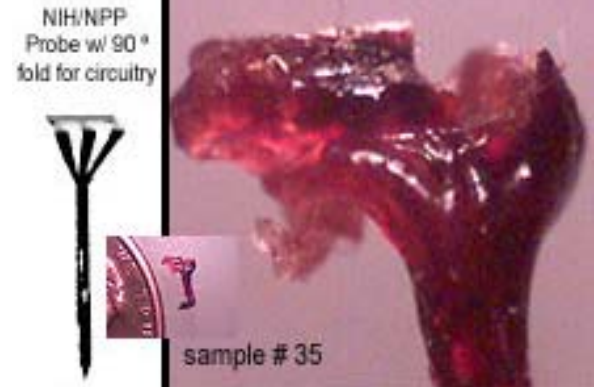
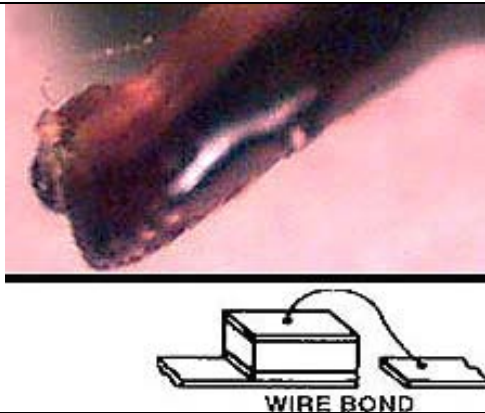
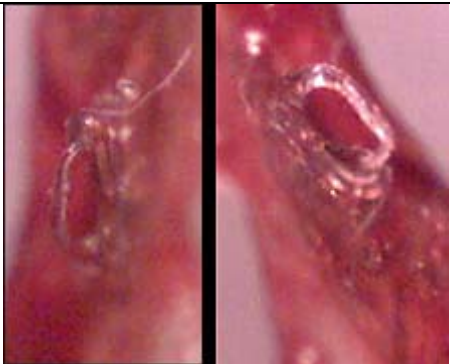
## **Evidence Detail**

The following will provide detail as to each instance of evidence that is listed above in "evidence summary". From a criminal/legal perspective, even though the assault occurred some time ago, the devices presented as evidence have remained active and in the control of the defendants up until the time they were recovered, and while implanted, were used violently and maliciously. Additionally, the devices were just recently recovered allowing discovery of their existence, so no statute of limitations is applicable, nor should apply. I wish that Joseph Schulman and Gerald Loeb were not guilty of this, as it would be much easier on everyone involved, however the evidence proves that they did this. Any skepticism or doubt that you may have regarding the legitimacy of my allegations should be weighed against the evidence. If you look at the evidence, it will prove the legitimacy of my allegations, so all that I ask is that you read on and try to comprehend what is presented. Even if the substantial amount of evidence presented here does not overcome all remaining doubt, it should at minimum prompt you to seek and find the answers needed through investigation to arrive at the truth. I had to endure an enormous amount of physical pain and frustration over the last 5 years (and especially as of late) in order to be able to present this evidence to you, and I would not waste your time or mine if I were not in a position to provide the evidence you require in order to take action. This is a call to action and if you received this document, it is because your assistance is needed. The evidence begins on the following page and each allegation listed above (1 – 13), is specifically addressed and the proof presented in detail with substantiated source or fact.

---

## **Article 1. - Microstimulator devices have been recovered from my person:**

1.1 Images: The devices below have all come from infected sites on my person and have been analyzed, numbered, documented and archived. Some of the images are shown below:

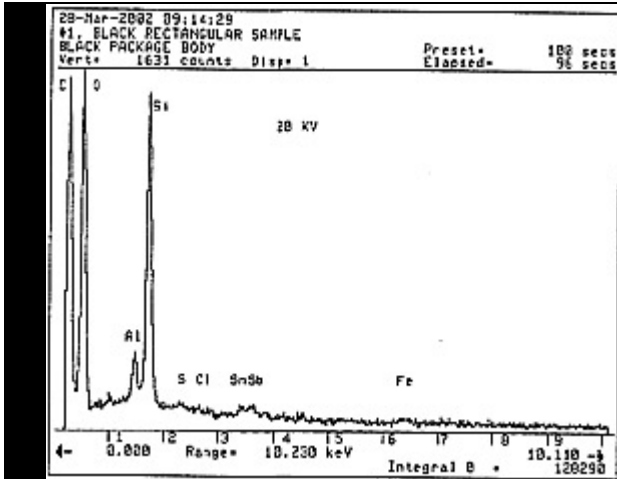
	 <p>NIH/NPP Probe w/ 90° fold for circuitry</p> <p>sample # 35</p>
<p>Image 1: sample # 50 at 60x magnification (main), and at 10x (inset) with penny for size reference. Circuitry is visible as is the (darker) hermetic feed-through w/ 90° bend for connection to electrode array or other component</p>	<p>Image 2: sample # 35 compares in size and shape with probe shown in NIH/NPP documentation. Integrated circuitry of active probe appears damaged from structural failure shown here at 60x. This type of probe has been used by Loeb (and Hambrecht) at the NINDS.</p>
 <p style="text-align: center;"><b>WIRE BOND</b></p>	
<p>Image 3: Sample # 70 at 60x. A typical wire bond similar to illustration. The substrate appears to have an epoxy or polymer coating.</p>	<p>Image 4: Sample # 95 at 60x magnification. Unknown feature that is definitely fabricated and structured to serve a purpose. Two views shown.</p>

Summary of Article 1: The photos clearly show something foreign that is not organic mixed with the biological tissue.

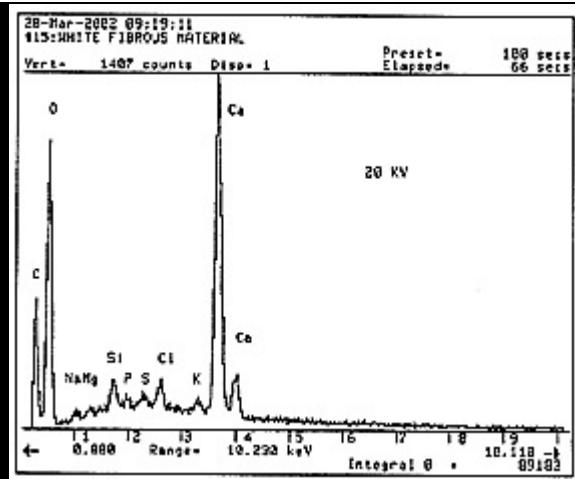
**Article 2. - Devices have been analyzed using Energy Dispersive Spectroscopy (EDS):**

Energy Dispersive Spectroscopy is a common method used for materials composition analysis and is well known within the scientific community. I had samples analyzed at Photometrics in Huntington Beach CA., which has one of the newer era systems that features light detection ability etc., and confirms the presence of Silicon in devices which I have recovered. Silicon is the primary and most common material in semiconductor chips and is not normally present in the body, and in fact is harmful because it contains trace amounts of arsenic. Additionally, other materials were identified such as Chlorine, which is used in the chip manufacturing process, Tin, and Aluminum. Some of the data from EDS performed at Photometrics appears below and complete information is attached to the end of this document.

2.1: EDS Data



The data for this sample reads left to right as follows (the higher the “peak” the higher the concentration of indicated material): Carbon (C), Oxygen (O), Aluminum (Al), Silicon (Si), Sulfur (S), Chlorine (Cl), Tin (Sn), Antimony (Sb), and Iron (Fe)



Sample #15, L to R: Carbon (C), Oxygen (O), Sodium (Na), Magnesium (Mg), Silicon (Si), Phosphorus (P), Sulfur (S), Chlorine (Cl), Potassium (K), Calcium (Ca) and (Ca) again. The sample that produced this result is shown below at left



This is a photo of sample #15 that showed Silicon and Chlorine being present. This device doesn't look as much like semiconductor as others, but it is and presents an opportunity to address the oxidization that occurs. The oxidization that builds up on the implants is a result of an activated process that occurs when certain metals interface with the electrolytes present in human tissue, and the longer the device is implanted, typically, the thicker the oxidation layer. This is documented in papers and QPR's from Loeb. The presence of the oxidization and some remaining blood and tissue on this device, which was recovered from my nasal cavity, make it difficult to visually see semiconductor components, however if you look closely the circuitry is indeed visible. Note the inset photo which reveals the actual size of this sample by showing a penny next to it on the lab slide for reference.

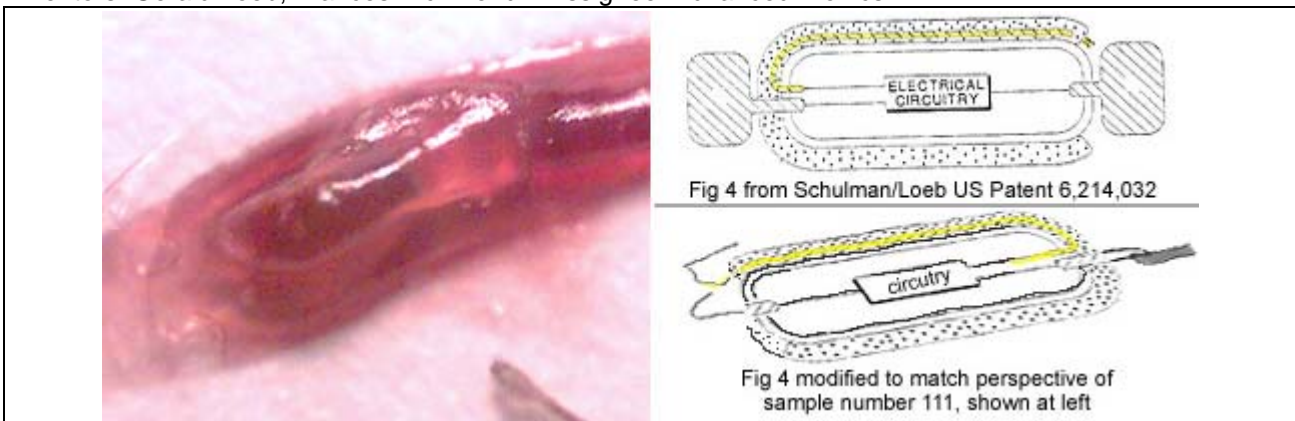
Summary of Article 2: EDS offers proven statistical data that confirms the devices recovered from my person contain semiconductor material and combined with the visual images provide absolute certain proof that these devices are foreign and should not be naturally present in my body. Moving forward, what follows will focus on proving the identity of the individuals responsible for placing them there.

**Article 3. - A number of recovered devices show fabrication and design methods which are unique to Joseph Schulman and/or Gerald Loeb, and are detailed in their U.S. Patent Office filings:**

The examples below directly compare images of my recovered devices to images from The US Patent Office outlining inventions patented by Schulman, Loeb and others within Alfred E. Mann Foundation. It should be noted that my recovered devices have been implanted for 5 years and obviously will are not in the pristine condition of a line-drawing illustration, but the design and construction of the recovered devices is similar or in some cases, identical. From a research perspective, a neural prosthesis needs to remain implanted for over 40 years to effectively serve a disabled individual, and because not enough research (none in fact, except for simulations) has been done regarding this, it would make sense for Schulman and Loeb to implant a variety of devices as well as materials because the continual real-time feedback (or lack there-of) would allow them to determine which were most effective and also remained biocompatible over time. I have

recovered devices appear to utilize a hardened epoxy coating on them as well as others that appear to have a rubbery type silicone coating. Schulman and Loeb's patents made during the time of the microstimulator contract describe much more than a typical "BION" type microstimulator and specifically address alumina, silicone and epoxy coatings instead of glass or titanium, describe how additional circuitry can upgrade a device from one which requires an external coil to one that is fully implantable, the creation of electrodes as small as 15µm using a YAG/UV laser, and one patent even describes how an implantable system can have it's components "daisy chained" together by microwire (which is slightly thinner than human hair) rather than completely assembled inside a glass package. The devices recovered from my person do not all conform to one design or configuration, and most conform to patent filings of Loeb and Schulman. Some of the comparisons to those U.S. Patent Office filings follow:

3.1 Sample #111 vs. US Patent 6,214,032 "System for implanting a microstimulator"  
 Inventors: Gerald Loeb, Frances Richmond Assignee: Advanced Bionics

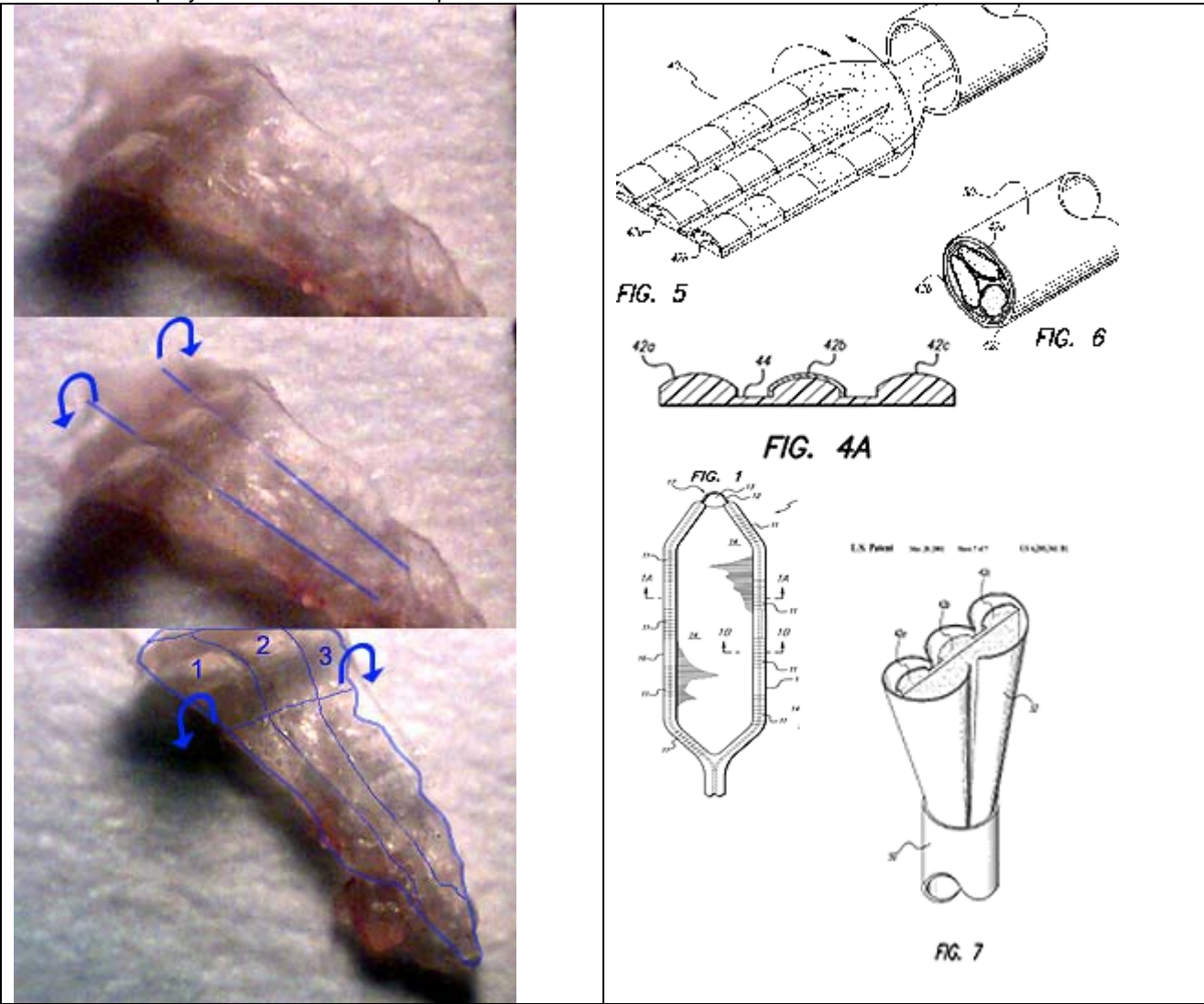


The image on the left shows my recovered device #111. Patent 6,214,032 on the right (top) illustrates an invention of Frances Richmond and Gerald Loeb (this same illustration appears in several patents held by Loeb and Schulman) in which the yellow highlighted path represents encapsulated wire containing electrode sites that are in addition to the electrodes shown at each end. The lower illustration is modified to show the invention in the same perspective as my sample #111. In this photo of my sample #111, the circuitry in the middle of the device is dark and difficult to identify, however the wire bond and path of the wire are clearly shown and it's construction is consistent with that of Loeb/Schulman's filings with the U.S. Patent Office and further, the documentation shows that "parent case" provisional applications were filed May 29, 1998 for U.S., and also Internationally on Feb.19, 1997 (within 30 days of my assault).

*Continued Next Page....*

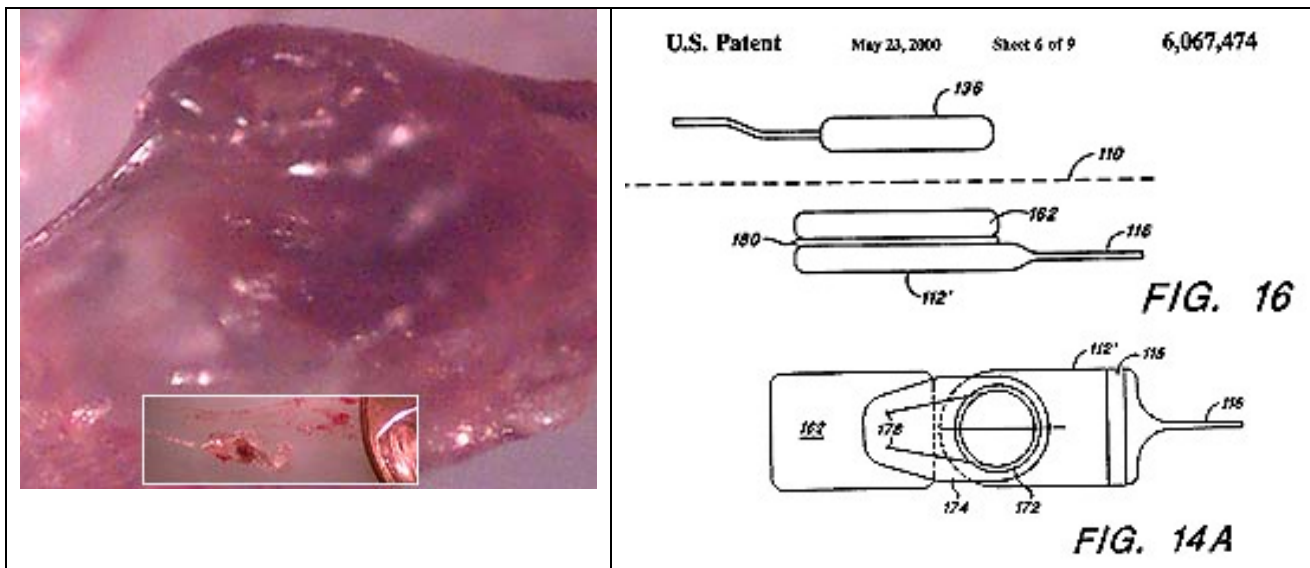
3.2: Sample 110 vs. US Patent 6,205,361 "Implantable expandable multicontact electrodes"  
 Inventors: Mann/Advanced Bionics

This invention describes an integrated circuit on flexible substrate (so that it will conform to the shape of the scalp, or muscle etc) with attached electrodes, and is designed to be "folded" into a syringe/hypodermic needle and injected into the target location. Upon being injected, the electrode array unfolds or "expands" to an open position. Although this device lists Mann as the inventor, and Advanced Bionics as the assignee, Gerald Loeb played a role in it's development as Advanced Bionic's "Chief Scientist".



The series of images to the left show my sample #110, and marked-up versions with the blue illustrations showing how my sample #110 compares to U.S. Patent 6,205,361 for a foldable/expandable array of electrodes. My sample #110 is consistent with above invention in that: (1) It has three defined areas at one end, and tapers to one section; and (2) it appears to be on a flexible substrate; and (3) it utilizes a wire perimeter for support

3.3 Sample #122 vs. US Patent 6,067,474 “Implantable device with improved battery (capacitor) recharging and powering configuration” - Inventors: Gerald Loeb, Frances Richmond Assignee: Advanced Bionics This patent lists Gerald Loeb (as well as F.J. R., also a principal investigator involved with the NIH microstimulator contract) as the inventors. In sample #122 recovered from my person, a coil (or “antenna”) can clearly be seen on top of the stack and is represented in the patent drawings as the innermost circle referenced as #172



My sample #122 is shown on the left. Patent #6,067,474 is shown to the right and illustrates a microstimulator that uses multiple circuits in a stacked configuration (with a coil on top, #172 in patent illustration) and the stacking is done to save space, however, such stacking complicates assembly and lowers yield. Schulman and Loeb claim that no such stacking of chips was performed, nor required by the “BION” microstimulator, however my sample #122 at left, US Patent # 6,067,474, at right, combined with Schulman reporting “unexplainable” low yields throughout his contract (as low as 30% in QPR#3) should be more than suspicious and immediately prompt inquiry followed by an investigation. And if the above wasn’t enough, the parent case documentation, U.S. Provisional Application Ser. No. 60/054,480, was filed Aug. 1, 1997 which is during the course of NIH contract #N01-NS5-2325, and not long after my assault.

**Summary of Article 3:**

Devices which have been recovered from my person and proved to contain semiconductor material, emulate and resemble devices which are unique to Joseph Schulman and Gerald Loeb.

**Article 4. - Schulman and Loeb file US Patent applications for patient monitoring systems and methods for long range bi-directional telemetry:**

Joseph Schulman and Gerald Loeb have each filed patents as well as later additions and newer patents for a “Patient Monitoring System” and “System of Implantable Devices for Monitoring and/or Affecting Body Parameters” and I contend that the “affecting” part is painful and has been applied by Schulman and Loeb maliciously through the recovered implanted devices shown in articles 1 and 3 above. What is defined in these systems is:

- Telemetry using multiple modes or frequencies so that one carrier can transmit a short distance with it being processed to another capable of longer range telemetry.
- Real time interaction that includes stimulation and recording from the human nervous system
- Audio and video conferencing are integrated into the system that displays the EEG, EMG and other data allowing Schulman, Loeb and Troyk to simultaneously interact with the system and each other
- A closed loop mode that has a “default” stimulus parameter loaded into memory for each electrode so that if the subject being monitored breaks the telemetry link, he will still be subject to electrical stimulation because power can be derived and stored into the capacitor from the Tantalum capacitor-electrode/bodily tissue electrolyte interface.

Because the illegal and unethical activity of monitoring an unwilling subject involves a database, which one can assume is backed up and stored somewhere (perhaps even online for remote retrieval) and also involves the program to be housed on a server that can be accessed by Schulman Loeb and Troyk, this presents considerable evidence that could be obtained by University personnel or law enforcement agencies, yet unobtainable to me. Because Schulman, Loeb and Troyk participate in this unethical/illegal research both at home and at work, this creates considerable liability for their employers and those who fund this research.

**4.1 - U.S. Patent 6,315,721: "System of implantable devices for monitoring and/or affecting body parameters"**  
Inventors: Schulman; Joseph H. (Santa Clarita, CA); Assignee: Alfred E. Mann Foundation

(This application is a division of U.S. patent application Ser. No. 09/048,826, now U.S. Pat. No. 6,208,894, and claims the benefit of U.S. Provisional Application Ser. No. 60/042,447 filed Mar. 27, 1997 and U.S. patent application Ser. No. 09/030,136, now U.S. Pat. No. 6,185,452, which claims the benefit of U.S. Provisional Application Ser. No. 60/039,164 filed Feb. 26, 1997.)

*"...a system implemented using multiple communication channels, e.g., a first sonic channel at a first carrier frequency and a second sonic channel at a second carrier frequency, is also considered to be within the scope of the present invention..."*

**4.2 – U.S. Patent 6,208,894: "System of implantable devices for monitoring and/or affecting body parameters"**  
Inventors: Schulman; Joseph H. (Santa Clarita, CA); Assignee: Alfred E. Mann Foundation

(This application claims the benefit of U.S. Provisional Application Ser. No. 60/042,447 filed Mar. 27, 1997 and U.S. Patent Application Ser. No. 09/030,106 filed Feb. 25, 1998 which in turn claims the benefit of U.S. Provisional Application Ser. No. 60/039,164 filed Feb. 26, 1997.)

*"...A system for monitoring and/or affecting parameters of a patient's body comprised of a system control unit (SCU) and one or more other devices implanted in the patient's body, i.e., within the envelope defined by the patient's skin. In accordance with a preferred embodiment, the system operates in closed loop fashion whereby the commands transmitted by the SCU are dependent, in part, on the content of the data signals received by the SCU..."*

*... a preferred SCU is also implemented as a device capable of being injected into the patient's body... Wireless communication between the SCU and the other implanted devices can be implemented in various ways, e.g., via a modulated sound signal, AC magnetic field, RF signal, or electrical conduction....In accordance with a further aspect of the invention, the SCU is remotely programmable, e.g., via wireless means, to interact with the implanted devices according to a treatment regimen ... While the invention herein disclosed has been described by means of specific embodiments and applications thereof, numerous modifications and variations could be made thereto by those skilled in the art without departing from the scope of the invention set forth in the claims. For example, a system including multiple SCUs, e.g., one external and one internal, is considered to be within the scope of the present invention. Additionally, while the use of a single communication channel for communication between one or more SCUs and the other implanted devices has been described, a system implemented using multiple communication channels, e.g., a first sonic channel at a first carrier frequency and a second sonic channel at a second carrier frequency, is also considered to be within the scope of the present invention ... Transponders, e.g., are devices which can be used to extend the interbody communication range between stimulators and sensors and other devices, e.g., a clinician's programmer and the patient control unit. ... The clinician's programmer and/or the patient control unit and/or other external control devices can also communicate with the implanted devices, as described in the parent application, ... Alternatively, such external devices can communicate with the SCU via a transceiver coupled to the programmable controller. Since, in a preferred operating mode, the signal transmitter and signal receiver operate using sound means, a separate transceiver which operates using magnetic means is used for communication with external devices. However, a single transmitter/receiver can be used in place of transceiver if a common communication means is used.."*

**4.3 – U. S. Patent: 6,175,764: "Implantable microstimulator system for producing repeatable patterns of electrical stimulation"** Inventors: Loeb; Gerald E. (Kingston, CA); Richmond; Frances J. R. (Kingston, CA)

Assignee: Advanced Bionics Corporation (Sylmar, CA) Appl. No.: 490921 Filed: January 25, 2000

(This application is a Divisional Application of U.S. application Ser. No. 9/077,662, filed May 29, 1998, now U.S. Pat. No. 6,051,017 which application is incorporated herein by reference, and which application was filed in the U.S. under 35 U.S.C. .sctn.371 based on



international application number PCT/US97/02576, filed Feb. 19, 1997; which international PCT application claimed priority to U.S. Provisional Applications Nos. 60/011,870; 60/012,019; 60/011,868; and 60/011,869; all filed Feb. 20, 1996.)

*“...Improved implantable microstimulators are covered with a biocompatible polymeric coating... The microstimulator systems include external control for controlling the operation of the microstimulators. The control include memory for programming preferred stimulation patterns for later activation by the patient or caregiver.... can be implanted non-surgically by injection. ...upon an external command, or at predetermined intervals, power and command signals sent from controller cause the various microstimulators to emit a series of electrical current pulses (i.e., a pulse train) at the desired frequency and amplitude sufficient to cause the muscles to lift the body for the duration of the pulse train. ...also provides storage and production means for a program of output currents and stimulation pulses that may then be produced autonomously by the implanted device without requiring the continuous presence of extra corporeal electronic components, i.e., without the need for an external control device.*

4.4 – U.S. Patent 6,164,284: “System of implantable devices for monitoring and/or affecting body parameters” Inventors: Schulman; Joseph H. (Santa Clarita, CA)

(This application claims the benefit of U.S. Provisional Application No. 60/042,447 filed Mar. 27, 1997 and U.S. patent application Ser. No. 09/030,106 filed Feb. 25, 1998 entitled “Battery-Powered Patient Implantable Device” which in turn claims the benefit of U.S. Provisional Application No. 60/039,164 filed Feb. 26, 1997)

*“...it is preferably that at least a portion of the program storage be an alterable form of memory, e.g., RAM, EEPROM, etc., whose contents can be remotely altered as described further below. However, it is additionally preferable that a portion of the program storage be nonvolatile so that a default program is always present. The rate at which the program contained within the program storage is executed is determined by clock, preferably a real time clock that permits tasks to be scheduled at specified times of day...”*

4.5 - U.S. Patent Application 20010054071: “Audio/video conference system for electronic care giving”  
Loeb, Gerald E. December 20, 2001

4.6 - U.S. Patent 5,791,344: “Patient monitoring system”  
Inventors: Joseph Schulman, A.E. Mann Foundation Appl. No.: 582756 Filed: January 4, 1996

4.7 – U.S. Patent 5,551,016 “Monitoring system and interface apparatus therefor”  
Inventors: Loeb; Gerald E. Assignee: Queen's University at Kingston Appl. No.: 084928

*“In monitoring systems for acquiring data about a subject, such as are used, for example, in the medical, scientific and engineering fields, determination of temporal relationships between data acquired from multiple monitoring devices is facilitated by means of an interface unit which interconnects the monitoring devices with tape recorders for storing the data and a computer for processing the data.”*

Summary Article 4: These systems allow Schulman, Loeb and Troyk to interact with the implanted devices via radio frequency, and enables this interaction from any location that allows them (Internet access) to connect to a remote server.

---

#### Article 5. - Timing 1:

U.S. Provisional Applications No. 60/039,164 was filed on Feb. 26, 1997 and U.S. Provisional Application Ser. No. 60/042,447 on Mar. 27, 1997 –

These patent filings pertain to the patient monitoring systems and other technology applicable to a long term human study and include methods capable of long range bi-directional telemetry.

Summary Article 5: The above patent filings occurred during the same quarter as my assault

---

## Article 6. - Timing 2:

Schulman and Loeb report the first functional microstimulators in a QPR just prior to my assault, and report “tightening up” the accountability and traceability of prototype devices in a QPR following my assault:

*“...we were able to demonstrate the feasibility of using a single coil in the microstimulator both to receive power and data and to act as the tank circuit and antenna for outgoing RF transmission at the same frequency. The next silicon breadboard is a demonstration of the feasibility and achievable modulation rates for incoming and outgoing data transmission based on the suspended carrier mode of operation. This will be done by a chip that records in a shift-register the sequence of detected carrier half-cycles during incoming RF transmission. The chip then telemeters out that information as a sequence of AM encoded bits..... This design has passed intensive simulation and is now in layout. Release to the foundry is anticipated in August, with chips likely to be available for evaluation by early October...”* (QPR #5 March 10, 1996 - June 9, 1996)

*“In the next quarter, we intend to finalize and validate the sealing process and provide the necessary active implants to support preclinical chronic animal tests.....”* (QPR #7 September 10, 1996 - December 9, 1996)

*“During this quarter, we tightened up the traceability and documentation of BION prototype production units.....”* (QPR #10 June 10 1997 – Sept 9 1997)

Summary Article 6: This proves that Schulman had achieved functional implants with back telemetry prior to my assault and that he expressed intent to chronically implant an animal during the following quarter which corresponds with the date of my assault. Additionally, it shows that following my assault, there was concern for the accountability of pre-production devices and chips.

---

## 7. Schulman has an FCC license (K6BWA) and has 224.840MHz repeaters operating to the North, South and East of my residence, all within a 22-mile range

An FCC database search reveals that Joseph Schulman holds an “Advanced” class radio license, call sign K6BWA, and that he upgraded his license as recently as Oct 2001. Evidence shows that repeaters (224.840 MHz) have been arranged in which licenses attributed to Danny Schulman and Barbara Schulman are involved and their location could facilitate access to the repeater data by Gerald Loeb, James T. Bennett, and even Phil Troyk. Because Joe Schulman license is used for a repeater in "Hollywood Hills" (yet lives 35 miles North from this location), and Danny Schulman's (lives in Los Angeles) is used for one located in "Altadena", which is 9 miles from Gerald Loeb, there seems to be some confusion as to where this hardware is and who can access it. Additionally, Robert Mancini (WB6PRR, note the “W” in call sign) apparently lives in Oak Lawn, IL (14 miles from Troyk @ IIT), but his license is attributed to another repeater in Altadena (why two both in Altadena?), this is also more than suspicious. Barbara Schulman will not even admit she is a radio operator, as attempts to contact here by email result in replies from Joe Schulman. Because Joe Schulman's license is attributed to a repeater located in Hollywood Hills, so the repeater at his home or AEMF in Santa Clarita is operated by Joe under his wife's license. One interesting observance is that neither “Sand Canyon”, nor “Hollywood Hills” is a real city, which again, is more than suspicious. Businesses bearing the name “Hollywood Hills Whatever” (such as Plumbing or Cleaners), are located in zipcode 90027 (which is Los Angeles, 4 miles from zipcode center to James Bennett's residence) and “Sand Canyon” is a road in Santa Clarita that runs past the residence of Joseph Schulman. The tables of information that follow show:

The Repeater Network (Frequency / Call Sign / Location of the Hardware / Status <open or private>)

The FCC Licenses (Callsign / Assignee / Address)

Physical Map of Repeaters (Shows locations of repeaters in relation to my residence)

Private Repeater / Auxiliary Network:

Freq (MHz)	Call Sign	Location of Repeater Hardware	Status / Mode
224.820 -	KJ6JY	SANTIAGO PEAK (CONDOR)	OPEN
224.840 -	K6BWA	HOLLYWOOD HILLS	PRIVATE
224.840 -	KA6UFC	SAND CANYON	PRIVATE
224.840 -	KS6E	SHERMAN OAKS	PRIVATE
224.840 -	KS6K	SILVERLAKE, L.A.	PRIVATE
224.840 -	N6EHD	ALTADENA	PRIVATE
224.840 -	WA6MTM	YORBA LINDA	PRIVATE
224.840 -	WB6PRR	ALTADENA	PRIVATE
224.860 -	N6ENV	LA COUNTY FIRE CAMP 49	OPEN

Information above has been confirmed from three sources:  
[www.220sma.org/smarpt.htm](http://www.220sma.org/smarpt.htm)  
[http://www.gsl.net/cora/222\\_Mhz\\_Repeater\\_Guide.htm](http://www.gsl.net/cora/222_Mhz_Repeater_Guide.htm)  
<http://www.geocities.com/CapeCanaveral/Hangar/6847/>

Identity of above call signs from FCC:

FCC Callsign Lookup <a href="http://www.fcc.gov">www.fcc.gov</a>		
Callsign	Licensee Name	City (according to FCC license)
K6BWA	SCHULMAN, JOSEPH H	SANTA CLARITA
N6EHD	SCHULMAN, DANNY I	LOS ANGELES
KA6UFC	SCHULMAN, BARBARA L	SANTA CLARITA
KS6K	LEIDNER, JOEL D	LOS ANGELES
KS6E	FREEDMAN, MARVIN	LOS ANGELES
WB6PRR	MANCINI, ROBERT F	ALTADENA

Physical Locations:

The map shows the Los Angeles metropolitan area with a red dot representing the author's residence in Santa Clarita and several yellow dots representing repeater locations. The repeaters are located in Santa Clarita, Hollywood Hills, Altadena, and West Hollywood. Major highways like I-5, I-10, and I-405 are visible.

Legend:  
 ● = My Residence (Red)  
 ● = Repeater (Yellow)

Joseph Schulman's repeater / auxiliary station locations (\*yellow) are shown in relation to my residence (\*red) in the map above. Schulman's RF repeaters are located to the North, South, East, and South East of my residence, all within a 22 mile radius. There is reason to believe Schulman arranged this repeater network through his membership in the ARRL (American Radio Repeater League).

Summary of Article 7: Joe Schulman has repeaters surrounding my residence and their locations do not match that of the licensee's. Additionally, one of the licensee's is located 14 miles from Phil Troyk at the Pritzker Institute/IIT. Other repeaters are located near Loeb and Bennett. The repeater locations are more than just coincidence. Enforcement is needed from the FCC and law enforcement.

\*Update - I just spoke to Marvin Freedman KS6E at his home. Marvin Freedman told me that “he hasn’t been active as a HAM for years” and that “I’m not repeating from Sherman Oaks” and “I’m looking at my equipment,... it’s unplugged, sitting here in Los Angeles”. He also told me that he and Joe Schulman were “friends”. This suggests that Joe Schulman could be using Marvin Freedman’s FCC license without his consent”.

Joe Schulman is using his “old friend’s” FCC license for a repeater that is located on or near my Mother’s property in Encino. He is also using his own license, K6BWA for a repeater that is being operated by James Bennett located at 2288 Bronson Hill Dr. Los Angeles (aka “Hollywood Hills”), and he is using a license that he obtained in his wife’s name (his wife would not admit to me that she was a radio operator and insisted I speak to her husband about it) for a repeater at his residence in Santa Clarita (aka “Sand Canyon”). In fact, of the six repeaters listed in the table below, 3 are in non-existent cities (Hollywood Hills, Sand Canyon and Silverlake are not real cities with no US zip code). Joe Schulman is using this radio signal to control biomedical sensors which is part of his work at the Alfred E. Mann Foundation ( <http://www.aemf.org/> ) which developed sensors called the “BION”, (<http://ami.usc.edu/Projects/Bion/media.asp> or

<http://npp.ninds.nih.gov/ProgressReports/MicrostimulatorsAndMicrotransducersForFunctionalNeuromuscularStimulation%20NS52325/MicrostimulatorsAndMicrotransducersForFunctionalNeuromuscularStimulation%207NS52325.pdf>

Marvin Freedman lives in Los Angeles and can be contacted below:  
Marvin Freedman KS6E 310-470-1116

---

#### **8. Professor Wise from Univ. of Michigan reports in a QPR that because of bandwidth and impedance issues, 200MHz should be chosen for remote, wireless stimulation using a neural prosthesis**

As if articles 1 through 4 were not enough evidence to prompt action, further evidence of the repeaters use is the frequency Joseph Schulman’s repeater/auxiliary stations are operating at (224.840 MHz). The significance of this is best explained by Professor Kensall D. Wise, University of Michigan, respected for his many years and considerable contributions to the NINDS/NPP:

*“...the most important issue of an implanted biotelemetry system is power dissipation. To minimize the power of a transmitter, we have to select the lowest practical RF carrier frequency, since RF circuits dissipate power that is proportional to their operating frequency. The FCC has authorized unlicensed use of the 38-41MHz, 88-108MHz, and 174-216MHz Very High Frequency (VHF) bands for Industrial, Scientific and Medical (ISM) telemetry purposes. Other bands in the microwave region (900MHz and above) are not considered because of high tissue absorption at those frequencies. On the other hand, a higher carrier frequency is desired for wider bandwidth in data transmission. As the result of these trade-offs, 200MHz is chosen as the carrier frequency...”* (recent QPR #5, (Oct/Dec 2001) Contract NO1-NS-0-2329 “Thin-Film Intracortical Recording Electrodes)

Ken Wise has been involved with the NPP and using micromachined Silicon electrodes to stimulate and record from the nervous system since as early as 1992, and is obviously very accomplished and knowledgeable. In the above, he establishes 200MHz as the preferred carrier frequency because higher frequencies use too much power, and lower frequencies do not penetrate the impedance of the tissue as well as higher frequencies.

Summary Article 8: If someone wanted to use wireless radio frequency to stimulate and record from an unwilling human participant, a frequency near 200 MHz would be desired

---

**9. Schulman reported acquiring an ISDN “Connectix” phone as part of contract #N01-NS5-2325 claiming it would be used for file sharing and later, a company he founded called “NeuroDyne” markets a new system which used such an ISDN video-phone to enable multiple physicians to tele-conference while analyzing a patients biofeedback signals**

In QPR #5 (March 10, 1996 - June 9, 1996), Schulman reports:

*“In order to facilitate communication between the Mann Foundation (Schulman) and Queens University (Loeb), an ISDN telephone line and high speed modem have been installed with service to the Internet. The Connectix VideoPhone system is in use and remote file access and terminal sessions will soon be available between engineering and documentation computers at both locations...”*

This ISDN video phone was actually used to provide real-time bio-feedback data from Schulman in Valencia CA, to Loeb at Queens University, and allow them to conference and interact while monitoring me and the biofeedback from their microstimulators during this unethical and illegal research. They also based parts of their “patient monitoring system” patents on this. The fact that Schulman and Loeb used the Connectix Phone for more than “remote file access” is substantiated by this press release from NeuroDyne, a corporation founded by (not a coincidence) it’s President, Joseph Schulman:



NeuroDyne/E-Tech Press Release:

*August, 1999: A new Internet technology will allow a patient to carry on a face-to-face conversation with a doctor who is a few miles or even thousands of miles away while his physiological measures, such as EKG, EEG, EMG, GSR etc. are displayed in real-time on the doctor's screen for assessment... the ability for healthcare colleagues and specialists several thousand miles apart to conference concerning a patient, while interacting via audio, video and active monitoring of the patient's physiology.... Optionally, a Video Camera, VoxPhone software, and an Internet connection, allow two individuals virtually anywhere in the world to carry out a private conversation or to support a conference with up to five individuals. Under the joint venture, the VoxPhone interface will be expanded to allow the transfer of real-time data from NeuroDyne Medical's biomonitors instrumentation in addition to voice and video information. This physiological information can then be displayed graphically on both computer screens as well as being saved for further analysis.....”*

Summary Article 9: It is becoming obvious (it isn’t already), that Schulman and Loeb are guilty of my allegations. They implanted these microstimulators using a syringe leaving virtually no externally visible evidence and they expect you and others not to believe it because “it sounds crazy”. You don’t have to believe me but you do have to believe the evidence. The evidence is far too much and too convincing to be “amazing coincidence”. All of the evidence presented is documented, factual and proven. No speculation or fabricated conclusions. Just fact. This is currently page 15 of about 50 and the evidence in the following pages don’t make Schulman and Loeb look any more innocent. That is because they are not.

---

**10. Schulman reports a “faulty” diode and resistor required that they be removed in-house and replaced, however the patents of Schulman, Loeb and Troyk allow diodes and resistors to be eliminated by integrating a chip based on the suspended carrier telemetry protocol resulting in a smaller device package size.**

Schulman reports vendor errors have required a diode and resistor to be cut from the chip (removed) and then replaced. It is specifically indicated that these components were “defective” and that they would be

replaced with two new (off the shelf) surface components and that doing so required in-house post-processing. The legitimate reason behind elimination of the resistor and diode involves Schulman's want and need for a smaller implant, so placing tiny additional circuitry with the implanted chip would be needed and thus, also would need to be explained. I contend that the components were not defective and that they were not replaced with two comparable components as indicated. The reason Schulman removed them is because the diode and resistor would not be necessary after stacking an additional, small circuitry, (based on the Suspended Carrier telemetry protocol) onto the existing ASIC as described above. This additional circuitry was for the mixed signal demodulation required for long-range RF bi-directional telemetry and also performed DAC, clock, state machine and other functions. Schulman lays it on thick in QPR's, however contradicting (but accurate) explanation exists in their patent filings where they are forced to be honest in order to protect their invention:

QPR #3 conceals the reason for removing the diode and resistor by claiming the components are defective:

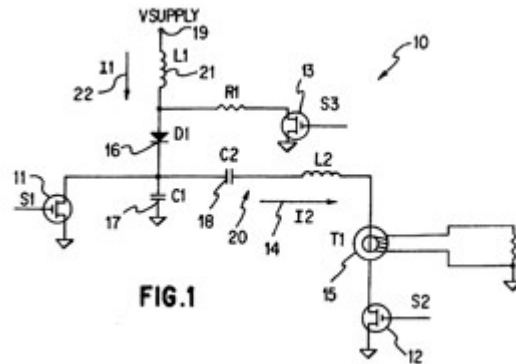
*"...we were dealing with a chip that had a defective diode on board, which made the chip useless. Luckily, the position of the diode in the circuit and physically on the chip was such that it could be effectively removed from the chip and replaced by an external diode. Moreover, the resistor, essential for the data demodulation, was also conveniently placed and could be replaced by an external device. Having two additional devices in a small package required redesign of the package which in turn demanded repositioning of the bonding pads using a relatively inexpensive gold-bumping process..."* (QPR #3, Sept 10, 1995 – Dec 9, 1995)

In the patent below, Schulman describes how additional circuitry can be used to upgrade an implant device from a system that requires an external coil and driver circuitry, to one that is fully implantable. One doesn't have to look far past contract #N01-NS5-2325 when wondering how Joseph Schulman came up with such a concept. Other descriptions in U.S. Patent 6,067,474 have allowed me to assess that Schulman and Loeb refer to the system used for their illegal and unethical research as a "proximity system" and that it was developed as part of NIH microstimulator contract N01-NS5-2325 and substantiated by its filing date.

*"Appropriate switching circuitry is included with the battery (or capacitor) in the second device to convert the dc power of the battery to ac power for transmission to the first device. This ac power may be modulated, as desired, to also transmit information, e.g., control signals, from the second device to the first device. Yet a further embodiment of the invention, hereafter referred to as the "proximity system" embodiment, resides in a fully implantable system that includes first and second implantable devices. The first device houses electronic circuitry for performing a desired function. The second device houses a replenishable power source, and may also include additional circuitry. Thus, one use of this proximity system embodiment allows a second device.. that has heretofore been included in an external device, to be implanted proximate an implant device of an existing system, thereby effectively upgrading the existing system to a fully implantable system."* (6,067,474: "Implantable device with improved battery recharging and powering configuration" Inventors: Joseph Schulman (U.S. Provisional Application Ser. No. 60/054,480, filed Aug. 1, 1997)

U.S. Patent Office filing contains accurate information:

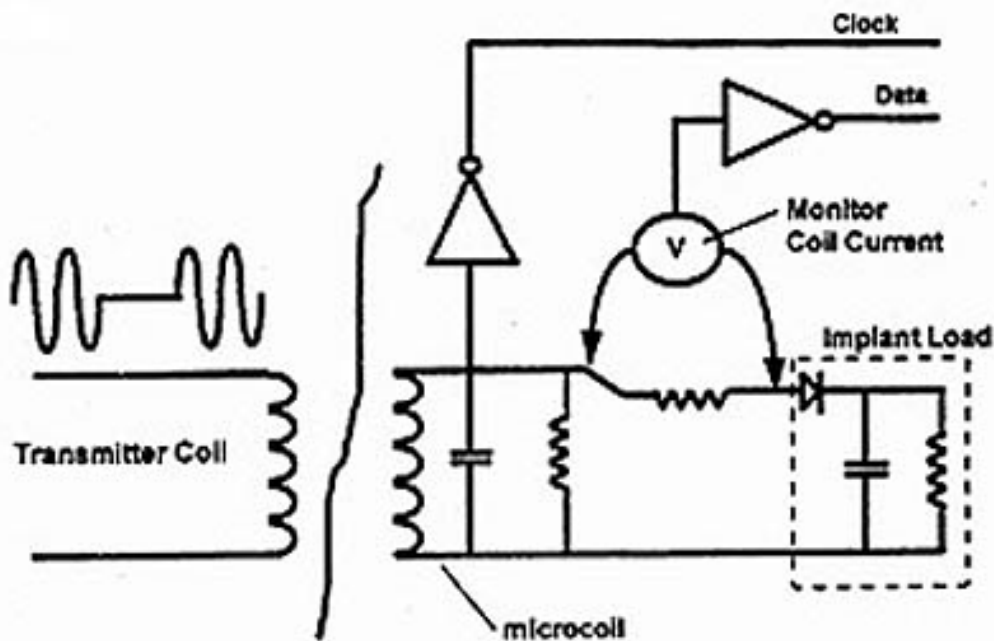
Patent 5,697,076: "Suspended carrier modulation of high-Q transmitters" (the only patent legitimately indicating that it was made under government support) December 9, 1997, Inventors: Troyk; Philip R.; Heetderks; William; Schwan; Martin; Loeb; Gerald Assignee: Illinois Institute of Technology (Chicago, IL); United States of America as represented by the Department of Health and Human Services (Washington, DC) Appl. No.: 432605 Filed: May 1, 1995



“...When switch 13 is closed, current 22 rises to  $(V_{supply}/R1)$  with an  $R1/L1$  time constant. Thus, with the proper choice of  $R1$ , if switch 13 is closed long enough in advance of the closing of switch 12 (restart), the proper initial condition for current 22 can be achieved. When switch 12 is closed and Class E operation resumes, switch 13 opens, causing current 22 to flow into diode 16 instead of  $R1$ .

...Elimination of the  $R1-S3$  branch may be possible with an appropriately sized choke inductor 21. In this case, the current 22 is allowed to go to zero, even for “short” off periods.”

The above diagram is from their patent and obviously shows accurate information. The diagram that follows however, from NIH QPR #10 contains less “precise” information and further conceals how suspended carrier was being used during contract #N01-NS5-2325 :



**This diagram is pictorial only, and does not precisely show the method used to sense the rectifier currents**

“Not precisely” isn’t even truthful because the truth would read “not even close”

The patent filing below shows how diode and resistance are present in the circuit diagram, but are not represented by actual components:

U.S. Patent 6,035,237: "Implantable stimulator that prevents DC current flow without the use of discrete output coupling capacitors"

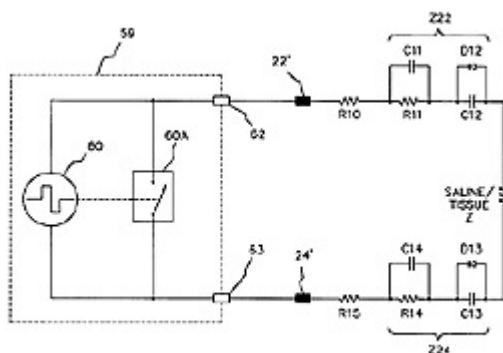


FIG. 10

*"As illustrated in FIG. 10, the coupling capacitor is actually formed by using the electrode-saline interface that results when a conductive electrode comes in contact with saline body fluids. In FIG. 10, Z22 represents the complex impedance between electrode 24 and the saline. Capacitor C12 and diode D12 (as well as capacitor C13 and diode D13) represent the passivation or oxide insulation layer that exists at the surface of each stimulating electrode."*

In this invention, the result of the oxide insulation layer functions as a capacitor and diode, so although the circuit layout shows a diode, there is only oxide build-up that causes some resistance and capacitance and no true hardware components.

There is no mention of chip stacking in the text body of contract QPR's and when Schulman finally reports using a chip allowing elimination of the diode and resistor (which ironically doesn't happen until the final QPR in 1999), the additional circuitry is showed next to the existing circuitry in a planar configuration. This poses the following question: If no chip stacking occurred, what is meant in the text header contained in QPR #2 following subject header "Work at the Alfred E. Mann Foundation" that reads as follows:

**3. Electronic assembly, stacked chips, side by side chips**

*"As reported in the 1st Progress report of this Contract, a plan was developed to rework the microstimulator integrated circuit wafers which had been fabricated at the end of the first microstimulator NIH contract. That circuit was not fully functional due to problems associated with the on-the-chip demodulation / rectifier diode....."*

Additionally, if no chip stacking was used for the BION microstimulator, what was the cause for the abnormally low yield of microstimulators which was consistently around 30%? The "Evidence Detail" section of this document has shown a stacked chip device recovered from my person and Schulman's patents also describe a stacked chip configuration, most of which were initially filed during the microstimulator contracts period. Although Schulman eventually revealed the method of additional circuitry that was capable of eliminating the diode and resistor, the added chip was shown in a planar configuration rather than stacked which requires more substrate area. He concealed this throughout the duration of the contract and only partially revealed it in ironically, the final QPR of the contract in 1999. His U.S. Patent applications prove that he was aware of and practicing this invention as early as 1995 and the QPR's during the four years of the contract never mention fixing the chip design so as not to necessitate the addition of replacement components which suggests that this method of adding additional circuitry was occurring for the duration of the contract beginning with QPR1 in 1995 when he reported a need to cut "faulty" diodes from the chip. The final QPR from 1999 offers an illustration which shows one simple chip added next to (not stacked) the existing circuitry:



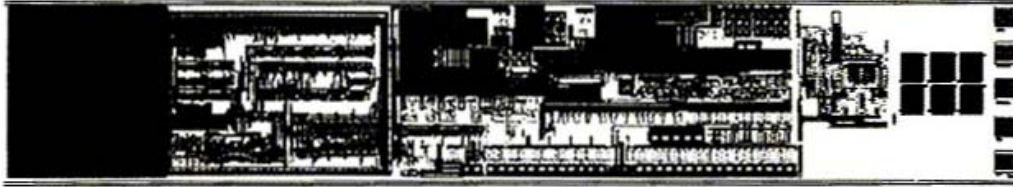


Figure 10 - 2MHz microstimulator with integrated rectifier and telemetry front end. (Front End)

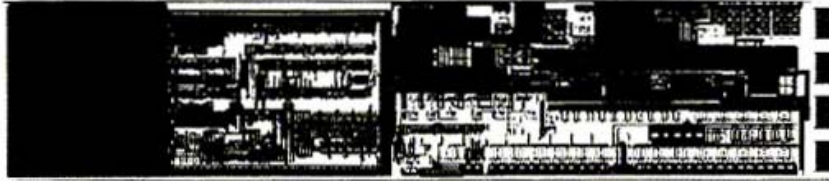


Figure 11 - Original 2MHz microstimulator that used external diode rectifier

*“...we revised the rectifier/data demodulator of MOS8 for integration with existing 2MHz microstimulator circuitry. The existing external diode, amplitude demodulator, and clock recovery circuit were replaced by a new ASIC cell that uses the techniques of MOS8. (Images 10 and 11, above) Presently these chips are under test...”*

It is reported that the above ASIC “cell”, which finally solves the 4 year old resistor/diode problem “uses techniques” of MOS8. I contend that this “cell” exists on a ASIC design layout that was submitted during the first year of the contract and that Schulman waited until the final QPR#16 to reveal it. This cell could have been present on (and cut from) any number of the early wafer/chip runs that he falsely reported being “unusable”.

It is reported multiple times that a “faulty” diode needed to be replaced and is shown again in this instance:

*“A major effort was expended dealing with the new microstimulator development and in correcting silicon foundry errors via chip microsurgery ...a new corrected layout was sent to the foundry for another run of wafers... when the wafers returned, it was observed that they did not work ...the design rules provided by the foundry allowed the shorts ...by cutting a trace on the integrated circuit, the external diode could be reliably removed. A diode-resistor combination could then substitute for the defective on-chip components.” - Schulman/Loeb/Troyk QPR#1 – March 10, 1995 – June 9, 1995*

Again, patent documentation makes argument against Schulman’s reporting, and here he describes how removal of components followed by adding additional small circuitry can actually result in a smaller overall package:

Patent 6,035,237: “Implantable stimulator that prevents DC current flow without the use of discrete output coupling capacitors”

*“The implantable stimulator of claim 3 wherein said first and second circuit means are realized using CMOS transistors configured to function as prescribed circuit elements on the same semiconductor chip ...while such DSP circuitry may utilize a significant number of CMOS transistors, configured into appropriate processing and logic circuitry, the overall space required by such DSP or other circuitry on the chip, or a supplemental chip, particularly given the smaller trace sizes associated with modern CMOS devices (0.8 micron and smaller) could still be less than using discrete coupling capacitors (resistor).*

The repositioning of the bond pads was done to create electrical contacts for attaching the additional suspended carrier/long-range telemetry circuitry which also allowed the elimination of diode/resistor. This activity is not revealed in NIH QPR's, because Schulman claims that the additional bond pads were needed to attach the diode and resistor which was cut off of the original chip. Further evidence can be found in U.S. Patents 6,315,721 and 6,164,284 which describe the following:

*"...The SCU 302 is primarily comprised of (1) a housing, preferably sealed and configured for implantation beneath the skin of the patient's body as described in the parent application in reference to the implanted devices, (2) a signal transmitter in the housing for transmitting command signals, (3) a signal receiver in the housing for receiving status signals, and a programmable controller, e.g., a microcontroller or state machine..."*

The "state machine" mentioned above is included in the additional circuitry that Schulman used which allowed the elimination of the resistor and diode

*"...For example, sensor circuitry can be coupled to the electrodes ...and provided the sensed data to the controller circuitry. Preferably, the sensor circuitry includes a programmable bandpass filter and an analog to digital (A/D) converter that can sense and accordingly convert the voltage levels across the electrodes into a digital quantity. Alternatively, the sensor circuitry can include one or more sense amplifiers to determine if the measured voltage exceeds a threshold voltage value or is within a specified voltage range. Furthermore, the sensor circuitry can be configurable to include integration circuitry to further process the sensed voltage. The operation modes of the sensor circuitry is remotely programmable via the devices communication interface..."*

The programmable bandpass filter is also part of the additional circuitry that Schulman integrated after eliminating the diode and resistor during his NIH contract and the additional functionality of the added circuitry enabled bi-directional telemetry over greater distances than with a normal inductively coupled link. In order to sense and monitor as described above in patent text, and also the ability to do so with only 2-5% modulation and switch in 4 clock cycles or less which is described in Schulman's QPR's, the "sensor circuitry" mentioned above must be implanted with (onto or next to) the microstimulator. This is in contrast to a typical inductive link where this sensing and monitoring takes place between a coil attached outside the skin and an implanted device where the impedance of the tissue and the increased distance do not allow the low modulation and instantaneous switching. Although much of this evidence pertains to Schulman and Loeb, Phil Troyk worked very closely with them on the Suspended Carrier chip, the back telemetry and also the overall packaging. The fact that it is reported that Martin Schwan performed the removal of diodes and resistors at IIT also suggests that both Troyk and Schwan direct knowledge of Schulman and Loeb's intent with the device. In email correspondence, Phil Troyk responds to my inquiry about addition of the small circuitry by denying any intent for it to be added to the implanted package and claimed that it was to exist outside the skin, but then in an about face, claims that the idea was "one of the other team members" and that he "saw no benefit to it". This email correspondence is attached at the end of this document.

Summary Article 10: Respected Investigators, Professors and Scientists are capable of telling lies, they do tell lies, and Schulman has told more than his share. The diode and resistor were not defective. They just needed the additional circuitry integrated / added into or onto the implanted portion.

**11. In a table of data listing the specs for components of the "BION" microstimulator, Schulman indicates that the diode used in replacement of the "faulty" diode is a "THD9064", however there doesn't appear to be any such diode in existence.**

In QPR #2 (June 10, 1995 - Sept 9, 1995), a table of "Specifications" is present that provides details regarding each of the "BION" microstimulator components and indicates a mystery diode. Additionally, a table of "Suppliers" lists an undetermined vendor for an unknown component.

Suppliers:

1. Kimbel Borosilicate Glass

- 2. AVX Tantalum Corp
- 3. Vendor to be selected by A.E. Mann Foundation

Specifications:

Part	Abbrev	Specification
Glass Capillary	cap	0.079" OD
Glass Bead	bd	0.015" ID 0.065 OD
Iridium Ball	Ir ball	0.065" OD
Diode	D	THD 9064
Integrated Circuit Chip	IC chip	3 micron, double poly, CMOS

1. THD 9064 does not appear to be a commercially available diode. The fact that no such "THD9064" diode exists should alone prove that Schulman eliminated the diode rather than replaced it and the false data above is reason for Alan Price and John Krueger at the Office of Research Integrity to find Schulman guilty of misconduct and because he didn't act alone, the other Investigators should be subject to disciplinary measures as well.

2. A search for THD 9064 on two popular search engines (Hotbot and Google using the terms diode thd 9064 in various forms) returns a total of 3 results: (1) New Focus 12-GHz Amplified Free-Space Photoreceiver (which uses a MEMS class 25µm schottky diode), (2) Siemens SDA9064-5 Digital Deflection MicroController (frequency doubler), and (3) another microcontroller from ICS.

---

**12. Throughout contract #N01-NS5-2325, Joseph Schulman fabricates an absurd amount of "vendor error" and "faulty product" which was done to conceal the real reason for the excessively low yield of microstimulators which can be attributed to the error prone stacking process of the smaller unethical implants**

*"Yield of the microstimulators after winding is approximately 30%. The reason for this poor yield is not yet known. The integrated circuits have been pretested. In some of the defective stimulators, measurements show that the external diode and resistor are not properly connected. In others the precise cause is unknown. Presently, test fixtures are under development, at the Mann Foundation, to permit testing of the electronic module at all assembly levels." (Contract N01-NS5-2325 QPR#3)*

This atrocious yield of 30% is factual evidence and should have flagged concern upon NIH review. For those not totally familiar regarding ASIC fabrication process, a brief explanation may help. When Schulman says the circuits have been pretested, this is pretty understandable as testing a single IC after production is a simple process. The window of consistency for yield of chips from foundry runs industry wide (in process sizes ranging from .05µ to 3.0µ, CMOS or BiCMOS) is always in excess of 96% and is typically approaches 99%, which means there is very little "vendor error". Schulman indicates that after he gets his hands on them, yield falls to 30% and struggles to explain, but indicates that the problem is "unknown", but "some" of the problem is that the diode and resistor replacing the "faulty" components that were removed at IIT are not "properly connected". The "BION" microstimulator documented in this contract does not stack circuits in a vertical or 3D fashion and attaching basic components to bond pads in a 3.0µ or 1.5µ CMOS process is not immensely difficult, especially when you take into account the experience of Mann personnel who have prior experience with smaller chip processes used in cochlear implants and at IIT who is a leader in MEMS scale and Nano scale fabrication. There is another reason for 70% of the finished product being non-functional and the problem is not "unknown", it is deception that Schulman must conceal because the only legitimate explanation for this poor yield is that the chips were being stacked in order add functionality (long range telemetry) and save space and a quantity were being misappropriated as well.

Here a “clerical error” causes open pads, however open pads such as these would be beneficial if your intent was to dice and stack the chip:

*“Among the chips included on the wafer run was a test chip for the suspended carrier front end. Due to a clerical error in the process of combining the chips onto one reticule, that test chip had open circuits at many of the pads, making testing of the chip essentially impossible. These opens and any other problems which can be fixed with metal layer changes will be corrected on the wafers which have been held in the early stages of processing. As this report is being prepared for submittal, the corrections in the metal layers have been submitted for two more wafer releases.” (QPR #15)*

In this quarter, Schulman blames Slicex, a vendor in Salt Lake for two different errors and claims an entire run of chips are useless

*“Prior to delivery of the chips it was discovered that an unfortunate error existed in the layout at the upper-most cell level. A long strip of via/contact had been placed, by Slicex, across the entire length of the repeater chip rendering it inoperative. Equally unfortunate was the substitution of cells from an earlier revision of the layouts in the test chip. Therefore neither of these chips were functional upon delivery. Fortunately, some of the circuits used in the Repeater chip design had been included in another MOSIS submission as part of work on ITT's contract: Multichannel Transcutaneous Cortical Stimulation System. Therefore we were able to use that chip, MOS2, to test the power supply regulator, and rectifier/data decoder Both of these circuits are key components for the telemetry of the proposed micromodules” (QPR #10)*

E-mail correspondence with Tom Wolf, President of Slicex, reveals conflicting information:

*Mr. Wolf writes: “...We brought Dr. Huber in primarily to address the latch up problem as AEM (Alfred E Mann Institute) was pushing the IC process beyond its ability... Dr. Schulman wanted to use an entire reticule which meant we placed 5 IC's in a stack, and then 2 IC's rotated 90 degrees at the end. At this point, the industry did not have good tools for debugging multiple IC's on a reticule (I am not sure if this is possible even today). But at that point, the only way to check was with the human eye. Shortly after the IC came back and was deemed acceptable by Schulman, I visited Schulman at AEM. He expressed his satisfaction with our work and then introduced me to Ron Liebel of MiniMed. He told Ron that he should use us and that we were great, just expensive. After which, we helped MiniMed with their artificial pancreas chip. At this point Dr. Schulman stated he had lots of new work for us, but we never saw any more from him. Just his sister companies, MiniMed and ABC (ABC = Advanced Bionics Corp) Now at ABC, we did several million dollars worth of business. I don't understand that Schulman could say we didn't understand anything, when we designed 1 IC for him that worked and 17 for ABC. Especially since according to Schulman, the only IC that had worked first time for AEM up until that point in time was the one done by SliceX....”*

In email correspondence between Joseph Schulman and myself, a different scenario is portrayed:

*“I WAS CONCERNED BECAUSE THEY ALL SAT QUIETLY AND DIDN'T ASK ANY QUESTIONS, AND WHEN I ASKED THEM IF THEY WERE SURE THEY UNDERSTOOD ALL THE SCHEMATICS WE WERE DISCUSSING THEY WOULD ALWAYS ANSWER YES. FINALLY WHEN THEY PRODUCED THE CHIPS, IT WAS OBVIOUS THAT THEY DIDN'T UNDERSTAND ANYTHING AND THUS HAD MANY ERRORS IN THE LAYOUT. WHEN I POINTED OUT THE ERRORS TO THEIR MANAGEMENT, THEY PROMPTLY RETURNED THE FUNDS WE PAID THEM. ANOTHER COMPANY THAT WORKED WITH THEM LOST OVER A MILLION DOLLARS DUE TO THEIR SCREWUPS...”*

Mr. Wolf mentions that Schulman was stacking a full reticule as well as “pushing the process beyond it's ability” which indicates that the design was sacrificing reliability in favor of a smaller chip size. If this chip were just being used for the BION, space savings should not have been a concern as this chip was in a 1.5µm process which is considerably smaller than the 3.0µm chip that the project began using initially and as shown below in another section, there is enough room inside the BION glass capsule for circuitry and this is definitely not an issue. Although Mr. Wolf does admit some troubles with the full reticule, he indicates that he

is certain that Slicex is not responsible for an entire run of chips being non-functional and that they provided Schulman and AE Mann with a working design.

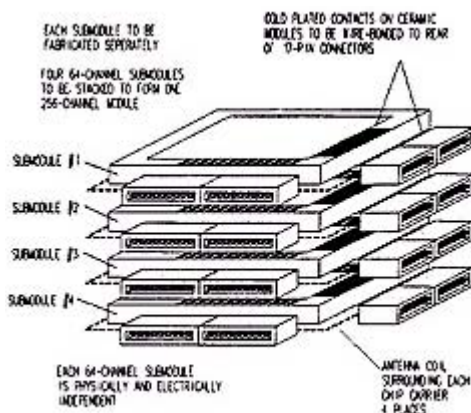
Summary Article 12: There are far more instances than what I report here. Schulman was keeping some of the chips for the smaller devices and was forced to cover this up by saying yield was poor and vendors made mistakes.

### 13. Other Evidence

Much further evidence exists in Joseph Schulman and Gerald Loeb's U.S. Patent Office filings and the Quarterly Progress Reports from contract #N01-NS5-2325. This evidence shows fraudulent reporting, that technology and alternative implantable device invention discoveries occurred during the contract period, and also provides insight regarding motive and other factual evidence. Most evident however is that many of the inventions and methods involve integrating or stacking additional circuitry allowing smaller implant sizes and the ability to eliminate external components and increase the range and options for bi-directional telemetry, and also making it possible to pursue the unethical long term research involving an unwilling subject.

Previously, in article 12, Mr. Wolf from Slicex reports that *"Dr. Schulman wanted to use an entire reticule which meant we placed 5 IC's in a stack, and then 2 IC's rotated 90 degrees at the end"*. The following shows an illustration from Philip Troyk's separate NIH contract for a visual prosthesis as well as text from those QPR's:

*"...the NPP is highly motivated to initiate the development of a fully implantable cortical stimulation system which could be used to provide inputs and computer control for hundreds, to over one thousand, implanted cortical electrodes. This project used the combined capability four organizations, the Illinois Institute of Technology, ....., and the A.E. Mann Foundation. The device specified and designed in this contract has significantly higher packaging and functional density (is smaller) than any other implantable neural prosthesis previously developed."*



Stacked visual prosthesis chips with portion rotated 90 degrees

Individuals at the Mann Foundation have expressed a desire to create a "universal family" of microstimulators that could be used for a visual, auditory or motor prosthesis. Such a universal device would have to be much smaller than the "BION" developed under contract #N01-NS5-2325 and be flexible so as to conform and stretch in response to it's implanted environment. The evidence presented here suggests that Schulman began development of this universal device using NIH grant funding from #N01-NS5-2325 and fundamentals learned are applied today by Mann Corporate ventures Advanced Bionics and Second Sight

*“Work at the Alfred E Mann Foundation is increasingly directed towards a medium to large scale manufacturing of microstimulators. Methods are being developed for reliable and reproducible manufacturing procedures that can utilize batch-processes rather than production steps on an individual basis” (Schulman, QPR #3)*

Later, from R. Greenberg, NC retinal chip investigator and now President of Mann founded “Second Sight”

*“Also, by splitting the chips up into smaller components, and utilizing techniques such as solder bumping to connect the chips with flexible electrode substrates, we shall keep the sizes to a minimum....” (Robert Greenberg Mann Foundation Multiple-Unit Artificial Retina Chipset System To Benefit The Visually Impaired)*

Most recently from USC media:

*“Health & Medicine – USC News;  
...developing an eye chip that would be placed in the retina and restore some vision to people with degenerative diseases. The intraocular retinal prosthesis is created through microelectronics. The eye chips will be manufactured by Second Sight, a company that was founded by Alfred E. Mann, the Chair of the Board of Directors of the Alfred E. Mann Institute for Biomedical Research at USC  
(<http://www.usc.edu/student-affairs/dt/V144/N26/04-doctor.26c.html>)*

Schulman defines smaller devices in patent applications with descriptions far smaller than the “BION”. At 25 mils total thickness, this is approx 1mm, and thus many times more compact than the BION.

*“...A thin hermetically sealed electrical feedthrough suitable for implantation within living tissue permits electrical connection between electronic circuits sealed within an hermetically sealed case and electrical terminals or contacts on the outside of the case ...For example, using currently known processing techniques, the frame and lid can be as thin as approximately 4 mils. The height of the cavity can be as small as about between 5 to 10 mils depending on the connections of the electronics contained within the cavity. The insulating layers, as indicated above, can be on the order of less than 1 mil thick, it is thus feasible using presently available processing techniques to construct an implantable device less than 25 mils thick, yet still providing an hermetically sealed case and hermetic feedthroughs.” (U.S. Patent 5,750,926: “Hermetically sealed electrical feedthrough for use with implantable electronic devices” Inventors: Joseph Schulman Filed: Aug 16, 1995)*

6,067,474: “Implantable device with improved battery recharging and powering configuration”

*“Although the preferred power source for use within the fully implantable systems described herein is a rechargeable battery, it is to be understood that other power sources may also be employed. For example, an ultracapacitor (also known as a supercapacitor) may be used.. Thus, for this type of application, where recharging must occur on a regular basis, and when appropriate discharge circuits are employed to control the rate of discharge or energy withdrawal, the ultracapacitor provides a viable alternative to a rechargeable battery for use within the implantable system.”*

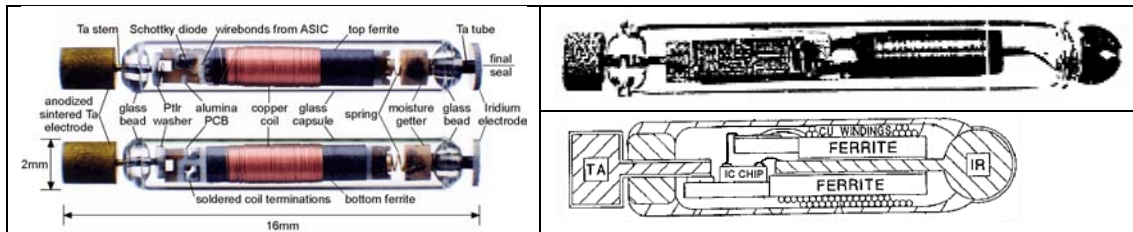
Patent 5,522,865 “Voltage/current control system for a human tissue stimulator”

*Multiple Chip Control: “In the preferred embodiments thus far described, the ICS comprises a single chip. However, multiple chips of the same or similar circuitry may be usefully employed in a human tissue stimulator. In such an embodiment, a circuit at the input of each chip permits the interconnection of several chips into one functional unit by making one of the chips a master device which receives data and clock signals and then distributes such signals to all the slave chips. In this manner a large number of chips may be connected together forming a system with a large number of output channels”*

Chips too big?

*“To provide an easier fit in the small Microstimulator package, the chips must be lapped from a thickness of about 500 microns down to under 200 microns. While this can be done on individual chips, it is much easier (and more economical) to do it on an entire wafer before it is diced into individual chips. As mentioned in the previous report, an additional passivation layer (probably silicon nitride) is needed to protect the chips from damage due to the attached ferrites.” (QPR #6)*

Schulman reports above that post-processing is needed to get the chips thin enough to fit into the microstimulator package, however there appears to be ample room within the glass housing for the tiny 1.5µ and 3.0µ process chips of the BION. A photo of an early and more recent BION is shown below:




---

## Summary

Joseph Schulman and Gerald Loeb have both acted irresponsibly, selfishly and have caused me considerable pain, suffering and other damages during the last 5 years. They have anticipated that no proof would exist of their crimes and that nobody would believe a story of this nature. This will no longer be possible as the evidence and cause for investigation is presented here and now. Loeb's plans for the future at USC-AMI show a new improved version of the "Connectix" Video Phone mentioned as Article 9 in the "Evidence Detail" section above, however in this embodiment the server for this criminal misconduct will remain on campus allowing Schulman and Loeb to record data from and stimulate their unfortunate research subjects who are unwilling recipients of microstimulators, whether at home, on campus, on vacation, or at a National conference or event by remotely accessing the university "Telemedicine" server from their location. I'm certain that Mr. Steven Sample, President of USC did not anticipate this kind of trouble when welcoming the Alfred Mann Institute to USC, and had he known, he may have reconsidered, because this kind of human research misconduct far exceeds any that has existed, even at Johns Hopkins, whose Biomedical Dept has also received Mann funding. Loeb's future plans and ulterior motive for developing tele-medicine technology are a liability for USC and should not be allowed to continue regardless of revenue or finances, because this is a losing proposition no matter how you look at it.

Loeb's newest invention for USC is as follows: United States Patent Application 20010054071  
 Inventors: Loeb, Gerald E.; (Los Angeles, CA)

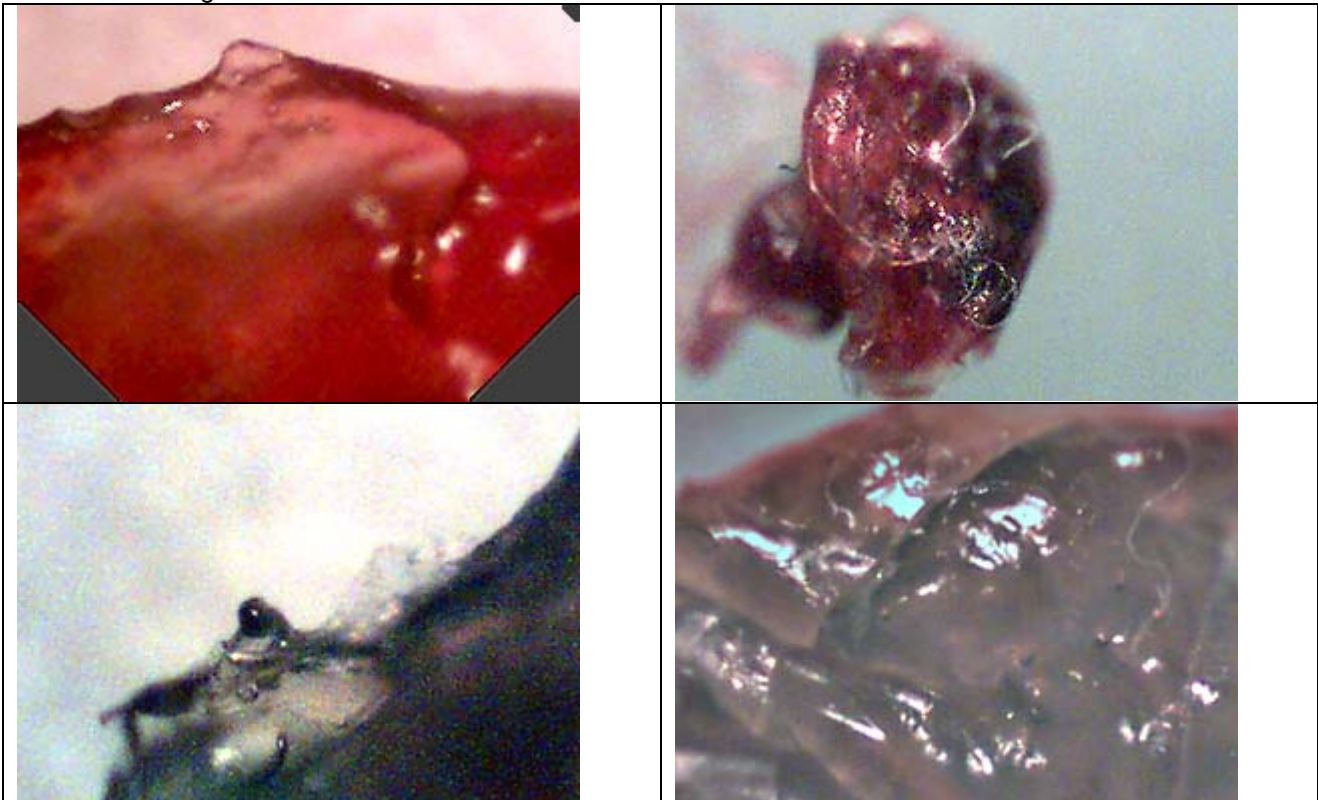
*“...audio/video-call system allows communication using audio, video and/or data. The system includes two communication systems--one for the user and one for the caregiver. Each communication system includes a console coupled to a television. The console has a digital processor, storage, software, and connections to telephone equipment, channel tuner, a video camera, and steering for the video camera. A remote control issues commands to the console to initiate or accept calls to the other communication system. The phone line is used to carry the audio portion of the call while the network is used to carry any video and/or data portion of the call. In some embodiments, a passkey allows the caregiver's communication system to control the user's communication system without any interaction by the user. In other embodiments, more than one video camera is used for a communication system. A microphone can be coupled to the remote control to collect the audio channel for the call...”*

In conclusion, I hope that everyone who reads this will do what he or she can to ensure this behavior is discontinued and that disciplinary measures are pursued. I cannot take the law into my own hands. I need your assistance resolving this. I am willing to cooperate in any fashion that will assist authorities in realizing the truth regarding this situation and this would include having these devices analyzed using techniques or methods that are available which I cannot currently arrange due to financial reasons. Thank you for your assistance regarding this manner.

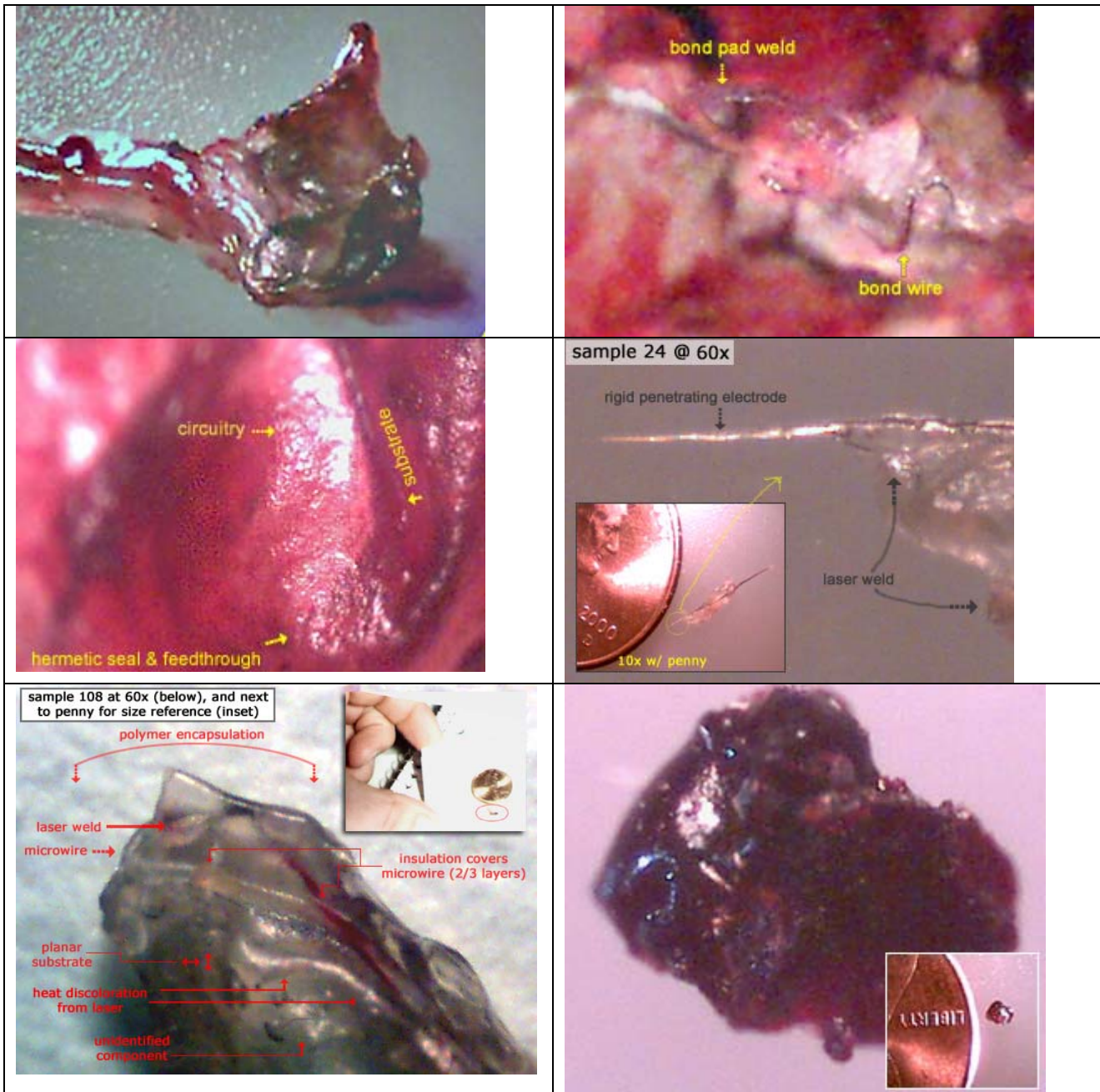
#### Attachments

1. Additional recovered device images
2. Glossary
3. Recovered device materials analysis documentation and data from Photometrics
4. N01-NS5-2325, titled "Implantable Microstimulators and Transducers for FES"
5. N01-NS-7-2365 titled "Multi-Channel Transcutaneous Cortical Stimulation System".
6. E-mail correspondence

More images of devices recovered from my person are presented here. Despite the overwhelming evidence presented above, these devices will make the strongest case against Joseph Schulman and Gerald Loeb and cannot be ignored.







There is a reason why these devices look the way they do. The first reason for their appearance is that they have been implanted for 5 years. The second is that if they were to be recovered, Schulman and Loeb would not want you to recognize them, and the third reason is explained by Jerry Loeb himself:

*“Therefore, modification of the microstimulators' chemical nature and/or superficial physical contours to avoid, prevent and/or discourage an immunological response by the body, would be advantageous.”*

---

## Glossary

**Glossary links on the Internet:**

Everything: <http://whatis.techtarget.com/whome/0,,sid9,00.html>

Semiconductor: <http://semiconductor glossary.com/>

Medical: [http://www.medicinenet.com/script/main/Alphaldx.asp?li=MNI&p=A\\_DICT](http://www.medicinenet.com/script/main/Alphaldx.asp?li=MNI&p=A_DICT)

AEMF: Alfred E. Mann Foundation

ASIC: Application Specific Integrated Circuit

ARPA: a way universities have of saying "DARPA" (since when has "defense" become a dirty?)

BiCMOS: Complimentary Metal Oxide Semiconductor implementing bipolar transistors

BION: "Bionic Neuron"; brand name that Schulman stole from Europe and used for the microstimulator that he stole from Najafi

CMOS: Complimentary Metal Oxide Semiconductor

DAC: Digital Analog Conversion

DARPA: "Defense Advanced Research Projects Agency" "...circumvents normal red-tape by reporting directly to the Secretary of Defense, established to develop imaginative, innovative and often high-risk research ideas that will go well beyond the normal evolutionary developmental approaches" [www.darpa.mil](http://www.darpa.mil)

DSP: Digital Signal Processing

EDS: Energy Dispersive Spectroscopy; analysis for materials characterization

EEPROM: A form of "read only memory"

FES: Functional Electrical Stimulation; the use of electrical stimulation only to replace lost function in the human body [Hambrecht, 1992]. While this may be the most accepted definition, it is not the most accurate. "FES" is more accurately defined as the use of ES (electrical stimulation) to produce a functional movement and/or replace a physiological function [Reswick, 1973]

FET: Field Effect Transmitter

FPGA: Field Programmable Gate Array

IC: Integrated Circuit

Micron: Short for "micron" (short for *micrometer*) is one-millionth of a meter. It can also be expressed as: One thousandth of a millimeter, or One 25-thousandth of an inch

MOSIS: a low-cost prototyping and small-volume production service for CMOS/VLSI circuit development, originally a DARPA-funded program <http://www.mosis.org/>

MCM: Multi-Chip Module; the stacking of chips vertically or even horizontally to use less chip real-estate and realize a smaller package. (Related; Hybrid, stacking, gold bumping; solder bumping; summit; mumps)

NIH: National Institutes of Health

NINDS: National Institute of Neurological Disorders and Stroke; division of the NIH

NPP: Neural Prosthesis Project; a project headed by William Heetderks within the NIH/NINDS  
<http://npp.ninds.nih.gov/>

QPR: Quarterly Progress Report

SCU: System Control Unit

$\mu\text{m}$ : Symbol for "micron" (short for *micrometer*) is one-millionth of a meter. It can also be expressed as: One thousandth of a millimeter, or One 25-thousandth of an inch

Wafer: Foundation that chips are constructed from

**PHOTOMETRICS, INC.**  
THE MATERIALS CHARACTERIZATION LAB  
15801 GRAHAM ST., HUNTINGTON BEACH, CA 92649  
(714) 895-4465 • FAX (714) 893-4682

## Packing List

Invoice No.: 29984  
Date: 3/28/2002  
Terms: Net 30 Days

**ACCOUNTS PAYABLE**  
Dave Larson

*Dave Larson*  
*Dave Larson*

5676 Lindley Ave.  
Encino, CA 91316

5676 Lindley Ave.  
Encino, CA 91316

Purchase Order Number: cash  
Release Number:  
Client Number: DLAC

Shipped Via: Visit/PRR/cc  
Shipped On: 3/28/2002

Qty.	Description	Unit Price	Total Price
1.0	Hour(s) Energy Dispersive X-ray Spectroscopy	165.00	165.00

**PHOTOMETRICS, INC.**  
15801 GRAHAM ST.  
HUNTINGTON BEACH, CA 92649

*paid cash*  
*3/28/02*

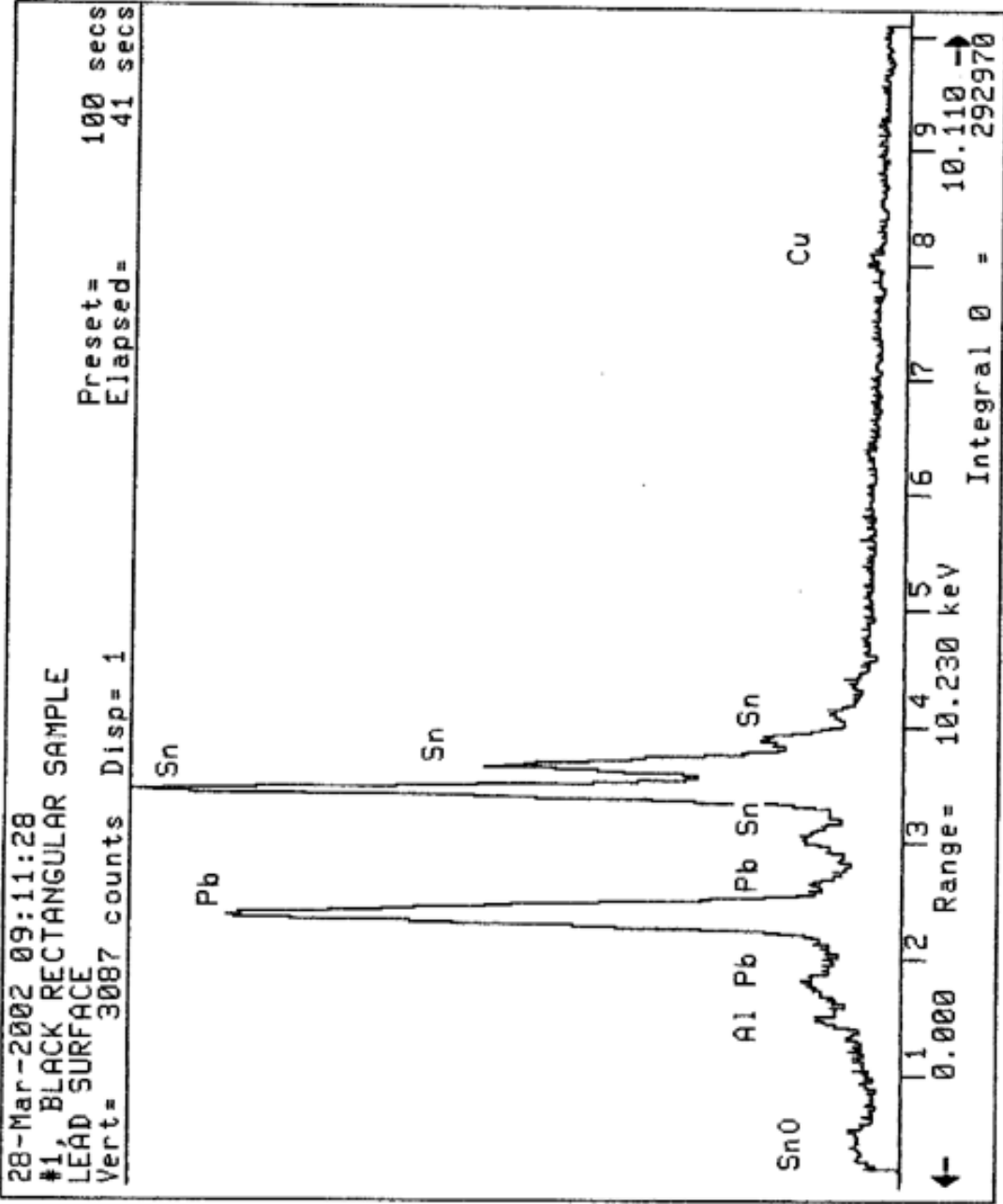
Received by \_\_\_\_\_

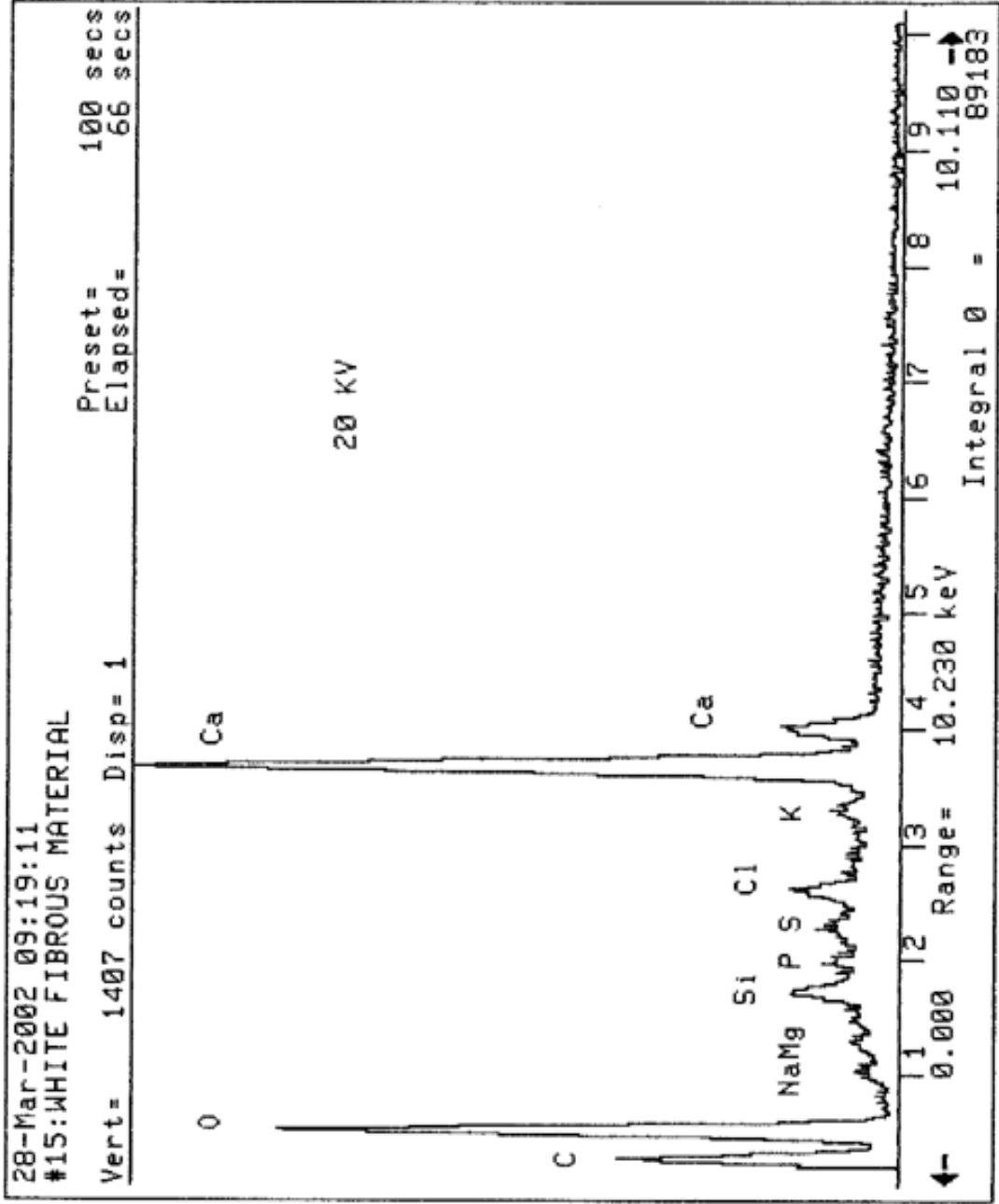
**Total:** 165.00

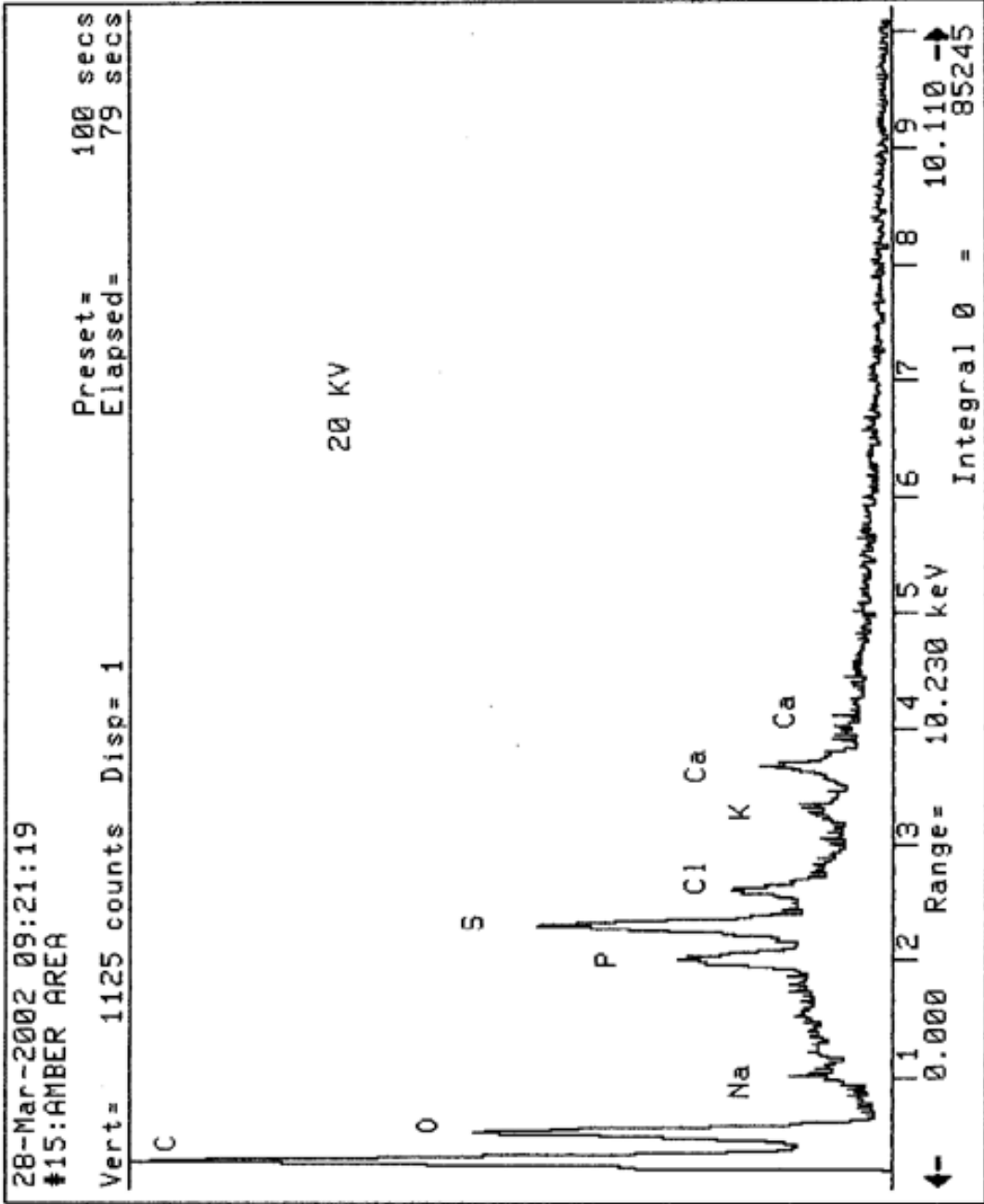
***Please Forward to Accounts Payable***

PROBLEM SOLVING WITH SURFACE SCIENCE FOR RESEARCH, PRODUCTION AND QA/QC











## Microstimulators and Microtransducers for Functional Neuromuscular Stimulation

Principal Investigator	Affiliation	Contract Number	Link
Joe Schulman, Ph.D.	A.E. Mann Foundation	N01-NS5-2325	

---

### ARTICLE C.1 BACKGROUND

The Neural Prosthesis Program (NPP) of The National Institute of Neurological Disorders and Stroke is committed to the development of safe, reliable, and effective systems for functional neuromuscular stimulation (FNS) in spinal cord injured individuals. Multiple implantable microstimulators which selectively stimulate paralyzed muscles in a controlled fashion may permit an individual to use his or her own muscles as the motors to produce limb movement. Multiple implantable microtransducers that sense contact, grasp force, and limb position from either implanted transducers or intact sensory receptors may provide sensory feedback from an otherwise insensate limb. To produce a useful system, these sensory and motor prostheses are being developed together as standard and compatible building blocks of an integrated FNS system.

The NPP is supporting research and development of a system consisting of an extracorporeal coil and a family of implantable microstimulators and microtransducers (Loeb et. al., Injectable microstimulator for functional electrical stimulation., Med. & Biol. Eng. & Comput. 29:NS13-NS19, 1991; Troyk and Schwan, Closed-loop class E transcutaneous power and data link for microimplants., IEEE Trans. on BME 39:589-599, 1992). The system is designed to power and control up to 64 individual transducers and stimulators through a single external coil. During the past five years significant progress has been made in developing addressable, implantable microstimulators and an extracorporeal coil and transmission system. This work has progressed to the point where first generation devices are available for implantation. Work was also begun during the past 2 years on the development of implantable addressable transducers. This contract will continue the engineering development of the microstimulators and transducers. In addition, in-vivo evaluation of these devices will be initiated in an animal model.

This RFP represents a competitive renewal of an ongoing contract. Copies of progress reports from current contracts related to this work and a bibliography of Neural Prosthesis Program publications are available from the Contracts Management Branch, Room 901 Federal Bldg., 7550 Wisconsin Ave., Bethesda, MD 20892.

---

### ARTICLE C.2. STATEMENT OF WORK

Independently, and not as an agent of the Government, the Contractor shall exert its best efforts to develop and test a system for functional neuromuscular stimulation (FNS) consisting of implantable receiver-stimulators and transducer-telemeters and an easily donned extracorporeal transmitter.

Specifically, the Contractor shall:

A. Design, fabricate, and test implantable receiver-stimulators, implantable transducer-telemeters, and an extracorporeal transmitter for FNS.

1. The implantable receiver-stimulators, consisting of power receiver, control receiver, stimulus and control circuitry, electrodes, and encapsulation shall meet the following specifications:

- a. They shall be made as small as possible while still being compatible with the other requirements described below. The target volume of each stimulator shall be 60 cubic mm or less.
- b. An individual stimulator shall initiate a stimulus pulse only in response to a properly addressed command from the transmitter. There shall be at least 64 different stimulator addresses that can be selected.
- c. The stimulators shall derive their power and commands by an inductive link from a transmitter coil with field specifications as outlined below.
- d. The stimulus current waveform shall have two phases of opposite polarity. The duration of the first phase shall be variable from 0 to 200 microseconds in at least 200 steps on a pulse by pulse basis.
- e. The stimulus current level of the first phase of a stimulus pulse shall be adjustable and shall include levels of 1 and 10 milliamps under control of the transmitter. The current of the second phase shall guarantee long

term charge balance at the electrodes under all pulsing conditions within specifications. Stimulator compliance voltage shall be at least 10 volts.

f. Stimulating electrodes shall be made of corrosion resistant and biocompatible materials (e.g. platinum, platinum-iridium, iridium, or tantalum pentoxide) and shall be stable in physiologic saline at 37 degrees C for at least 1 year at the maximum charge delivery rate.

g. The stimulator shall be encapsulated in a manner that permits stable operation in physiologic saline at 37 degrees C for at least 1 year with materials that are known to be biocompatible.

2. The implantable transducer-telemeters consisting of transduction unit, power receiver, control circuitry, digitizer, telemetry circuitry, and encapsulation shall meet the following specifications:

a. The transducer-telemeters shall not interfere with the operation of the receiver-stimulators and vice-versa.

b. The transducer-telemeter package shall occupy a volume of no more than 125 cubic mm (not including magnets, electrodes, or other passive elements which can be external to the package if they are needed for the design). Biocompatibility and life expectancy shall be as specified for the microstimulator in part 1.

c. The transducer-telemeters shall derive power and commands from the same transmitter coil as the microstimulators and shall transmit information to this coil upon command.

d. The transducer-telemeters shall be addressable with at least 64 unique addresses using an addressing scheme that is compatible with the microstimulator addressing.

e. The telemeter's digitizing and telemetry circuits shall be designed to provide at least 8 bits of digital data per sample. Signal conditioning for sensing biopotentials suitable for closed-loop control and/or sensory feedback shall be incorporated in telemeters with addresses 1 through 16.

f. A joint angle transducer-telemeter shall be designed to provide wrist joint angle information for use as a command signal with a resolution of at least 5 degree as the wrist moves through its physiological range of flexion and extension. The error of angle measurement (including drift, hysteresis, and errors due to wrist movement other than pure flexion and extension) shall not exceed 15 degrees.

3. The extracorporeal transmitter shall meet the following specifications:

a. The transmitter field shall permit specified operation of the stimulators and transducers anywhere within a cylindrical shaped volume of physiologic saline at least 9 cm. in diameter and at least 16.0 cm in length.

b. The transmitter field shall permit specified operation with up to 10 degrees of axial misalignment between the transmitter coil and the receiver coils in the stimulators and transducers.

c. The transmitter, under the control of a computer, shall be capable of powering and controlling at least 64 stimulators and/or transducers in any combination.

d. The transmitter coil shall be designed to be donned and doffed as easily as an item of clothing.

e. The transmitter shall operate reliably in the presence of electromagnetic interference commonly found in a home or work environment.

4. The Contractor shall make its best effort to deliver to the Project Officer one set of 32 receiver-stimulators each with a different address, one set of 4 joint angle transducer-telemeters each with a different address, and one transmitter with specifications as outlined above by the end of the contract period.

B. Select a suitable animal model and evaluate implanted microstimulators in this animal model.

1. Evaluate the tissue reaction to implanted stimulators placed intramuscularly and subdermally for periods of at least 3 months.

a. Evaluate the tissue reaction to active and inactive devices.

b. Evaluate the tissue reaction to any components used in making the device that might be exposed to tissue should the encapsulation fail.

2. Investigate the threshold stability and motor recruitment stability of microstimulators over a period of at least 3 months.

3. Investigate the function of implanted transducer-telemeters over a period of at least 3 months.

C. Cooperate with other investigators in the Neural Prosthesis Program to integrate this work into practical FNS systems.

## Multichannel Transcutaneous Cortical Stimulation System

Principal Investigator	Affiliation	Contract Number	Link
Phil Troyk, Ph.D.	Illinois Inst. of Technology	N01-NS7-2365	

---

### BACKGROUND

The Neural Prosthesis Program (NPP), National Institute of Neurological Disorders and Stroke, National Institutes of Health develops implanted devices that interface directly with the nervous system to replace or supplement function in neurologically disabled individuals.

Significant advances have been made in demonstrating the feasibility of bypassing peripheral sensory organs such as the ear and the eye to restore lost sensory functions. Recently, a blind individual had an array of 38 microelectrodes with percutaneous leads implanted into her visual cortex for a period of three months. During this time she was able to see and describe visual phosphenes produced by electrical stimulation through the microelectrodes. Critical engineering parameters, such as phosphene thresholds as low as 4 microamperes and a two point resolution of 500 microns, were determined which now permit the design of a permanent implant controlled by electromagnetic signals passing across the skin. (Ref. Hambrecht, F.T., 1995, Visual prostheses based on direct interfaces with the visual system. In Brindley, G.S. and Rushton, D.N. eds., Baillière's Clinical Neurology: Neuroprostheses, Baillière Tindall, London, pp 147-165.) Because such a permanent implant would not require any leads passing through the skin, there should be little risk of infection and the implant could be left in place indefinitely.

Specifically, a transcutaneous stimulation system consisting of a computer controlled transmitter and a group of implantable receiver-stimulator modules, each with 256 stimulus channel outputs, is needed. Research and development are required to assure that the implanted portion of this system will be small enough to fit safely and comfortably beneath the scalp and that the stimulus outputs are flexible enough to provide the range of stimulus parameters necessary for producing patterns of phosphenes by intracortical microstimulation. This transcutaneous transmission system will interface not only with discrete wire microelectrodes but also with silicon microstimulating microelectrodes presently being developed by other investigators in the NPP. The extracorporeal portion of the system will include a computer controlled transmitter for sending power and control signals across the skin to the implanted receiver-stimulator modules.

This contract research is a new project not previously supported by the NPP. A bibliography listing publications resulting from NPP studies related to this project is available, free of charge, from the Neural Prosthesis Program, NIH, Federal Building, Room 916, Bethesda, MD 20892-9170. (fax: 301-402-1501, e-mail: fh2@cu.nih.gov)

---

### STATEMENT OF WORK

I. Independently, and not as an agent of the government, the contractor shall exert its best efforts to design and fabricate a transcutaneous transmission system suitable for use in a human visual prosthesis. The system shall consist of: 1.) a group of appropriately packaged, implantable, 256 channel receiver-stimulator modules each with high density connectors suitable for connection to 256 cortical microelectrodes; 2.) an external transmitter interfaced to an external computer which can be used to control the implanted receiver-stimulator modules; 3.) a reverse telemetry system for monitoring key voltages in the receiver-stimulator modules. The contractor will not be required to furnish the microelectrodes nor perform any animal or human testing.

Specifically the Contractor shall:

- A. Design the overall system such that it can be expanded in modules of 256 channels up to 1024 channels.
- B. Design the receiver-stimulator module to meet the following target specifications. (Note: Priorities and

limits will have to be assigned to some of the stimulus parameters under certain operational conditions to avoid conflicts. These rules of operation will be established shortly after the beginning of the contract in a joint meeting between the contractor and the Project Officer.)

1. Capable of passing truly simultaneous biphasic current pulses through at least 16 of any of the 256 microelectrodes that will be connected to its output with the capability of passing interleaved biphasic current pulses through any or all of the non-simultaneously pulsed microelectrodes.
2. Capable of stimulating each channel at repetition rates of 10 to 250 Hz. The repetition period should have a resolution of 250 microseconds.
3. Phase durations of each phase of a biphasic pulse pair controllable over the range of 50 to 750 microseconds (50 microsecond resolution/phase)
4. Output compliance voltages of at least +/- 5 volts.
5. Amplitude of each phase controllable over the range of 0 to +/- 64 microamperes (0.5 microampere resolution) unless limited by the compliance voltage.
6. Have an anodic bias supply adjustable over the range of 0 to 0.75 volts referenced to a standard calomel electrode (SCE). In a functional system, the actual voltage of this bias must be translated to correspond to the use of a large surface area platinum reference electrode that will be used in place of a SCE. Each channel must be connected to the anodic bias supply through large bias resistors (e.g. 10 megohm)
7. Train length on each channel of 1 to 255 biphasic pulse pairs.
8. Train delay times for each channel referenced to a reference timing pulse (e.g, a reference pulse for channel #1) adjustable over the range of 0 to 12700 microseconds (100 microsecond resolution).
9. Maximum total output current of each module at least 1024 microamperes.
10. Have built-in safety features that include the ability to sense the failure of any of the output drivers or other modes of operation that could result in charge imbalance and tissue damage.
  - a. Since it is assumed that space will not permit output coupling capacitors on each channel, a desired feature would be the ability to disconnect a shorted output driver from its power supply.
11. The modules should be capable of independent operation, i.e. should any of the modules fail, such failure should not affect the operation of the remaining modules.
12. Have a reverse telemetry system for monitoring the following voltages within each module on demand
  - a. The voltage waveform developed across any remotely selected microelectrode during stimulation.
  - b. The anodic bias supply.
  - c. The positive and negative power supply voltages
13. Contained in a hermetic package
  - a. Suitable for implantation between the scalp and the skull with dimensions no greater than 5 mm thick, 30 mm wide, and 30 mm long, not including any receiving antennae which can be external to the package.
  - b. With 4 separate 66 contact, high-density, low disconnect force, connectors in the package walls that interface with appropriate connectors terminating the leads from the microelectrodes or on dummy loads during in-vitro testing.

c. With smooth, rounded edges and attachment sites for immobilizing the package to the skull in an appropriate manner.

C. Design the external transmitter portion of the system to supply power and full control of up to four receiver-stimulator modules.

1. The transcutaneous transmission signal should be capable of reliably operating through human scalp tissue with thicknesses ranging from 2 mm to 10 mm.

2. The transmitter shall be under the control of a computer whose output can be altered by keyboard entries, by software or by a remote interface that will eventually be controlled by an image sensing device such as a television camera. (The contractor does not need to supply the image sensor, nor any interface electronics between the image sensor and the computer, only the remote interface that will allow full control of the receiver-stimulator modules.)

3. Although the external portions of the systems to be supplied under this contract do not need to be "patient portable", the design should not ignore the fact that future generations of the system will be portable.

D. Fabricate and test, in-vitro, a complete system with 1024 channels.

1. Document the ability of the system to deliver the specified design stimuli at the specified rates without significant cross talk between channels.

2. Test functional receiver-stimulator packages with integral connectors for hermeticity and proper electrical operation in an accelerated aging environment of heated saline solution for a period of at least 6 months.

E. Before the end of the first year of the contract, furnish the Project Officer with 5 empty but hermetically sealed receiver-module packages with at least one 66 contact connector for in-vivo testing by other investigators in the Neural Prosthesis Program (NPP).

F. Before the end of the second year of the contract, furnish the Project Officer with at least 5 fully functional, 256 channel systems, complete with mating connectors so that other investigators in the NPP can attach microelectrode leads and test the complete system in animals.

G. Before the end of the third year of the contract, furnish the Project Officer with at least 2 fully functional, 1024 channel systems, complete with mating connectors for attachment to microelectrode leads for human testing by other NPP investigators.

H. Should failures occur during the above mentioned in-vitro or in-vivo testing, the remaining required deliverables will be reduced, upon mutual agreement between the contractor, the Project Officer and the Contracting Officer, to allow resources to be applied to the redesign and fabrication of relevant parts of the system.

I. The contractor shall coordinate his work, through the Project Officer, with other investigators in the NPP.

---

**Dave Larson**

---

**From:** "Philip R. Troyk, Ph.D."  
**To:** "Dave Larson" <lars121@pacbell.net>  
**Sent:** Sunday, September 16, 2001 5:49 PM  
**Subject:** Re: NPP Quarterly Report #

This meant that the electronic transmitter circuitry would be small enough to physically mount on the extra-corporal transmitter coil, not that both coils would be implanted. This was a goal of one of the other team members. I was never convinced that there was any functional advantage to this.

P. Troyk

At 05:03 PM 9/16/01 -0700, you wrote:

Dr. Troyk,

I am hoping you can help me with a question regarding some of the research performed by yourself and other colleagues as part of NIH contract N01-NS5-2325. The text below is taken from a quarterly report and mentions an interest in reducing coil driver circuitry so that it could potentially be located on the transmission coil itself. This in essence would mean both coils would be implantable correct? I did not see any mention of this interest in subsequent quarterly reports. Do you know if any efforts were made to achieve this? Thank you in advance for your assistance.

"We are working on a re-design of the coil drivers in which the driver circuitry is sufficiently small that it can be located outside the bedside controller and close to, or even on the transmission coil" (Microstimulators And Microtransducers For Functional Neuromuscular Stimulation. Quarterly Progress Report #6 - June 10, 1996 - September 9, 1996).

D. Larson  
Los Angeles, CA.

**Dave Larson**

---

**From:** "Philip R. Troyk, Ph.D." <troyk@iit.edu>  
**To:** "Robert Arzbaecher"  
**Cc:** "Dave Larson" <lars121@pacbell.net>  
**Sent:** Monday, September 10, 2001 11:59 AM  
**Subject:** Re: Fwd: Email to Dr. Troyk

As far as I know little to no imaging work has been done in the living subject at resolutions that would be high enough to be informative.

Phil

At 06:53 PM 9/9/01 -0700, Robert Arzbaecher wrote:

Date: Wed, 05 Sep 2001 17:57:11 -0700  
From: Dave Larson <lars121@pacbell.net>  
Subject: Email to Dr. Troyk  
To: arzbaecher@iit.edu  
Organization: Home  
X-Mailer: Microsoft Outlook Express 5.50.4807.1700

Dr. Arzbaecher,

My attempt to send this email to Dr. Troyk was unsuccessful. Would you be so kind as to forward this to him? Also, I copied you on the original as I am sure you may have some insight as well. Thanks again.

- DL

Dr. Troyk,

I have followed the development of the NINDS NPP for the last couple of years and admire your more than significant contributions to this project. I am quite familiar with the published data, however there is one aspect of the research that I have not found in the papers and I am hoping you can help.

Papers outlining NPP background and statement of work often refer to various electrode designs as well as leadless implanted receiver-stimulator modules for chronic implantation offering between 256 and 1024 individual recording/stim sites. Testing of these devices is also mentioned both in primates and a small number of human based experiments.

I am curious if any research effort has been made to examine the devices and surrounding tissue using radiology and diagnostic imaging in the living subject. There is only mention of histological inspection for evidence of substrate breakage, insulation damage etc. I would think that the ability to check for tissue or electrode damage as well as migration could be accomplished in-vivo using high field MRI or radiography and would prove especially useful during the longer term, chronic studies. Can you provide any insight regarding this?

Regards,

DL

9/16/01

-----Original Message-----

From: Dave Larson [mailto:lars121@pacbell.net]

Sent: Friday, January 25, 2002 10:42 PM

To: hajimiri@caltech.edu

Subject: CMOS ASIC Question.....

Ali,

Hello and happy new year. You probably don't remember me as I have only met you once on campus. I occasionally depend on the help of Steve Potter and others in Biomedical Imaging Center for help with Physiology studies etc., however I find myself in need of help regarding Integrated Circuits and Sensors.

1. If developing ASIC for implantable microstimulator for functional nerve stimulation (FES), mention is made in various papers of both CMOS and Bi-CMOS process. Can you provide a general or "approximate" idea as to what kind of yields from wafer are typically achieved in this process? Assume 3.0 $\mu$  or 1.2 $\mu$  process through MOSIS run?

The yield is usually very high (better than 99%) on those processes.

2. When testing chips and assemblies, what factors are used in determining the amount of difference in carrier levels? I have seen reference made to the lower level at 80% or 90% of the upper level. I am trying to determine what factors would require a greater depth of modulation to be desired.

Consider the following: If one were to multiplex a single electrode driver to multiple electrode sites and assuming a nominal stimulation rate of approximately 30 Hz, 10 to 100 electrodes could share a common driver provided that temporal and phase modulation of individual electrodes were not required for control. In such a scheme, creative circuit techniques would be needed to insure electrode charge recovery. Could this configuration require a greater depth of modulation?

I quite frankly don't follow this question entirely. If the question is how many electrodes can you control using one wireless driver, the answer is quite a few, if you use the right coding schemes. In general this depends on a large number of factors, such as bandwidth, the sophistication of the electrode function and the frequency of stimuli.

In an ASIC design that functioned appropriately at 5 volt compliance, and then encountered latch-up problems when redesigned for 20 volt compliance, how difficult do you think this problem would be to correct? If the process plays a significant role, consider the problem for both 3.0 and 1.2 $\mu$  CMOS through MOSIS run.

Latch up is primarily determined by the substrate doping level, and is usually avoided by following the design rules, including, but not limited to, close proximity of substrate contacts with the active devices. In general it is not advisable (at all) to use a process qualified for 3 or 5 volts with higher voltages. Even if the circuits work, you will have reliability and longevity problems.

Many thanks for your assistance and best wishes.

Dave Larson



Mr. Larsen: In going through my efiles as part of my year-end cleanup, I did not have a clear record that the following response was actually sent to you with regard to your September inquiry. Hence, I am sending it to you now. My apologies if it duplicates an earlier transmittal.

Donna J. Dean, Ph.D.  
Acting Director  
National Institute of Biomedical Imaging and Bioengineering  
Building 31, 1B37, MSC 2077  
National Institutes of Health  
Bethesda, MD 20892-2077  
Phone 301-451-6768  
Fax 301-480-4515  
deand@nibib.nih.gov  
<http://www.nibib.nih.gov>

Dear Mr. Larson,

I have asked some people at NIH for an informed opinion on your query and am pleased to transmit the following response.

There are several parts to this inquiry, but basically the concern is the possibility that a clinical researcher could implant a clinically undetectable (with standard imaging methods) probe without approval to do so. A second part is if NIH is considering this possibility.

With regard to unapproved implantation, there is always the chance that a researcher could perform such an implant in conjunction with other clinical research activities without approval or without including this in the reviewed research protocols. Patient rights and informed consent are major concerns for clinical research, and the NIH requires stringent reviews of clinical procedures and on-site monitoring of clinical research for their funded grants. However, a knowledgeable researcher absolutely intent on implanting a sub-millimeter size probe may be able to find a way to do it during experimental activities.

Information obtained in such a manner could not likely be published in a reputable journal, since the methodology would have to be part of the article, which makes it less tempting if the unethical clinician is seeking acclaim. Also, the fact that such work could not then be replicated by others would heighten the likelihood of scrutiny and ultimately, exposure as fraud.

Is possible unethical research of concern to the NIH? Certainly. There is a Bioethics Office that explicitly deals with such issues, and the Office of Extramural Research coordinates clinical research requirements for NIH grantees for all the NIH research institutes and centers. On-site reviews of NIH-funded clinical research are conducted by NIH staff to evaluate compliance with patient rights issues and adherence to planned experimental protocols. The issue of unauthorized implants is one of many causes for concern in the general realm of unethical clinical research, and the NIH endeavors to deal with such issues before and during clinical investigations.

By way of information, all micro devices that are capable of sending signals into or out of the body are readily detectable using relatively standard x-ray imaging methods. You are correct that we use extensive histological analysis to evaluate the tissue response to implanted micro devices. This is not because we cannot detect the devices using x-rays but because there is a need to do safety studies that involve detailed light and electron microscopic analysis of the tissue response at a cellular and subcellular level.

The question of informed consent applies to all research involving human volunteers and is clearly one of increasing importance in light of recent shortcomings. Informed consent as outlined in the Belmont Report and subsequent guidelines as well as trust between the physician and human volunteers is essential for ethical research and should be a high priority at NIBIB as well as at the other institutes.

-----Original Message-----

**From:** Dave Larson [mailto:lars121@pacbell.net]

**Sent:** Tuesday, September 18, 2001 4:10 PM

**To:** Dean, Donna (OD)

**Cc:** Leonard, Laurie (NINDS); Wheelles, Timothy (OD)

**Subject:** NIBIB Information

I have a question and I am hoping you can provide some insight. Current research in the NIH/NINDS Neural Prosthesis Project (NPP) has led to the development of implantable sensors and probes (Professors Troyk, Najafi, Banks etc., PI on NIH NPP contracts) that are just below the size of feature detection in typical clinical radiology (MRI/CT etc). Continuing research on animal models and a select few human cases show that these probes can not be imaged and the only means of examination is a histological method after the research subject dies with no in-vivo examination due to the sub-millimeter size. The MRI capability of a clinical MRI system (1.5T) is in no way capable of imaging an object below 500x500 microns, however there is development of new high field MRI as high as 7.0T and research at Caltech has used fields as high as 11.7T in rodents. My question Donna pertains to the potential that exists for an unethical researcher to implant probes in an unwilling human in order to gain long-term research data that is otherwise unattainable in animal models. The work of Professors Loeb and Troyk also involve passive, radio frequency techniques that make remote sensing and stimulation of the probes attainable. If this were to happen, there would be no way to detect the probes in a clinical situation and begin any treatment for the potential victim/subject. A horrible thought, yes, but it appears to be a potential cause for concern at this moment. Do you think the establishment of the NIBIB will address this, or has anyone in the NIH addressed this issue at all? Many thanks for your help.

David A. Larson  
Los Angeles, CA

Dr. Kerns received a Ph.D. in Electrical Engineering from the California Institute of Technology and has been kind enough to take an interest in my situation and I am grateful because his knowledge of implantable devices and experience with integrated circuits and sensors has been very beneficial. His prior professional experience includes: Senior Analog Designer, Analog Devices, Inc. (A leading manufacturer of high performance analog and digital IC's) Staff Scientist, Tanner Research (IC design, simulation and layout software development), and Dr. Kerns has also worked for the Jet Propulsion Laboratory and Fermilab.

Date: Tue, 19 Feb 2002 12:16:27 -0500 (EST)

From: Kernsd@██████████

Subject: Re: Hello?

Dave,

Your "wirebond" pic does look very much like a single wirebond from a substrate to the top of a die, but I would have expected at least two bonds to a device capable of wireless communication and local stimulation. My opinion is certainly not the last word.

It looks like you're making good progress in pushing forward the analysis of the object.

- Doug K

"flip-chip" is an assembly process for connecting

an IC chip to other electronic devices. The process essentially consists of soldering the chip face-down (hence the name "flip-chip") onto the other stuff. Flip-chip assembly is full of technical hassles, but it's one of the most compact assembly techniques yet invented. The truly most-compact I've ever seen is "chip stacking." That has hassles that make flip-chip seem like a walk in the park, but there are still people who care so much for compact stuff that they'll pay the outrageous prices.

I hope that's helpful.

- Doug K

----- Original Message -----

> From: <Kernsdou[REDACTED]>  
> Sent: Sunday, January 20, 2002 1:47 PM  
> Subject: Re: semiconductor technical stuff....

Chip stacking is the assembly technique of gluing several silicon chips together, typically adding metal interconnections between the layers of the stack by lapping the edges and doing the usual evap/resist/litho/etch process that's used to make the chips in the first place. The hassles one encounters are usually of two types:

- (a) one of the chips in the stack is flawed, so it doesn't work correctly, and it's difficult to discover the flaws prior to assembling the stack. The end result is that the stack containing the flawed chip has to be thrown away, even though most of it is composed of good pieces. Sometimes a flaw can be caused by the stacking process, so even if you start with chips that are all known to be good, the stack may still be flawed after assembly.
- (b) The interconnection between chips in a stack may have flaws, so there's a missing connection or a shorted connection. Again, the stack may have to be thrown away, even if all the chips were good.

> I believe that the implanted devices have on-chip  
>> memory allowing "event-related" or closed-loop stimulation. This would allow the  
>> use of naturally occurring events to trigger a stimulus response  
>> from the implant. I have noticed that when I'm on vacation or in the  
>> Faraday cage that the stimulus is consistent in its properties and is  
>> event related which is in contrast to the stimulus that I am typically forced to  
>> endure which is more spontaneous and random in its occurrence as well as its amplitude and duration. This leads me to believe that even if I am able to disrupt the signal, I will still be subject to  
>> closed-loop, event related stimulus. I believe this is made possible by an electrode made from tantalum which is able to act as a capacitor which stores a  
>> charge derived from an electrolytic/voltammetry condition resulting from  
>> interaction with the body fluid, or perhaps the implantable devices  
>> can continue to store a charge from a carried signal even if the data  
>> comm is non-functional. I'm trying to assess what wireless communication method  
>> would be feasible for command of the implants and what possible  
>> counter-measures may exist. I'll assume  
>> that the individual responsible for this is licensed by the FCC and has his own  
>> little piece of the spectrum unless a programmable micro-controller would allow  
>> variety of frequencies to be used. I'm wondering if a frequency in the  
>> vicinity of 220.000 MHz as the carrier signal and amateur radio relay used to  
>> establish a network is being used. If this were the case, what equipment would I  
>> need to purchase that would be capable of jamming this signal and what kind of trouble can I create by attempting to disrupt the signal? Can a specific  
>> frequency be targeted in a manner that would not cause problems elsewhere, or is  
>> the idea to simply stomp on everything in the vicinity with noise and power?  
>> Thanks for your insight Doug.  
>>  
>> Sincerely,  
>>  
>> Dave Larson

Hi David,  
EDS would NOT be able to identify an epoxy polymer.  
FTIR would be able to identify it as an epoxy, but probably not a specific type. However, if it is not epoxy and is some other polymer FTIR would be able to determine that.  
With EDS, epoxies, urethanes, acrylics.... all look like they are carbon and oxygen.  
If you are interested in the FTIR let me know, I will have Greg Ansten call you.  
If you would like to schedule an appointment for EDS let me know

Thanks

Paul Reidel  
PhotoMetrics  
(714) 895-4465

<lars121@pacbell.net>; Mon, 10 Dec 2001 14:41:21 -0500 (EST)  
Date: Mon, 10 Dec 2001 14:42:45 -0500  
From: Stuart Cogan <scogan@eiclabs.com>  
Subject: Re: Implantable Microstimulators  
Dave,

We have only a little experience evaluating tissue.

If the tissue samples are in an appropriate form we can look at them with the electron microscope and probably identify the chemical nature of the substances that may be foreign bodies. This we have done with specially prepared histology slides with some success.

If you want to discuss this further, please feel free to call me.

Regards

Stuart Cogan  
EIC Laboratories, Inc.  
1 781 769-9450

<lars121@pacbell.net>; Fri, 01 Feb 2002 17:52:42 -0500  
Date: Fri, 01 Feb 2002 17:56:00 -0500  
From: wise@umich.edu  
Subject: Re: Supply Voltage

The level shifter problem was just a design error. The output bus has a fair amount of capacitance and the driver was too small to drive it quickly. The best solution was to resize the driver, which we have done. But with the existing probes, another solution that helped a bit was to increase the voltage. This isn't desirable long term because it puts more stress across the encapsulating dielectrics. We would like the probes to operate as low in voltage as possible to minimize our packaging problems; however, ultimately, we are limited by the water window and the spreading resistance drops in the solution. They make it hard to operate below 5V for small sites. So the voltage is really a trade off between device site size and the needed stimulating current levels. Hope this helps a little.  
... Ken D. Wise, Professor.

>Professor Wise:

>

>Hello, and greetings from California. I have recently had the  
>opportunity to read published papers regarding micromachined stim  
>electrode arrays and one your papers mentions level shifters that  
>operate slower than which is desired. You acknowledge that an  
>adjusted (higher I assume) voltage is a solution, but only for  
>present day use. Why would adjusted voltage not be a solution for a  
>"standardized" device? Thank you for your insight and any help you  
>may be able to provide.

>

>Regards,

>

>Dave L

>Northridge CA

> There is a new 8.0T system at OSU which can image particle sizes of  
> 200umx200umx2000um, so new advances in imaging are becoming attainable

Petra Schmalbrock Ph.D.  
Associate Professor  
The Ohio State University, Dept. Radiology  
170 Means Hall, 1654 Upham Dr, Columbus OH 43210  
Phone 614-293-4139  
FAX 614-293-8129