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CERTIFICATE OF COMPLIANCE (SAR EVALUATION)

SAFE ENVIRONMENT ENGINEERING 25061 West Avenue Stanford, #30 Valencia, CA 91355

Attn: David M. Lamensdorf, President

Dates of Tests: November 1-2, 2001 Test Report S/N: SAR.211010614.PZE Test Site: PCTEST Lab, Columbia, MD USA

FCC ID PZELLP2E-20

APPLICANT SAFE ENVIRONMENT ENGINEERING

EUT Type: Personal Safety Monitor (PSM)

Tx/Rx Frequency Range: 403 ~ 512 MHz

Max. RF Output Power: 2.0 W

Max. SAR Measurement: 3.53mW/g AMPS Body SAR; 4.23mW/g AMPS Mouth SAR

Trade Name/Model(s): Life line LLP2E-20

FCC Classification: Licensed Non-Broadcast Transmitter Held to Face (TBF)
FCC Rule Part(s): §2.1093; FCC/OET Bulletin Supplement C [July 2001]

Application Type: Certification

This wireless portable device has been shown to be capable of continued compliance for localized specific absorption rate (SAR) for uncontrolled environment/general population exposure limits specified in ANSI/IEEE Std. C95.1-1992 and had been tested in accordance with the measurement procedures specified in ANSI/IEEE Std. C95.3-1992 and IEEE Std. 1528 (Draft Aug.2000).

I attest to the accuracy of data. All measurements reported herein were performed by me or were made under my supervision and are correct to the best of my knowledge and belief. I assume full responsibility for the completeness of these measurements and vouch for the qualifications of all persons taking them.

NVLAP accreditation does not constitute any product endorsement by NVLAP or any agency of the United States Government.

PCTEST certifies that no party to this application has been denied the FCC benefits pursuant to Section 5301 of the Anti-Drug Abuse Act of 1988, 21 U.S.C. 862.

Randy Ortanez President







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SAR MEASUREMENT REPORT

1.1 Scope

Environmental evaluation measurements of specific absorption rate¹ (SAR) distributions in simulated human head and body tissues exposed to radiofrequency (RF) radiation from wireless portable devices for compliance with the rules and regulations of the U.S. Federal Communications Commission (FCC).²

Company Name: SAFE ENVIRONMENT ENGINEERING

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Held to Face (TBF)

Modulation(s):

§2.1093, OET Bulletin 65, FCC Rule Part(s): Supplement C (July 2001)

Max. RF Output Power: 2.0W

3.53mW/g AMPS Body SAR 4.23mW/g AMPS Mouth SAR Max SAR Measurement:

Application Type: Certification

Dates of Tests: November 1-2, 2001 PCTEST Engineering Lab. Columbia, MD, U.S.A. Place of Tests:

Report Serial No.: SAR.211010614.PZE



Fig. 1 SAR Test Setup



Specific Absorption Rate (SAR) is a measure of the rate of energy absorption due to exposure to an RF transmitting source (wireless portable device).

IEEE/ANSI Std. C95.1-1992 limits are used to determine compliance with FCC ET Docket 93-62.

2.1 INTRODUCTION

The FCC has adopted the guidelines for evaluating the environmental effects of radiofrequency radiation in ET Docket 93-62 on Aug. 6, 1996 to protect the public and workers from the potential hazards of RF emissions due to FCC-regulated portable devices.[1]

The safety limits used for the environmental evaluation measurements are based on the criteria published by the American National Standards Institute (ANSI) for localized specific absorption rate (SAR) in IEEE/ANSI C95.1-1992 Standard for Safety Levels with Respect to Human Exposure to Radio Frequency Electromagnetic Fields, 3 kHz to 300 GHz. (c) 1992 by the Institute of Electrical and Electronics Engineers, Inc., New York, New York 10017.[2] The measurement procedure described in IEEE/ANSI C95.3-1992 Recommended Practice for the Measurement of Potentially Hazardous Electromagnetic Fields - RF and Microwave[3] is used for guidance in measuring SAR due to the RF radiation exposure from the Equipment Under Test (EUT). These criteria for SAR evaluation are similar to those recommended by the National Council on Radiation Protection and Measurements (NCRP) in Biological Effects and Exposure Criteria for Radiofrequency Electromagnetic Fields, "NCRP Report No. 86 (c) NCRP, 1986, Bethesda, MD 20814.[5] SAR is a measure of the rate of energy absorption due to exposure to an RF transmitting source. SAR values have been related to threshold levels for potential biological hazards.

2.2 SAR Definition

Specific Absorption Rate (SAR) is defined as the time derivative (rate) of the incremental energy (dU) absorbed by (dissipated in) an incremental mass (dm) contained in a volume element (dV) of a given density (ρ). It is also defined as the rate of RF energy absorption per unit mass at a point in an absorbing body (see Fig. 2).

$$S A R = \frac{d}{d t} \left(\frac{d U}{d m} \right) = \frac{d}{d t} \left(\frac{d U}{\rho d v} \right)$$

Figure 2. SAR Mathematical Equation

SAR is expressed in units of Watts per Kilogram (W/kg).

SAR = $\sigma E^2/\rho$ where: σ = conductivity of the tissue-simulant material (S/m) ρ = mass density of the tissue-simulant material (kg/m³) E = Total RMS electric field strength (V/m)

NOTE: The primary factors that control rate of energy absorption were found to be the wavelength of the incident field in relations to the dimensions and geometry of the irradiated organism, the orientation of the organism in relation to the polarity of field vectors, the presence of reflecting surfaces, and whether conductive contact is made by the organism with a ground plane.[5]

3.1 SAR MEASUREMENT SET-UP

These measurements are performed using the DASY3 automated dosimetric assessment system. It is made by Schmid & Partner Engineering AG (SPEAG) in Zurich, Switzerland. It consists of high precision robotics system (Staubli), robot controller, Pentium III computer, near-field probe, probe alignment sensor, and the generic twin phantom containing the brain equivalent material. The robot is a six-axis industrial robot performing precise movements to position the probe to the location (points) of maximum electromagnetic field (EMF) (see Fig. 2).

A cell controller system contains the power supply, robot controller, teach pendant (Joystick), and remote control, is used to drive the robot motors. The PC consists of the Micron Pentium III 500 MHz computer with Windows NT system and SAR Measurement Software DASY3, A/D interface card, monitor, mouse, and keyboard. The Staubli Robot is connected to the cell controller to allow software manipulation of the robot. A data acquisition electronic (DAE) circuit performs the signal amplification, signal multiplexing, AD-conversion, offset measurements, mechanical surface detection, collision detection, etc. is connected to the Electro-optical coupler (EOC). The EOC performs the conversion from the optical into digital electric signal of the DAE and transfers data to the PC plug-in card.

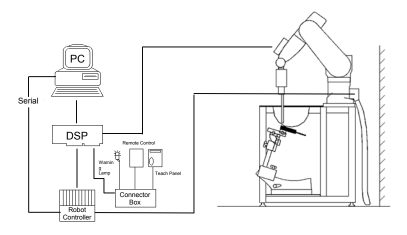


Figure 3. PCTEST SAR Lab II Test Measurement Set-up

The DAE3 consists of a highly sensitive electrometer-grade preamplifier with auto-zeroing, a channel and gain-switching multiplexer, a fast 16 bit AD-converter and a command decoder and control logic unit. Transmission to the PC-card is accomplished through an optical downlink for data and status information and an optical uplink for commands and clock lines. The mechanical probe mounting device includes two different sensor systems for frontal and sidewise probe contacts. They are also used for mechanical surface detection and probe collision detection. The robot uses its own controller with a built in VME-bus computer. The system is described in detail in [6].

4.1 DASY3 E-FIELD PROBE SYSTEM

4.2 ET3DV6 Probe Specification

Construction Symmetrical design with triangular core

Built-in optical fiber for surface detection System

Built-in shielding against static charges

Calibration In air from 10 MHz to 2.5 GHz

In brain and muscle simulating tissue at Frequencies of 450 MHz, 900 MHz and

1.8 GHz (accuracy ± 8%)

Frequency 10 MHz to > 6 GHz; Linearity: \pm 0.2 dB

(30 MHz to 3 GHz)

Directivity \pm 0.2 dB in brain tissue (rotation around probe axis)

 \pm 0.4 dB in brain tissue (rotation normal probe axis)

Dynamic 5 : W/g to > 100 mW/g; Range Linearity: \pm 0.2 dB

Surface \pm 0.2 mm repeatability in air and clear liquids

Detection over diffuse reflecting surfaces.

Dimensions Overall length: 330 mm

Tip length: 16 mm Body diameter: 12 mm Tip diameter: 6.8 mm

Distance from probe tip to dipole centers: 2.7 mm

Application General dosimetry up to 3 GHz

Compliance tests of mobile phones

Fast automatic scanning in arbitrary phantoms



Figure 4. Photograph of the Probe and the Phantom



Fig. 5. ET3DV6 E-field Probe

The SAR measurements were conducted with the dosimetric probe ET3DV6, designed in the classical triangular configuration [6] and optimized for dosimetric evaluation. The probe is constructed using the thick film technique; with printed resistive lines on ceramic substrates. The probe is equipped with an optical multifiber line ending at the front of the probe tip. It is connected to the EOC box on the robot arm and provides an automatic detection of the phantom surface. Half of the fibers are connected to a pulsed infrared transmitter, the other half to a synchronized receiver. As the probe approaches the surface, the reflection from the surface produces a coupling from the transmitting to the receiving fibers. This reflection increases first during the approach, reaches a maximum and then decreases. If the probe is flatly touching the surface, the coupling is zero. The distance of the coupling maximum to the surface is independent of the surface reflectivity and largely independent of the surface to probe angle. The DASY3 software reads the reflection during a software approach and looks for the maximum using a 2nd order fitting. The approach is stopped at reaching the maximum.

5.1 E-FIELD PROBE CALIBRATION PROCESS

5.2 E-Probe Calibration

Each probe is calibrated according to a dosimetric assessment procedure described in [7] with an accuracy better than +/- 10%. The spherical isotropy was evaluated with the procedure described in [8] and found to be better than +/-0.25dB. The sensitivity parameters (NormX, NormY, NormZ), the diode compression parameter (DCP) and the conversion factor (ConvF) of the probe is tested.

The free space E-field from amplified probe outputs is determined in a test chamber. This is performed in a TEM cell for frequencies bellow 1 GHz, and in a waveguide above 1 GHz for free space. For the free space calibration, the probe is placed in the volumetric center of the cavity and at the proper orientation with the field. The probe is then rotated 360 degrees.

E-field temperature correlation calibration is performed in a flat phantom filled with the appropriate simulated brain tissue. The measured free space E-field in the medium correlates to temperature rise in a dielectric medium. For temperature correlation calibration a RF transparent thermistor-based temperature probe is used in conjunction with the E-field probe.

$$SAR = C \frac{\Delta T}{\Delta t}$$

where:

 $\Delta t = \exp \text{osure time (30 seconds)},$

C = heat capacity of tissue (brain or muscle),

 ΔT = temperature increase due to RF exposure.

SAR is proportional to $\Delta T/\Delta t$, the initial rate of tissue heating, before thermal diffusion takes place. Now it's possible to quantify the electric field in the simulated tissue by equating the thermally derived SAR to the E- field;

$$SAR = \frac{\left|E\right|^2 \cdot \sigma}{\rho}$$

where:

 σ = simulated tissue conductivity,

 ρ = Tissue density (1.25 g/cm³ for brain tissue)

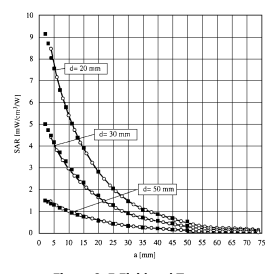


Figure 6. E-Field and Temperature measurements at 900MHz [6]

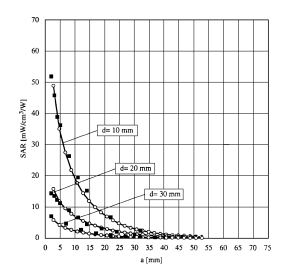


Figure 7. E-Field and temperature measurements at 1.8GHz [6]

5.3 Data Extrapolation

The DASY3 software automatically executes the following procedures to calculate the field units from the microvolt readings at the probe connector. The first step of the evaluation is a linearization of the filtered input signal to account for the compression characteristics of the detector diode. The compensation depends on the input signal, the diode type and the DC-transmission factor from the diode to the evaluation electronics. If the exciting field is pulsed, the crest factor of the signal must be known to correctly compensate for peak power. The formula for each channel can be given as [9]:

with
$$V_i$$
 = compensated signal of channel i (i=x,y,z)
 $V_i = U_i + U_i^2 \cdot \frac{cf}{dcp_i}$ U_i = input signal of channel i (i=x,y,z)
 U_i = input signal of channel i (i=x,y,z)
 U_i = crest factor of exciting field (DASY parameter)
 U_i = diode compression point (DASY parameter)

From the compensated input signals the primary field data for each channel can be evaluated:

E-field probes: with
$$V_i = \text{compensated signal of channel i (i = x,y,z)}$$

 $Norm_i = \text{sensor sensitivity of channel i (i = x,y,z)}$
 $\mu V/(V/m)^2 \text{ for E-field probes}$
 $E_i = \sqrt{\frac{V_i}{Norm_i \cdot ConvF}}$
ConvF = sensitivity of enhancement in solution
 $E_i = \text{electric field strength of channel i in V/m}$

The RSS value of the field components gives the total field strength (Hermetian magnitude):

$$E_{tot} = \sqrt{E_x^2 + E_y^2 + E_z^2}$$

The primary field data are used to calculate the derived field units.

$$SAR = E_{tot}^2 \cdot \frac{\sigma}{\rho \cdot 1000}$$
 with $SAR = local specific absorption rate in W/g = total field strength in V/m $\sigma = conductivity in [mho/m] \text{ or [Siemens/m]}$ $\rho = equivalent tissue density in g/cm^3$$

The power flow density is calculated assuming the excitation field to be a free space field.

$$P_{pwe} = \frac{E_{tot}^2}{3770}$$
 with $P_{pwe} = \text{equivalent power density of a plane wave in W/cm}^2$ = total electric field strength in V/m

6.1 PHANTOM & EQUIVALENT TISSUES

6.2 Generic Twin Phantom

The Generic Twin Phantom is constructed of a fiberglass shell integrated in a wooden table. The shape of the shell is based on data from an anatomical study designed to determine the maximum exposure in at least 90% of all users [10][11]. It enables the dosimetric evaluation of left and right hand phone usage as well as body mounted usage at the flat phantom region. A cover prevents the evaporation of the liquid. Reference markings on the Phantom allow the complete setup of all predefined phantom positions and measurement grids by manually teaching three points in the robot. See Figure 8.



Filling Volume Volume Approx. 20 liters

Dimensions 810 x 1000 x 500 mm (H x L x W)



Fig. 8 Generic Twin Phantom

6.3 Brain & Muscle Simulating Mixture Characterization

The brain and muscle mixtures consist of a viscous gel using hydroxethylcellullose (HEC) gelling agent and saline solution (see Table 1). Preservation with a bacteriacide is added and visual inspection is made to make sure air bubbles are not trapped during the mixing process. The mixture is calibrated to obtain proper dielectric constant (permittivity) and conductivity of the desired tissue. The mixture characterizations used for the brain and muscle tissue simulating liquids are according to the data by C. Gabriel and G. Hartsgrove [12].

MIXTURE %	FREQUENCY (Brain)	FREQUENCY (Muscle)	FREQUENCY (Brain)	FREQUENCY (Muscle)	
	800 - 850 MHz	800 - 850 MHz	1850 -1910 MHz	1850 -1910 MHz	
WATER	41.45	52.40	47.00	40.40	
SUGAR	56.00	45.00	51.90	58.00	
SALT	1.450	1.400	0.000	0.500	
BACTERIACIDE	0.100	0.200	0.100	0.100	
HEC	1.000	1.000	1.000	1.000	

Table 1. Composition of the Brain & Muscle Tissue Equivalent Matter

6.4 Device Holder for Transmitters

In combination with the Generic Twin Phantom V3.0, the Mounting Device (POM) enables the rotation of the mounted transmitter in spherical coordinates whereby the rotation points is the ear opening. The devices can be easily, accurately, and repeatably positioned according to the FCC and CENELEC specifications. The device holder can be locked at different phantom locations (left head, right head, flat phantom).

Fig. 9. Device Holder

^{*} Note: A simulating human hand is not used due to the complex anatomical and geometrical structure of the hand that may produced infinite number of configurations [11]. To produce the worst-case condition (the hand absorbs antenna output power), the hand is omitted during the tests.

7.1 SYSTEM SPECIFICATIONS

7.2 Robotic System Specifications

Specifications

POSITIONER: Stäubli Unimation Corp. Robot Model: RX60L

Repeatability: 0.02 mm

No. of axis: 6

Data Acquisition Electronic (DAE) System

Cell Controller

Processor: Pentium III

Clock Speed: 450 MHz

Operating System: Windows NT

Data Card: DASY3 PC-Board

Data Converter

Features: Signal Amplifier, multiplexer, A/D converter, and control logic

Software: DASY3 software

Connecting Lines: Optical downlink for data and status info.

Optical uplink for commands and clock

PC Interface Card

Function: 24 bit (64 MHz) DSP for real time processing

Link to DAE3

16 bit A/D converter for surface detection system

serial link to robot

direct emergency stop output for robot

E-Field Probes

Model: ET3DV6 S/N: 1560

Construction: Triangular core fiber optic detection system

Frequency: 10 MHz to 6 GHz

Linearity: \pm 0.2 dB (30 MHz to 3 GHz)

Phantom

Phantom:Generic TwinShell Material:FiberglassThickness: $2.0 \pm 0.1 \text{ mm}$

Measured Tissue403-470 MHzParameters(Muscle)Dielectric Constant: ε56.7Conductivity: σ0.94

8.1 MEASUREMENT PROCESS

8.2 System Verification

Prior to assessment, the system is verified to the $\pm 5\%$ of the specifications at 835MHz and 1900MHz by using the system validation kit. (Graphic Plots Attached)

Validation Kit	Muscle	Targeted SAR _{1g} (mW/g)	Measured SAR _{1g} (mW/g)
D835V2, S/N: 406	iviuscie	2.11	2.16

8.3 Dosimetric Assessment Setup

The evaluation was performed with the following procedure:

- 1. The SAR value at a fixed location above the ear point was measured and was used as a reference value for assessing the power drop.
- 2. The SAR distribution at the exposed side of the head was measured at a distance of 3.9mm from the inner surface of the shell. The area covered the entire dimension of the head and the horizontal grid spacing was 20mm x 20mm. Based on this data, the area of the maximum absorption was determined by spline interpolation.
- 3. Around this point, a volume of 32mm x 32mm x 34mm was assessed by measuring 5 x 5 x 7 points. On this basis of this data set, the spatial peak SAR value was evaluated with the following procedure:
 - a. The data at the surface were extrapolated, since the center of the dipoles is 2.7mm away from the tip of the probe and the distance between the surface and the lowest measuring point is 1.2mm. The extrapolation was based on a least square algorithm [14]. A polynomial of the fourth order was calculated through the points in z-axes. This polynomial was then used to evaluate the points between the surface and the probe tip.
 - b. The maximum interpolated value was searched with a straight-forward algorithm. Around this maximum the SAR values averaged over the spatial volumes (1g or 10g) were computed using the 3D-Spline interpolation algorithm. The 3D-spline is composed of three one-dimensional splines with the "Not a knot" condition (in x, y, and z directions) [14][15]. The volume was integrated with the trapezoidal algorithm. One thousand points (10 x 10 x 10) were interpolated to calculate the average.
 - c. All neighboring volumes were evaluated until no neighboring volume with a higher average value was found.
- 4. The SAR value, at the same location as procedure #1, was re-measured. If the value changed by more than 5%, the evaluation is repeated.

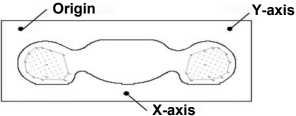


Fig 10. SAR Measurement Points in Area Scan

9.1 TEST POSITIONS OF THE PHONE

9.2 Handset Test Positions

The test device was placed in a normal operating position with the "test device reference point" located along the "vertical centerline" on the front of the device aligned to the "ear reference point" (See Fig. 11). The "test device reference point" was than located at the same level as the center of the earpiece region. The test device was positioned so that the "vertical centerline" was bisecting the front surface of the handset at it's top and bottom edges, positioning the "ear reference point" on the outer surface of the head phantom on each ear spacer.

9.3 EAR Reference Point

The test device was initially positioned with the earpiece region pressed against the ear spacer of a head phantom. The device was positioned parallel to the cheek for maximum RF energy coupling. The "test device reference point" was aligned to the "ear reference point" on the head phantom and the "vertical centerline" was aligned to the "phantom reference plane". (see Figure 12). While maintaining these three alignments, the body of the test deice was gradually adjusted to both of the following positions for SAR evaluation [5]:

EARPIECE Y-axis

Figure 11.
Ear Reference Point

A. Cheek / Touch Position

For Cheek/Touch Position, the test device was brought toward the mouth of the head phantom by pivoting against the "ear reference point" of the head phantom. The test position was established:

- --When any point on the display, keypad or mouthpiece portions of the test device was in contact with the head phantom, or
- --When any portion of a foldout, sliding or similar keypad cover opened to its intended self-adjusting normal use potion was in contact with the cheek or mouth of the head phantom.

When the test device lost contact with the phantom at the pivoting point, rotation continued until the device touched the cheek of the head phantom or broke it's last contact from the ear spacer.

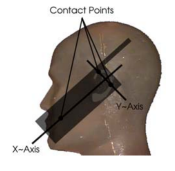


Figure 12.
Phantom Reference Plane

B. Ear / Tilt Position

With the test device aligned in the "Cheek/Touch Position":

- --If the earpiece of the test device was not in full contact with the head phantom's ear spacer in the Cheek/Touch Position and the peak SAR location for the "Cheek/Touch" position was located at the ear spacer region or corresponded to the earpiece region of the test device, the device was returned to the "initial ear position" by rotating it away from the mouth until the earpiece was in full contact with the ear spacer, otherwise
- --The test device was moved (translated) away from the cheek perpendicular to the line that passes through both the "ear reference points". While in this position, the handset was tilted away from the mouth with respect to the "test device reference point" by 15°. After the tilt, the test device was then moved (translated) back toward the head perpendicular to the line spacer. If the antenna touched the head phantom first, then the positioning process was repeated with a tilt angle less than 15° so that the device and its antenna touched the phantom simultaneously.

10.1 BODY-WORN TEST SETUP

10.2 Ear-Microphone Jack

Portable transmitting devices that have an ear-microphone jack must be evaluated for RF exposure in a body-worn configuration. The testing is performed with the use of the flat phantom filled with muscle equivalent tissue. The test device is positioned with the keypad facing away from the flat phantom, and the ear-microphone (headset) wire attached to the phone jack, simulating the device placed in a shirt pocket or attached to a body holster. The SAR tests are then performed in both the antenna in and antenna out positions using the low, middle, and high channels to investigate the worst case SAR value (see Figure 13). Please note that body-worn configurations which have not been SAR tested may result in operating conditions that could exceed FCC RF exposure limits, therefore, users are cautioned to use tested and/or approved accessories.



Figure 13. Ear-Microphone Jack

A. Shirt Pocket Configuration

The shirt pocket configuration is used for devices designed to be body-worn, and small enough to be placed inside a shirt pocket. To simulate the worst-case configuration, the EUT is placed in a torso position on the flat phantom with the keypad facing away from the phantom, and the headset wire connected to the phone to simulate hands-free operation in a shirt-pocket configuration (see Figure 14).



Figure 14.
Shirt Pocket Configuration

B. Body Holster Configuration

The body holster configuration is used for body-worn devices that have a body holster / beltclip accessory. Typically, a holster (carrying case) or beltclip is provided or available as an accessory item for supporting headset and body-worn operations. SAR may vary depending on the body separation distance provided by the type of holster/beltclip and batteries supplied for a phone. In some cases, the antenna may become closer to the user's body than next to the head. The design of the holster/beltclip permits the phone to be positioned only with the keypad facing away from the phantom. Proper use of the holster/beltclip restricts the antenna to a specified distance away from the surface of the body. For this test the EUT is placed into the holster/beltclip and the holster/beltclip is positioned against the torso of the flat phantom in a normal operating position. The headphone wire is then connected to the phone to simulate hands-free operation in a body holster/beltclip configuration (see Figure 15).



Figure 15.
Body Holster Configuration

C. Other Configurations

If other operating configurations are possible (i.e.: pants pocket, car adapter kit, etc), it will be indicated to users in the instruction manual regarding untested conditions and the possibility of exceeding FCC RF exposure limits for such use or the use of third-party accessories. If there is a high potential for exceeding limits in certain unintended configurations, a warning statement will be included in the manual, warning the user to avoid such operating conditions.

11.1 ANSI/IEEE C95.1 - 1992 RF EXPOSURE LIMITS

HUMAN EXPOSURE	UNCONTROLLED ENVIRONMENT General Population (W/kg) or (mW/g)	CONTROLLED ENVIRONMENT Occupational (W/kg) or (mW/g)
SPATIAL PEAK SAR * (Brain)	1.60	8.00
SPATIAL AVERAGE SAR ** (Whole Body)	0.08	0.40
SPATIAL PEAK SAR *** (Hands / Feet / Ankle / Wrist)	4.00	20.00

Table 2. Safety Limits for Partial Body Exposure [2]

NOTES:

- * The Spatial Peak value of the SAR averaged over any 1 gram of tissue (defined as a tissue volume in the shape of a cube) and over the appropriate averaging time.
- ** The Spatial Average value of the SAR averaged over the whole-body.
- *** The Spatial Peak value of the SAR averaged over any 10 grams of tissue (defined as a tissue volume in the shape of a cube) and over the appropriate averaging time.

Uncontrolled Environments are defined as locations where there is the exposure of individuals who have no knowledge or control of their exposure.

Controlled Environments are defined as locations where there is exposure that may be incurred by persons who are aware of the potential for exposure, (i.e. as a result of employment or occupation).

12.1 MEASUREMENT UNCERTAINTIES

Measurement uncertainties in SAR measurements are difficult to quantify due to several variables including biological, physiological, and environmental. However, we estimate the measurement uncertainties in SAR to be less than 15-25 % [17].

According to ANSI/IEEE C95.3, the overall uncertainties are difficult to assess and will vary with the type of meter and usage situation. However, accuracy's of \pm 1 to 3 dB can be expected in practice, with greater uncertainties in near-field situations and at higher frequencies (shorter wavelengths), or areas where large reflecting objects are present. Under optimum measurement conditions, SAR measurement uncertainties of at least \pm 2dB can be expected.[3]

According to CENELEC [18], typical worst-case uncertainty of field measurements is \pm 5 dB. For well-defined modulation characteristics the uncertainty can be reduced to \pm 3 dB.

Uncertainty Description	Error	Distribution	Weight	Std. Deviation	Offset			
Probe Uncertainty								
Axial isotropy	±0.2 dB	U-Shaped	0.5	±2.4 %				
Spherical isotropy	±0.4 dB	U-Shaped	0.5	±4.8 %				
Isotropy from gradient	±0.5 dB	U-Shaped	0	±				
Spatial resolution	±0.5 %	Normal	1	±0.5 %				
Linearity error	±0.2 dB	Rectangle	1	±2.7 %				
Calibration error	±3.3 %	Normal	1	±3.3 %				
SAR Evaluation Uncertainty								
Data acquisition error	±1 %	Rectangle	1	±0.6 %				
ELF and RF disturbances	±0.25 %	Normal	1	±0.25 %				
Conductivity assessment	±10 %	Rectangle	1	±5.8 %				
Spatial Peak SAR Evaluation Uncertainty								
Extrapolated boundary effect	±3 %	Normal	1	±3 %	±5 %			
Probe positioning error	±0.1 mm	Normal	1	±1 %				
Integrated and cube orientation	±3 %	Normal	1	±3 %				
Cube Shape inaccuracies	±2 %	Rectangle	1	±1.2 %				
Device positioning	±6 %	Normal	1	±6 %				
Combined Uncertainties				±11.7 %	±5 %			

Table 3. Breakdown of Errors [19]

13.1 SAR TEST DATA SUMMARY

		Relative HUMIDITY (%)	22.4 60.7 99.3
Mixture Type:	450MHz Muscle		
Dielectric Constant:	56.7	Measured Depth of Simulating Tissue:	15.5 cm
Conductivity:	0.94	Measured Tissue TEMPERATURE (°C)	22.2

13.2 Measurement Results (AMPS Body SAR w/ Holster)

FREQUI	ENCY	Modulation	POWER* (dBm) Battery		Separation	Antenna	SAR	
MHz	Ch.				(dBm)		Battery	Distance (cm)**
403.100	Low	AMPS	2.0	2.0	Standard	2.0 [with Holster]	Fixed	2.13
438.050	Mid	AMPS	2.0	2.0	Standard	2.0 [with Holster]	Fixed	2.43
469.650	High	AMPS	2.0	2.0	Standard	2.0 [with Holster]	Fixed	3.53
ANSI / IEEE C95.1 1992 - SAFETY LIMIT Spatial Peak Controlled Exposure/Occupational Exposure Limits					8.0 W/	Body kg (mW/g) d over 1 gram)	

NOTES:

- All modes of operation were investigated and the worst-case are reported. 1.
- 2. Battery condition is fully charged for all readings. Standard battery is the ONLY battery option.
- * Power Measured X Conducted EIRP □ ERP 3.
- **SPEAG** 4. SAR Measurement System X IDX
- Phantom configuration SAR Configuration 5. Head \times Body Hand
- ** Test Configuration X Holster Without Holster 6.

Spacing = 2.0cm from flat phantom to the back top & base of radio, and 4.0cm from flat phantom to antenna. The worst-case spacing of 2.0cm is noted and specified in the User's Manual on the RF Exposure Warning page.

Randy Ortanez President



Figure 16. Body SAR Test Setup

13.1 SAR TEST DATA SUMMARY (Continued)

		Relative HUMIDITY (%)	22.4 60.7 99.3
Mixture Type:	450MHz Muscle		
Dielectric Constant:	56.7	Measured Depth of Simulating Tissue:	15.5 cm
Conductivity:	0.94	Measured Tissue TEMPERATURE (°C)	22.2

13.3 Measurement Results (AMPS Mouth SAR)

FREQU	ENCY	Modulation	POWER*		Separation	Antenna	SAR	
MHz	Ch.		(dBm)		Battery	Distance (cm)**	Position	(W/kg)
403.100	Low	AMPS	2.0	2.0	Standard	2.0	Fixed	2.35
438.050	Mid	AMPS	2.0	2.0	Standard	2.0	Fixed	3.08
469.650	High	AMPS	2.0	2.0	Standard	2.0	Fixed	4.23
ANSI / IEEE C95.1 1992 - SAFETY LIMIT Spatial Peak Controlled Exposure/Occupational Exposure Limits				8.0 W/	uth/Face /kg (mW/g) ed over 1 gram			

NOTES:

- 1. All modes of operation were investigated and the worst-case are reported.
- Battery condition is fully charged for all readings. Standard battery is the ONLY battery option. 2.
- * Power Measured X Conducted **EIRP** 3.
- X 4. SAR Measurement System **SPEAG** IDX
- Phantom configuration X Flat Phantom
- SAR Configuration 5. Head \times Body \times Mouth ** Test Configuration Holster \times Without Holster 6.

Spacing = 2.0cm from flat phantom to the front top & base of radio, and 2.2cm from flat phantom to antenna. The worst-case spacing of 2.0cm is noted and specified in the User's Manual on the RF Exposure Warning page.





Figure 17. Body SAR Test Setup

14.1 SAR TEST EQUIPMENT

14.2 Type / Model	Calib. Date	S/N
Stäubli Robot RX60L	Feb. 01	599131-01
Stäubli Robot Controller	Feb. 01	PCT592
Stäubli Teach Pendant (Joystick)	Feb. 01	3323-00161
Micron Computer 450 MHz Pentium III, Windows NT	Feb. 01	PCT577
SPEAG EDC3	Feb. 01	321
SPEAG DAE3	Feb. 01	330
SPEAG E-Field Probe ET3DV6	Feb. 01	1560
SPEAG Dummy Probe	Feb. 01	PCT583
SPEAG Generic Twin Phantom	Feb. 01	PCT587
SPEAG Light Alignment Sensor	Feb. 01	205
SPEAG Validation Dipole D835V2	Feb. 01	PCT613
SPEAG Validation Dipole D1900V2	Feb. 01	PCT593
Muscle Equivalent Matter (450MHz)	Nov. 01	PCTMEM32
Robot Table		PCT586
Phone Holder		PCT588
A/B Power Indicator		PCT589
Remote Power Switch		PCT590
Phantom Cover		PCT591
HP Spectrum Analyzer	Dec. 00	PCT200
IFI TEM Cell Model: CC110EXX	Jan. 01	A427-0697
(DC - 2000 MHz)		
Microwave Amp. Model: 5S1G4 (800MHz - 4.2GHz, 5 Watts)	Jan. 01	22332

NOTE:

The E-field probe was calibrated by SPEAG, by temperature measurement procedure. Dipole Validation measurement is performed by PCTEST Lab. before each test. The brain simulating material is calibrated by PCTEST using the dielectric probe system and network analyzer to determine the conductivity and permittivity (dielectric constant) of the brain-equivalent material.

The following list of equipment was used to calibrate the brain equivalent material:

Power Meter Gigatronics 8651A Signal Generator HP-8648D (9kHz ~ 4GHz)

Power Amp Amplifier Research 5S1G4 (5 Watts, 800MHz ~ 4.2GHz)

Network Analyzer HP-8753E (30kHz ~ 3GHz)

Dielectric Probe Kit HP85070B

15.1 CONCLUSION

The SAR measurement indicates that the EUT continues to comply with the RF radiation exposure limits of the FCC. These measurements are taken to simulate the RF effects exposure under worst-case conditions. Precise laboratory measures were taken to assure repeatability of the tests.

Please note that the absorption and distribution of electromagnetic energy in the body are very complex phenomena that depend on the mass, shape, and size of the body, the orientation of the body with respect to the field vectors, and the electrical properties of both the body and the environment. Other variables that may play a substantial role in possible biological effects are those that characterize the environment (e.g. ambient temperature, air velocity, relative humidity, and body insulation) and those that characterize the individual (e.g. age, gender, activity level, debilitation, or disease). Because innumerable factors may interact to determine the specific biological outcome of an exposure to electromagnetic fields, any protection guide shall consider maximal amplification of biological effects as a result of field-body interactions, environmental conditions, and physiological variables.[3]

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