

PCTEST ENGINEERING LABORATORY, INC. 6660 - B Dobbin Road • Columbia, MD 21045 • USA Telephone 410.290.6652 / Fax 410.290.6654 http://www.pctestlab.com (email: randy@pctestlab.com) **CERTIFICATE OF COMPLIANCE (SAR EVALUATION)**



APPLICANT NAME & ADDRESS:

HYUNDAI Curitel Inc. San 136-1, Ami-Ri, Bubal-Eub Ichon-Si, Kyungki-Do, KOREA 467-701 Attn: Mr. Ki-Soo Kim, QM Dept. System Quality Assurance Office

DATE & LOCATION OF TESTING:

Dates of Tests: August 20, 2002 Test Report S/N: SAR.220811425.PP4 Test Site: PCTEST Lab, Columbia MD

FCC ID:	PP4HWP-2100
APPLICANT:	SAMSUNG ELECTRONICS CO., LTD.

EUT Type:	WLL Telephone (CDMA)
Tx Frequency:	824.70 – 848.31 MHz (CDMA)
Rx Frequency:	869.70 – 893.31 MHz (CDMA)
Max. RF Output Power:	0.338 W ERP CDMA (25.283 dBm) / 25.0 dBm Conducted
Max. SAR Measurement:	1.30W/kg CDMA Body SAR
Trade Name/Model(s):	HWP-2100
FCC Classification:	Licensed Non-Broadcast Transmitter Held To Ear (TNE)
FCC Rule Part(s):	§2.1093; FCC/OET Bulletin 65 Supplement C [July 2001]
Application Type:	Certification
Test Device Serial No.:	Identical prototype

This wireless portable device has been shown to be capable of 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 FCC/OET Bulletin 65 Supplement C (2001) and IEEE Std. 1528-200X (Draft 6.4, July 2001).

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 youch for the qualifications of all persons taking them.

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.

Alfred Cirwithian Vice President Engineering





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1. INTRODUCTION / SAR DEFINITION

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.[6] 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.

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. 1.1).

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

Figure 1.1 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/m ³)
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.[6]

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2. SAR MEASUREMENT SETUP

Robotic System

Measurements are performed using the ALIDX-500 automated dosimetric assessment system. The ALIDX-500 is made by IDX Robotics, Inc. (IDX) in the United States and consists of high precision robotics system (CRS), robot controller, Pentium 4 computer, near-field probe, probe alignment sensor, and the Left and Right SAM phantoms containing the head/brain equivalent tissue, and the flat phantoms for body/muscle equivalent. 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. 2.1).

System Hardware

The Robot table consists of the power supply, robot controller, safety computer, teach pendant (Joystick), six-axis robot arm, and the probe. The cell controller consists of DELL Dimension 4300 Pentium-4 1.6 GHz computer with Windows 2000 system and SAR Measurement software, National Instruments analog card, monitor, keyboard, and mouse. The robot controller is connected to the cell controller to communicate between the two computers. The probe data is connected to the cell controller via data acquisition cables.

System Electronics

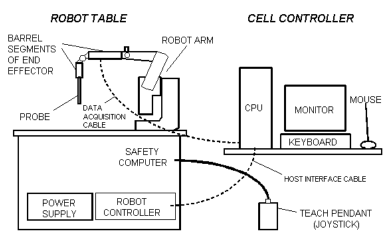


Figure 2.1 SAR Measurement System Setup

When the Robot is in the home position, the Y-axis of the coordinate system parallels the line of intersection between the tabletop and the long axis of the Robot's Large Shoulder. The Teach Pendant may be used to establish the X,Y coordinate directions by depressing the 0-X and 0-Y MOTOR/AXIS switches while in axis mode.

The robot is first taught to position the probe sensor following a specific pattern of points. In the first sweep the sensor enclosure touches the inside of the phantom head. The SAR is measured on a defined grid of points that are concentrated on the surface of the head closest to the antenna of the transmitting device (EUT).

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3. ALIDX-500 E-FIELD PROBE SYSTEM

Probe Measurement System



Fig 3.1 IDX System The near-field probe is an implantable isotropic E-field probe that measures the voltages proportional to the $|E|^2$ (electric) or $|H|^2$ (magnetic) fields. The probe is enclosed in a hollow glass protective cylinder 9-mm. outer diameter, 0.5 mm. thickness and 30 cm. in length. The E-probe contains three electrically small array of orthogonal dipoles strategically placed to provide greater accuracy and to compensate for near-field spatial gradients. The probe contains diodes that are placed over the gap of the dipoles to improve RF detection. The electrical signal detected by each diode is amplified by three DC amplifiers and are contained in a shielded container in the robot end effector so its performance is not affected by the presence of incident electromagnetic fields (see Fig. 3.1).

Probe Specifications

	Probe specifications	
Frequency Range:	10 kHz – 3.0 GHz	
Calibration:	In air from 10 MHz to 3.0 GHz	
	In brain and muscle simulating tis MHz, 1900MHz and 2450MHz	ssue at Frequencies of 835
Sensitivity:	3.5 mV/mW/cm ² (air – typical)	Rev.E-10-1 E&OE.
DC Resistance:	300 kohm	
Isotropic Response:	0.25 dB	
Dynamic Range:	10 mW/kg – 100 W/kg	
Resistance to Pull:	25 N	\checkmark
Probe Length:	290 mm	Figure 3.2
Probe Tip Material:	Glass	Triangular Probe Configuration
Probe Tip Length:	40 mm	
Probe Tip Diameter:	7 ± 0.2 mm	Triangular Probe Isotropic Characteristics
Application:	SAR Dosimetry Testing	(plane perpendicular to probe axis) 달 8.00
	HAC (Hearing Aid Compatibility)	net nn e 4.00
	Compliance tests of mobile phones	110 8.00 6.00 4.00 2.00
		Δ

Proberotation Figure 3.3

0 30 60 90 120 150 180 210 240 270 300 330

Probe Characteristics

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4. Probe Calibration Process

Dosimetric Assessment Procedure

Each E-Probe/Probe amplifier combination has unique calibration parameters. A TEM calibration procedure is conducted to determine the proper amplifier settings to enter in the probe parameters. The amplifier settings are determined for a given frequency by subjecting the Probe to a known E-field density (1mW/cm²) using an RF Signal generator, TEM cell, and RF Power Meter. The SAR measurement software is used for Probe calibration.

Free Space Assessment

The free space E-field from amplified probe outputs is determined in a test chamber. This calibration can be performed in a TEM cell if the frequency is below 1 GHz and in a waveguide or some other methodologies above 1 GHz for free space. For the free space calibration, we place the probe in the volumetric center of the cavity and at the proper orientation with the field. We then rotate the probe 360 degrees until the three channels show the maximum reading. The power density readings equates to 1mW/cm².

Temperature Assessment *

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:

 Δt = exposure 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;

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

where:

 σ = simulated tissue conductivity,

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

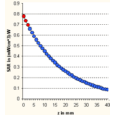


Figure 4.1 E-Field and Temperature measurements at 900MHz

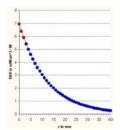


Figure 4.2 E-Field and temperature measurements at 1.9GHz

* NOTE: The temperature calibration was not performed by PCTEST. For information use only.

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5. PHANTOM & EQUIVALENT TISSUES

SAM Phantom



Figure 5.1 SAM Phantoms

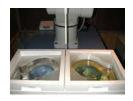


Figure 5.2 Head Simulated Tissue

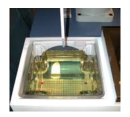


Figure 5.3 Body/Muscle Simulated Tissue

The Left and Right SAM Phantoms are constructed of a vivac composite integrated in a corian stand. 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 [7][8]. 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 Fig. 5.1)

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 6.1). 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 head tissue dielectric parameters recommended by the IEEE SCC-34/SC-2 have been incorporated in the following table. Other head and body tissue parameters that have not been specified in P1528 are derived from the issue dielectric parameters computed from the 4-Cole-Cole equations. The mixture characterizations used for the brain and muscle tissue simulating liquids are according to the data by C. Gabriel and G. Hartsgrove [9].(see Table 5.1)

Ingredients	Frequency (MHz)									
(% by weight)	4:	50	8	35	9	15	19	00	24	50
Tissue Type	Head	Body	Head	Body	Head	Body	Head	Body	Head	Body
Water	38.56	51.16	41.45	52.4	41.05	56.0	54.9	40.4	62.7	73.2
Salt (NaCl)	3.95	1.49	1.45	1.4	1.35	0.76	0.18	0.5	0.5	0.04
Sugar	56.32	46.78	56.0	45.0	56.5	41.76	0.0	58.0	0.0	0.0
HEC	0.98	0.52	1.0	1.0	1.0	1.21	0.0	1.0	0.0	0.0
Bactericide	0.19	0.05	0.1	0.1	0.1	0.27	0.0	0.1	0.0	0.0
Triton X-100	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	36.8	0.0
DGBE	0.0	0.0	0.0	0.0	0.0	0.0	44.92	0.0	0.0	26.7

 Salt: 99'% Pure Sodium Chloride
 Sugar: 98'% Pure Sucrose

 Water: De-ionized, 16 MΩ' resistivity
 HEC: Hydroxyethyl Cellulose

 DGBE: 99'% Di(ethylene glycol) butyl ether, [2-(2-butoxyethoxy)ethanol]
 Triton X-100 (ultra pure): Polyethylene glycol mono [4-(1,1,3,3-tetramethylbutyl)phenyl] ether

Table 5.1 Composition of the Brain & Muscle Tissue Equivalent Matter

Device Holder



Figure 5.4 Device Positioner

In combination with the SAM Phantom, the EUT Holder (see Fig. 6.2) enables the rotation of the mounted transmitter in spherical coordinates whereby the rotation point is the ear opening. Device positioning is accurate and repeatable according to the FCC and CENELEC specifications. The device holder can be locked at different phantom locations (left head, right head, flat phantom).

* Note: A simulating human hand is not used due to the complex anatomical and geometrical structure of the hand that may produce infinite number of configurations [8]. To produce the worst-case condition (the hand absorbs antenna output power), the hand is omitted during the tests.

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6. TEST SYSTEM SPECIFICATIONS

Automated Test System Specifications

Positioner

Robot:	CRS Robotics, Inc. Robot Model: F3
Repeatability:	± 0.05 mm (0.002 in.)
No. Of axes:	6

Data Acquisition Electronic (DAE) System

Pentium 4
1.6 GHz
Windows 2000 [™] Professional
NI DAQ Card (in CPU)
IDX Flexware
Data Acquisition Cable
RS-232 Host Interface Cable
6000 samples/sec



Figure 6.1 ALIDX-500 Test System

E-Field Probes

Model:	E-010	S/N: PCT001 & PCT002
Construction:	Triangu	lar core absolute encoder system
Frequency:	10 MH	z to 3.0 GHz

Phantom

Phantom:	SAM Phantoms (Left & Right)
Shell Material:	Vivac Composite
Thickness:	$2.0\ \pm 0.2\ mm$

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7. DOSIMETRIC ASSESSMENT & PHANTOM SPECS

Measurement Procedure

The measurement procedure consists of the process parameters, probe parameters, EUT product data, and measurement scans (teach points). The measurement procedure is a set of predefined points to be scanned and measured by the probe, DC amplified and processed by the cell controller. The corresponding voltages determined by the electric and magnetic fields are extrapolated to determine peak SAR value.

The SAR Measurement System measures field strength by employing two different types of systematic measurement scans; a coarse scan and a fine scan. Coarse and fine scans measure field strength in a rectangular area within the XY plane (a plane parallel to the top of the Robot Table). The measurement area is divided into a grid of small squares defined by equally spaced grid lines. During an actual measurement process, the probe moves along grid lines systematically recording the field strength at grid line intersections. Typically, after a coarse scan is completed, a fine scan is conducted at the peak field strength value (hot spot) that was measured in the coarse scan. The fine scan has a greater resolution (smaller grid squares) than the coarse scan, and covers only a fraction of the measurement area in the coarse scan.

Specific Anthropomorphic Mannequin (SAM) Specifications

The phantom for handset SAR assessment testing is a low-loss dielectric shell, with shape and dimensions derived from the anthropometric data of the 90th percentile adult male head dimensions as tabulated by the US Army. The SAM Phantom shell is bisected along the mid-sagittal plane into right and left halves (see Fig. 7.1). The perimeter sidewalls of each phantom halves are extended to allow filling with liquid to a depth that is sufficient to minimized reflections from the upper surface. The liquid depth is maintained at a minimum depth of 15cm to minimize reflections from the upper surface. The SAM shell thickness is $2.0 \pm 0.2 \text{ mm}$.



Figure 7.1 Left and Right SAM Phantom shells

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8. **DEFINITION OF REFERENCE POINTS**

EAR Reference Point (ERP)

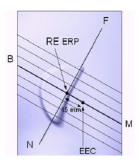


Figure 8.2 Close-up side view of ERPs

Figure 8.1 shows the front, back and side views of the SAM Twin Phantom. The point "M" is the reference point for the center of the mouth, "LE" is the left ear reference point (ERP), and "RE" is the right ERP. The ERPs are 15mm posterior to the entrance to the ear canal (EEC) along the B-M line (Back-Mouth), as shown in Figure 9.2. The plane passing through the two ear canals and M is defined as the Reference Plane. The line N-F (Neck-Front) is perpendicular to the reference plane and passing through the RE (or LE) is called the Reference Pivoting Line (see Figure 8.2). Line B-M is perpendicular to the N-F line. Both N-F and B-M lines are marked on the external phantom shell to facilitate handset positioning [5].



Figure 8.1 Front, back and side view of SAM Twin Phantom

Handset Reference Points

Two imaginary lines on the handset were established: the vertical centerline and the horizontal line. 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. 8.3). The "test device reference point" was than located at the same level as the center of the ear reference point. 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 both the left and right head phantoms on the ear reference point.

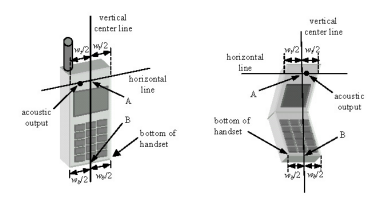


Figure 8.3 Handset Vertical Center & Horizontal Line Reference Points

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9. TEST CONFIGURATION POSITIONS

Positioning for Cheek/Touch

1. The test device was positioned with the handset close to the surface of the phantom such that point A is on the (virtual) extension of the line passing through points RE and LE on the phantom (see Figure 9.1), such that the plane defined by the vertical center line and the horizontal line of the phone is approximately parallel to the sagittal plane of the phantom.



Figure 9.1 Front, Side and Top View of Cheek/Touch Position

- 2. The handset was translated towards the phantom along the line passing through RE & LE until the handset touches the ear.
- 3. While maintaining the handset in this plane, the handset was rotated around the LE-RE line until the vertical centerline was in the plane normal to MB-NF including the line MB (reference plane).
- 4. The phone was hen rotated around the vertical centerline until the phone (horizontal line) was symmetrical was respect to the line NF.
- 5. While maintaining the vertical centerline in the reference plane, keeping point A on the line passing through RE and LE, and maintaining the phone contact with the ear, the handset was rotated about the line NF until any point on the handset made contact with a phantom point below the ear (cheek). See Figure 9.2)

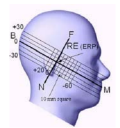


Figure 9.2 Side view w/ relevant markings

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9. TEST CONFIGURATION POSITIONS (Continued)

Positioning for Ear / 15° Tilt

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

- 1. While maintaining the orientation of the phone, the phone was retracted parallel to the reference plane far enough to enable a rotation of the phone by 15 degree.
- 2. The phone was then rotated around the horizontal line by 15 degree.
- 3. While maintaining the orientation of the phone, the phone was moved parallel to the reference plane until any part of the phone touches the head. (In this position, point A was located on the line RE-LE). The tilted position is obtained when the contact is on the pinna. If the contact were at any location other than the pinna, the angle of the phone would then be reduced. The tilted position was obtained when any part of the phone was in contact of the ear as well as a second part of the phone was in contact with the head (see Figure 9.3).



Figure 9.3 Front, Side and Top View of Ear/15° Tilt Position

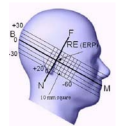


Figure 9.4 Side view w/ relevant markings

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9. TEST CONFIGURATION POSITIONS (Continued)

Body Holster /Belt Clip Configurations

Body-worn operating configurations are tested with the belt-clips and holsters attached to the

device and positioned against a flat phantom in a normal use configuration (see Figure 9.5). A device with a headset output is tested with a headset connected to the device. Body dielectric parameters are used.

Accessories for Body-worn operation configurations are divided into two categories: those that do not contain metallic components and those that do contain metallic components. When multiple accessories that do not contain metallic components are supplied with the device, the device is tested with only the accessory that dictates the closest spacing to the body. Then multiple accessories that contain metallic components are supplied with the device, the device is tested with each accessory that contains a unique metallic component. If multiple accessories share an identical metallic component (i.e. the same metallic belt-clip used with different holsters with no other metallic components) only the accessory that dictates the closest spacing to the body is tested.



Figure 9.5 Body Belt Clip & Holster Configurations

Body-worn accessories may not always be supplied or available as options for some devices intended to be authorized for body-worn use. In this case, a test configuration where a separation distance between the back of the device and the flat phantom is used. All test position spacings are documented.

Transmitters that are designed to operate in front of a person's face, as in push-to-talk configurations, are tested for SAR compliance with the front of the device positioned to face the flat phantom. For devices that are carried next to the body such as a shoulder, waist or chest-worn transmitters, SAR compliance is tested with the accessory(ies), including headsets and microphones, attached to the device and positioned against a flat phantom in a normal use configuration.

In all cases SAR measurements are performed to investigate the worst-case positioning. Worst-case positioning is then documented and used to perform Body SAR testing.

In order for users to be aware of the body-worn operating requirements for meeting RF exposure compliance, operating instructions and cautions statements must be included in the user's manual.

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

Uncontrolled Environment

UNCONTROLLED ENVIRONMENTS are defined as locations where there is the exposure of individuals who have no knowledge or control of their exposure. The general population/uncontrolled exposure limits are applicable to situations in which the general public may be exposed or in which persons who are exposed as a consequence of their employment may not be made fully aware of the potential for exposure or cannot exercise control over their exposure. Members of the general public would come under this category when exposure is not employment-related; for example, in the case of a wireless transmitter that exposes persons in its vicinity.

Controlled Environment

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). In general, occupational/controlled exposure limits are applicable to situations in which persons are exposed as a consequence of their employment, who have been made fully aware of the potential for exposure and can exercise control over their exposure. This exposure category is also applicable when the exposure is of a transient nature due to incidental passage through a location where the exposure levels may be higher than the general population/uncontrolled limits, but the exposed person is fully aware of the potential for exposure and can exercise control over his or her exposure by leaving the area or by some other appropriate means.

	HUMAN EXPOSURE LIMITS		
	UNCONTROLLED ENVIRONMENT	CONTROLLED ENVIRONMENT	
	General Population	General Population	
	(W/kg) or (mW/g)	(W/kg) or (mW/g)	
SPATIAL PEAK SAR ¹	1.60	8.00	
Brain	1.00	8.00	
SPATIAL AVERAGE SAR ²	0.08	0.40	
Whole Body	0.08	0.40	
SPATIAL PEAK SAR ³	4.00	20.00	
Hands, Feet, Ankles, Wrists		20.00	

Table 10.1. Safety Limits for Partial Body Exposure [2]

³ 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.

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¹ 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.

² The Spatial Average value of the SAR averaged over the whole body.



11. MEASUREMENT UNCERTAINTIES

а	b	с	d	e=	f	g	h =	i =	k
				f(d,k)			cxf/e	cxg/e	
Uncertainty		Tol.	Prob.		Ci	Ci	1 - g	10 - g	
Component	Sec.	(± %)	Dist.	Div.	(1 - g)	(10 - g)	u _i	u _i	V _i
·						·	(± %)	(± %)	
Measurement System									
Probe Calibration	E1.1	6.0	Ν	1	1	1	6.0	6.0	∞
Axial Isotropy	E1.2	4.88	R	$\sqrt{3}$	0.5	0.5	1.4	1.4	∞
Hemishperical Isotropy	E1.2	9.6	R	$\sqrt{3}$	0.5	0.5	2.8	2.8	∞
Boundary Effect	E1.3	11.0	R	$\sqrt{3}$	1	1	6.4	6.4	∞
Linearity	E1.4	4.7	R	$\sqrt{3}$	1	1	2.7	2.7	∞
System Detection Limits	E1.5	1.0	R	$\sqrt{3}$	1	1	0.6	0.6	∞
Readout Electronics	E1.6	1.0	R	1	1	1	1.0	1.0	∞
Response Time	E1.7	0.8	R	$\sqrt{3}$	1	1	0.5	0.5	∞
Integration Time	E1.8	1.7	R	$\sqrt{3}$	1	1	1.0	1.0	∞
RF Ambient Conditions	E5.1	1.2	R	$\sqrt{3}$	1	1	0.7	0.7	∞
Probe Positioner Mechanical Tolerance	E5.2	0.4	R	$\sqrt{3}$	1	1	0.2	0.2	∞
Probe Positioning w/ respect to Phantom	E5.3	2.9	R	$\sqrt{3}$	1	1	1.7	1.7	∞
Shell									
Extrapolation, Interpolation & Integration	E4.2	3.9	R	$\sqrt{3}$	1	1	2.3	2.3	∞
Algorithms for Max. SAR Evaluation									
Test Sample Related									
Test Sample Positioning	E3.2.1	10.6	R	$\sqrt{3}$	1	1	6.1	6.1	11
Device Holder Uncertainty	E3.1.1	8.7	R	$\sqrt{3}$	1	1	5.0	5.0	8
Output Power Variation - SAR drift	5.6.2	5.0	R	$\sqrt{3}$	1	1	2.9	2.9	∞
measurement									
Phantom & Tissue Parameters									
Phantom Uncertainty (Shape & Thickness	E2.1	4.0	R	$\sqrt{3}$	1	1	2.3	2.3	∞
tolerances)									
Liquid Conductivity - deviation from	E2.2	5.0	R	$\sqrt{3}$	0.7	0.5	2.0	1.4	∞
target values									
Liquid Conductivity - measurement	E2.2	10.0	R	$\sqrt{3}$	0.7	0.5	4.0	2.9	∞
uncertainty									
Liquid Permittivity - deviation from	E2.2	5.0	R	$\sqrt{3}$	0.6	0.5	1.7	1.4	∞
target values									
Liquid Permittivity - measurement	E2.2	5.0	R	$\sqrt{3}$	0.6	0.5	1.7	1.4	∞
uncertainty									
Combined Standard Uncertainty (k=1)			RSS				14.4	14.0	
Expanded Uncertainty (k=2)							28.8	28.0	
(95% CONFIDENCE LEVEL)									

The above measurement uncertainties are according to IEEE Std. 1528-200x (July, 2001)

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12. SYSTEM VERIFICATION

Tissue Verification

Table 12.1 Simulated Tissue Verification

	MEASURED TISSUE PARAMETERS											
Date(s)	05/06/02	835MHz Brain		835MHz Muscle		1900MHz Brain		1900MHz Muscle				
Liquid Temperature (°C)	22.7	Target	Measured	Target	Measured	Target	Measured	Target	Measured			
Dielectric Constant: ε		41.50	40.56	55.20	54.96	40.00	N/A	53.30	N/A			
Conductivity: σ		0.900	0.880	0.970	1.000	1.400	N/A	1.520	N/A			

Test System Validation

Prior to assessment, the system is verified to the $\pm 10\%$ of the specifications at 835MHz by using the system validation kit(s). (Graphic Plots Attached)

Table 12.2 System Validation

	SYSTEM	DIPOLE VALIDATION TA	ARGET & MEASURED	
System Validation Kit:	835MHz	Targeted SAR _{1g} (mW/g)	Measured SAR _{1g} (mW/g)	Deviation (%)
D-835S, S/N: 103	Brain	2.375	2.24	- 5.5





Figure 12.1 Dipole Validation Test Setup

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13. SAR TEST DATA SUMMARY

See Measurement Result Data Pages

Procedures Used To Establish Test Signal

The handset was placed into simulated call mode (Cellular CDMA mode) using manufacturers test codes. Such test signals offer a consistent means for testing SAR and are recommended for evaluating SAR [4]. When test modes are not available or inappropriate for testing a handset, the actual transmission is activated through a base station simulator or similar equipment. See data pages for actual procedure used in measurement.

Device Test Conditions

The handset is battery operated. Each SAR measurement was taken with a fully charged battery. In order to verify that the device was tested at full power, conducted output power measurements were performed before and after each SAR measurement to confirm the output power. If a conducted power deviation of more than 5% occurred, the test was repeated.

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SAR DATA SUMMARY

Mixture Type: 835MHz Muscle

14.1 MEASUREMENT RESULTS (CDMA Body SAR)											
FREQU	IENCY	Modulation	Begin / End POWER [‡]			Separation	Antenna	SAR			
MHz	Ch.	Modulation	(dł	3m)	Battery	Distance (cm) ¹¹	Position	(W/kg)			
824.70	1013	CDMA	25.0	25.0	Internal	2.5	Fixed	1.30			
835.89	363	CDMA	25.0	25.0	Internal	2.5	Fixed	1.27			
848.31	777	CDMA	25.0	25.0	Internal	2.5	Fixed	1.27			
ANSI / IEEE C95.1 1992 - SAFETY LIMIT Spatial Peak Uncontrolled Exposure/General Population						1.6 W	Muscle /kg (mW/g) ed over 1 gram				

NOTES:

1. The test data reported are the worst-case SAR value with the antenna-head position set in atypical configuration. Test procedures used are according to FCC/OET Bulletin 65, Supp.C [July 2001].

ERP

IDX

Body

Flat Phantom

Without Holster

Base Station Simulator

 \times

X

 \times

- 2. All modes of operation were investigated, and worst-case results are reported.
- 3. Battery is fully charged for all readings.
 - [‡]Power Measured
- SAR Measurement System 4. Phantom Configuration
- 5. SAR Configuration
- 6. Test Signal Call Mode
- 7. ^{‡‡}Test Configuration
- 8. Tissue parameters and temperatures are listed on the SAR plots.
- 9. Liquid tissue depth is 15.1 cm. \pm 0.1

Alfred Cirwithian Vice President Engineering

1	- 4 1	(1) and
	THE R	

□ EIRP

Right Head

Hand

Figure 14.1 Body SAR Test Setup

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- I ⊂ Conducted
- DASY3
- □ Left Head
- □ Head
- Manu. Test Codes
- With Holster



15. SAR TEST EQUIPMENT

Equipment Calibration

Table 15.1 Test Equipment Calibration

EQUIPMENT SPECIFICATIONS

Туре		Calibration Date	Serial Number
CRS Robot F3		February 2002	RAF0134133
CRS C500C Motion Controller		February 2002	RCB0003303
CRS Teach Pendant (Joystick)		February 2002	STP0132231
DELL Computer, Pentium 4 1.6 GHz, W	/indows 2000 [™]	February 2002	
E-Field Probe E-010		February 2002	PCT002
Right Ear SAM Phantom (P-SAM-R)		February 2002	
Left Ear SAM Phantom (P-SAM-L)		February 2002	
IDX Robot End Effector (EE-103-C)		February 2002	07111223
IDX Probe Amplifier		February 2002	07111113
Validation Dipole D-835S		February 2002	PCT640
Brain Equivalent Matter (835MHz)		August 2002	PCTBEM101
Muscle Equivalent Matter (835MHz)		August 2002	PCTMEM201
Microwave Amp. Model: 5S1G4, (800M	1Hz - 4.2GHz)	January 2002	22332
Gigatronics 8651A Power Meter		January 2002	1835299
HP-8648D (9kHz ~ 4GHz) Signal G	enerator	January 2002	PCT530
Amplifier Research 5S1G4 Power A	mp	January 2002	PCT540
HP-8753E (30kHz ~ 3GHz) Networ	k Analyzer	January 2002	PCT552
HP85070B Dielectric Probe Kit		January 2002	PCT501
Ambient Noise/Reflection, etc.	<12mW/kg/<3%of SAR	January 2002	Anechoic Room PCT01

NOTE:

The E-field probe was calibrated by IDX, by temperature measurement procedure. PCTEST Lab re-calibrated the E-field probe, by using TEM Cell and wave guide 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.

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16. CONCLUSION

Measurement Conclusion

The SAR measurement indicates that the EUT complies 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. The tested device complies with the requirements in respect to all parameters subject to the test. The test results and statements relate only to the item(s) tested.

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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17. REFERENCES

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