A COMPUTER CODE FOR

HIGH ALTITUDE EMP

THESIS

GNE/PH/74-1

Terry C. Chapman Capt USAF

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resulting from a nuclear burst above 60 km altitude with a gamma yield up to 60 tons. Either the direct or the ground reflected wave can be calculated. With special care, bursts up to 1 kt of gamma yield can be used.

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A COMPUTER CODE FOR

HIGH ALTITUDE EMP

THESIS

Presented to the Faculty of the School of Engineering of the Air Force Institute of Technology

Air University

in Partial Fulfillment of the Requirements for the Degree of Master of Science

by

Terry C. Chapman, B.S.

Capt USAF

Graduate Nuclear Engineering

January 1974

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Preface

It is my pleasure to recognize several people who contributed in various ways to make this work possible.

I want to thank my advisor, Maj Carl T. Case. His patience and helpful suggestions were important factors to the successful conclusion of this work. In addition, I want to point out that the theoretical portion of this work is based largely on a series of lectures he gave while teaching the Electromagnetic Waves (EE 6.30) course during the summer quarter of 1973. His unusual ability to present difficult topics in a way that is easily understood by the student was the largest and most important contribution to the success of this work.

I want to thank Dr. Charles J. Bridgman for his suggestions and helpful comments made during the early stages of the work.

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Last, but not least, I want to gratefully acknowledge the large contribution of my wife, Karen. She offered moral support, punched computer cards, typed drafts, and in many other ways contributed to the successful conclusion of this work.

Terry C. Chapman

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Abstract

A relatively inexpensive computer code is developed to calculate the peak value of the electric field contained in an electromagnetic pulse generated by the gamma rays from a high altitude nuclear burst. The code is based on the Karzas and Latter theory for the production of Compton electrons and their interaction with the earth's magnetic field.

The code can be used to calculate the peak value of the electric field at a target anywhere on or above ground level, resulting from a nuclear burst above 60 km altitude with a gamma yield up to 60 tons. Either the direct or the ground reflected wave can be calculated. With special care, bursts up to 1 kt of gamma yield can be used.

A COMPUTER CODE FOR HIGH ALTITUDE EMP

I. Introduction

The effects of a nuclear environment on aerospace systems is an important factor in systems analysis. During the past few years several students have worked with Professor Bridgman at the Air Force Institute of Technology (AFIT) on a computer code to determine survivability of a system with known nuclear vulnerabilities from a variable nuclear threat. The AFIT survivability code capabilities include blast, thermal, x-ray, gamma ray, and neutron effects. The high altitude EMP code presented in this report is intended to be used in conjuction with the AFIT survivability code.

The EMP (electromagnetic pulse) from a nuclear weapon is usually considered to be a radiating electromagnetic wave of short duration containing many frequencies. However, the nuclear generated EMP was not studied seriously until a considerable time after the first nuclear explosion. At present there is a significant amount of work being done to model EMP generation and effects. For example the Air Force Weapons Laboratory (AFWL) and several civilian companies under contract to the USAF are working in the field.

There are several different types of EMP with distinctions made between the mechanisms which generate them. Kinsley (Ref 1) presents a comprehensive discussion of the various types of EMP. For example, a nuclear burst on the ground produces an EMP with different characteristics

than those from a high altitude burst. Also, nuclear burst products interacting directly with a system can produce an EMP within the system or even within the circuits of the system. This report considers only the EMP generated by high altitude burst gamma rays interacting with the atmosphere.

The high altitude EMP code developed in this report is based on the theory of Karzas and Latter (Ref 2). Briefly, the theory develops a model for the interaction of Compton electrons with the geomagnetic field. The Compton electrons are produced by prompt gamma radiation from the burst in a reasonably well defined region in the atmosphere. Several simplifications are made before arriving at the final equations.

Since several of the simplifications and assumptions used are implicit in the presentation of the theory, it is appropriate to list them here. Only one group of monoenergetic unscattered gamma rays are considered to produce Compton electrons. Each gamma which interacts is assumed to produce one and only one Compton electron initially traveling precisely in the radial direction. No angular distribution of Compton electrons is allowed. All Compton electrons are assumed to have the same energy. Curvature of the Earth's magnetic field is ignored. The electromagnetic fields are not self-consistent, that is, only the geomagnetic field is considered to affect the motion of the Compton electrons. Cascading of secondary electrons and recombination of ions is ignored. The low

frequency portion of the pulse is not considered. The Earth is assumed to be flat and the finite conductivity of ground is not considered. The burst is assumed to be far from the absorption region. Only gamma ray effects are considered.

Although the final model is somewhat restricted by these assumptions and simplifications, the end result is a relatively inexpensive computer code which gives a peak value of the electric field at any target point on or above the ground, which is an upper bound on the actual peak value.

Section II of this report develops the theory and derives the equations used in the code. Section III describes the calculational procedures used in the code. Section IV presents a sample of typical results and a study of input parameter variation. Section V is a discussion of the code's limitations and uses, with recommendations for possible improvements. Appendix A is a code user's guide. Appendix B is the detailed flow charts for the entire code. And finally Appendix C is a listing of the complete code.

II. Theory

Overview

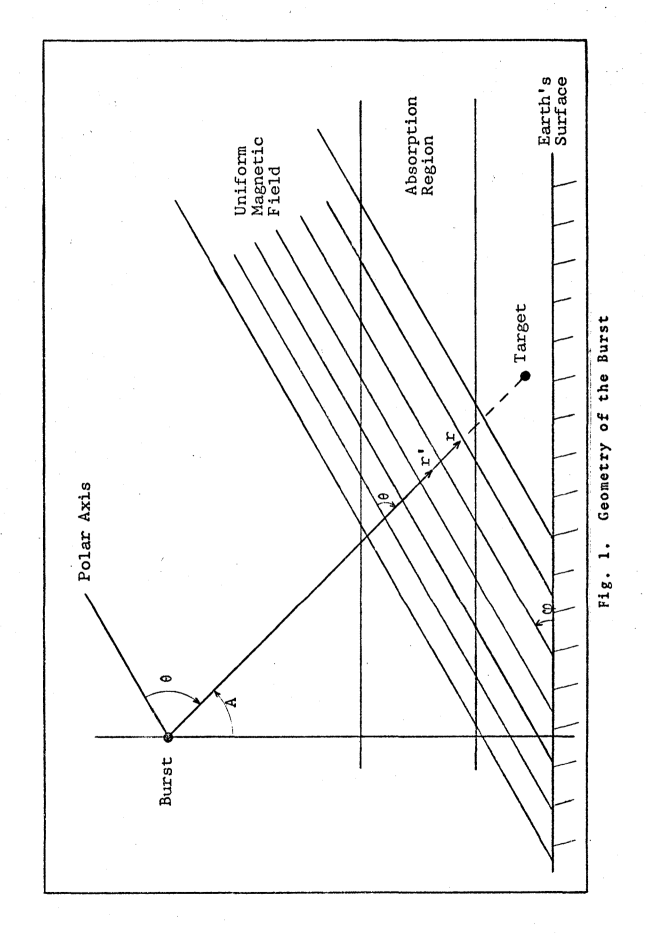
The EMP problem is a problem in classic electromagnetic theory. A solution of Maxwell's equations is a solution of the problem. In this case it is necessary to model the current and charge densities generated by the gamma rays in the absorption region to obtain the sources and conductivity needed to solve Maxwell's equations.

Expressions for the current sources and conductivity are obtained in four steps. The transport of the gammas from the burst to the absorption region is used to obtain the number density of reacting gammas. This result is used with the models for the current and charge densities to obtain preliminary expressions. Then after considering the relativistic motion of the Compton electrons, the preliminary expressions are transformed to spherical coordinates.

After presenting Maxwell's equations in a convenient form, they are transformed to spherical coordinates and retarded time. A high frequency approximation is then made to arrive at the final equations.

Electron Current and Density

<u>Gamma Transport</u>. Consider the geometry shown in Fig. 1. The nuclear burst occurs at the origin at time, t = 0. The gamma rays move to point r' in time t' and at that point and time interact to create Compton electrons. It is assumed



that each gamma creates one and only one Compton electron traveling in the radial direction with the maximum Compton recoil energy.

The gamma ray emission rate can be taken as

$$\frac{dN(t)}{dt} = \frac{Y}{E} f(t)$$
 (1)

where

N(t) = number of gamma rays emitted
Y = gamma ray yield of burst
E = mean energy of the gamma rays
f(t) = time dependence of the yield

and

$$\int_{-\infty}^{\infty} f(t) dt = 1$$
 (2)

The number density of gammas, g(r), which interact at a point, r, can be taken as

$$g(\mathbf{r}) = \frac{\gamma}{E} \frac{\exp\left\{-\int_{0}^{\mathbf{r}} \frac{d\mathbf{r}'}{\lambda(\mathbf{r}')}\right\}}{4\pi r^{2} \lambda(\mathbf{r})}$$
(3)

where

 λ = mean free path for production of Compton

electrons.

Electron <u>Currents</u> and <u>Densities</u>. The rate of production of primary (Compton) electron density, n_{pri} , is

$$\frac{dn_{pri}}{dt} = g(r)f(t - r/c)$$
(4)

Following the Karzas-Latter approach (Ref 2) it is assumed that the electrons maintain their initial speed, V_0 , throughout their range, R, and then abruptly stop. Also, it is assumed that the secondary electrons are made at a uniform rate during the lifetime, R/V_0 , of the Compton electrons. Therefore, the rate of production of secondary electron density, n_{sec} , is

$$\frac{dn_{sec}}{dt} = \left\{ \frac{E_{pri}/33ev}{R/V_0} \right\} n_{pri} = \frac{qV_0}{R} n_{pri}$$
(5)

where

Epri = the initial energy of the Compton electrons
R = the range of the Compton electrons in air
q = Epri/33ev
33ev = average ionization energy per molecule for
air

 V_0 = the speed of the Compton electrons R/V₀ = the lifetime of the Compton electrons

Now consider the current resulting from the Compton electrons. The differential current is the charge times the velocity times the differential density of electrons. Hence

$$d\vec{J}^{c} = -e\vec{V}(t-t')g(r')f(t' - r'/c)dt'$$
 (6)

7.

where

 $\vec{V}(t-t')$ = velocity of the Compton electrons at time t which were created at time t'.

Putting (6) into integral form gives

$$\vec{J}^{c} = -e \int_{t-R/V_{0}}^{t} g(r')f(t' - r'/c)\vec{V}(t-t')dt'$$
(7)

Now let

$$\tau' = t - t' \tag{8a}$$

$$\tau = t - (r/c)$$
(8b)

$$X(\tau') = X(\tau) = \mathbf{r} - \mathbf{r}' \qquad (8c)$$

Also note that

$$(\mathbf{r} - \mathbf{r}') << \mathbf{r} \text{ or } \mathbf{r}' \tag{9}$$

for distant explosions (see Fig. 1). So,

$$g(\mathbf{r}) \simeq g(\mathbf{r}') \tag{10}$$

Using Eqs (8), (9), and (10) in Eq (7) gives

$$\dot{J}^{c} = -eg(r) \int_{0}^{R/V_{0}} \dot{V}(\tau') f\left(\tau - \tau' + \frac{X(\tau')}{c}\right) d\tau' \qquad (11)$$

Using similar arguments,

$$n_{\text{pri}} = g(\mathbf{r}) \int_{0}^{R/V_{0}} f\left(\tau - \tau' + \frac{X(\tau')}{c}\right) d\tau' \qquad (12)$$

And putting Eq (12) into Eq (5) yields

$$n_{sec} = \frac{qV_0}{R} \int_{-\infty}^{\tau} n_{pri}(\tau') d\tau'$$

$$= g(\mathbf{r}) \frac{qV_0}{R} \int_{-\infty}^{\tau} \left[\int_{0}^{R/V_0} f\left(\tau' - \tau'' + \frac{X(\tau'')}{c}\right) d\tau'' \right] d\tau' \quad (13)$$

<u>Relativistic Electron Motion</u>. Equations (11), (12), and (13) contain $r(\tau)$ and $X(\tau)$ which are not yet defined in an easily obtained form. The equation of motion for a Compton electron is

$$\frac{d}{dt} (m\gamma \vec{V}) = -e\vec{V}X\vec{B}_0$$
(14)

where

m = electron rest mass

$$\gamma = [1 - (V/c)^2]^{-1/2}$$

 \vec{B}_0 = earth's magnetic field = $B_0 \hat{U}_z$

Again it is assumed that V_0 is constant throughout the electron's lifetime.

With $\omega = eB_0/m\gamma Eq$ (14) becomes

$$\frac{d}{d\tau} \vec{V}(\tau) = -\vec{V}(\tau) \hat{XU}_{z} \omega \qquad (15)$$

Breaking Eq (15) into its rectangular components

$$\frac{\mathrm{d}V_{\mathbf{x}}}{\mathrm{d}\tau} = -\omega V_{\mathbf{y}} \tag{16a}$$

$$\frac{dV}{d\tau} = \omega V_{\rm X}$$
 (16b)

$$\frac{\mathrm{d}V}{\mathrm{d}\tau} = 0 \tag{16c}$$

A solution for this set of equations is

$$V_{\rm x} = V_{\rm \perp} \cos \omega \tau$$
 (17a)

$$V_y = V_{\perp} \sin \omega \tau$$
 (17b)

$$V_{z} = V_{||}$$
(17c)

where V_{\perp} is the initial velocity component perpendicular to \vec{B}_0 and V_{\parallel} is the initial velocity component parallel to \vec{B}_0 and both are constants with respect to τ .

<u>Transformation to Spherical Coordinates</u>. It is convenient to transform the above solution to a spherical coordinate system with its origin at the burst point. The transformation from rectangular to spherical coordinates is

 $V_r = V_x \sin \theta \cos \phi + V_y \sin \theta \sin \phi + V_z \cos \theta$ (18a)

 $V_{\theta} = V_x \cos \theta \cos \phi + V_y \cos \theta \sin \phi - V_z \cos \theta$ (18b)

$$V_{\phi} = -V_{x} \sin \phi + V_{y} \cos \phi \qquad (18c)$$

Without loss of generality the coordinates can be chosen such that \vec{V} lies in the X-Y plane, hence $\phi = 0$, and the transformation becomes

$$V_{r} = V_{x} \sin \theta + V_{z} \cos \theta \qquad (19a)$$

$$V_{\rho} = V_{\chi} \cos \theta - V_{z} \sin \theta$$
 (19b)

$$V_{\phi} = V_{y}$$
(19c)

Note that

$$V_{\perp} = V_0 \sin \theta$$
 (20a)

$$V_{||} = V_0 \cos \theta$$
 (20b)

Putting Eqs (17) and (20) into Eq (19) gives

$$V_{r} = V_{0} [\sin^{2} \theta \cos \omega \tau + \cos^{2} \theta]$$
 (21a)

$$V_{\theta} = V_{0} [\cos \theta \sin \theta \cos \omega \tau - \sin \theta \cos \theta]$$
 (21b)

$$V_{\phi} = V_0 [\sin \theta \sin \omega \tau]$$
 (21c)

Now $X(\tau)$ can be written as

$$X(\tau) = \int_{0}^{\tau} V_{r} d\tau = V_{0} [\sin^{2} \theta \frac{\sin \omega r}{\omega} + \tau \cos^{2} \theta] \quad (22)$$

Equations (21) and (22) substituted into Eq (11) give

$$J_{\mathbf{r}}^{\mathbf{c}} = -eg(\mathbf{r})V_0 \int_0^{\mathbf{R}/V_0} f(\mathbf{T})[\cos^2\theta + \sin^2\theta \cos \omega \tau']d\tau' \quad (23)$$

$$J_{\theta}^{c} = -eg(r)V_{0} \int_{0}^{R/V_{0}} f(T)[\sin \theta \cos \theta (\cos \omega \tau' - 1)]d\tau' (24)$$

$$J_{\phi}^{c} = -eg(r)V_{0} f_{0}^{R/V_{0}} f(T) [\sin \theta \sin \omega \tau'] d\tau' \qquad (25)$$

where

$$\Gamma = \tau - (1 - \beta \cos^2 \theta)\tau' + \beta \sin^2 \theta \frac{\sin \omega \tau'}{\omega}$$
 (26a)

with

$$\beta = V_0/c \tag{26b}$$

Equations (23), (24), (25), and (26) provide the Compton currents within the absorption region in a form which can be used in the final field equations. In addition to the Compton currents, an expression for the conductivity within the absorption region is needed.

Equations (21) and (22) substituted into Eq (13) give

$$n_{sec}(\tau) = \frac{qV_0}{R} g(\mathbf{r}) \int_{-\infty}^{\tau} \left[\int_{0}^{R/V_0} f(T') d\tau'' \right] d\tau'$$
(27)

where

$$T' = \tau' - (1 - \beta \cos^2 \theta)\tau'' + \beta \sin^2 \theta \frac{\sin \omega \tau''}{\omega}$$
(28)

Consider the equation of motion for secondary electrons. Neglecting the $\vec{V}X\vec{B}_0$ term, which is small compared to the other terms (Ref 2) it is

$$m \frac{d\vec{V}}{dt} = -e\vec{E} - m\vec{V}\nu_c \qquad (29)$$

where

$$v =$$
 electron collision frequency.

These secondary electrons have velocities in the thermal region and are assumed to reach their maximum velocity instantly. In this case, Eq (29) becomes

$$\vec{V} = \frac{-e}{m\nu} \vec{E}$$
(30)

The current source from the secondary electrons is

$$\dot{J}^{sec} = -e\vec{V}(\tau)n_{sec}(\tau) = \frac{n_{sec}(\tau)}{mv_c}e^{2t}$$
(31)

or in terms of conductivity

$$\vec{J}^{sec} = \sigma(\tau)\vec{E}$$
(32)

where

$$\sigma(\tau) = \frac{n_{sec}(\tau)}{mv_c} e^2$$
 (33)

Equations (32) and (33) provide the needed expressions for the conductivity.

Electromagnetic Fields from High Altitude Currents

<u>Maxwell's Equations</u>. Now that the Compton currents and the conductivity due to secondary electrons have been obtained, consider the field equations.

Maxwell's equations are

$$\vec{\nabla} \mathbf{x} \vec{E} = - \frac{\partial \vec{B}}{\partial t}$$
(34a)

$$\vec{\nabla} \mathbf{x} \vec{B} = \mu_0 \vec{J} + \frac{1}{c^2} \frac{\partial \vec{E}}{\partial t}$$
(34b)

$$\vec{\nabla} \cdot \vec{E} = \frac{q_v}{\varepsilon_0}$$
(34c)

$$\vec{\nabla} \cdot \vec{B} = 0 \tag{34d}$$

where

 \vec{J} = total current density q_y = total charge density

Continuity of charge requires

$$\frac{\partial q}{\partial t} + \vec{\nabla} \cdot \vec{J} = 0$$
 (35)

It is convenient to combine the above equations into equations containing only \vec{E} in one and only \vec{B} in the other. Doing so gives

$$\left(\nabla^{2} - \frac{1}{c^{2}} \frac{\partial^{2}}{\partial t^{2}}\right) \stackrel{?}{E} = \mu_{0} \frac{\partial \vec{J}}{\partial t} + \frac{1}{\varepsilon_{0}} \vec{\nabla} q_{\nu}$$
(36)

$$\left(\nabla^{2} - \frac{1}{c^{2}} \frac{\partial^{2}}{\partial t^{2}}\right)\vec{B} = -\mu_{0}\vec{\nabla}x\vec{J}$$
(37)

<u>Transformation to Spherical Coordinates and Retarded</u> <u>Time</u>. Equations (36) and (37) will now be transformed to spherical coordinates and retarded time. Consider the transformation

$$\tau = t - r/c \qquad (38a)$$

$$\mathbf{r'} = \mathbf{r} \tag{38b}$$

$$\theta^{\dagger} = \theta$$
 (38c)

$$\phi' = \phi \qquad (38d)$$

This is a spherical coordinate system where time is measured at each radial point in terms of the arrival of the bomb gamma rays at that point.

Using Eq (38) it is easily shown that

$$\frac{\partial}{\partial t} = \frac{\partial}{\partial \tau}$$
(39)

$$\frac{\partial}{\partial \mathbf{r}} = \frac{\partial}{\partial \mathbf{r}'} - \frac{1}{c} \frac{\partial}{\partial \tau}$$
(40)

$$\frac{\partial}{\partial \theta} = \frac{\partial}{\partial \theta^{\dagger}}$$
(41)

$$\frac{\partial}{\partial \phi} = \frac{\partial}{\partial \phi}$$
(42)

Thus the operator

$$\frac{\partial}{\partial t}$$

 $\frac{\partial}{\partial \tau}$

transforms to

and the operator

transforms to

$$\vec{\nabla} - \hat{\mathbf{U}}_{\mathbf{r}} \frac{1}{c} \frac{\partial}{\partial \tau}$$

Similarly, the operator

$$\nabla^2 - \frac{1}{c^2} \frac{\partial^2}{\partial t^2}$$

transforms to

$$\nabla^2 - \frac{2}{c} \frac{\partial}{\partial \tau} \frac{1}{r} \frac{\partial}{\partial r} r$$

Equation (36) now becomes

$$\left[\nabla^{2} - \frac{2}{c}\frac{1}{r}\frac{\partial}{\partial\tau}\frac{\partial}{\partial\tau}r\right] \stackrel{*}{E} = \mu_{0}\frac{\partial\vec{J}}{\partial\tau} + \frac{1}{\epsilon_{0}}\vec{\nabla}q_{v} - \hat{U}_{r}\frac{1}{c}\frac{\partial q_{v}}{\partial\tau}$$
(43)

and Eq (35) becomes

$$\frac{\partial \mathbf{q}}{\partial \tau} = -\vec{\nabla} \cdot \vec{\mathbf{j}} + \hat{\mathbf{U}}_{\mathbf{r}} \frac{1}{c} \frac{\partial}{\partial \tau} \cdot \vec{\mathbf{j}} = -\vec{\nabla} \cdot \vec{\mathbf{j}} + \frac{1}{c} \frac{\partial \mathbf{J}_{\mathbf{r}}}{\partial \tau}$$
(44)

Using Eq (44) in Eq (43) and rearranging gives

$$-\nabla^{2}\vec{E} + \hat{U}_{\mathbf{r}} \frac{1}{c\varepsilon_{0}} \vec{\nabla} \cdot \vec{J} + \frac{1}{\varepsilon_{0}} \vec{\nabla} q_{\mathbf{v}}$$
$$+ \frac{\partial}{\partial\tau} \left[\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial\tau} (r\vec{E}) + \mu_{0} (\vec{J} \cdot \hat{U}_{\mathbf{r}} J_{\mathbf{r}}) \right] = 0 \qquad (45)$$

Similarly, Eq (37) becomes

$$-\nabla^2 \vec{B} + \mu_0 \vec{\nabla} x \vec{J} + \frac{\partial}{\partial \tau} \left[\frac{2}{rc} \frac{\partial}{\partial r} (r\vec{B}) \right]$$

$$+ \frac{\partial}{\partial \tau} \left[\frac{\mu_0}{c} \left(\hat{U}_{\phi} J_{\phi} - \hat{U}_{\theta} J_{\theta} \right) \right] = 0$$
 (46)

<u>High Frequency Approximation</u>. Again, following the Karzas-Latter model, note that the variation of currents with distance is slow compared to variations with time and that the fields resulting from the transverse currents are rapidly varying in character, as are the currents themselves. Therefore, only the $\partial/\partial \tau$ terms are kept in the transverse field equations. Since the radial components do not propagate outside of the absorption region, they are not considered further in this report.

The transverse equations become

$$\frac{\partial}{\partial \tau} \left[\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (r E_{\theta}) + \mu_0 J_{\theta} \right] = 0$$
 (47)

$$\frac{\partial}{\partial \tau} \left[\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rE_{\phi}) + \mu_0 J_{\phi} \right] = 0$$
 (48)

$$\frac{\partial}{\partial \tau} \left[\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rB_{\theta}) - \frac{\mu_0}{c} J_{\phi} \right] = 0$$
 (49)

$$\frac{\partial}{\partial \tau} \left[\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rB_{\phi}) + \frac{\mu_0}{c} J_{\theta} \right] = 0$$
 (50)

These equations are called the Karzas-Latter high frequency approximation for the EMP fields, and they are useful in the range $0 < \tau < 100$ shakes.

Integration with respect to time yields

$$\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rE_{\theta}) + \mu_0 J_{\theta} = 0 \qquad (51)$$

$$\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rE_{\phi}) + \mu_0 J_{\phi} = 0$$

$$\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rB_{\theta}) - \frac{\mu_0}{c} J_{\phi} = 0 \qquad (53)$$

$$\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rB_{\theta}) + \frac{\mu_0}{c} J_{\theta} = 0 \qquad (54)$$

Recall that the total current density is

$$\vec{J} = \vec{J}^{\text{pri}} + \vec{J}^{\text{sec}} = \vec{J}^{\text{c}} + \sigma(\tau)\vec{E} \qquad (55)$$

so Eqs (51) and (52) become

$$\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rE_{\theta}) + \mu_0 J_{\theta}^c + \mu_0 \sigma(\tau) E_{\theta} = 0$$
 (56)

$$\frac{2}{c} \frac{1}{r} \frac{\partial}{\partial r} (rE_{\phi}) + \mu_0 J_{\phi}^c + \mu_0 \sigma(\tau) E_{\phi} = 0$$
 (57)

With the aid of a computer, it is now possible to obtain numerical solutions for the above equations which will yield a slightly high estimate of the peak value of the EMP pulse resulting from a high altitude burst.

Below the absorption region the Compton currents and the conductivity are zero. In this case, Eqs (56) and (57) have the following solutions:

$$E_{A} = C_{1}/r \tag{58}$$

$$E_{\phi} = C_2/r \tag{59}$$

where C_1 and C_2 are determined by the values of E_{θ} , E_{ϕ} , and r at the bottom of the absorption region.

III. Code Description

General Approach

Equations (56), (57), (58), and (59) were chosen as the simplest ones to solve numerically. Of course, Eqs (24), (25), (27), and (33) are used to obtain the Compton currents and conductivity needed to solve Eqs (56) and (57).

The B - field equations are not solved since

$$\mathbf{E} = \mathbf{c}\mathbf{B} \tag{60}$$

can be used to obtain B once E is found. This relationship is based on the assumption that the EMP pulse is a spherical wave propagating in free space, below the absorption region.

The function used for the time dependence of the weapon yield is the one recommended by Pomranning (Ref 3).

$$f(\tau) = (1/N) \frac{(\alpha+\beta) \exp (\tau-\tau_0)}{\beta + \alpha \exp [(\alpha+\beta)(\tau-\tau_0)]}$$
(61)

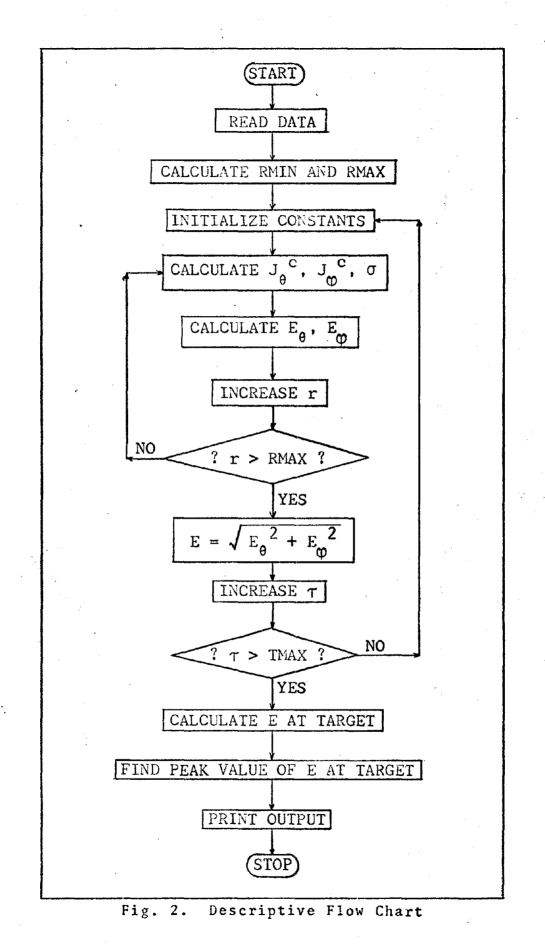
where N is chosen such that

$$\int_{0}^{\infty} f(\tau) d\tau = 1$$
 (62)

and $\alpha > \beta$.

This function rises like $e^{\alpha \tau}$ for small τ , falls like $e^{-\beta \tau}$ for large τ , and has a single maximum at τ_0 .

Figure 2 presents a flow chart which is descriptive of the approach taken solving the equations. The top of the absorption region is assumed to be at 50 km altitude and the bottom of the absorption region is assumed to be



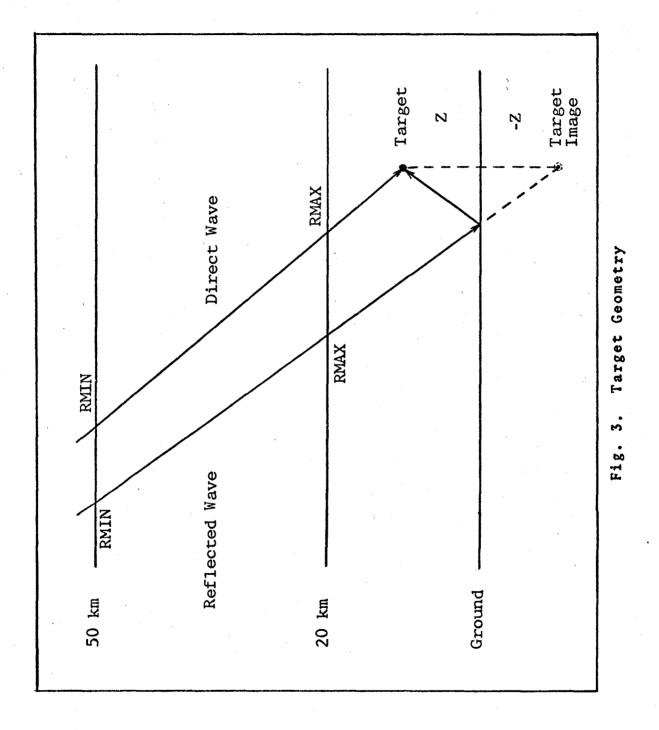
at 20 km altitude. Calculations by Latter and LeLevier (Ref 4) indicate that 20 km to 50 km is the altitude where most of the prompt gamma ray energy is deposited.

Figure 3 depicts the target geometry. The value for RMIN is determined by the intersection of the line of sight with the 50 km altitude. The value for RMAX is determined by the intersection of the line of sight with the 20 km altitude. If the target is in the absorption region the target altitude determines RMAX for the direct wave calculation. These two values of r are the limits on the mesh in the r direction. The line of sight is divided into the desired number of steps along r for the integration on r in the absorption region.

The retarded time direction of the mesh is divided into 0.1 shake steps up to 10 shakes and then 1.0 shake steps on up to 100 shakes. Calculation can be stopped at any desired TMAX from 10 to 100 shakes, which is the upper limit of the usefulness of the high frequency approximation.

If the ground reflected wave is to be calculated, the mirror image of the target, below ground, is used to find the line of sight from the burst to the target. (Refer to Fig. 3.)

At r = RMIN all of the fields are assumed to be zero. For each τ , equations (57) and (58) are integrated over r from RMIN to RMAX and the value of E at the bottom of the absorption region is stored. At each step in r, equations (24), (25), and (27) are numerically integrated. Then



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equations (58) and (59) are combined into

$$E = \frac{(RMAX)(E_{RMAX})}{r_{target}}$$

(63)

to find E at the target.

The E array is then searched to find the peak value before printing out the results.

Inputs

The code uses a right handed Cartesian coordinate system with the ground in the X-Y plane, the \vec{B}_0 vector in the Y-Z plane, and \hat{j} pointing towards the equator. For example, in the northern hemisphere, \hat{i} is magnetic west, \hat{j} is magnetic south, and \hat{k} is altitude. The origin of the coordinate system is always at ground zero, directly below the burst. Note that this coordinate system is not the same as the Cartesian systems used earlier.

Referring to the above coordinate system the target coordinates, (X,Y,Z), are read in using units of meters. If the reflected wave is to be calculated the altitude is read in as a negative number, (X,Y,-Z).

The height of the burst is read in using units of kilometers. The gamma yield of the burst is read in using units of kilotons.

The magnitude of the Earth's magnetic field is read in using units of webers per square meter. The dip angle (ϕ in Fig. 1) of the magnetic field is read in using units of degrees.

NDELR, the desired number of steps to be used in the integration over r in the absorption region, is read in as any integer in the closed interval [50, 500].

TMAX, the retarded time where calculations are to be stopped, is read in, using units of shakes, as any integer in the closed interval [10, 100].

Preliminary Calculations

Before starting the numerical integrations, the code performs several preliminary calculations. The input data is converted to MKS units. The reflected wave is used whenever Z is greater than 49 km or less than 0. The target coordinates are transformed to a spherical coordinate system with the burst at the origin and the polar axis parallel to \vec{B}_0 . The line of sight intersections with the absorption region are determined. And finally, the constant angles required by the code, θ and A, (see Fig. 1) are calculated.

Calculation of Compton Currents and Conductivity

The two Compton currents, J_{θ}^{c} and J_{ϕ}^{c} are calculated at each r, τ mesh point by numerically integrating equations (24) and (25). The step size used is 0.1 times the Compton lifetime, R/V_{0} . The integration itself is done using the 4^{th} order Runge-Kutta method (Ref 5). It should be noted that both the mean free path for Compton interaction and the Compton lifetime are exponentially scaled from sea

level values using a 7 km scale height. However, the Compton lifetime is not allowed to be greater than 100 shakes, since this is the maximum time of interest.

Monoenergetic gammas of energy 1.5 Mev are assumed. The most energetic Compton electrons resulting from 1.5 Mev gammas have a speed of 2.88 (10)⁸ m/sec. Therefore $V_0 = 2.88 (10)^8$ m/sec.

Since the integration on τ " in equation (27) is also over the Compton lifetime, this integration is carried out simultaneously with the Compton current integrations. Again, the 4th order Runge-Kutta method is used. It is broken into two parts, one for $-\infty < \tau' < 0$ and the other for 0 < $\tau' < \tau$. In this case, $-\infty$ is defined to be the time when the first gamma ray reached the top of the absorption region, since no secondaries can be produced before that time.

The integration on τ' in equations (27) is also broken into two parts, one for $-\infty < \tau' < 0$ and the other for $0 < \tau' < \tau$. In the first case, integration is started at $\tau' = 0$ and proceeds to $\tau' = -(r-RMIN)/V_0$ in steps of $\Delta \tau' = -\Delta r/V_0$. In the second case, integration is started at $\tau' = 0$ and proceeds to $\tau' = \tau$ in steps of $\Delta \tau' = \Delta \tau$. In both cases, simple step integration is used. That is

 $\int f(\tau^{\dagger}) d\tau^{\dagger} = \sum_{all i} (\Delta \tau_{i}^{\dagger}) [f(\tau_{i}^{\dagger})]$ (64)

The integration over τ' is carried out parallel to the integration of (56) and (57) over r (using space as a pseudo retarded time) and simultaneously with the increase in τ as the space integrations are repeated for each new τ .

This rather involved approach to solving equation (27) is necessary to save running time. A direct approach, with separate integrations, would at least triple or quadruple the total running time required for execution of the code.

Integration of the Field Equations

For each τ , equations (56) and (57) are integrated from r = RMIN to r = RMAX in steps of $\Delta r = (RMAX-RMIN)/NDELR$ using the 4th order Runge-Kutta method. Then the magnitude of E is found from the two components and the result is stored in the E array. τ is increased by $\Delta \tau$ and the whole process is repeated until τ reaches TMAX.

On completion of the iterations, each member of the E array is multiplied by $RMAX/r_{target}$ (equation 62). Then the E array is searched to find the peak value.

Outputs

There are several output options available in the code. The basic output it:

- 1. Gamma yield and altitude of burst.
- 2. Target coordinates from ground zero.
- 3. Distance from burst to target.
- 4. A message indicating whether the direct or the reflected wave is being calculated.

5. The time period covered by the calculation.

6. The time when the peak value occured.

7. The peak value of E at the target.

8. The τ and E arrays.

In addition, a linear and a log-log plot of $E(\tau)$ can be obtained. Also, a listing of the values of E at the bottom of the absorption region for each τ can be obtained. Either or both of these two options can be added to the basic output.

IV. Results and Input Parameter Variation

The output from a typical run is shown in Fig. 4. The $E(\tau)$ calculated during the run is shown in Fig. 5. The input data for this run was:

X	= 0 meters	(65a)
Y	= 0 meters	(65b)
Z	= 0 meters	(65c)
HOB	= 100 km	(65d)
Υ _Υ	= .001 kt	(65e)
^B 0	$= 2(10)^{-5} \text{ wb/m}^2$	(65f)
Dip Angle	= ^{20°}	(65g)
NDELR	= 50	(65h)
TMAX	= 20 shakes	(65i)

The CDC 6600 Computer required 191 sec and 33000_8 words of central memory to execute this run.

The peak value of E, 6400 V/m, obtained in this run compares favorably with Karzas-Latter's order of magnitude estimate of 10^4 V/m (Ref 2) from similar input data.

In order to gain a better knowledge of the operating capabilities of the code, the effect of varying input parameters one at a time was studied. The basic set of parameters used was:

THE TARGET IS AT CCORCINATES 0. Which is 1.000E+05 meters from the burst	•	•
DIRECT WAVE IS BEING CALCULATED		
ITERATION TERMINATED AFTER 20.0 SHAKES		•
PEAK OCCURRED AT 2.1 SHAKES		
* * * * * * * * * * * * * * * * * * *	* * * * 02 * * H * 	$(X_{i})^{(i)}$

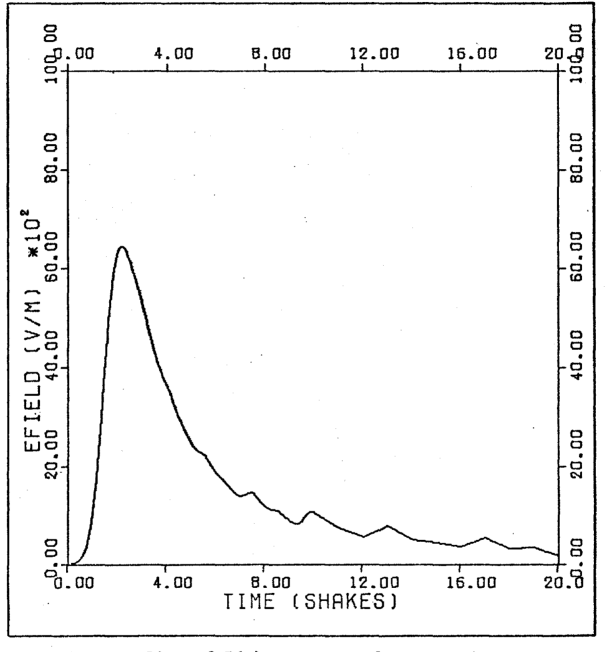


Fig. 5. Plot of $E(\tau)$ at target from a typical run

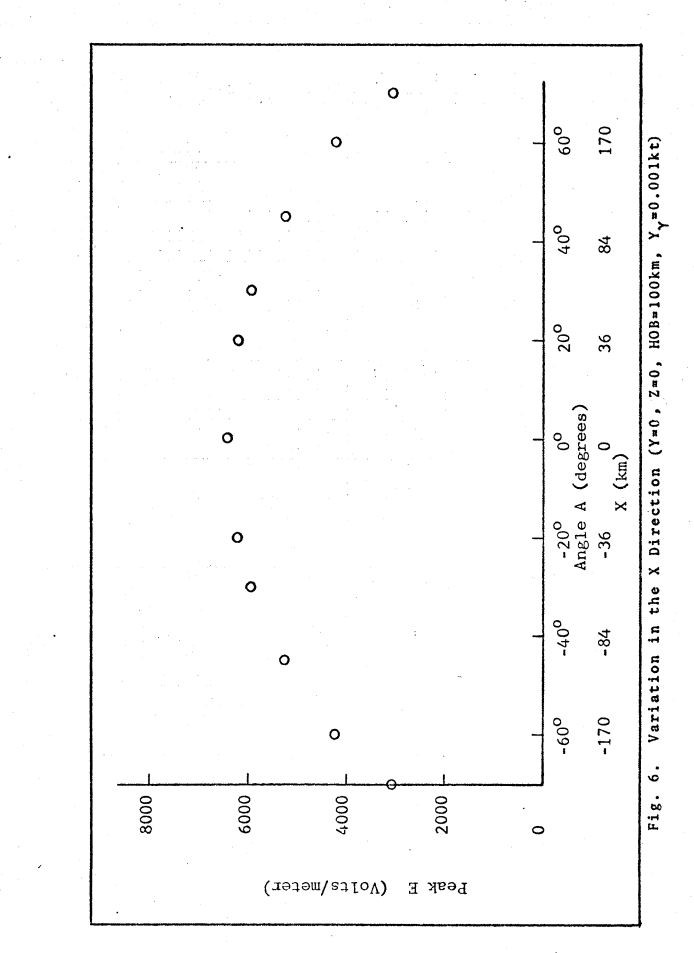
X	= 0 mete	rs		(66a)
Y	= 0 mete	rs	~	(66b)
Z	= 0 mete:	rs		(66c)
HOB	= 100 km			(66d)
Y,	= .001 k	t		(66e)

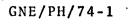
Each of the above parameters was systematically varied while holding the others constant. The other inputs were held constant at the values shown in equations (65).

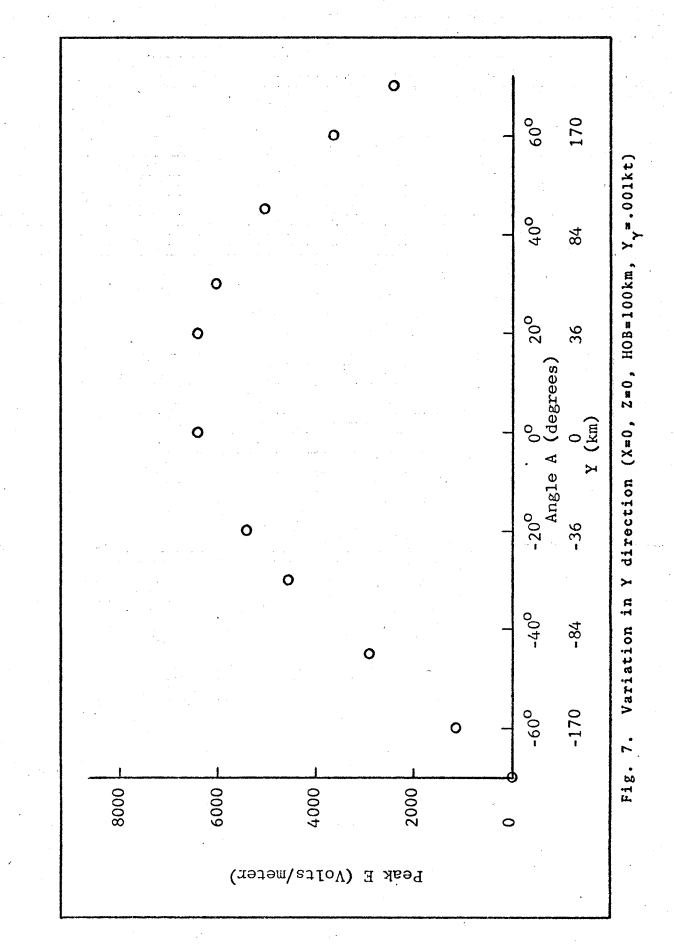
The results of the variation in X are shown in Fig. 6. Since the X axis is perpendicular to the magnetic field the symmetry about X = 0 is expected. The decrease in peak value of E for increasing distance from ground zero is due to the increasing distance from the burst.

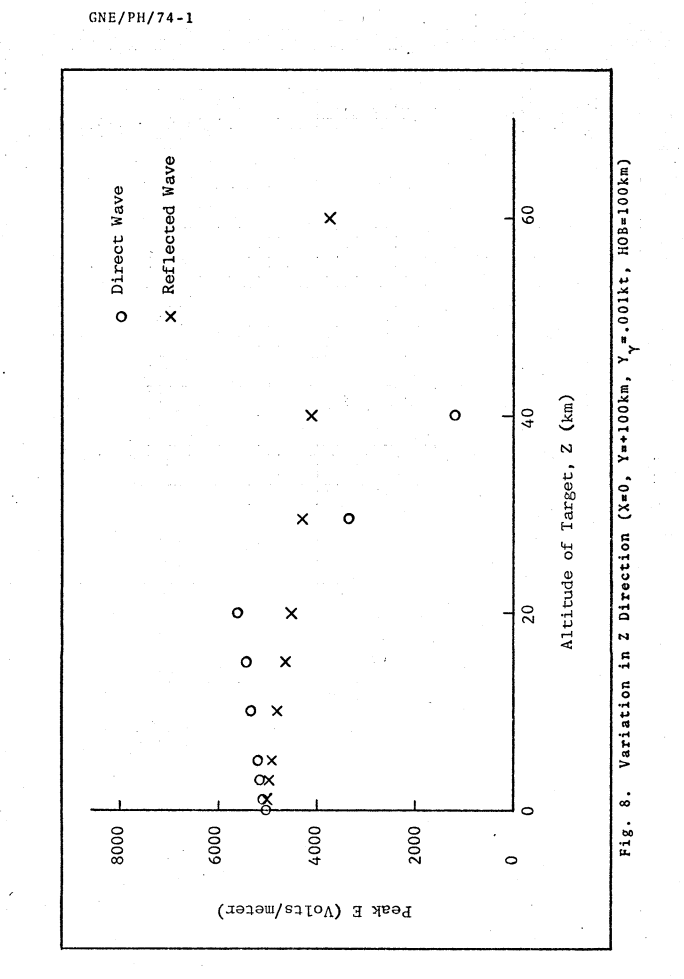
The results of the variation in Y are shown in Fig. 7. Here the peak values of E depend on the angle between \vec{r} and \vec{B}_0 , θ . When $\theta = 180^\circ$ (A = -70° and Y = -275 km) the peak E drops to zero. The maximum peak E is skewed toward A = 20° (θ = 90° and Y = 36 km). The maximum is not exactly at A = 20° because of the increased distance from the burst. These characteristics are expected since an electron moving perpendicular to the magnetic field would feel the strongest acceleration from it while an electron moving parallel to the magnetic field would feel no acceleration at all.

The results of variation in Z are shown in Fig. 8. In this case, both the direct and the reflected waves were calculated at each point below the top of the absorption









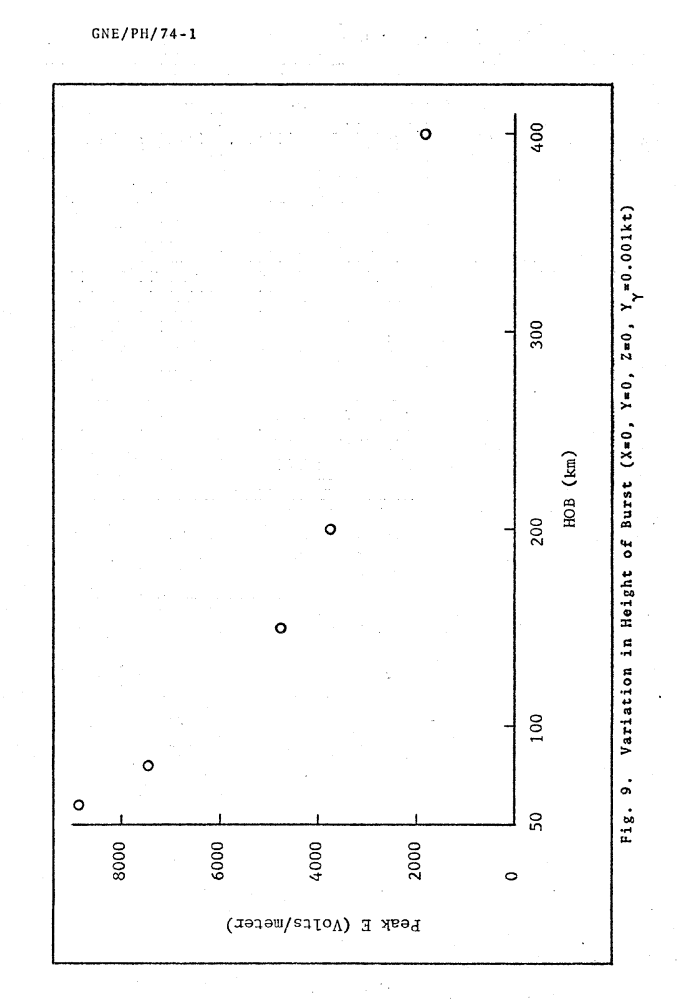
region. Note that Y = 100 km for these runs. As expected, the direct wave falls off rapidly as the target altitude passes through the absorption region, since less of the absorption region contributes to the wave with each increase in altitude. The crossover point where the reflected wave becomes the largest occured at 25 km in this case. Above ground zero the crossover point was 29.4 km. The altitude of the crossover point is both yield and geometry dependent. It is necessary for the user to calculate both waves whenever there is any doubt which one is the largest.

The reflected wave calculation assumes 100% reflection from the ground and no attenuation in the absorption region or the ionosphere. These assumptions are reasonable if it is recalled that only the high frequency component is being considered and that it requires at least

$$\frac{40 \text{ km}}{3(10)^8 \text{ m/sec}} = 133 \text{ }\mu \text{ sec}$$
(67)

for the wave to leave the absorption region, reach the earth, be reflected, and return to the absorption region. This length of time is enough for a significant number of the free electrons to recombine and reduce the effective conductivity of the absorption region.

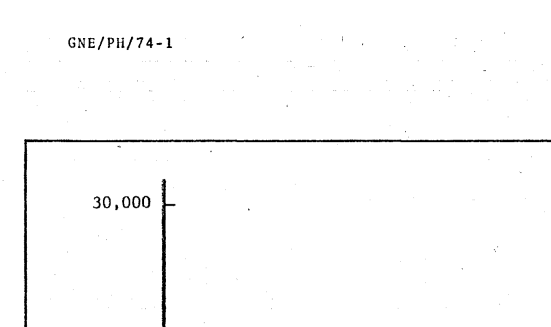
The results of variation in HOB are shown in Fig. 9. For all values of HOB attempted below 60 km the code went unstable. Infinite values for E were obtained which resulted in abnormal termination of the calculations. This is



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expected since the burst is assumed to be distant from the absorption region (equations 9 and 10).

The results of variation in gamma yield are shown in Fig. 10. For all gamma yields attempted above 60 tons the code went unstable, giving infinite values for E. However, the instability always occured at times later than the natural peak value of E. For example, with 80 tons of gamma yield, the natural peak occured at 1 shake and the instability occured at 10 shakes. By using the natural peak value and ignoring the instability, reasonable values for peak E were obtained up to 1 kt of gamma yield.



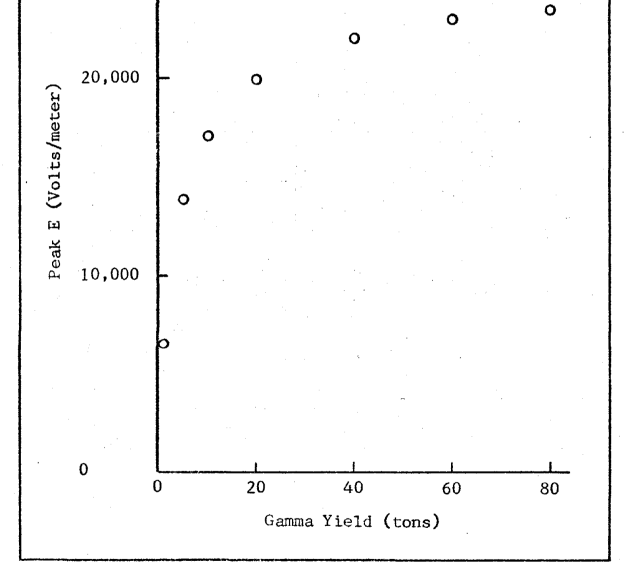


Fig. 10. Variation in Gamma Yield (X=0, Y=0, Z=0, HOB=100km)

V. Discussion and Recommendations

Limitations

Most of the limitations of the code are inherent in the model upon which it is based. Approximations such as a flat earth, a uniform magnetic field, and constant speed Compton electrons can be improved only by changing the model. In addition, the effect of the self generated electromagnetic fields on the motion of the Compton electrons is ignored, as is recombination of both primary and secondary electrons. The possibility of a single gamma ray interacting to produce more than one Compton electron is not allowed. In the absorption region the contribution of the non-propagating radial component of the electric field is neglected. Also, the model is not easily adapted to multi-group gamma transport, or to multiple burst calculations.

The code calculates only the effect of the gamma rays. The user must keep in mind that X-ray generated EMP becomes important for bursts above 100 km.

The code does not account for the increase in altitude of the absorption region for slant angles (angle A in Fig. 1) greater than 60° which is indicated by Latter and LeLevier (Ref 4).

Since 97% of the running time of the code is used for numerical iteration it is not practical to adapt the code to run more than one target at a time. Two targets would

merely double the running time, so it is simpler to just make two runs. Typical requirements are 200 seconds running time with 33000_8 words of central memory on the CDC 6600 computer using NDELR = 50 and TMAX = 20 shakes.

Uses

The code can be used to calculate the peak value of the E field at a target, anywhere on or above ground level, resulting from a nuclear burst above 60 km altitude with a gamma yield up to 60 tons. Either the direct or the ground reflected wave can be calculated. With special care, bursts up to 1 kt of gamma yield can be used.

Recommendations

In the interest of accuracy, the targets should be located such that the slant angle, A, is between -60° and $+60^{\circ}$.

By accepting a much longer running time the accuracy and hopefully, the stability of the code could be improved by using a smaller step size in the integration of the Compton current equations. Reducing the step size from one tenth of the Compton lifetime to one shake would require approximately ten times as much running time as the code presently requires. This possibility should be investigated further to determine the optimum step size for obtaining the best relationship between accuracy and running time.

Another possibility for increasing the accuracy and stability of the code is to reduce the step size in r. The present code has the capability of dividing the absorption region into 500 steps in r along the line of sight. Of course, the running time required for 500 steps is ten times that required for 50 steps. A modification of the code to allow more than 500 steps would increase the amount of computer core required as well as increasing the running time. This provides another area for investigation to determine the best trade off point between accuracy and running cost.

These two possibilities could be investigated with minor modifications to the present code. However, the computer time required would be considerable.

In addition, there are numerous possibilities for improvements in the model itself. Some of the more important ones are;

Using multigroup gamma transport.

Using multigroup Compton electrons.

Allowing angular distribution of Compton electrons.

Using self consistent electromagnetic fields.

Including the low frequency components. Each of these would require major modifications to the present code.

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Appendix A

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EMP Code User's Guide

EMP Code User's Guide

The code is run the same as any other Fortran Extended program, but due to the running time it should be converted to binary form before execution. The plotting subroutine requires an on-line plotter and both linear and log plotting library subroutines.

The input data is read in the following order: Data card #1, using FORMAT (7F10.0, 215), contains;

> X,Y,Z The target coordinates in meters HOB The height of the burst in kilometers (60 km \leq HOB)

GAMYLD The gamma yield in kilotons

(GAMYLD < 1 kt)

BFIELD	The	Earth's	magnetic	field	in	wb/m ²	
DITTTD	INC	Laith S	magnette	TTGTU	II	WUIM	

BANGLE The magnetic field dip angle in degrees

NDELR The number of steps in r taken through

the absorption region (50 < NDELR < 500)

OUT The output control parameter

Data card #2, using FORMAT (13), contains;

ITER The time period covered by the iterations in shakes ($10 \le ITER \le 100$) (ITER = TMAX) Data card #3, using FORMAT (4F10.0), contains;

Α	Pomranning constant α in inverse shakes
В	Pomranning constant β in inverse shakes
RN	Pomranning constant N in shakes
то	Pomranning constant τ_0 in shakes

Default values are provided for BANGLE, BFIELD, and NDELR. They are 40°, 0.00002 wb/m^2 , and 50 respectively. If these default values are desired, zero must be punched in their respective card fields.

The ground reflected wave at the target is obtained by reading in the target altitude, Z, as a negative number. For any target within the absorption region, both the direct and the ground reflected wave should be calculated to determine which one is the strongest.

For values of GAMYLD between 0.06 kilotons and 1.0 kilotons the code will most likely go unstable. This instability occurs after the real peak has been calculated, but the peak value printed out may not be the real peak. Since execution is terminated when the field becomes greater than 1E15 V/m, the array search can result in a false peak value. In this case, the arry itself (or the plot) can be used to determine the real peak value.

Increasing NDELR makes the step size in r through the absorption region smaller and the calculation becomes more accurate. However, total running time varies directly with changes in NDELR. For example, using NDELR = 100 instead of NDELR = 50 will approximately double the running time required for NDELR = 50.

There are four output options provided. Option 0 prints out the informative messages, the calculated peak value at the target, the E array, and the τ array. Option 1 adds a linear plot of the first 20 shakes and a log-log

plot of 100 shakes of E as a function of τ at the target. Option 2 includes both Option 0 and Option 1 and adds a printout of E and σ as a function of τ at the bottom of the absorption region. Option 3 deletes the plots from Option 2. The last two options are primarily for debugging since a partial printout is made for each completed iteration even if execution is terminated before the iterations are completed. The first two options are best for production runs.

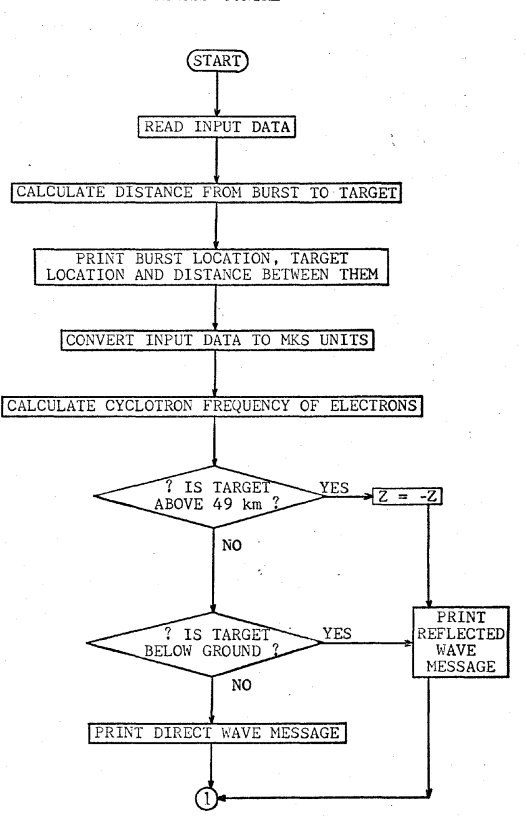
The only requirements on the Pomranning constants are N must be chosen such that equations (61) and (62) are satisfied, all of them must be positive, and $\alpha > \beta$.

Increasing ITER also increases the running time. For ITER = 10 shakes, running time is approximately 180 seconds on the CDC 6600 computer. For ITER = 100 shakes, running time is approximately 340 seconds. A good compromise, which gives nice looking plots, is ITER = 20 shakes with a running time of approximately 200 seconds.

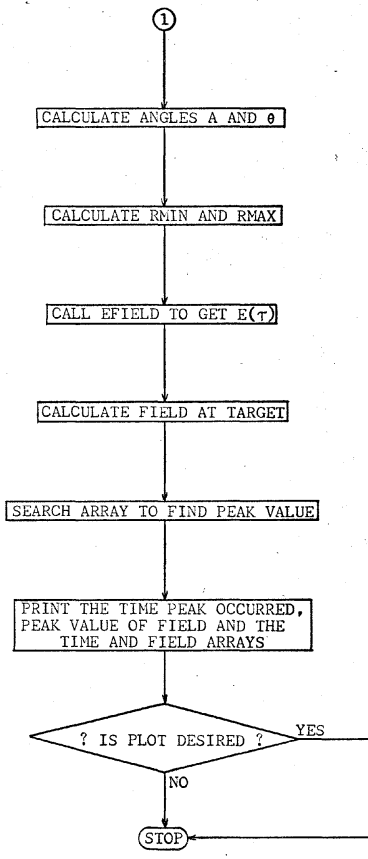
In binary form, the code requires 33000₈ words of core on the CDC 6600 computer.

Appendix B

EMP Code Flow Charts



PROGRAM CONTRL

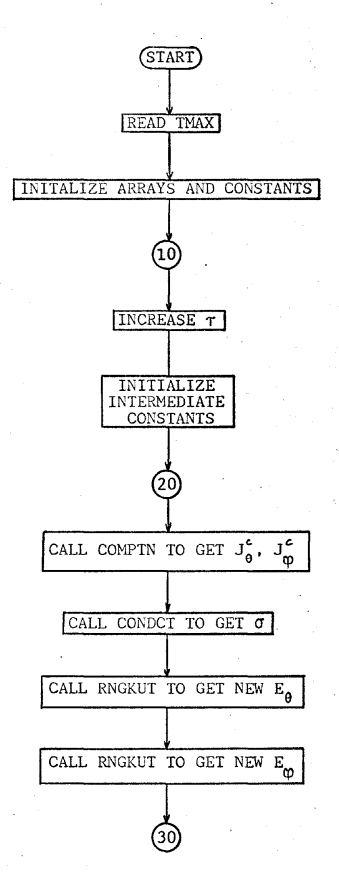


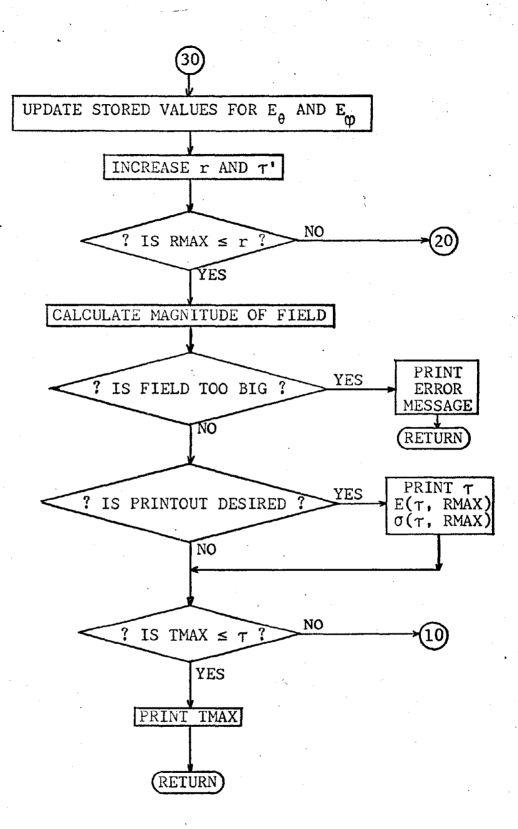
50

CALL

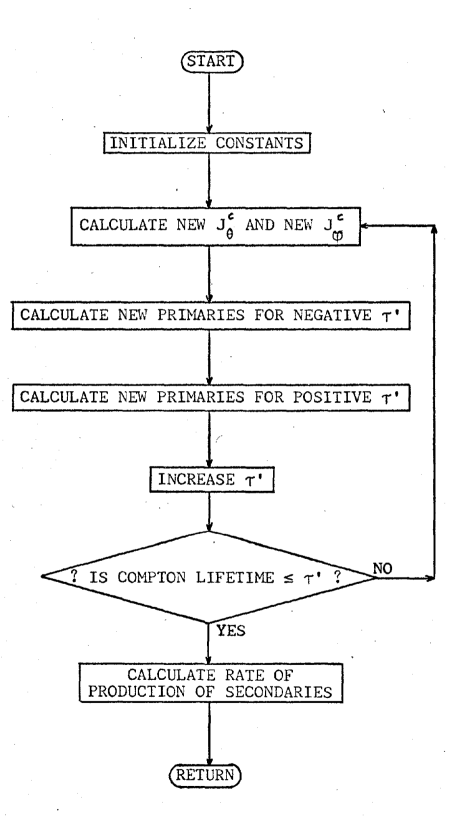
ELGPLT

SUBROUTINE EFIELD

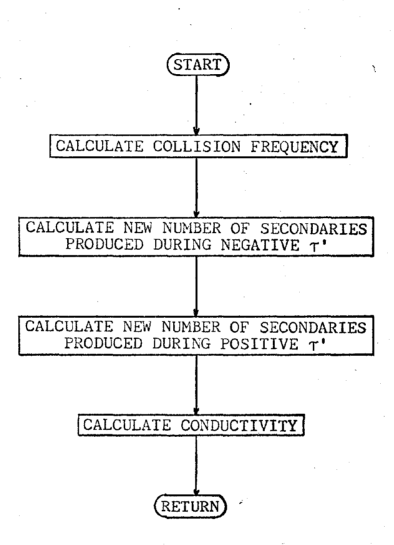




SUBROUTINE COMPTN

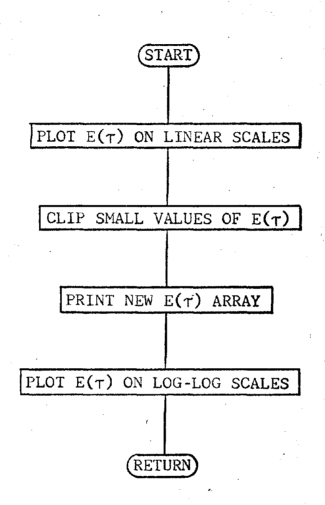


SUBROUTINE CONDCT

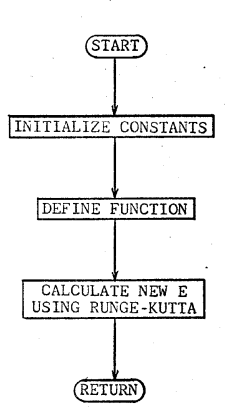


SUBROUTINE ELGPLT

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SUBROUTINE RNGKUT



Appendix C

EMP Code Listing

FROGRAM CONTRL (INPUT.CUTFLI.FLOT)			
, , , ,	CNTL		
THIS PROGRAM CONTROLS THE SUBROUTINES	Z		
	N1		
COMMON OUT		06.0	
FULL COL			
こう しきしつ うみ しょうそう しょうしき しょうき しこう	ż		
THE TARGET LCCATI	NI		
FOR THE NORTHERN HEVISPHE	z	0	
X IS MAGNETIC WEST V IS MACHETIC SCITU			
IS ALTITUDE	z	υM	
	N	1	
HOE IS HEIGHT CF BURST IN KILOMETERS > 50KM	N 1	5	
	Z	S	
GAMYLD IS GAMMA YIELC CF EURST IN KILCTCNS	z	\sim	
	N1	ω	
IS THE PAGNITUCE OF EARTHS PAGNETIC FIELD IN THE	Z	σ	
PSORPTION REGION PELCE THE BURST IN WEBERS/SQUARE MET	E :	•	
	z	 t	
BANGLE IS THE DIP ANGLE CF THE MAGNETIC FIELD IN DEGREES	Zī	\sim r	
CID TO THE NUMBER OF CIERS SETUES BUT AND DAY		0.5	
	- 1 Z 2	t 1	
<pre><=NUELK<=>0</pre>	- F Z 2	υu	
S THE OUTPUT CONTECL PARAMETE	ż		
T=O ==> PRINT PEPK VALLE AND ARRAY	L Z	œ	
T=1 ==> PRINT PEAK VALLE AND MAKE FLC	Ľ	σ	
2 ==> PRINT EVERYTFING AND MAKE FL	N	0	
T=3 ==> PRINT EVEFYTHIN	Z	-	
	Z	2	
EAD 1091,X,Y,Z,HOB,GAYYLC,EFJE	N1	cut	
(X*X+Y*Y+(HOE*10002)**2	Z1	4	
RINT 2006, GAMYLD, HOB, X, Y,	z	5	
	CNTL	9	

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TL 3	CNTL 380 CNTL 390 CNTL 400	NTL 41	NTL 43	NTL 44	NTL 40	NTL 47	NTL 49	NTL 50	NTL 51	NTL 52 NTL 53	NTL 54	NTL 55	NTL 56	NTL 57	NTL 59	NTL 60	NTL 61	NTL 63	NTL 64	NTL 65	NTL 66	NTL 67	NTL 68	N - L 0 0 N - L 0 0	NTI 74	TL 72	 - -
												•		NO													
SET UP DEFAULT VALUES	IF(BANGLE.EQ.0.) BANGLE=40. IF(BFIELC.EQ.0.) BFIELC=0.00002	F(NDELR.EQ.0) NDELR=5	CONVERT DATA TC MKS L'NITS	08=408 4100	AMYLD=2.61625E25*GAMY	BANGLE=0.017453295*BANGLE CMFGA=1.6F-10*AFTFLF//3.605*C.145-31)		PRINT TYFE OF CALCULATION		LCT=49000. Z.GT.REFLCT)PRINT 2	F(Z.LT.0.0) PRINT 2008	F(Z.LE.REFLCT.AND.Z.G		WAVE CALCULATION ASSUMES 100% REFLECTI TOPOD TWAGE SE TADGET DELOW COMMON	ET Z = -Z IF REFLECTEC VAVE IS TO EE USED		(Z.6T.REFLCT) Z=-Z	IF(Z.GT.FOB-1000.) FRINI ZUU/ IF(Z.GT.FOB-1000.) Z=-Z		DETERMINE ANGLES		=SQRT (X+X+Y+		いつう このつうち ーちー ロビ	DETERMINE DMIN DNAY		

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ar be	CNTL107 CNTL108
RAIN=5.E4 A=(ZRMN-H0E)/ RMIN=TA*X RMIN=TA*X RMAX=TA*X PIN=SGRT(XRMIN RMAX=Z F(Z.LT.2.E.E.) Z B=(ZRMAX-H0B)/ RMAX=TB*X RMAX=TB*X CALCULATE E ALL EFIELD(E,T CALCULATE E ALL EFIELD(E,T 100E2) CALCULATE E F(R.LE.RMAX) G CALCULATE E F(R.LE.RMAX) G CALCULATE E CALCULATE E F(R.LE.RMAX) G CALCULATE E CALCULATE E CALCULATE E F(R.LE.RMAX) G CALCULATE E CALCULATE E CALCULATE E F(R.LE.RMAX) G CALCULATE E CALCULATE E F(R.LE.RMAX) G CALCULATE E F(R.LE.RMAX) G CALCULATE E CALCULATE E F(R.LE.RMAX) G CALCULATE E F(R.LE.RMAX) G F(R.LE.RMAX)	PRINT 2010, TIME(IT) Print 2005, Big

RINT 200	NTL109
FRINT 2002, (TIME(I), I=1,15()	TL1
FINT 2003	NTL111
RINT 200	NTL112
	NTL113
IF DESIRED, MAKE PLCI	NTL114
	NTL115
F(OUT.LE.D.CR.OUT.GE.	NTL116
CALL ELGPLT (E,TIME, BIG)	NTL117
F	NTL118
1 FCRMAT (7F10.0,21	NTL119
CRMAT (//5X, "PEAK OCCUERED AT", F5. 1," SHAKE	NTL120
9 FCRMAT (5X,"DIRECT MAVE IS BEING CALCULATED"////	NTL121
8 FORMAT (FX, "REFLECTED WAVE IS BEING CALCULATED"///	NTL122
7 FCRMAT (5X,"TARGUT IS ABOVE AESCRPTION REGION SO	NTL123
EING CALCULATED"////	NTL124
CRMAT ("1 THE GLEST WITH GAMMA YIELD CF", 1PE10.	NTL125
/5×,"IS AT AN ALTITUCE CF", 1PE10.3," KILOMETE	NTL126
/5X,"THE TARGET IS AT CCORDINATES", 3(5X,1PE1	NTL127
/5X,"WHICH IS",1PF10.3," METERS FROM THE BU	NTL128
FCRMAT (//5X,* * * * * * * * * * * * * * * * * *	NTL129
1 /5X,"* PEAK EFIELC 21 TARGET IS ",1PE10.3," V	NTL130
5×°** * * * * * * * * * * * * * * * * *	NTL131
4 FCRMAT(19(10(3X,1PE10.3)/	NTL132
CRMAT (///5X,"EFIELD VALLE	NTL133
2 FCRMAT (19(10(4X, F5.1, 4X)/	NTL 134
1 FCRMAT ("1 TIMES USED (IN	NTL135
Z	NTL13

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hi hi. Lithir	11 1 1 1 3 3	FLD 5	FL) ô	۲ 1 8 8	6 (1:	FLJ 10		FLJ 13	7.0 14	FLJ 15		FLD 17	7L) 18	FLO 19	FLJ 20	FLD 21	FLJ: 22	FL) 23		FLJ 25	1 2 2 2 2 2	72 0.1	FLJ 28	ت ال 29	FLJ 30	FLD 31		1 L) 00 1 L) 00 1 L) 00		FLJ 36
			•		TESER OUT	MMON OUT, AP, BP, RNP, TOP	T DUT/CJIJ-/0 F SJ/VHS NI NUTIVCJI BU JMIT SI GJI	CHANGE IT TO NJMBER OF TIME STEPS		100, ITER	E2=100+(ITER-10)				AD 101, AP, BP, RNP, I OP	R44T(4F10.0) ·	51 J=1,100	J)=0.0 \$ TIME(J)=0.1*J	NTINUE	71 J=101,190 //	J)=0.0 \$ TIME(J)=10.+(J-130.)		51 L=1,NDELR \$ STORE2(L)=0.		HE=0. \$ DELRN=NDE.? \$ T=0. \$ DF=1. \$ EPHI=0.	LTAR=(RMAX-RMIN)/JELRN \$ R=RMI4+DE_TAR \$ RNP=1.E+3*RNP		E LOOP IS FOR CALCULATION IN RETARDED TIME	NSIDE LOOP IS FOR INTESRATION IN & AT EACH TIME STEP	
1 SS S	ں ں ں	0		C) 12		•) (, c		X	₽ 3		c) i		N			1					C			0	0	s 0	0	G

410 33 410 33 410 33	2010000 2010000			С 53 54 54	1055 10780			 	LCC 680 LCC 680 LCC 710 LCC 710 Z20	•
	LE LE LE LE LE	u u u u	. LL. LL. LL	ևևև	_ LL_ LL_ LL_ L	سا سا س		ᆠᇉᄔ		
00 00 00 00 00 00 00 00 00 00 00 00 00	A A L L A L L A L L A L L A L L A L L A L L A L A L L A L	P=TP+CTP CNTINLF	FIND MAGNITUDE CF EFIELC E(I)=SGRT(ETHE**2+EPHI**2)	CHECK	IF(E(I).GT.1.E15) GC TO 52 IF (I.EG.100) DT=10. F=RMIN+DELTAR	IF DESIRED, PRINT OUTFLT	IF (OUT-1.LE.D) GO TO 21 PRINT 5,I,TIME(I),E(I),SIGMA 1 CONTINUE	PRINT MESSAGE AFTER TERMINATION OF TIME LOOP	2 FRINT 201,TIME(ITER) Return Print message After Afncemal termination of time Loop	
		3	00	ပပပ	> 0	000	. N	ა თ <i>ღ</i>	4	>

	• • •
730 750 750 750 750 750 8810 8820 8820 8820 8820 8820 8820 882	60000000000000000000000000000000000000
<pre>FRINT 301 FRINT 201,TIME(IT) IF(IT.LT.10) RETURN SET LAST 5 VALUES OF EFIELD TO 0.0 TC AVOID INCORRECT FEAK E(IT)=E(IT-1)=E(IT-2)=E(IT-3)=E(IT-4)=E(IT-5)=0.0 E(IT)=E(IT-1)=E(IT-3)=E(IT-4)=E(IT-5)=0.0 FCRMAT (I3) FCRMAT (I3) FCRMAT (I3) I =",I4," TIME =",F6.1," SHAKES E(T,RMAX) =", 1FE10.3," VOLTS/METER SIGMA =",1PE10.3," MHO/METER") I FCRMAT(//J5X,"ITERATION TERPINATED AFTER",F5.1," SHAKES"/) FCRMAT(//J5X,"ITERATION TERPINATED AFTER FF5.1," SHAKES"/) FCRMAT(//J5X,"ITERATION TERPINATED AFTER FF5.1," SHAKES"/) FCRMAT(//J5X,"ITERATION TERPINATED AFTER FF5.1,") FCRMATED AFTER FF5.1," FCRMATED AFTER FF5.1," FCRMATED AFTER</pre>	<pre>SUBROUTINE CCNDGT(SIGMA,FFI,CTP,DT,HOB,R,A,STORE1,STORE2,K,hCELR, IFPI2) CALCULATES SIGPA AFTEF FINDING NSECCNDARY FROM NPRIPARY SICRE1 CCNTAINS INTEGFAL FOR NEGATIVE TAU STCRE1 CCNTAINS INTEGFAL FOR NEGATIVE TAU STCRE2 CCNTAINS INTEGFAL FOR NOSTITIVE TAU CCLISN=4.E12*EXF((R*COS(A)-+CB)/7000.) STORE1=STOPE1-PRI*CT*(1.0E-9) STORE1=STOPE1-PRI*CT*(1.0E-9) STORE2(K)=STCRE2(K)+PRI2*CT*(1.0E-9) STORE2(K)=STORE1 STORE1=STORE1 STORE2(K)=STORE1 STORE2(K)=STORE1 STORE1=STORE1 STORE2(K)=STORE1 STORE1=STOPE1-PRI*CT*(1.0E-9) STORE2(K)=STORE2(K)+PRI2*CT*(1.0E-9) STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STORE1 STORE2(K)=STORE1 STOR</pre>
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SUBROUTINE CCMPTN(JTHETA, JFHI,T,R,A,THETA,CMEGA,HOB,GAMYLC,TF,FRI, 1PRI2)	τ Σ Σ	10
	2	
ALCULATE THE TWO CCPFCNENTS	Σ	
CURPENT AT GIVEN T AND R	ž	
ALCULATE NUMBER OF FRIMARY	μ	
	Å	
THETA IS THETA COMPCHENT CF COMPTCN	Σ	
PHI IS FHI COMPONENT CF COMPTON CURREN	μ	
MAX IS COMFTON LIFEIIPE	Ē	0
ATH IS ALTITUCE SCALFC COM	μ	-
IS NEGATIVE FETARCEC	Ξ	2
IS POSITIVE RETARCEE TIME	ž	3
RI IS NUMBER OF PRIMARY ELECTRONS GENERATED DURING	Ч	t.
RIZ IS NUMBER CF FRIMARY ELECTRONS GENERATED DURING	μ	5
PRIME IS VARIAFLE CF INTEGRATI	Σ	9
	۲ ۲	2
INITIALIZE CONSTANTS	Σ	Ø
	ž	σ
AL JIHETA, JPHI	μ	
JPHI=0. \$ TFRIPE=(5.E-9) \$ TMAX=CLIFE(R,A,HOB) \$FRI=0	μ	
=TMAX/10.\$ PATH=309.*EXF((FOB-R*COS(A))/7000.) \$TPRIME=01\$W=	μ	\sim
I2 =	Σ	M
	Σ	4
RUNGE-KUTTA INTEGRATICN CF COMPTON CURRENT	Ξ	മ
) 2	0
0 31 k=1,10	⊢ Σ	~
K1=DT*CMTHET(HOB, R, A, THETA, CMEGA, PATH, T, TFRIME, GAMYLD)		CC (
K2=DT*CMTHET(HOB,R,A,TFEJP,CMEGA,PATH,T,TFRIME+W,GAMY	E I	σ
K3=RK2	μ Σ	0
K4=DT *CMTHET (HOB * R)		
HE A=J HE A+(PK1+2。+ (FK2+FK3)+FK4)/6。 ///////////////////////////////////	= ;	NI
KS=U *UYPHI (HUB9K9A)HELP9CHEGA9PAHA) 9 FKLME9G Ke-dikomput (uco o a turta cmrca datu i trotusiji	- +	n .:
	- +- : 2	t 13
RK3=DT*CPPHI (HOB, R,A,THETA,CMEGA,PATH,T,TPRIME+DT,GAMYLD)	- }	360

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C MTN C MTN C MTN 400

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440 450 460 470

JPHI=JPHI+(RK5+2.*(RK6+RK7)+RK8)/6. RUNGE-KUTTA INTEGRATION OF FRIMARIES RKP1=DT*RKCMTN(R,THETA,OMEGA,TP,TPRIME) RKP2=DT*RKCMTN(R,THETA,OMEGA,TP,TPRIME+W) RKP2=DT*RKCMTN(R,THETA,OMEGA,TP,TPRIME+W) RKP4=DT*RKCMTN(R,THETA,OMEGA,TP,TPRIME+DT) PRI=PRI+(RKP1+2.*(RKP2+RKP3)+RKP4)/6. RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+W) RP2=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+W) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+W) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+W) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+W) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=PTRIME+DT RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=PTRIME+DT RP1=PTRIME+DT RP1=PTRIME+DT RP1=PTRIME+DT RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=PTRIME+DT RP1=PTRIME+DT RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=PTRIME+DT RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=PTRIME+DT RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=TRRIME+DT RP1=DT*RKCMTN(R,THETA,OMEGA,T,TPRIME+DT) RP1=TRRIME+DT RP1=TRRIME+

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F + 10	FF102 FF103 FF104	CMFF1060 CMFF1060 CMFF1070 CMFF1080 CMFF1080 CMFF1090			·	TH101 TH102 TH102 TH103	CMTF1050 CMTF1060 CMTF1070 CMTF1080 CMTF1090 CMTF1090
		•					•
FUNCTION CMPHI(HOB,P,A,THETP,OMEGA,PATH,T,TPRIME,GAMYLD)	CALCULATES F(T,P) FOF RUNGE-KUTTA INTEGRATION OF PHI COMPONENT OF COMPION CURRENT	SCLVE=TOFT(T.TPRIME,THETA,CPEGA) SCLVE=FOFT(SCLVE) SCLVE=FOFT(SCLVE) SOLVE=SOLVE*(-4.608E-11)*GCFF(R,A,HOB,FATF,GAMYLD) CMPHI=SOLVE*SIN(THETA)*SIN(CPEGA*TPRIME) RETURN \$ END				FUNCTICN CMTHET(HOE,R,A,THETP,OMEGA,PATH,T,TPRIME,GAMYLD) Calculates F(T,F) FCF FLNGE-KUTTA INTEGRATION OF THETA COMPONENT CF CCMFTON CURRENT	SOLVE=TOFT(T,TPRIME,THETA,CMEGA) SCLVE=FUFT(SOLVE) SCLVE=FUFT(SOLVE) SOLVE=SOLVE*(-4.608E-11)*GCFF(R,A,HOB,FATH,GAMYLD) CMTHET=SCLVE*SIN(THETA)*CCS(THETA)*(COS(CMEGA*TPRIME)-1.) RETURN \$ END
•	0000	2				0000)
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SUBROUTINE RNGKUT (E1,E,F,F,SIGMA,COMPTJ)	RNK11010
U)	FNK11030
THE RUNGE-KUTT	FNK71040
	RNKT1050
CATA (C=3.0E8),(RMLC=12.56637E+7)	FNK11060
FUN (R	FNK11070
K1=H*EFUN(P,E)	FNK11080
H+A) ND:	RNK71090
K3=H*E	FNKT1100
K+=H*EFUN (R+H	RNKT1110
1=E+(RK1+2.*()	FNK11120
ETURN \$ E	FNKT1130
	•

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RKC+1020 RKC+1030 RKC+1030 RKC+1030 FKC 1060 FKC 1070 FKC 1080 FKC 1010

CALCULATES F(T) FOR RUNGE-KUTTA INTEGRATION OF FRIMARY ELECTRONS

FUNCTION RKOMIN(R, THETA, CREGA, TP, TPRIME)

SCLVE=TOFT(TF,TPRIME,THETP,CMEGA) RKCMTN=FCFT(SOLVE) RETURN \$ END

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FLNCTICN GOFF (R,A,HOF,FATF,GAMYLD)	GCFF1010
	GOFF1020
SOLVES VIRGIN TRANSFCRI DND USES REACTION RATE TO	6CFF1030
CALCULATE THE NUMBER CENSITY OF RACIAL ELECTRONS	GCFF1040
	COFF1050
SCLVE= (.0226275/COS(A))*(-1.+EXP(R*COS(A)/7000.))*EXP(-HOE/7000.)	CCFF1060
DENOM=12.556537*R*PATH*1.5	GOFR1070
GCFR=EXP (-SOLVE) *GAMYLD/CENCP	GCFF1080
RETURN \$ END	GOF F1090

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CLIF1010 CLIF1020 CLIF1030 CLIF1040 CLIF1050 CLIF1050 CLIF1050 CLIF1060 CLIF1090

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CALCULATES COMPTON LIFETIME AT RADIUS = P MAX ACCEFTABLE LIFETIPE = 100 SHAKES FCR THE KARZAS-LATTER PIGH FREQUENCY AFFRCX FE=1.041667E-8*EXP((PCE-R*COS(A))/7000.)

FUNCTION CLIFE (R, A, HOB)

00000

CLIFE=1.041667E-8*EXP((FCE-R*COS(A))/7000.) IF(CLIFE.6T.1.E-6) CLIFE=(1.E-6) Return \$ END

FUNCTION TOFT (T,TPRIME,THETA, OMEGA)		FT 10
T(T) IS TIME TRANSFORMED TO KIRZAS-LATTER FORM		
=0.358		FT 10
FIRST=T-(1B*(COS(THETA)**2))*TP2IME SECOND=B*(SIN(THETA)**2)*SIN(OME3A*TPRIME)/04E3A	· ·	FT 10 FT 10
JFT=FIRST+S		FT108
RETJRN & END		FT10
	,	

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FUNDTION FOFT(T) F(T) IS THE

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F(T) IS THE POMRANNING MODEL FOR TIME DEPENDENCE Df nuclear meapin yield in retarded time

INTESER OUT . 20MMON OUT,A,B,RN,TD TSHAKE=1.E8*T DENOM=(B+A*EXP((A+B)*(TSHAKE-TO)))*RN FOFT=(A+B)*EXP(A*(TSHAKE-TO))/JENOM RETJRN \$ END

FJFT1100 FJFT1100

FJF1093

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	SUBROUTINE ELGPLT(EFLOT, TIME, BIG)			
	CO DELIGE TO DIFERS OF SECTOR DELIGION OF FUCEFU DE			
່ວເ	THE FIXAT ZU SHAKES (F E 1) IS FLUTIEU UN LINEAR SUAL	2		
2	IMENSION EPLOT (19		 	1 5
	5 \$ SMALL = 0.0	•	a	
	HECK = 0.0	•	٩	
	0 6 I=1,11			
	(I) = EFLCT		u	
9	CNTINU			C
	ALL PLOT (0		o.	-
	ALL PLOT (2.		Ω.	N
	ALL SCAL		D.	\mathbf{m}
	ALL SCALE (Ω.	4
	ALL LINE (T .		a.	5
	ALL AXIS(0.,0.,13HTIME (SHAKES),-13,5.,0.,T (111),	(112))	α.	9
	ALL AXIS(C.,0.,12HEFIELE (V/M),12,5.,50.,E (1	(112)	Ľ	~
	ALL AXIS(0.,5.,2H ,2,5.,0.,T (111),T (112)		Ъ	8
	ALL AXIS(5.,0.,2H ,-2,5.,9		Q_	σ
	ALL FLOT(10.0,		с Б	С
с С			L L	44
с С	VALUES OF E(T) AFE C		<u>с</u>	2
ပ	NC E(T) IS PLOTTED CA LCG-LOG SCAL		1	3
с С			Ц Ц	÷.
	0 3 I=1,19		Ľ	5
	F((CPLCI(I)/PIG).LI.(10.**(+MAG)		Ц	Q
~	FLOT(I)=BIG*(10.**(-MAG)) 2 CH		ц Б	~
m	CNT TNU		ر م	Ø
	F COMECK.		ر. س	ഗ
	RINT 2005		٦	0
	PINT 2006, (EPLOT(I)		Ľ	
	MALL=EIG* (10.** (-MAG		a	N
t t	CNTINU	•	o.	4
	ALL PLOT (0.		а.	ណ
	ALL FLOT		О.	Q
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Ц Ц Ц	п Г Г	EPL	0	0	0	\mathbf{O}	0	n	е С	Ц П П П	3EPL	ц П Ц	БРГ
			((2))	$\widehat{\mathbf{a}}$					(/*		•		
			Π []	01 0					* *		IS", 1FE1		
			VII,	, EP L		~			PPE0		IS.	.•	
			.91)	T (191), EPLCT (19	_	T (192)			CLIP		E(T)		•.
			IME (1	0T (1	92))	0T (1			z	2	MUM E		
			F	, EPLO		EPLO.			111	-	MIXAN		
				90.	31), TIME	191),			HA I	LPE1	2		
			13,	۰ ۵	0., TIME (191), T	01 (:			ECI	ax •	.0.3/"		
		6	-, (S	,12	IMF (, EPL			<u>о</u> F	(10(E10.		
		6 0 1 1	(SFAKES),	SV/M	1. • · I	.06.			:101	1/19	., 1P		
~	0	- 0 J	5	L L		• ມາ				: 	S H		
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Vita

Terry C. Chapman was born on 21 August 1943 in Vancouver, Washington. He graduated from high school in Manitou Springs, Colorado in 1961. He attended the University of Colorado in Boulder where he was elected to Tau Beta Pi, Engineering Honor Society and Sigma Pi Sigma, Physics Honor Society. He received the degree Bachelor of Science, Engineering Physics and a commission in the U. S. Air Force from the University of Colorado in 1969. After attending communications training, he was assigned to Kelly AFB, Texas as a communications operations officer in 1970. He entered the Air Force Institute of Technology in 1972.

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This thesis was typed by Ladonna Stitzel.