Three-Dimensional Simulation of Multistage Depressed Collectors on Micro-Computers

Lalit Kumar, P. Spätzke, R. G. Carter, and D. Perring

Abstract—A three-dimensional (3-D) package for simulation of asymmetric and crossed-field multistage depressed collectors for microwave tubes has been developed. This package is based upon the 3-D finite-difference code KOBRA3-INP. The main features of the package are a user-friendly input interface, post-processors for collector analysis and calculation of secondary electron trajectories, and versatile output graphics. Both PC and mainframe versions of the package have been developed. The results of simple benchmark tests and those of simulation and analysis of asymmetric and crossed-field collectors including the effects of secondary electrons are presented. It is found that the asymmetric hyperbolic electric field collector shows very low backstreaming. It is shown that the representation of trajectories in energy space gives a better insight into the behavior of individual trajectories than plotting in coordinate space. The package will be useful for designing novel types of depressed collector.

I. INTRODUCTION

MULTISTAGE depressed collectors are widely used in microwave linear beam tubes [1], high power gyrotron devices [2], and particle accelerators [3] to recover the energy of the spent charged particle beam and thus enhance the overall efficiency of the device. The use of depressed collectors also has the additional advantages of reducing the collector cooling requirements in high power devices and the X-rays generated in high voltage devices. In some systems in which the overall power consumption is a primary concern, such as satellite communications, mobile transmitters, large TV transmitters, and high energy accelerators, a 100 percent recovery of the spent beam energy would be highly desirable. Significant advances have therefore been made in the design of depressed collectors to approach this goal. Detailed reviews of the development of depressed collectors for microwave tube have been made by Kosmahl [1] and Kumar and Carter [4].

Among all the types of collector geometry used so far the axi-symmetric geometries developed by Kosmahl and his coworkers at NASA [5], [6] and Neugebauer and Mihran at the General Electric Company [7] have been most widely investigated and used in the industry. This is because they are simple to construct and do not require a magnetic field. These collectors have achieved efficiencies in excess of 85% in travelling-wave tubes (TWT's) with basic efficiencies of about 17% [6]. This performance was achieved as a result of major and consistent efforts devoted to collector development and the optimization of their design using 2.5-D computer simulations [6]. Backstreaming of electrons into the circuit region is inherent in the design of depressed collectors [5]. While this may be tolerated to some extent in ECM tubes it is highly undesirable in radar and communication tubes because it degrades the tube performance and may cause oscillations. Some other types of collectors including the hyperbolic electric field (HEF) collector with off-axis electron beam [8], the tilted electric field (TEF) collector [9], and crossed-field collectors [10], [11] have demonstrated reasonably high efficiencies and promise of reduced backstreaming. Further work on the optimization of these designs has been hindered hitherto by the lack of analytical techniques or computer codes suitable for modeling them. A fully three-dimensional electrostatic code with the facility of electron trajectory tracing in the presence of an external magnetic field is required for simulating asymmetric and crossed-field collectors. A facility for electron trajectory sorting and collector efficiency calculations is also necessary for a collector simulation code. Secondary electrons play a significant role in determining the performance of a collector so provision for calculation of their trajectories is also desirable.

To satisfy this requirement, a 3-D package for simulating depressed collectors has been developed at Lancaster University under an ESTEC project [12], [13]. Results of simple benchmark test runs of the package have demonstrated its accuracy for depressed collector problems. The usefulness of the package for asymmetric and crossed-field collectors has been shown by the results of simulations of Hechtel’s hyperbolic electric field (HEF) collector [8] and of a tilted electric field (TEF) collector with a transverse magnetic field for which some experimental data was available.

II. DEVELOPMENT OF THE 3-D PACKAGE LKOBRA

The collector simulation package (LKOBRA) is based upon the 3-D code KOBRA3 which was originally written for the
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simulation of ion sources [14]. LKOBRA incorporates the IBM-PC version, KOBRA3-INP\(^1\), unchanged [15]. Upgrades of KOBRA3-INP can thus be readily substituted for the version (1.13) which was used for the simulations reported here. Some improvements in KOBRA3-INP were made by INP to make it more suitable for collector simulation on the basis of feedback from the work carried out at Lancaster.

LKOBRA also contains pre- and post-processors which have been developed to facilitate interactive creation of input data files, collector analysis, and calculation of secondary electron effects. LKOBRA runs interactively on an IBM-PC or compatible system with at least 640 kbytes of memory. The disk storage requirement of the package is about 1.5 Mbytes and a typical problem may require about 5 Mbytes of disk space.

For large problems the extended-DOS, WINDOWS, or mainframe versions of KOBRA3 can be used. The output graphics of the mainframe version are inferior to those of the PC version but LKOBRA can be used on a PC for pre- and post-processing the data of problems which are run using the mainframe version. The main segments of the package are described below.

A. Input Interface

The LKOBRA input interface facilitates the interactive creation of the input data file for a given problem in the format required by KOBRA3 [15]. Its main features are a set of pre-defined shapes for generating plane-by-plane cross-sectional data for asymmetric geometries and a beam model for generating the starting conditions of the trajectories at the collector entrance. It also has conversion of units from millimeters to meters and degrees to radians built into it. The basic shapes which have been pre-defined in this program are shown in Fig. 1. It is possible to combine these shapes to define more complex shapes and to cut holes through some of them. The beam model in the interface is useful for generating sensible initial conditions for the trajectories when 3-D spent-beam data is not available. However, if 3-D spent-beam data is available from the output of a large-signal analysis program or from experimental measurements, then the starting conditions of the trajectories can be added to the input data file by the user. For instance the spent-beam output of the 2.5 D large-signal program [16] can be directly appended to the input data file. A beam of arbitrary cross-section can also be simulated by choosing appropriate starting positions of the beamlets.

The beam model requires as basic data of the electron beam its radius, current, and voltage, an estimated kinetic energy distribution, and the maximum angular spread of the beam. It assumes the initial beam cross section to be circular and divides it into a number of rings. Each ring is then divided into a number of sectors such that the areas of all the sectors in the beam cross section are equal. The current density in the beam is assumed to be uniform and thus all the sectors represent equal amounts of current. The given spent-beam energy distribution is mapped in a pseudo-random manner onto the sectors so that the number of sectors to which a particular energy is assigned is proportional to the percentage of the total current in this energy range. A specified number of trajectories can be started from the centre of each sector with their initial angles distributed randomly within a given range \(-\theta\) to \(+\theta\). This randomization avoids, to some extent, a preferential distribution of energy or starting angles.

B. The Mesh Generator

The KOBRA3-INP mesh generator discretises the three-dimensional box containing the problem geometry into cubical elements. In the DOS version, the mesh array size is limited to 16,000 mesh points but in the extended DOS version up to 120,000 mesh points can be used on a machine with 5 Mb of random access memory. Arbitrary electrode shapes can be simulated but the feature size is limited by the mesh spacing used. The coordinates of the corners of the polygonal cross section of the electrodes in each mesh plane must be provided for arbitrary shapes. Simple rectangular or circular geometries can be input using geometrical data. Local mesh refinement is possible in all three coordinate directions in any desired mesh interval. Neumann (open circuit) boundaries are allowed only on the sides of the problem box.

C. The Poisson Solver

The KOBRA3-INP Poisson solver uses a seven-point finite difference (FD) method for the solution of Poisson's equation

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\(^1\)KOBRA3-INP is distributed by INP, Junkernstrasse 99, 62 Wiesbaden, Germany. Copies of LKOBRA can be obtained (subject to ESTEC approval) from Dr. R. G. Carter at Lancaster University.
in three dimensions. In this method the potential at any mesh node is given, in terms of the potentials at the six neighboring nodes in the coordinate directions, by the expression

\[
\phi_i = \left( \frac{h_1 \phi_1 + h_2 \phi_2}{h_1 h_2 (h_1 + h_2)} + \frac{h_3 \phi_3 + h_4 \phi_4}{h_3 h_4 (h_3 + h_4)} + \frac{h_5 \phi_5 + h_6 \phi_6}{h_5 h_6 (h_5 + h_6)} - \phi_0 \right) \left( \frac{1}{h_1 h_2} + \frac{1}{h_3 h_4} + \frac{1}{h_5 h_6} \right)
\]

where \( h_i \) is the distance of the \( i \)-th node from the central node and \( \phi_i \) is the potential at the \( i \)-th node (see Fig. 2). Thus all the six nodes are allowed to have different distances from the central node. Where an electrode intersects the mesh line joining the \( i \)-th node to the central node \( \phi_i \) is taken to be the electrode potential and \( h_i \) the distance of the point of intersection from the central node. The accuracy of this method is of the order of \((h_{\text{max}})^2\), where \( h_{\text{max}} \) is the width of the coarsest mesh. The space charge term on the right hand side is determined by tracing the charged-particle trajectories. The system of FD-equations for all the internal nodes of the problem is solved using an iterative procedure. The successive over-relaxation (SOR) method is used to speed up the convergence of the solution.

D. The Trajectory Solver

The KOBRA3-INP trajectory solver solves the relativistic Lorentz force equation for charged particles in electric and magnetic fields

\[
\frac{d\vec{v}}{dt} = \frac{q}{m} \left( \vec{E} + \left( \frac{\vec{v} \cdot \vec{E}}{c^2} \right) \vec{v} + \vec{v} \times \vec{B} \right).
\]

The starting velocity of a trajectory can be specified by all the velocity components in m/sec or any one component by the energy in keV and the other two components by the angle with the axis or by the emittance of the beam. The position, velocity components and total energy of each particle at each point of its trajectory are stored in a data file.

E. Post-Processor for Collector Analysis

The LKOBRA post-processor sorts out the trajectories landing on different electrodes and the backstreaming trajectories. It also calculates the backstreaming power, the power dissipated and recovered at each collector stage and the collector efficiency. KOBRA3 stores the Hamiltonian (total energy) at each trajectory point in the trajectory data file. The Hamiltonian and the velocity components of the trajectory at its starting- and end-points are used to determine the input power carried by the trajectory, and the power dissipated by it on the electrode at its end-point. The total power entering the collector \( (P_{\text{ent}}) \) is calculated from

\[
P_{\text{ent}} = \sum_{i=1}^{N_t} I_i V_i
\]

where \( V_i \) is the voltage corresponding to the initial kinetic energy of the \( i \)-th trajectory, \( I_i \) the current associated with the \( i \)-th trajectory, and \( N_t \) the total number of trajectories. The power \( P_j \) recovered at the \( j \)-th collector stage is determined by

\[
P_j = -V_j I_j M_j
\]

where \( V_j \) is the potential of the \( j \)-th electrode with respect to ground and \( M_j \) is the number of trajectories arriving at this electrode. The total power recovered \( (P_{\text{rec}}) \) is calculated by summing the powers recovered \( (P_j) \) at all the collector stages. The collector efficiency \( \eta_c \) is found by using the simple relation

\[
\eta_c = \left( \frac{P_{\text{rec}}}{P_{\text{ent}}} \right) \times 100%.
\]

Backstreaming trajectories are identified by the positions of their end-points which lie at or before the entrance plane of the collector.

F. Post-Processor for Secondary Electron Effects

The LKOBRA post-processor facilitates the automatic calculation of secondary electron trajectories. It determines the starting conditions of a secondary trajectory for each primary trajectory and writes it to the input data file. All trajectories are assumed to carry equal amounts of current. The current density distribution is therefore represented by the distribution of the trajectories. Trajectories of multiple species of particles can be calculated simultaneously. The trajectory equations are solved using a method similar to the Runge-Kutta method with step size control. The maximum number of trajectories is 30,000 but this may be limited by the disk space available for storing the trajectory data.

The starting velocity of a trajectory can be specified by all the velocity components in m/sec or any one component by the energy in keV and the other two components by the angle with the axis or by the emittance of the beam. The position, velocity components and total energy of each particle at each point of its trajectory are stored in a data file.
secondary trajectories are then traced using the KOBRA3-INP trajectory solver. The starting points of the secondary trajectories are located at the end-points of the primary trajectories. This is not likely to introduce any significant error since these end-points are generally close to the electrode surfaces. A total kinetic energy of 50 eV is assigned to each slow secondary electron which is generally considered as the maximum energy of true secondary electrons. The initial direction of the secondaries is assumed to be normal to the electrode surface since slow electrons are preferentially emitted in that direction. The normal to the electrode surface is determined using the coordinates of the trajectory end-point and the mesh information. In the present version it is not possible to include the space-charge of the secondaries. This is because the secondary electron trajectories are calculated separately from the primary trajectories and KOBRA3 deletes the space-charge of the previous iteration on restarting the trajectory calculations. It is also not possible to calculate the secondary trajectories simultaneously with the primaries because KOBRA3 does not allow trajectories to have different currents. The influence of the space-charge of the primaries will, however, be present in the potential distribution which is used to determine the electric field for the secondary electron trajectory calculations. Further improvements in KOBRA3 will be carried out to overcome these problems.

G. Post-Processor for Resetting Primary Trajectories

An option has been provided in LKOBRA for restoring all the data related to the primary trajectories after the secondary electron effects processor has been run. However, the secondary electron data is lost on running this processor.

H. Program Output

KOBRA3-INP provides printed and graphical output in forms which are useful for collector simulations. The 3-D display of electrode geometry, mesh and trajectories individually or together can be viewed from any angle. The geometry and mesh can be displayed in any particular mesh plane or between any two mesh planes. The trajectories can be sorted according to ranges of serial number, initial energy, charge or mass of the particles. 3-D plots of potential surfaces in any mesh plane are available. A 2-D plot option provides plots of geometry, equipotentials and electric field components in any mesh plane. It is also possible to display curves of potential, total energy and kinetic energy components (individual or combined) versus trajectory path length. Sorting of trajectories can also be done in these plots in the same way as in the 3-D plots. Such plots provide very useful insights into the behavior of trajectories in different energy ranges.

The various KOBRA3 program units interact with other units through disk files. This modular structure made it possible for us to develop LKOBRA as an extension of KOBRA3 and made it convenient to control the flow of the program interactively. The package can also be run in batch mode for a given number of potential and trajectory calculation cycles.

III. VALIDATION OF THE PACKAGE

To validate the package and test its performance for 3-D depressed collector simulation, some simple benchmark tests were performed. The benchmark tests were a beam spread test, to check the accuracy of the space-charge field calculation, and an electron spectrometer test to check the trajectory calculation in given electro- and magneto-static fields. The results of the benchmark tests are discussed in the following sections.

A. Beam Spread Test

The expansion of a cylindrical electron beam in a field-free region under the influence of space-charge forces is given by the universal beam spread curve [17]. According to this a uniform electron beam of radius \( r_0 \) doubles its radius in a distance

\[
L = 0.01207 \frac{r_0}{(\mu_{perv})^{0.5}}
\]

where \( \mu_{perv} \) is the micropervance of the beam. The beam spread test was performed for a beam 1 mm in radius carrying a current of 100 mA at 1 kV (3.1623 \( \mu_{perv} \)). To simulate this problem 100 trajectories representing an electron beam with the above parameters were started at one end of a box with a square cross section. The trajectories were distributed uniformly over the beam cross section. The axis of the beam coincided with the axis of the box. The potential on all sides of the box was held constant at 1 kV. The distance at which the beam radius became twice its initial value was found to be 6.67 mm which agrees well with the theoretical value of 6.78 mm given by (6).

B. Electron Spectrometer Test

The accuracy of the trajectory calculation in itself and in accounting for the bending effect of a magnetic field on an electron can be checked by a very simple electron spectrometer test. In this test a number of trajectories with energies in the range of 0.6–1.3 kV in steps of 0.1 kV were started from a single point in the \( z \)-direction in a 14.5 cm cube having a constant potential throughout and a magnetic field of 9.43 Gauss in the \( z \)-direction. The agreement between the theoretical and computed values of the \( z \)-position of the end-points of the trajectories was found to be better than 1%.

IV. SIMULATION OF AN ASYMMETRIC HYPERBOLIC ELECTRIC FIELD (HEF) COLLECTOR

A four-stage HEF collector, similar to one described by Hechtel [8], was simulated in order to study its general behavior and to demonstrate the capabilities of LKOBRA. In Hechtel's collector the electrodes were hyperbolic in shape and the electron beam was launched off-axis. For the simulation we approximated the hyperbolic shapes of the electrodes by cones with a total collector length of 31.8 mm. The beam
Table I

<table>
<thead>
<tr>
<th>V/V₀</th>
<th>0.35</th>
<th>0.65</th>
<th>0.75</th>
<th>0.85</th>
<th>0.95</th>
<th>1.05</th>
<th>1.15</th>
</tr>
</thead>
<tbody>
<tr>
<td>H/V/V₀</td>
<td>0.107</td>
<td>0.271</td>
<td>0.373</td>
<td>0.150</td>
<td>0.017</td>
<td>0.046</td>
<td>0.056</td>
</tr>
<tr>
<td>N</td>
<td>20</td>
<td>55</td>
<td>75</td>
<td>50</td>
<td>5</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>

voltage was 24.54 kV and the beam current 3.43 A, which gives a beam perveance of 0.892 µperv. The potentials of the four collector stages were 24.54 kV (body), 12.54 kV, 6.54 kV, and 0.54 kV. The spent beam energy distribution used is shown in Fig. 3 (curve b) and the beam diameter was 1.4 mm. The starting conditions of the trajectories representing the electron beam were generated using the beam model described in Section II-A. The beam cross section was divided into 41 segments of equal area. Five trajectories were started from the center of each segment with their incident angles randomly distributed in the range −10° to +10°. The total number of trajectories was 205, and these were distributed among various energy ranges by the program (Table I). A self-consistent solution of the problem was obtained in five iterations. A tolerance of 0.1% in collector efficiency was used as a criterion for self-consistency. The average run-time of this problem on an IBM compatible microcomputer (33 MHz, 80386 processor with 80387 Maths co-processor) running under MS-DOS was 105 sec. for the potential solution (50 iterations) and 27 min. for the trajectory calculation for each iteration. The total disk memory requirement for this problem was 10 Mbytes. The complete details of the simulation are given in [13] and [18].

Plots of the electron trajectories in the HEF collector are shown in Fig. 4. Note that this figure and all subsequent figures were produced using the output facilities of KOBRA3-INP. An alternative representation of trajectories is available in KOBRA3-INP in which any component of the energy, the total energy, or the potential along the trajectory can be plotted as a function of its path length. Fig. 5 shows the axial and transverse kinetic energy components and the potential energy versus the path length of the trajectories. In the plot of axial energy versus path length (Fig. 5a) the continuously falling curves correspond to trajectories which hit the electrodes directly. The curves having well defined minima correspond to those trajectories which turn back after attaining zero axial velocity and then possibly land on an electrode. The height of a curve after the minimum indicates the axial kinetic energy which will be dissipated by this trajectory. This height should be minimized for all trajectories to achieve the greatest collector efficiency. The spread in the initial kinetic energy of the trajectories belonging to a particular kinetic energy class occurs because part of the total kinetic energy is associated with transverse velocity components. Two trajectories around 23 kV have very short path lengths because they are being intercepted by the body electrode.

The maximum transverse kinetic energy in Fig. 5(b) is 2 kV which is about 8% of the highest beam energy. Fig. 5(c) is a plot of the potential versus the path length of the trajectories. The end-points of the curves in this figure indicate approximately the potentials of the electrodes on which the electrons are collected. The curves in this plot achieving the highest (body) potential indicate the electrons reaching the body electrode or backstreaming. As in this case no backstreaming trajectories are visible in the trajectory plots, these curves show the trajectories reaching the body electrode only. In this plot also as in the plot of axial kinetic energy (Fig. 5a), the electrons hitting the electrodes directly and those landing after undergoing a reflection can be identified.

Table II shows the results of the computer analysis of the HEF collector. These results demonstrate the capabilities of LKOBRA for the simulation of depressed collectors of this type.

A. Secondary Electron Effects

To study the effects of secondary electron emission in the HEF collector the slow secondary electron trajectories were also simulated. A 3-D plot of the secondary electron trajectories in this collector is shown in Fig. 6. This shows the trajectories originating from the lower potential electrodes

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Fig. 3. Spent beam energy distribution (a) calculated for a high power TWT by a ring model [8], and (b) used for simulation of Hechtel’s HEF collector.

Fig. 4. 3-D representation of the electrode geometry and trajectories in the HEF collector.
Fig. 5. Energies of 205 trajectories in the HEF collector plotted against distance along each trajectory: (a) axial kinetic energy, (b) transverse kinetic energy, and (c) potential energy.

Table II

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Computed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power entering the collector</td>
<td>63.65 kW</td>
</tr>
<tr>
<td>Current at stage 1 (body)</td>
<td>0.25 A</td>
</tr>
<tr>
<td>Current at stage 2</td>
<td>2.27 A</td>
</tr>
<tr>
<td>Current at stage 3</td>
<td>0.75 A</td>
</tr>
<tr>
<td>Current at stage 4</td>
<td>0.20 A</td>
</tr>
<tr>
<td>Power recovered at stage 1</td>
<td>0.6 kW</td>
</tr>
<tr>
<td>Power recovered at stage 2</td>
<td>26.7 kW</td>
</tr>
<tr>
<td>Power recovered at stage 3</td>
<td>13.6 kW</td>
</tr>
<tr>
<td>Power recovered at stage 4</td>
<td>4.8 kW</td>
</tr>
<tr>
<td>Power dissipated at stage 1</td>
<td>4.3 kW</td>
</tr>
<tr>
<td>Power dissipated at stage 2</td>
<td>11.3 kW</td>
</tr>
<tr>
<td>Power dissipated at stage 3</td>
<td>4.6 kW</td>
</tr>
<tr>
<td>Power dissipated at stage 4</td>
<td>0.6 kW</td>
</tr>
<tr>
<td>Total power recovered</td>
<td>65.1 kW</td>
</tr>
<tr>
<td>Collector efficiency</td>
<td>70.8%</td>
</tr>
</tbody>
</table>

Fig. 6. 3-D plot of the secondary electron trajectories in the HEF collector with a cut-away view of the electrodes.

(to the right hand side of the diagram) reaching electrodes at higher potentials. Many more trajectories (not visible in Fig. 6) are captured by the electrode from which they were emitted and thus do not cause any performance degradation.
Fig. 7 The geometry of TEF collector in the x-y plane with the finite difference mesh superimposed.

Fig. 8 (a)-(d) Plots of the equipotential lines in the 3-stage TEF collector in transverse planes at 0.8 mm axial intervals in the vicinity of the gap between the second and third stages.

Only one or two of the secondary electron trajectories appear to be backstreaming. The behavior of the slow secondaries suggests that this type of collector will have low or zero backstreaming from this cause. The fast secondaries, whose number is usually much smaller than that of the slow ones, may cause some additional backstreaming. These trajectories, however, could not be studied with the present version of the program.
TABLE III

<table>
<thead>
<tr>
<th>V/V_n</th>
<th>0.55</th>
<th>0.65</th>
<th>0.75</th>
<th>0.85</th>
<th>0.95</th>
<th>1.05</th>
<th>1.15</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
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<td>18</td>
<td>51</td>
<td>12</td>
<td>15</td>
<td>9</td>
<td>6</td>
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</table>

V. SIMULATION OF A TILTED ELECTRIC FIELD (TEF) COLLECTOR

A three-stage TEF-collector of simple geometry for which some experimental data was available was simulated [18]. The electric field configuration in this collector was similar to that of Okoshi's soft-landing TEF collector [9] but in this case the magnetic field was transverse instead of being parallel to the axis. The tilted electric field sorts the electrons according to their energy and the transverse magnetic field (prominent in the third stage) is expected to reduce the backstreaming. This collector was used in a helix travelling-wave tube with a beam voltage of 4.8 kV and a net collector current of 53.2 mA. The collector stage voltages were 4.8 kV (body), 2.4 kV, and 1.2 kV. The experimentally measured spent-beam kinetic energy distribution is given in Table III. The beam radius was 0.63 mm.

To simulate the problem on a PC, coarse mesh widths of 1.9 mm in the x-direction and 0.9 mm in both the y- and z-directions were used. Local mesh refinement was used in the regions where the inter-electrode gap or feature sizes were small. Fig. 7 shows the geometry of the 3-stage TEF collector with the finite-difference mesh superimposed showing the local mesh refinement in the x-y plane. A total of 14,440 mesh points was used to define the problem. To model the beam the circular beam cross section was divided into six rings which together contained 61 segments of equal area. Three trajectories were started from each segment with their initial angles randomly distributed in the range of -2° to +2°. This smaller range of angles was chosen arbitrarily considering the fact that the perveance of the beam is very low (~0.16 μPerv). The 183 trajectories used were assigned kinetic energies such that the distribution approximated that given in Table III. The simulation of this collector was carried out both without and with a uniform transverse magnetic field in the y-direction. A self-consistent solution of the problem was obtained in six iterations. The average run time of the problem on the micro-computer specified in Section IV was 73 sec. for the potential solution (50 iterations) and 45 minutes for the trajectory solution in each iteration. The total disk memory requirement of the problem was about 10 Mbytes. The results of simulation of the TEF collector without a magnetic field will be described first followed by the effects of the magnetic field.

A series of interesting equipotential plots in four successive planes at 0.8 mm intervals in the vicinity of the second inter-electrode gap is shown in Fig. 8. These plots show how the potential distribution near the axis changes from nearly circularly symmetric to almost planar and then again circularly symmetric as one goes across the gap from the second electrode to the third. The 3-D potential plot shown in Fig. 9 for the y-z plane (at z = 13.5 cm) is useful for giving an overview of the potential distribution in any mesh plane and thus an understanding the behavior of the trajectories.

![3-D plot of the potential in a transverse plane (z = 13.5 cm) in the 3-stage TEF collector.](image)

A 3-D plot of the electron trajectories in the TEF collector is shown in Fig. 10 with the cross-section of the electrodes in the mid-plane. Some backstreaming trajectories are visible on the left side of the figure. Many of the trajectories also appear to be reflected after penetrating some distance into the third stage. The series of trajectory plots in Fig. 11 shows groups of trajectories belonging to each of the first four classes of energy in the electron beam. The trajectories in the other three classes of energy hit the far end of the third stage like those in the fourth energy class. Slow electrons having energy 0.55 V₀ (2640 eV) are collected at the second stage at 2400 V.
Fig. 11. Sorting of the trajectories by initial energy in the TEF collector without a magnetic field: (a) 0.55 \( V_0 \), (b) 0.65 \( V_0 \), (c) 0.75 \( V_0 \) and (d) 0.85 \( V_0 \).

A. Effect of a Magnetic Field

The shapes of the inner surfaces of the first pair of electrodes were found to have a marked influence on the trajectories of these electrons in some other simulations. Electrons with energy 0.65 \( V_0 \) (3120 eV) are collected at this stage after being reflected by the retarding field at the second gap (Fig. 11(b)). Fig. 11(c) is interesting because it shows that most of the trajectories having a kinetic energy equal to 0.75 \( V_0 \) (3600 V) are not getting collected at the third stage which is at a potential of 3600 V below that of the first stage (body). This is because a part of their total kinetic energy has been converted into transverse energy and their remaining axial energy is not sufficient to take them to the walls of the third stage. Increasing the voltage of the third stage to about 3400 V would, perhaps, have made it possible for this group of electrons to land on it so increasing the collector efficiency. The faster electrons above 0.85 \( V_0 \) reach the third stage directly with a deflection in the vertical \( (y) \) direction depending upon their starting positions and energies.

A comparison between the experimental results and the computer analysis without a transverse magnetic field is given in Table IV. Good agreement is seen in the collector efficiencies with a maximum difference in efficiencies of 0.6%. The distributions of currents among the collector stages however differ by as much as 32%. Similar differences between experimental and computed results have been described by other authors [6]. The simulation also predicts a small amount of backstreaming of secondaries (body). This is because a part of their total kinetic energy has been converted into transverse energy and their remaining axial energy is not sufficient to take them to the walls of the third stage. Increasing the voltage of the third stage to about 3400 V would, perhaps, have made it possible for this group of electrons to land on it so increasing the collector efficiency. The faster electrons above 0.85 \( V_0 \) reach the third stage directly with a deflection in the vertical \( (y) \) direction depending upon their starting positions and energies.

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A. Effect of a Magnetic Field

A dipole magnet of 0.2 A \( \cdot \) m\(^2\) magnetic moment oriented along the y-axis and located toward the end of the third stage was applied to the experimental TEF collector (Fig. 7). This type of magnetic field could not be simulated by the DOS version of KOBRA3 but it would be possible using the extended DOS version which is undergoing evaluation tests. In order to see the influence of a transverse magnetic field on the trajectories in the TEF collector a uniform magnetic field of 8.0 Gauss was applied in the y-direction. This is roughly equal to the field produced on the axis of a magnetic dipole having a moment of 0.2 A \( \cdot \) m\(^2\) at a distance of 20 mm from its center. The complete problem was run again until convergence. The primary trajectories showed little influence of this field by bending of the trajectories in the y-z plane. The effect of the magnetic field on the secondary trajectories was significant. Fig. 12 shows the secondary electron trajectories in the TEF collector without and with a transverse magnetic field. A large number of secondary trajectories are backstreaming in the case without the magnetic field (Fig. 12(a)). In the case with the magnetic field the secondary electron trajectories starting from the backwall of the last stage and from other electrodes were bent in such a fashion that they were collected at the electrode of their origin and backstreaming of secondaries was completely eliminated (Fig. 12b). There was also some redistribution of current among the electrodes. The collector stage currents were: 6.0, 37.2 and 13.7 mA for stages 1 to 3, respectively. The agreement of this current distribution with the experimental distribution appears to be better than that of the current distribution without the magnetic field (Table IV). The collector efficiency was reduced to 72.9% which is only 0.6% less than the experimental value. Thus we can conclude that the prediction of the collector performance has been improved by taking the magnetic field into consideration.
the conventional method of trajectory plotting. The package is expected to serve as a useful design tool for novel types of depressed collectors.

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REFERENCES


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