The General Electric Pacemaker was developed in conjunction with Adrian Kantrowitz, M. D., Maimonides Hospital of Brooklyn, Brooklyn, New York.
INTRODUCTION

The General Electric Cardiac Pacemaker is a miniature implantable energy pulse generator for the correction of heart block or Stokes-Adams seizures. The miniature transistorized implantable pacemaker powered by long life mercury batteries was developed by the Electronics Laboratory of the General Electric Company in conjunction with Dr. Adrian Kantrowitz of the Maimonides Hospital, Brooklyn, New York.

The rate of the pacemaker is about 70 pulses per minute. This has been found adequate in the majority of cases where average physical activity is maintained. For greater activity or periods of stress, an external control may be applied to temporarily raise the rate to any desired level between 75 and 120 pulses per minute. Removal of the external control automatically returns the pacemaker to its original rate. For special circumstances, the pacemaker may be built with higher or lower rates. It may be kept indefinitely at a higher rate with the external control but the minimum rate cannot be lowered. Battery life is not diminished in the implantable pacemaker by continued use of the external control.

The research leading to the development of the pacemaker established that the heart is an energy-sensing organ requiring five micro-joules of energy for stimulation. This threshold energy must be delivered with a minimum of approximately two volts but is also dependent on sufficient energy produced by voltage and current flow for a certain pulse length. The minimum required threshold of five micro-joules is observed to increase to approximately 15 micro-joule threshold over a long period of time after implanting.

To achieve reliable heart stimulation, it is desirable to approximately double the voltage and energy required by the heart. The pulse height is established, therefore, at about 3.7 volts. The pulse energy decreases with pulse rate when the external control is used, so the energy chosen is 35 micro-joules at 120 pulses per minute. This results in a pulse energy input of 65 micro-joules at the base rate of 70 pulses per minute, or approximately four times the required energy.

This energy is supplied by five mercury batteries. The batteries are chosen for the high standards of reliability established by the manufacturer and are carefully handled prior to assembly; therefore, failure due to premature battery rundown should be rare.
All other components in the pacemaker are also of high established reliability. Each part is chosen because of its ability to meet very rigid Government reliability standards. Each part is then carefully tested above, below, and at body temperature. The components are then arranged in matched sets to assure that all active elements meet rigid specifications. The pacemaker is then assembled, the desired frequency is obtained by changing circuit time constants, and the output is measured. Four separate tests are conducted on the assembled pacemaker during the soldering, encapsulating, sealing in silicone rubber, and packaging for shipment.

Physically, the pacemaker is a silicone rubber encased device weighing approximately five ounces. It is 2.5 inches long, 2.25 inches wide, and 1.0 inches thick. Two lead wires, each 21.9 inches long, made of No. 316 stainless steel sutures are sealed into the unit for attachment to the heart. The lead cable 0.015 inch in diameter is made of 49 individual wires in 7 bundles. Each wire is less than two-thousandths of an inch in diameter. The lead wires are insulated by two layers of silicone rubber to provide electrical isolation and a Dacron web is placed between the two layers which adds to the tensile strength of the lead. Two centimeters of these leads are left uninsulated to act as electrodes. Surgical needles are connected to the ends of the leads to facilitate attachment to the heart.
The technique of installing the General Electric implantable pacemaker is not difficult and many variations will undoubtedly be employed. In experience gained at the University of Michigan Medical Center and St. Joseph Mercy Hospital in Ann Arbor the following methods have proved to be most reliable.

An interim technique for pacing the heart must be employed during the period of operation prior to the time the implantable pacemaker electrodes have been secured in the myocardium. This is most conveniently accomplished using an endocardial electrode of bi-polar design. The catheter electrode is introduced by way of the left basilic vein and its tip positioned in the right ventricular outflow tract. The heart is then paced by electrical pulse delivered through the endocardial electrode by means of any standard external pacemaker device. If the endocardial electrode remains in position longer than twenty-four hours the chance of perforation of the right ventricle by the tip of the electrode is increased. For this reason the implantation of the permanent pacemaker is carried out promptly after positioning the endocardial catheter electrode.

Interim pacing by way of skin electrodes while effective introduces interference due to contraction of chest wall musculature.

The pacemaker is implanted under general intratracheal anesthesia providing for pulmonary ventilation throughout the operation.

Two methods of implantation have been employed successfully. The abdominal wall implantation (Figure 1) of the electronic package which has been the most widely employed technique and the left axillary implantation technique (Figure 4) which has been reserved for special circumstances which will be dealt with later.

ABDOMINAL IMPLANTATION TECHNIQUE

The patient is placed in a dorsal recumbent position (supine), with the left arm abducted on an arm board. The entire left chest, left axilla and abdomen are prepared and draped. The pocket to accommodate the pacemaker unit is first developed in the left upper quadrant. A three inch transverse incision is made over the left rectus
ABDOMINAL WALL IMPLANT — FIGURE 1
muscle at the level of the umbilicus or slightly higher. The incision is developed to the level of the rectus fascia and this surface cleared for an area slightly larger than the face of the pacemaker. This should allow for approximation of the wound edges over the pacemaker without tension. The resulting bulge is well tolerated and the waistline rides above the prominence.

A left submammary incision is made and the pleural space entered through the fifth intercostal space. An extracostal tunnel is developed from the lateral end of the submammary incision extending down over the costal margin to communicate with the lateral aspect of the pocket in the left upper quadrant. A short incision in the anterior rectus fascia is necessary to complete this tunnel communication. This more lateral course of the tunnel provides greater protection of the leads as they are subject to less deformity and flexion during bending of the torso. The pacemaker unit is then placed in the abdominal pocket. The needles swedged on the electrode leads are sheathed in their protective tubing prior to pulling the leads through the tunnel. The leads are most easily drawn through the tunnel by looping a piece of umbilical tape around the leads distal to the proximal heavy insulation. Care should be exercised not to pull the tape so tight that the lead becomes kinked. The polyethylene tubes sheathing the electrode needles are not removed until just before implanting the electrode in the myocardium. Care is taken to avoid grasping the leads in the jaws of a hemostat. Under no circumstances should the bare electrodes be grasped by an instrument. Pulling the leads through the tunnel by grasping the needles may result in detachment of the needles at the swedge.

**Exposure of the Heart**
Exposure of the pericardium is accomplished by a rib spreader which opens the fifth interspace for approximately four inches, care is taken to avoid catching the leads between the rib and the retractor. The pericardium is opened vertically along its anterior lateral aspect well anterior to the phrenic nerve. The avascular portion of the left ventricle is selected for electrode implantation.

**Method of Implantation**
The leads are led from the extracostal tunnel through the fifth intercostal space at the anterior axillary line. The thirteen-inch length of heavy insulation provides adequate protection to the leads as they enter the thorax between the fifth and sixth ribs (Figure 2). A gentle curve is fashioned to bring the leads to the midline of the mediastinum whence they are lead cephalad to the base of the heart and brought
down to the surface of the left ventricle. This route is important as it
minimizes the flexion to which leads are subjected to by myocardial
contractions. Care should be taken to avoid sharp bending of the leads
at the taper where the heavy tubing insulation gives way to progres-
sively thinner insulation. One of the individual electrodes is then led
into the avascular portion of the left ventricle at the base of the heart
directed toward the apex (Figure 3). The tip of the curved needle
should exit from the myocardium approximately fifteen millimeters
from its point of introduction. Care is taken to avoid entering the
ventricular cavity with this suture pass.

After the needle is withdrawn from the myocardium the bare electrode
is grasped with the fingers and the electrode drawn along the needle
path in the myocardium until the insulated portion of this single elec-
trode enters and exits from the myocardium along the needle path. *
Pulling the electrode into position by means of a needle holder ap-
plied to the needle may result in separation of the needle from the
electrode at the swedge. The needle is then reintroduced into the
myocardium close to the point of exit of the electrode and passed
in a reverse direction toward the base of the heart making a parallel
path with the initial suture pass. The bare electrode is withdrawn
from the heart until a small metal flange comes into view.

A 3-0 Dacron ligature is then used to ligate the bare electrode to the
insulated proximal electrode placing the knot below the level of the
metal flange. The needle is then cut off the electrode distal to the
metal flange. The same procedure is followed implanting the second
electrode parallel to the first so that the distance between the bare
electrodes is one centimeter. The second electrode should be placed
so that the last suture pass is adjacent to the last suture pass of the
previous electrode positioning providing for a zone of myocardium
free of injury between the bare electrodes (Fig. 3).

If the heart is being paced by an endocardial electrode this pace-
maker should be turned off as the second electrode is implanted in
the myocardium. If the myocardial implantation is too close to the
apex of the ventricle stimulation of the phrenic nerve may result. The
thoracotomy wound is closed in the usual manner employing thora-
cotomy tube drainage for as long as may be required. Prior to clos-
ing the incision in the left upper quadrant the thin silicone-impregnated
dacron flaps on either side of the pacemaker unit are sutured with
four nonabsorbable sutures to the rectus fascia to prevent migration
or rotation of the unit during the healing stage.

* See diagram and footnote, p. 8.
Approximately 1.5 cm. of the distal end of the insulated lead is permanently positioned in the myocardium. Failure to observe this detail may result in early lead breakage at this point.
AXILLARY IMPLANTATION ALTERNATE TECHNIQUE, FIGURE 4

Younger patients with greater activity potential prefer the left axillary implantation site for the pacemaker package as this location interferes less with body motion and bending. Wearing apparel is more comfortable than with the abdominal implantation which requires men to wear loose clothing and suspenders. Belts may be uncomfortable for some patients with abdominally placed pacemaker. In obese patients there may be an increased tendency for the pacemaker to migrate in the abdominal implantation site or rise on edge and turn over, stressing the leads or even pulling them out of the myocardium. This has happened in spite of suturing the suture pad adequately to the rectus fascia. The above circumstances have led to the use of the axillary implantation site with encouraging results.

In the axillary implantation the operation is carried out with the patient in the same position, supine, with the left arm abducted. The anterior chest and left axilla are prepared and draped. A submammary incision, five to six inches in length, is developed over the fourth or fifth intercostal space and the pleura entered through the most appropriate space as determined by the surgeon. The pocket to accommodate the pacemaker package is developed extracostally by undermining the lateral aspect of the wound along the axis of the intercostal incision. Care is taken to maintain absolute hemostasis during the development of this pocket otherwise hematoma formation will be impressive in the loose areolar tissue of the axilla.

The pocket is developed as far posteriorly as the midaxillary line. The pacemaker package is then placed in the axillary pocket and the leads which are of shorter dimension (fifteen inches) on axillary models are led through the intercostal space to the midline of the anterior mediastinum. The leads are then swept cephalad and curved to the left being brought down in a gentle arc to the base of the left ventricle where implantation is carried out in the usual manner. Use of the standard abdominal implantation model for axillary implantation requires the placement of an extra loop in the region of the anterior mediastinum to take up the extra slack in lead length.

Lead Handling
Care should be taken to avoid making any sharp turns or acute angles in manipulating the pacemaker leads. An ample length of lead material is provided with the unit to allow sufficient slack between the point of electrode implantation and the site of the pacemaker package.
AXILLARY IMPLANT — FIGURE 4
EXTERNAL RATE CONTROL

The General Electric External Rate Control (Figure 5) is used for increasing the basic rate of an implanted General Electric cardiac pacemaker to any rate to approximately 120 pulses per minute. The External Rate Control consists of an induction coil and the electronic control. The induction coil is plugged into the control and placed with GE monogram away from patient's skin directly above the implanted pacemaker. The electronic control can be placed in the patient's shirt pocket, other pocket, held to the body with a loose fitting strap or otherwise placed in a convenient location.

The External Rate Control is energized by switching the slide switch to "ON." The rate is adjusted by rotating the knurled round knob. The rate will increase from the basic rate at "A" position on the control knob to a maximum rate of "G" position, near 120 pulses per minute. Return to the basic pacemaker rate results from removing the coil from near the implanted pacemaker. The electronic control should be switched to "OFF" position when not in use to save battery energy.

The advantage of the External Rate Control unit is its ability to elevate the heart rate to meet increased physiological demands of the patient. Such demands are usually temporary and intermittent. An increase in pulse rate to approximately 80 to 85 per minute may be of considerable benefit to some patients in the early postoperative phase, particularly those patients who may manifest slight hypotension in this phase of their recovery. Similarly the Rate Control is of value in subsequent illnesses or operations. Younger patients or those with more activity potential because of an otherwise healthy state may welcome the External Rate Control as a means of increasing their level of activity at will. The Rate Control unit may serve as a means of checking the threshold of myocardial response. As the rate is increased the energy pulse is decreased. Failure of the heart rate to increase above 90 per minute may indicate a minimal reserve above threshold for pacing, thereby warning of eventual pacing difficulty.

Required care of the External Rate Control is minimal. With normal handling, it is rugged and highly reliable. The case is not waterproof. Battery life depends on initial charge of the battery and how many hours a day it is in use.

When the External Rate Control appears to operate poorly, first replace the battery with a new one. If the difficulty persists, return it to Pacemaker, General Electric Company, X-Ray Department, 4855 West Electric Avenue, Milwaukee, Wisconsin 53201.
GENERAL ELECTRIC EXTERNAL RATE CONTROL — FIGURE 5
GENERAL INSTRUCTIONS

Handling
The pacemaker is a reliable, rugged device, but certain precautions should be taken in its use.
Normal transportation handling will not damage the pacemaker electronic package. Care should be exercised to insure that the electrodes and lead wires do not become kinked, abraded or nicked. Out of the package, the pacemaker should not be dropped. In the event that it is dropped from a height greater than six inches, the unit should not be implanted.
Care should be taken to avoid contaminating the silicone rubber covering. Silicone rubber has a static attraction and affinity for surface contaminants such as fingerprints, dust, lint, talc, starch and many other materials which can evoke foreign body reactions. Handling with lint-free sterile surgical gloves is recommended. Should the pacemaker become contaminated, it should be thoroughly washed and rinsed in distilled water.

Sterilizing
Do not autoclave the pacemaker. Any cold or ethylene oxide method of sterilization may be used in which the pacemaker is not heated above 115°F. The pacemaker should be very carefully washed after sterilization, ventilated and then rinsed in sterile water. It should not be left in sterilizing solutions for periods in excess of 24 hours.

Checking Operation
To check operation of the pacemaker, either in or out of a patient, place a small radio, like a pocket transistor radio, on top of the pacemaker, tune between stations and turn to full volume. Hold the pacemaker leads shorted together. The pacemaker pulse will be heard as a small click at a rate below 60 pulses per minute. To simulate implanted conditions, connect a 300 ohm, 5% tolerance resistor and a 20 microfarad, 10% tolerance capacitor in series with the electrodes. This will approximately produce the implanted pulse rate.

How to Order
The General Electric cardiac pacemaker and external rate control is sold directly to hospitals and members of the medical profession. Vital Evaluation Information is requested to be returned to the General Electric Company. For convenience forms are furnished with each unit shipped.
PACEMAKER
HELCABLE ELECTRODES

INTERIM CHANGES in procedure
for implanting HELICABLE Electrodes
of the G-E Pacemaker
HELICABLE ELECTRODES

INTERIM CHANGES in Procedure for Implanting HELICABLE Electrodes of the G-E Pacemaker

1. REMOVE - The tubular Plug on the connector of the Pulse Generator should be removed before sterilization. This Plug is used to retard the discharge of the Pulse Generator during storage and shipment and prior to implanting.

2. STERILIZING - The electrodes may be sterilized by autoclaving. DO NOT AUTOCLAVE THE PULSE GENERATOR. For sterilization of the Pulse Generator see "Cardiac Pacemaker" booklet, page 13.

3. CONNECTOR - The integral connector allows the electrodes to be sterilized and implanted before attaching to the Pulse Generator. The electrodes are designed to be drawn connector end first through any tunnel preparations during implanting. This will tend to avoid twists and kinks. The protective cover on the connector should be removed just prior to engagement with the Pulse Generator.

4. HEART ATTACHMENT - Two suture passes are made for each electrode (See Revised Figure 3 on back cover of this Interim Changes). On the first pass the insulation on the heart attachment end is to be drawn into the myocardium to the place where it abruptly increases in cross-section. At the forward end of the insulation, nearest the suture needle, there is a cone on the electrode which should protrude from the myocardium 4 or 5 mm. After the second pass (this time bare electrode) is drawn, this cone will provide a strain-relief pad for the exposed loop of electrode.

5. FERRULES - One pure silver ferrule is required for each electrode. It is to be placed and attached prior to cutting off the needle. Be sure to crimp the ferrule snugly, using the tool furnished from General Electric. Each electrode contains 49 small tempered strands laid together, which, when severed, will flare, or unravel. The snugly crimped ferrule prevents this condition when cutting off the needle.
6. ATTACHMENT OF PULSE GENERATOR - Remove protective cover from electrode connector by rolling it off like a rubber glove. A double seal is provided. The pins and sockets are aligned by means of a key. The metal sleeve of the Pulse Generator fits over the metal sleeve of the electrode connector. The inner silicon tubing of the electrode fits over the metal sleeve of the Pulse Generator and will butt against the short length protruding from the Pulse Generator. The outer length of tubing on the electrode will fit over the short length protruding from the Pulse Generator and will butt against the Pulse Generator case. Grasp the electrode and the Pulse Generator firmly and, if aligned, they should engage with effort. Pinching the tubing around the connector as you apply, engaging pressure will assist in bringing them into place.

7. MODE CONTROL - The Mode Control is a pencil shaped magnetic device furnished with the Dual Mode pacemaker (not applicable to the "Single Basic Rate" pacemaker). It has the words NORMAL (black) and ACTIVE (red) engraved along the side, the letter "A" engraved on one end, and the G-E monogram engraved on the other end.

The Mode Control is used to change the rate of the Dual Mode pacemaker from NORMAL (approx. 70 pulses per minute) to ACTIVE (approx. 85 pulses per minute) or vice versa. It functions best when held within 4 centimeters of the side of the Pulse Generator opposite the suture pad, and at a level with and parallel to the edge to which the leads attach. Normally the pacemaker will remain at the rate set until changed.

**Abdominal Implant:** - Orient the Mode Control by holding it such that the selected word NORMAL or ACTIVE can be read by the patient (upside down to the attending person). Move it toward the implant site. When within the 4 centimeter range the Pulse Generator should be at the selected rate. Retract it in the same manner.

**Axillary Implant:** - Orient the Mode Control by holding it with the end bearing the G-E monogram pointed toward the patient's head for NORMAL rate or the end bearing the letter "A" pointed toward the patient's head for ACTIVE rate. Move and retract it as described above.

After implant you will want to check to make sure that the Mode Control functions properly.
HEART ATTACHMENT - FIGURE 3

This Figure Supercedes the figure of the same number in the text.