

FIG. 3. Large brass plates at eighty kilocycles.

mathematical equations for the lines on the circular plate satisfy Kirchhoff's equation³ except at the center; while the forty-five lines on the square plate may be calculated from the Ritz⁴ formulas.

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A METHOD OF PASSING AIR, GAS OR VAPOR OVER OR THROUGH MICRO-ORGANISMAL GROWTHS¹

THE apparatus herein described can be used to study toxic and stimulating effects of gases or vapors on colonies of micro-organisms. As illustrated, it is adapted to solid medium. Liquid medium may be used, in which instance gas or vapor bubbles through the medium.

The gas or vapor may be drawn through the culture with suction as with the water pump A, or it may be forced through by pressure. In the latter instance, if the gas is inflammable the culture tube is connected with bottle I and the gas burnt at its nozzle. When suction is used a trap such as B is interposed between culture and the source of suction. This contains water, except where pathogens are studied, when an antiseptic is substituted. The glass tube connecting bottle with culture tube projects just below the surface. By this means an idea of the rapidity of passage of gas or vapor can be gleaned from the number of bubbles. The water pump should be shut off slowly in order to avoid any possibility of soilage. The bottle B may receive connections from many culture tubes instead of the one.

At stand C will be noted culture tube D, which is bent on itself slightly. After planting, the plug of cotton is forced further into the tube so that the stopper with glass tubing and connecting rubber can be fitted. The shaded area illustrates solid medium. E is a rubber vaccine stopper into which is plunged a 23-25 gauge hypodermic needle so that it rests on the

medium. The needle connects with a bottle, such as H, which contains water except when the effect of a volatile liquid is to be studied. H in turn connects with a glass cleaning tube containing cotton which filters out organisms from air or gas. D and all parts distal to it

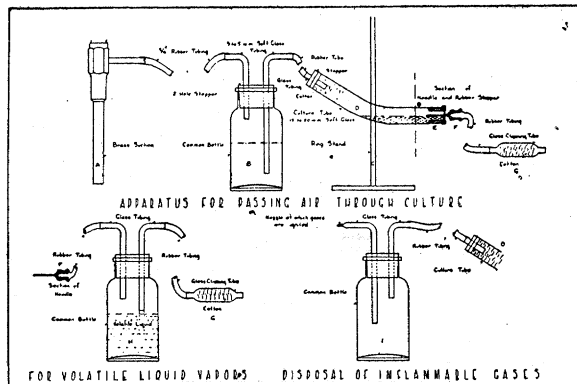


FIG. 1

must be sterile and adjusted, using sterile precautions. Without bottle H containing a liquid, excessive drying will be encountered.

This apparatus is easily constructed of stock laboratory material. It is inexpensive, easily adjustable and has many adaptations. A liquid medium may be used instead of solid. Gas or vapor may be passed through or over the culture only a short time in order to determine its noxiousness. Glass parts may be substituted for rubber where desired. When liquid medium is used frothing must be compensated for either by length of tube or other means. It is wise to have stop-cocks in the system so that any sudden change of gas pressure may be avoided. Hypodermic needles can be inserted, withdrawn and reinserted. Care must be taken that rubber of the vaccine cap is good and that all fittings are sound.

In conclusion, the variation of this method is largely in the use of the vaccine cap. In my experience I have found this a valuable adjunct. Many culture tubes can be run in the same suction or pressure set-up at one time.

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³ Rayleigh, "Sound," Vol. I, Chapter X.

⁴ Ritz, *Ann. der Phys.*, 28: 737 et seq., 1909.

¹ Contribution No. 58 from the Department of Biology and Public Health, Massachusetts Institute of Technology, Cambridge, Mass.