On the Structure and Mechanics of the Protozoan Flagellum

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The flagellum is an organelle of exceedingly wide distribution among both animals and plants. Among the Protozoa, it occurs not only in all of the Mastigophora, but also in many Sarcodina and Sporozoa during part of their life cycle. The swarm spores and gametes of certain algae and fungi possess flagella, as do the sperm cells of bryophytes, pteridophytes, and cycads. The tail of the typical animal spermatozoon is probably nothing more than a flagellum. The cilia of the ciliate Protozoa and of metazoan ciliated epithelium in all probability represent modified flagella. There is reason to believe that the bacterial flagellum is perhaps the simplest and most primitive manifestation of this organelle. In spite of the apparent diversity of form and function represented in these numerous examples, it seems highly probable that the organization and mode of functioning are fundamentally similar in all.

During the past century numerous investigators have studied the structure and mode of action of flagella, and their conclusions have varied widely. Certain facts have been established, but many points have remained open to question. It has been evident for some time that new techniques would be necessary for the settlement of such questions. Today the electron microscope, with resolution and accompanying magnification far beyond the limits of the ordinary compound microscope, provides such a new technique for the study of minute structures.

Having access to an electron microscope and to pure clone cultures of several flagellate Protozoa, I undertook an intensive study of flagellar structure, the major results of which are embodied in this paper. Flagellar action and flagellate locomotion I investigated by direct observation with the aid of a special technique, by the construction of working models, and by actual underwater swimming experiments.

The work was done at the suggestion and under the supervision and encouragement of Professor W. J. Kostir, to whom I am especially indebted. The electron microscope employed was one constructed by Professor A. F. Prebus, of The Ohio State University, and operated by him and his assistant, Mr. John Dankworth, to both of whom I here express my appreciation for invaluable assistance.

REVIEW OF PREVIOUS WORK ON THE SUBJECT

STRUCTURE OF THE FLAGELLUM

Until the 1880's or later, the flagellum was generally considered simply an elongate, homogeneous fiber, bearing no appendages, and characterized by weak staining (Batschli, 1883–7; Klebs, 1892; Danziga, 1901 b). The flagella were usually depicted as tapering to a point, but Batschli considered them commonly of equal diameter throughout their entire length, or tapering only slightly at the end.

In opposition to such views appeared the papers of Kunstler (1882, 1889), Loeffer (1889, 1890), and Fischer (1894, 1895). The former described (Kunstler, 1889, p. 408, translation) "... a dark axial line resembling a slit or canal, partic-
ularly when the filaments observed are well extended horizontally; in these cases one may often distinguish delicate transverse septa, dividing the axial cleft into rather short sections." (Fig. 1 a). The dark sections he considered to represent vacuoles which are separated from one another by septa of the transparent ground mass which surrounds them. He also described a cortical layer or sheath (Fig. 1 b, c) which was often separated from the ground substance and its enclosed slit.

To the presence of this enveloping membrane he attributed the weak stainability of the flagellum. The axial substance he considered the more contractile portion. Kunstler also first described an interesting external structure in the flagella of *Oxyrrhis* and *Cryptomonas*: "... the attenuated extremity often seems prolonged in a pale filament, thinner than the flagellum and of a different appearance (Fig. 2). The limits of the flagellum proper are distinguished clearly; suddenly, at its tip, the diameter and appearance change; there begins a transparent filament, hardly visible, very slender, sometimes ending in a slight swelling."

Loeffler (1889) independently described this last-mentioned structure, employing a new technique he was using for bacterial flagella. He suggested that the thick basal part might be a sheath, from which extends the delicate protoplasmic fiber in the manner of a pseudopod, slightly swollen terminally. The apical swelling, he postulated, might function as a tactile organ. Loeffler also discovered on the
flagellum of a monoflagellate (perhaps *Oikomonas*) two dense rows of extremely delicate hairs, projecting almost at right angles to the axis of the flagellum, one along each side. (See Fig. 3). He did not investigate the possibility that in life the hairs actually might cover the filament instead of being arranged in two rows, and in drying be flattened down laterally, producing the impression of two rows of hairs. At any rate, he had described and figured photomicrographs of two previously unknown appendages, and outlined his method in detail.

**Fig. 3.** *Monas* sp. The first published figure of a flimmer-flagellum, from a mordant-stain preparation designed for bacterial flagella. (Redrawn from Loeffler, 1889, Plate II.)

**Fig. 4.** *Euglena viridis.* a. "Cast-off flagellum ... still completely stretched out; untwisted, with the hairs on one side . . ." b. "A cast-off flimmer-flagellum rolling up, partially swollen and therefore with an apparent axial fiber." c. "A cast-off, rolling-up flagellum which had previously become uniformly swollen and consequently shows no such apparent structure" (as Fig. 4 b). From preparations of undiluted culture dried upon cover glasses in about one-half hour, and treated with a modification of Loeffler's mordant-stain technique. (Redrawn from Fischer, 1894, Plate XI.)

Fischer (1894), using a modification of Loeffler's technique, made a careful study of the flagella of *Euglena viridis*, *Monas guttula*, and species of *Bodo, Chlorogonium*, and *Polytoma*. He confirmed Loeffler's findings and discovered that in *Euglena* the flagellum bears but a single row of hairs (Fig. 4), while that of *Monas* has two opposite rows. He termed these flagella Flimmergeisseln, or ciliated flagella. Since "flimmer" appears to be a convenient word for the structures, and does not carry any obvious implications in the English language, I shall hereafter employ the term "flimmer-flagellum" in reference to any flagellum bearing hair-like projections along one, two, or all sides.

Whip-flagella, or Peitschengeisseln, Fischer found in *Bodo, Chlorogonium*, and *Polytoma*, and described as follows (p. 230, translation): "The whip-flagellum consists of a thick homogeneous stalk, previously thought to be the entire flagellum (the only part visible in unstained specimens) and, arising from its tip, a very delicate whip-thread 2-3 times as long as the stalk. This thread is slung about like a coach-whip by the strokes of the basal stalk."

Fischer observed in *Euglena* what he concluded to be an artifact arising from incomplete swelling: "... the central, not yet swollen part of the flagellum appears as a denser, more strongly stained axial fiber, while the outer, swollen part, seems a less dense, more weakly stained ground substance." (See Fig. 4 b). Further, "In *Polytoma* and *Bodo*, a relatively commonly seen granular structure of the basal part of the whip is likewise only a result of the technique. This flagellar structure,
which agrees with that described by Kunstler, is to be regarded in this light" (i. e., as an artifact). (For further discussion of the granular appearance, see Gelei, 1926.)

Fischer studied in detail the processes of flagellar disintegration, whereby he made his observations on the artificial nature of the granular or alveolar appearance of certain flagella. He figured in detail many examples of all the above-mentioned types, and discussed each matter thoroughly. In spite of the fact that he could demonstrate no natural inner structure in either whip- or flimmer-flagellum, he stated (p. 204): "... yet there is no doubt that the apparently homogeneous fiber must possess a very fine structure made up of definitely arranged particles (micelles), toward which point the twisting of the flagellum, its rolling up, and the arrangement of the flimmer-hairs."

Regarding the arrangement of the flimmer, Fischer assumed that they must beat in unison, since, on any given flagellum, all the flimmer project in the same direction.

For years flimmer-flagella were generally considered artifacts, perhaps the result of a final wriggle and smear, or of chemical action (Plenge, 1898; Dangeard, 1901b; Bütschli, 1902; Schuberg, 1905; Korschikov, 1923; Günther, 1928). On the other hand, most of these workers confirmed the existence of whip-flagella, and in many cases observed such structures in living specimens. During the latter part of the period covered by the above references, several investigators were, however, confirming the reports of Fischer and Losfler. Petersen (1918) demonstrated the two flagella of Synura uvella to be of different types (Fig. 5a): "... the one of whiplike form, the other a pinnate flagellum showing two opposite rows of little secondary filaments, and resembling a feather; it is to be presumed, however, that these [filaments] extend from all sides of the primary filament." (Translation from summary in French.) Petersen also found flimmer-flagella in Uroglena and Dinobryon. In a later paper (1929) he expanded his observations to a large number of forms, some possessing only whip-flagella, some only flimmer-flagella, and some having one of each. He also described a combination whip-flimmer-flagellum in Craspedomonads. (Fig. 5b). The distribution among the Protozoa of consistent flagellar types fits well into the accepted scheme of classification.
Mainx (1928), though failing to demonstrate flimmer-flagella in several species reported to possess them, confirmed their presence in *Euglena viridis* and figured excellent photomicrographs of the flimmer-flagellum of *Phacus pleuronectes*. His most exacting attempts to observe flimmer on the living flagellum failed. In consideration of their extreme regularity along one side of the flagellum, their presence with or without fixation, etc., he concluded that they could hardly be artifacts. He considered them, however, not motile, as Fischer had assumed, but rather simply as devices serving to increase the flagellar surface.

Petrova (1931) employed flagellar types in the solution of a taxonomic problem. Deflandre (1934), using a very different method of preparation, confirmed the presence of whip- and flimmer-flagella in many organisms, and considerably expanded the list of species studied.

Vlk (1931, 1938) has perhaps treated the problem most thoroughly, expanding and organizing the described types, and tabulating his results. For a more thorough treatment of this phase of flagellar structure, the reader is referred particularly to Vlk, 1938, or to Deflandre, 1934. Vlk even succeeded in observing flimmer (in two distinct rows) on the flagellum of active, living specimens of *Mallomonas acaroides*, a species having an unusually large flagellum. These flimmer are described as having a length about 6 times the diameter of the flagellum proper. He was able to demonstrate these living flimmer-flagella to members of the institute staff, including Professor A. Pascher.

Unfortunately, Kudo (1939), in his brief treatment of the flimmer-flagellum, gave all credit to Vlk but misrepresented Vlk's diagram of a two-row flimmer-flagellum (Fig. 6). The error is certainly unintentional, but misleading. This diagram, which Kudo used to illustrate a two-rowed flimmer-flagellum, was originally intended by Vlk (1938, p. 486) to represent a situation which *should* occur in twisting if the flagellum actually bears two distinct flimmer rows. But such a situation had never been observed. Instead, the flimmer of these forms usually appear somewhat as in Fig. 3 or Fig. 5 a. In other words, Vlk used it as a probable indication that the flimmer, in most organisms at least, are distributed on the surface of the flagellum like the hairs on a dog's tail, and not just in two lateral rows.
Recognition of the whip-flagella has been rather general, and they seem almost universally accepted today. Gerloff (1940) figures the whip-flagella in many species of *Chlamydomonas*. A considerable variation in relative length of lash and stalk may be noted (Fig. 7). Many authors (e.g. Awerinzew, 1907; Minchin, 1922) consider the lash or end-piece as a portion of an axial fiber extending beyond the enveloping sheath of protoplasm and membrane. Cilia are now commonly considered to have a similar structure (Klein, 1929). Klein seems to agree with Loeffler's hypothesis that the terminal swelling of the lash (Fig. 8) may have a sensory function.

Figure 8 (or 10 a) will also serve for a discussion of the generally accepted concept of the internal structure of a flagellum. Most textbooks and articles on the subject agree to the presence of an axial fiber surrounded by a protoplasmic sheath. Beyond that there is much less agreement. A few consider the axial core as the major contractile or vibratile portion, the sheath serving as a supporting structure (Rosskin, 1922; Calkins, 1933). The vast majority think differently. (Butschli, 1902; Schuberg, 1905; Awerinzew, 1907; Goldschmidt, 1907; Williams, 1907; Erhard, 1910; Hamburger, 1911; Minchin, 1922; Korschikov, 1923; Doñean, 1929; Klein, 1929; Kudo, 1939; Hyman, 1940).

The flagellum (or its axoneme) usually arises from a basal granule (blepharoplast), apparently essential to the continued function of the organelle. In certain forms the flagellum has two roots and basal granules (e.g. *Euglena*; Wager, 1899; Hall and Jahn, 1929). For information on other modes of attachment, see Prowazek, 1903; Goldschmidt, 1907; Schouteden, 1907.

Regarding the interrelationships among axoneme, surrounding plasm, and outer membrane, there are interesting reports, based upon studies of disintegrating flagella, which throw light upon the subject. Fischer (1894) made extensive
observations on the coiling or rolling-up of flagella, noting that the process might begin at either end of the flagellum or at any point in between. Korschikov (1923) confirms Fischer's description of flagellar "aggregation" (Figs. 4 b, c; Fig. 9), summarizing his observations as follows (translation from German summary): "The axial fiber and the outer membrane are the most resistant structural components of the flagellum. In the 'aggregation' of the flagellum, the intermediate substance is first destroyed. In the corresponding region the membrane becomes swollen. In the course of this process, the axial fiber is drawn into the swelling and forms there a ball or coil (Fig. 9). Afterward, however, when this blister of the flagellar membrane is destroyed, the axial fiber is freed and stretches again in length."

Awerinzew (1907) and Hamburger (1911), describing flagella undergoing destruction, state that the "protoplasmic" portion of the flagellum rounds into globules which proceed to flow or withdraw toward the base of the flagellum, either leaving the axoneme exposed and straightened out (Awerinzew) or telescoping it (Hamburger). (See Fig. 10.) All of these facts tend to indicate an elastic, supporting axoneme.

Ulehléa figures, but barely mentions, an interesting flagellar structure (Fig. 15), reminiscent of Kunstler's diagrams, but seen in living flagella.

![Fig. 11. a. "The uncoiled fibrils of the flagellum of Euglena." b. "Euglena with the fibrils of the flagellum branching out into a system of rootlets in the protoplasm of the body."]

"This structure is demonstrated with ease by subjecting a flagellum to slight pressure."

(Drawn from Dellinger, Figs. 1, 3.)

Roskin (1922, 1923) postulates and Korschikov (1923) asserts that the axial fiber consists of a great number of thin fibrils. Koltzoff (1903) stated that in certain pteropods the cilium "consists of several fibers which are encased in a common fluid plasma layer . . . the inner fibers running to the basal body . . . " Schmitt, Hall, and Jakus (1943) figure cilia (Frontonia) and flagella (Trichonympha) frayed into 9–11 fibrils.

Dellinger (1909), who figures fibrils in both cilia and flagella (Fig. 11), states: " . . . the flagella of Euglena, Chilomonas and Spirillum are composed of four spiral filaments." He also depicts the axial filament in the pseudopod (axopod) of Actinosphaeria (Heliozoa) as fibrillar. A similar structure is described in the heliozoan axopod by Rosskin (1925). For discussions of the possible homology between such pseudopods and flagella see Minchin (1922), Goldschmidt (1907).

The flagella of many bacteria entwine in a tuft and function as a unit, in some cases even becoming encased with a glutinous mass which keeps them bound together (Reichert, 1909; Ulehléa, 1911; Metzner, 1920; Pijper, 1941; Hutchinson and McCracken, 1943). This situation is perhaps analogous to that occurring in cirri or in the "cilia" studied by Gray (1922) and Carter (1924), in which a number of units capable of beating individually are united into a single organelle.
Sperm tails display a fibrillar axis and an enclosing sheath, and have been more exhaustively studied than flagella. Furthermore, they are probably homologous to the protozoan flagellum, although the details of the homology are not yet settled (Dangeard, 1901a; Alexeieff, 1924; Grassé, 1926; Duboscq and Grassé, 1933). Subsequent to the work of several investigators, Ballowitz stated in 1890 that many types of sperm tails frayed into 2-4 subfibrils. In his paper of 1908, he figures tails fraying into as many as 18 subfibrils. Koltzoff (1909) produces interesting figures of sperm tails, including such as Fig. 12, showing a sheath of helically wound gel fibrils. Recently the electron microscope has made possible more minute studies. Several articles (Baylor, Nalbandov, and Clark, 1943; Harvey and Anderson, 1943; Schmitt, Hall, and Jakus, 1943; Schmitt, 1944) figure micrographs of sperm tails showing, in frayed parts, 9-12 fibrils (Fig. 13). According to Schmitt, these fibrils are smooth, parallel, 300-500 A wide, and run the full length of the tail. Occasionally even finer subfibrils are seen. "In mammalian sperm tails the bundle of fibrils is surrounded by a sheath, the major component of which is a closely wrapped helical fibril, having a thickness also in the range of 300-500 A (Fig. 14). When the tails are fragmented by ultrasonic radiation, portions of this helix can be seen, appearing like miniature solenoids." (Schmitt, 1944, p. 36.)

MECHANICS OF THE FLAGELLUM

In the past, little actual correlation has been made between structure and function in flagella. Many authors have recorded types of flagellar movement and general movement, but relatively few have attempted to analyze the forces or mechanisms involved. First, let us enumerate the types of movements described. Úlehla (1911, pp. 727-728) classifies them as follows:

1. Monad type. Flagellum long, cylindrical, flexible; beats forward in loops (spirals) grading into flattened waves (i.e., undulations; a given point on the flagellum follows an elliptical path).

2. Chrysomonad type. Similar to above, but with shorter and stiffer flagellum.
(3) *Euglena* type. Long, twisted, ribbon-like flagellum; beats laterally in loop-shaped waves.

(4) *Bodo* type. (Trailing) flagellum with long end-piece, rather stiff; beats in flattened waves.

(5) *Clostridium* (a bacterium) type. Flagellum long, spiral, rather stiff; slow waves from base to tip.

(6) Chlorophycean type.
   a. Swarmer type. Flagellum short, cylindrical, rather strong, with the basal part more flexible; functions as an oar (in sculling?), swinging around as if contracting as a unit.
   b. *Pandorina* type. Flagellum longer, more flexible, often somewhat flattened; beats with a spiral lateral stroke.

Uhlela hardly considered the paddle-stroke as such, though this type of movement is described by many, being reported in flagella by Plehn (1904), Rosenbusch (1908), Doeflein (1916), Kofoid and Swezy (1920, 1923), Krijgsman (1925), Nigrelli (1929), Kirby (1943); in cilia by Gray (1922) and Carter (1924), among others. Such a lash-motion is mentioned also by Pütter (1903), Goldschmidt (1907), Schindera (1922), and Grassé (1926). In some such cases, it evidently occurs in flagella which ordinarily undulate. In all of these cases, the flagella concerned were located anteriorly.

![Fig. 16. See text for explanation.](image)

The second major type of movement executed consists of undulations, usually occurring in two planes (i.e., a spiral). Reports of this type may be found in Bütschli (1887), Kunstler (1889), Goldschmidt (1907), Laveran and Mesnil (1907), Friedrich (1909), Reichert (1909), Bancroft (1913), Kofoid and Swezy (1919), Schindera (1922), Grassé (1926), Entz (1928), Petersen (1929), Lowndes (1936, 1941, 1943, 1944, 1945).

The third type, not listed above, is simple conical gyration, reported by Keysselitz (1906), Metzner (1920), and Entz (1928), and by McDonald (1922), in cilia.

Primarily because of Krijgsman's detailed paper, which was based on studies of *Monas* sp. under darkfield, most textbooks today emphasize the paddle-stroke. It presents a clear-cut, diagrammatic, relatively simple-looking movement, familiar in our own swimming stroke and similar to that commonly depicted for cilia. The filament is held rigid during the effective stroke, but is flexible during the recovery stroke.

The concept of progression by a spiral turning of the flagellum has led a painful existence because of Bütschli's analysis of the situation as he saw it (Bütschli, 1887, p. 857 et seq.). Delage and Hérouard (1896, pp. 306–307) treat his explanation thus.
(translation): "Suppose that the flagellum assumes the form of an elongate helix (as indicated in Fig. 16), and that this helix begins to turn clockwise (as seen from the base). At any point such as $o$, the pressure on the water would create a force $ob$ perpendicular to the flagellum, which could be resolved into two others, the vertical one $oa$ determining its rotation about its axis. It is to be noted that this rotation is in the opposite direction to that of the flagellum. If the helix turns in the reverse direction, the animal will go backwards. In the case of a helix wound to the left, there would be likewise progression forward for a certain direction of rotation and backward for the opposite direction. . . . All that is very well, but Bütschli does not note that the movement he describes supposes an agency met in our mechanical instruments, but never among living beings. In order that the body and the flagellum might turn indefinitely in opposite directions about the point $p$, it would be necessary that the mode of union between them be that of a pin stuck through a card, capable of turning freely in its hole, its union with the card assured by the head which could not go through the hole." But this leaves Delage and Hérouard with the problem of explaining matters. They do so by going into great mathematical detail concerning the possibility of a forward component produced by conical gyration, concluding this to be impossible by direct action. However, by its indirect action, i.e., by the rotation produced in the body, they think to have found the answer; namely, that since the flagellum is firmly attached to the body it must rotate with the body, and being held in a corkscrew shape, will thus screw the body and flagellum through the water.

Reichert (1909) noted the forced aspect of this scheme and pointed out that, in his experience, the facts argued against such a mechanism. Further, he states (translation): "The necessity for their relatively cumbersome explanation of flagellar movement appears to have been a result of a misconception of Bütschli's idea. For Bütschli postulates that an opposing rotation of body and flagellum is not feasible in his view that the lines of contraction move about the flagellum. If the lines of contraction on the flagellar surface circle to the right, the body will turn to the left, and so will the flagellum, about its axis. . . . It is only necessary that the lines of contraction move correspondingly faster about the flagellum, so that its turning with the body is equalized or offset. Then the same effect would be achieved as if this turning of the flagellum did not occur at all."

Delage and Hérouard are not the only ones to misinterpret Bütschli's idea. Reichert explains it, and the physics involved, quite clearly. Then, in his summary he states: "Reversal of movement is accomplished quickly by polar-flagellated bacteria, for they simply reverse the flagellar rotation (as Reichert and Bütschli postulate that they should), or by bipolar-flagellated bacteria, whose flagella take turns (those at the other end begin beating). In peritrichous bacteria, on the other hand, reversal occurs slowly. Movement must first cease for a moment, and the flagella assume an opposite orientation to the body." But he is inconsistent, for he also states: "a. The flagella (in bacteria) are always wound in a right spiral (clockwise) and rotate always to the right (as seen from the rear), i.e., the screw-like lines of contraction wind around the flagella to the right. b. The body always rotates to the left. c. The flagella are usually directed backward during locomotion. In the spirilla this occurs too, whether the flagellum arises at the anterior or posterior pole." If they "always" rotate in one direction, how can they "simply reverse" this rotation? Further, if the flagellum is directed backward during locomotion, whether the flagellum be at either end of the body, does this not indicate that the flagellum is assuming a different orientation, rather than that it is merely rotating to the right or to the left? For, if it simply reversed its direction of rotation in order to back up, it would then pull the body instead of pushing it, and would not be directed backward. The error doubtless lies in the assumption that the organism moves backward by simply reversing the flagellar rotation. Indeed, the theory itself seems disputed by these facts.
Gray (1928, p. 35) gives a fine explanation of the mechanics involved in undulating flagella, the gist of it being that there is a series of waves passing along the flagellum in the same direction, requiring no recovery stroke. The propulsive power "... is equivalent to that which would be produced by projecting along the length of the flagellum a series of 'humps' of the same form as the waves, the velocity of the humps being made equal to the velocity of movement of each wave. If the waves pass from the base of the flagellum to its tip, the organism is driven forward in front of the flagellum; if the waves pass from the tip to the base the organism is drawn forward with the flagellum in front. If the waves pass along the flagellum in one plane there will be no force tending to rotate the animal on its axis: if, however, the waves pass round the flagellum as well as along it the organism will rotate." This is well and good, but it fails to fit a couple of facts; namely, that most flagellates move flagellum-foremost, and that the waves of transmission along the flagellum have only been demonstrated to progress from the base toward the tip (Lowndes, 1936, 1941 a and b, 1943, 1944 a and b, 1945 a and b). To date, Lowndes has made the most complete and accurate analysis of the situation in a number of flagellates. His conclusions are based upon observed facts and specific data, backed up by high-speed cinema photomicrography (for techniques, see Lowndes, 1935, 1944 a). Too much of previous explanations was based upon pure speculation and rationalization. Lowndes (1941 b) has demonstrated that the flagellum ordinarily pushes the organism, even though situated at the anterior end, by being directed backward. In this connection it is interesting to consult Ulehla's figures, many of which indicate the same. This appears true even for Peranema, the organism classically cited as having a "tractellum," or flagellum which pulls the organism forward, presumably by a wave starting at the tip. Lowndes shows that, in all cases studied, the wave invariably progresses from base to tip. This fact is an upsetting one to most of the previous theories. Among other workers reporting this type of action are Kunstler (1889), Friedrich (1909), Ulehla (1911), Bancroft (1913), and Schindera (1922). Of these, Ulehla and Bancroft give accounts almost precisely like that of Lowndes concerning the locomotion of Euglena viridis.

In his later papers, Lowndes elaborates a new concept, previously mentioned by him, but misunderstood by such workers as Barker (1943). In the summary of one of these papers (Lowndes, 1944 a), he states it as follows:

1. The primary function of the flagellum in a monoflagellate organism is to produce both rotation and gyration of the organism about a certain axis which constitutes the main direction in which the organism is swimming.

2. The mechanical principle by which the organism is propelled is simply that of the inclined plane which is caused to rotate. In other words it is that of the screw or propeller.

3. Since the disturbances or waves pass down the flagellum in the form of a spiral they produce two distinct components. It is the resultant of these two components which causes the tip of the organism both to rotate and gyrate.

4. So long as this rotation and gyration is maintained it will supply the necessary force for the propulsion of the organism.

5. The flagellum itself may or may not produce a forward component. If it is more or less swung out at right angles, as in Menoidium, it will produce no forward component but if it is swung back, as in Euglena, it will do so."

Hence, it is the rotation and gyration of the body of the organism which is considered to produce the major component of force. Others have observed, naturally, that in organisms bearing obvious helical external structures, rotation of the body would produce a component of force. For instance, Günther (1928) correlated the speed of several species of Euglena with spiral ridges on the periplast and flagellar length. Thus, of two species having flagella of approximately the same length, one having strong spiral ridges progresses at about three times the relative
speed of another having very insignificant ridges. But no one previous to Lowndes, so far as I know, had suggested the possibility that the gyration of the body itself might produce locomotion.

Propulsion by simple conical gyration is what Delage and Hérouard thought they had proved, by mathematics, to be impossible. Metzner (1920) showed this to be erroneous, both in theory and in practice. By means of experiments on wires rotating in water, he demonstrated that a simple conical gyration produces a pulling force which reaches maximum at an angle of 20–23 degrees from the axis of rotation. He even went so far as to state (translation): "Among flagellates, the simple conical gyration predominates; the organism ‘sucks’ itself along through the water by means of the flagellum." It should be noted that he considered only gyration of the flagellum, not gyration of the body. Such conical gyration of the flagellum is reported by Keysselitz (1906) and Petersen (1929). It is also described by Lowndes (1944 b), who performed experiments similar to those of Metzner. However, Lowndes contends that this mode of locomotion is incompatible with high speed, as the flagellum could not be maintained in such a forwardly-directed position. In rapidly moving organisms, the flagella must beat laterally to the rear.

In this same paper, and in his most recent one (1945 b), Lowndes renders untenable a number of misconceptions which arose long ago through misinterpretation of the original data, and have grown or continued from textbook to textbook. One such item regards diagrams of Monas swimming, taken from Krijgsman (1925). I shall not discuss these errors here. As Lowndes points out, Krijgsman's observations were excellent records of the situation he was studying, but that situation was not representative of the free swimming of the organism. When the animal is freely swimming, the mechanism is quite different from the paddle-stroke observed by Krijgsman, being similar to that described by Lowndes for Euglena, and the rate of progression is about ten times as rapid as that given by Krijgsman. Locomotion is primarily brought about by the rotation and gyration of the body of the organism. (See also Lowndes, 1945 a.)

As for the question of the precise nature of what goes on within the flagellum—it remains a question. For years, writers were occupied with a controversy as to whether the flagellum was an active unit or simply an appendage like a whip, manipulated from the base or cell body. The latter concept was eventually discarded as such, but lingers in part in the question whether or not the flagellum can execute any movement after losing connection with the cell. Various workers (e. g. Klebs, 1892; Entz, 1926) have reported the phenomenon; others have flatly asserted such to be impossible (Verworn, 1890; Korschikov, 1923). However, the work of many investigators has indicated that the flagellum is an active unit. Gray (1928) enumerates good reasons for assuming such, and Lowndes (1936, 1941 b, 1945 b) proves it with photographic, quantitative evidence. Granting, then, that the flagellum does possess the ability to move, by what mechanism does it effect this motion?

From the versatility of movements and also "... because its movements at times cannot be explained by simple mechanical laws," Krijgsman (1925) concluded that the flagellum "... must be of complicated construction." Engelmann (1881) and Ballowitz (1890) assert that movement is associated with fibrillar structure, which, as we have seen, is evidently present. Erhard (1910) and Lowndes (1941 b) regard the transmission of the stimulus as a surface phenomenon, not connected with the axoneme. Such statements are generalities, to be sure, but, if accepted, eliminate such theories as that of Gurwitsch (1904), which postulates a rapid streaming of protoplasm into and out of the flagellum, acting in antagonism to the elasticity of the axoneme. Heidenhain (1911) proposed a "theory of smallest waves" to explain movements: these waves may travel along one side of the filament, they may vary in length and effect, they may vary in extent, they
may vary in path, frequency, and rhythm. They are evidently quite versatile waves. So far as I know, they have been forgotten. Pütter (1903) discussed Engelmann's theory and decided that it required too many spiral fibrils of different pitch and direction. Verworn (1915) stated what he considered the basic principle involved (translation): "...a contractile side contracts from the cell body outwards and thereby stretches the opposite side, which latter in the expansion phase, by its elasticity, returns the filament to its rest position. According to the opposing relationship of the contractile to the passively stretched substance, there results a beat in one plane or in a complicated form." This does epitomize the principle employed by most of the theories. For a discussion of the theories, see Gray (1928).

Viewing the matter from a more generalized standpoint, but with fundamentals in mind, Gray (1928) and Schmitt (1944) liken the process involved to that of muscle, and consider the basic phenomenon to result from a change of distribution of water molecules "between polar groups in fibrous proteins and ionogenic groups in the environment" (Schmitt). In other words, if the proteins in the fibers along one side of the flagellum suddenly take on great numbers of water molecules at the expense of the opposite side of the filament, the filament (flagellum) will tend to bend toward the side losing the water. This sudden affinity for water by the protein might be due to a change in the degree of ionization of the protein molecule, caused by a local production of acid. Should the acid then be neutralized, the water would again be liberated (Gray). Or it might be that the protein fiber consists of folded molecular chains which fold up further upon the addition of water, thus contracting the fiber. At any rate, we get glimpses of the possibilities.

**Original Investigations**

**Structure of the Flagellum**

**Materials and Methods**

It was considered desirable to experiment briefly with the more productive methods described by other workers in the study of flagellar structure. Such techniques as those of Gicklhorn (1921), Kater (1929), Allen (1936), and Smyth (1944) are useful procedures for general study or mere observation of flagella, but contribute relatively little to a detailed investigation of the type here undertaken. On the other hand, the papers of Loeffler (1890), Fischer (1895), and Deflandre (1923) describe techniques of primary importance in the development of our knowledge of whip- and flimmer-flagella. McClung (1937, pp. 141–145) presents Loeffler's method and may be more readily available to persons attempting to duplicate some of the work done on flimmer- and whip-flagella. Vlk (1938) describes his modifications of the Loeffler technique. Petersen (1929) points out the possibilities for failure and sources of error, emphasizing the extreme delicacy required in preparation. Several authors (e.g. Batschli, 1902; Korschikov, 1923) have failed to achieve success with the mordant technique.

Such contributions as this paper may offer to the knowledge of flagellar structure are made possible through the use of the electron microscope. Further probings with the light microscope promise but little, as the possibilities have been rather carefully covered. For previously employed methods of preparation of specimens for electron microscope examination, see Marton (1941, 1943), Morton and Anderson (1942), Mudd and Anderson (1942), Richards and Anderson (1942), Prebus (1944), Williams and Wyckoff (1944, 1945 a, b), Claude and Fullam (1945), and Porter, Claude, and Fullam (1945). Some of the techniques described are adapted to specialized situations, but they bring out various points of importance in the preparation of biological material for electron microscope study. Among the most important of these, perhaps, are the removal of salts and other matter which might crystallize upon or otherwise obscure or mar the specimens, and the thinness
of film plus specimen, which should total less than 50 \( \mu \) in thickness. A good
discussion of the limitations of the electron microscope for biological work in
general is given by Richards and Anderson (1942). For basic principles and
general limitations of the instrument, see Hillier and Vance (1941), or such pop-
ularized books as Burton and Kohl (1942) or Hawley (1945).

Dellinger (1909) and Porter, Claude, and Fullam (1945), among others, consider
osmic acid the best killing and fixing reagent for retaining original appearances. I
have found it extremely useful in this work. Over a dozen different procedures
have been employed, the most successful of which have all involved the use of
osmic acid. Formalin as a fixing reagent has apparently resulted in excessive
clumping, but this has not been thoroughly investigated. Other major difficulties
in the preparations were due to too many or too few organisms on the film,
crystallization, loss of flagella in centrifuging, and, most of all, disruption of mem-
branes or films under electron bombardment.

The principal organisms studied were *Astasia klebsii* Lemmermann, *Euglena
gracilis* Klebs, *Ochromonas variabilis* Meyer, and *Chilomonas paramecium* Ehren-
berg which were obtained from pure cultures maintained in the Protozoology
Laboratory at The Ohio State University.

The film and screen upon which specimens were to be mounted was prepared as
follows (slight modification of method employed by Prebus):

1. 300 mesh/inch bronze screen is cleaned by immersing in dilute HCl and
washing a number of times in distilled water.
2. Rectangular pieces about 6 x 18 mm. are cut from the screen and kept
dust-free (e. g., in a Petri dish).
3. A vessel (e. g., a Pyrex pie plate) of about 10-inch diameter with a level
rim is filled to the brim with distilled water.
4. The water is saturated with amyl acetate.
5. A drop or two of 2% collodion (cellulose nitrate) in amyl acetate is allowed
to fall upon and spread over the surface of the water.
6. After a few seconds a film is left, subsequent to the evaporation of the
solvent. This film is then swept off by means of a glass rod, in order to clean the
surface of the water. This may be repeated if it seems advisable.
7. A drop of the collodion solution is allowed to fall an inch or less onto the
surface of the water.
8. A rectangular piece of screen, held in an artery clamp, is immersed in the
water near the rim of the vessel, moved over beneath the film, and raised directly
up. If parts of the film do not tear neatly around the edge of the screen, they
may be severed with a needle tip by running it along the edge of the screen.
9. Excess water is removed from the screen and film by touching the droplets
with a folded corner of filter paper.

The film thus mounted on the rectangle of screen may be used immediately or
allowed to dry before having specimens placed upon it. Immediate use seems
preferable.

Of the techniques employed in the preparation of specimens, the following
were most successful:

1. (a) Place 4.5 ml. of a pure culture of the desired organisms in a centrifuge
tube.
   (b) Add and mix 0.5 ml. of 2% osmic acid or 2% osmic acid in 1% chromic
   acid.
   (c) Add 5 ml. of distilled water and centrifuge 30 seconds.
   (d) Decant, then refill to 10 ml. mark with distilled water.
   (e) Re-suspend organisms by rolling tube between palms of hands.
   (f) Centrifuge 30 seconds.
   Repeat (d) to (f) inclusive once more.
(g) Decant, then add distilled water to raise meniscus to the 0.5 or 1.0 ml. mark. Again suspend organisms by rolling the tube vigorously between the palms.

(h) Place a drop of the liquid containing the concentrated, washed organisms upon film-screen and allow the organisms to settle for 30-60 seconds.

(i) Carefully remove some of the excess liquid from the top of the drop by touching to it the folded corner of a piece of filter paper.

(j) Allow to dry, then examine under compound microscope for promising specimens. An average of about 1 organism per "pane" or mesh of the film is the optimum concentration of specimens. If much more numerous, the film does not support them when under electron bombardment. If much less numerous, the specimens are too difficult to find in the electron microscope.

(k) Place screen, specimen side down, upon a rectangular piece of heavy onion-skin paper which is conveniently larger than the piece of screen. With a screen rectangle of 6 x 18 mm., a paper rectangle of about 12 x 30 mm. is of convenient dimensions. (See Fig. 17.)

(l) Flatten the screen against the paper and secure it there by means of small pieces of Scotch tape (thin, transparent, cellulose adhesive tape).

(m) Label one end of the paper for identification and reference to procedure, as in Fig. 17: "E-6, HB."

(n) Place in a Petri dish or between concavity slides to keep perfectly clean, and remove to the electron microscope laboratory.

(o) Using a die constructed for the purpose, punch out circular disks of screen about 2 mm. in diameter.

(p) Separate disks of screen from those of paper and examine under compound microscope (on a clean slide).

(q) Select promising disks for electron microscope examination.

2. The following method, since it does not involve centrifugation, subjects the flagella to less drastic treatment, but requires a greater initial concentration of organisms in the culture. Various modifications of this method were employed, some eliminating step (b).

(a) Fill depression of a concavity slide with an especially rich pure culture.

(b) Add several drops of Gicklhorn's stain (made by adding about 5 drops of concentrated NH$_4$OH to 50 ml. of 0.05% methylene blue solution).

(c) Let stand about 1 minute and add 1 drop of 2% osmic acid in 1% chromic acid solution.

(d) Allow organisms to settle for 30-60 seconds, then touch the surface of the liquid with a folded corner of filter paper to remove excess liquid.

(e) After removing a large part of the liquid as above, carefully replace this liquid with distilled water, by means of a pipette.

Repeat (d) and (e) at least 5 times, to wash the specimens of acid and of salts, extraneous material, etc.
(f) Observe under a compound microscope (16 mm. objective) while drawing organisms into a micro-pipette. They will be concentrated in certain areas at the bottom of the drop.

(g) Deposit by means of this pipette as many of the organisms as possible in a medium-sized drop upon a film-screen mount.

(h) Allow organisms to settle, then remove excess liquid as in (d). Continue as in (j) to (q) inclusive, technique No. 1.

Both of the above techniques were essentially developed by the author in the course of this work, as were various other less successful ones.

Results

Upon the development of satisfactory techniques, it was possible to demonstrate a number of details relative to the structure of the flagellum. Some of the more satisfactory and instructive of the electron micrographs obtained are shown in the appended plates. In viewing these photomicrographs, the reader should bear in mind that the objects and details perceptible are not rendered so by staining, as in preparations for the light microscope, but are visible because they have diffracted or absorbed varying proportions of the electrons passing through them. Thus a body appears dark because it is denser or thicker, or both, than the surrounding substance, and not because it has been differentially stained. From a careful examination of these plates, the following points may be observed:

1. Each flagellum is of approximately uniform diameter throughout its entire length (Plates 4; 6; 9; 10; 11; 12).

2. Each flagellum consists of a denser axial core (axoneme) and a less dense sheath surrounding the core (Plates 2 to 8 inclusive).

3. In the flagella of *Euglena* and *Astasia*, the axial core appears to consist of two closely approximated fibers of equal size (Plates 1 B; 2 A; 4; 5; 6; 9).

4. The sheath appears to contain or to consist of a coiled fiber which encircles the axial core in the form of a helix (Plates 1 B; 3 A; 12).

5. The flagella of *Euglena* and *Astasia* bear, along one side, what appears to be a single row of delicate filaments extending from the sheath. The length of the filaments is about 5 or 6 times the diameter of the flagellum, or 1.5 to 2.0 μ. (Plates 1 B; 4; 9).

6. The long flagellum of *Ochromonas* bears similar filaments along both (all ?) sides (Plates 11; 12).

7. The flagella of *Chilomonas* bear no such lateral filaments (Plate 10).

It is possible that the ground substance or intermediate substance which might be expected to occur between axial fiber and sheath, perhaps comprising a large part of the bulk of the living flagellum, has escaped in most of the specimens shown. Consideration of Plate 7 might lead us to this conclusion, assuming the darker upper portion of the flagellum to represent a region which, somehow, had not yet lost the inner plasm. Perhaps the swollen appearance in Plate 8 is due to an accumulation or exudation of such matter during the drying of the specimen. From Plate 7 we also get an indication that the lateral filaments might possibly be due to the escape (and subsequent coagulation) of plasm from a lateral series of minute pores. In view of the work of Vlk, who demonstrated flimmer upon a living flagellum, this explanation might be held to apply to the possible origin of flimmer in the living organism. In such a case, complete coagulation might not occur until death.

A second possible explanation for the appearance shown in Plate 7 is that a portion of the flagellum might have contracted, bringing the coils of the helix into closer approximation and producing the denser and thicker appearance seen in the upper portion of the flagellum shown. It is difficult to determine precisely what actually happened to produce this effect, and the interpretation of such an appearance must remain uncertain for the present.
Upon inspection of the plates, it will be noted that, as stated in (1) above, the flagella depicted are of approximately uniform diameter throughout their entire length. Little, if any, tapering occurs at either extremity. Emmel, Jakob, and Götz (1942) report the same condition in electron micrographs of *Leishmania donovani*.

Of the other points listed above, (2) simply substantiates the generally-accepted notion of flagellar structure. (4) serves to emphasize the similarity between the protozoan flagellum and the mammalian sperm tail, as described by Schmitt and others. (5), (6), and (7) corroborate, by an entirely new technique, the findings of a number of workers, extending somewhat the knowledge of the details.

Unfortunately, many of the best "shots" were lost, due to the rupture or the supporting film or the sudden curling up of flagella under the impact of the electron bombardment, before photographs could be taken. However, the fact that a flagellum, even though flattened out against the film, in a high vacuum (0.00001 to 0.0001 mm. Hg.), and dead for many hours, can retain within itself the potential ability to tear itself loose and curl up like a watchspring, may be significant. Similar coiling during the disintegration of flagella has been described by several investigators (e. g., Fischer, 1894), but never with violence like this.

The photographs included in this paper were, of course, selected to bring out various points. Plate 6 shows an entire flagellum, about 13 \( \mu \) in length, extending from the body of the euglena. The flagellum is seen to extend, not from the anterior tip of the organism, but from a point slightly lateral and posterior to the tip, where it emerges from the mouth of the gullet. Plate 6 also shows the naked axoneme, where the supporting film has torn, snapping the flagellum off near its base. Plate 2A probably represents a flagellum lying adjacent to the body, and extending away from it near the posterior tip of the body. Plate 2B gives an indication of the relative strength or durability of the axoneme and the sheath, the axoneme being apparently stronger and more elastic.

Plates 4 and 9 display especially well the single flimmer-row on the flagellum of *Euglena*, and represent flagella prepared by different techniques. Plates 11 and 12 show flimmer along both sides of the long flagellum of *Ochromonas*, which might, in life, occur either in two opposite rows or all over the surface of the flagellum like the hairs on a dog's tail. It will be noted that the flimmer do not occur in the neatly regular rows depicted by previous workers. This may be accounted for by the fact that these specimens have been centrifuged several times in preparation, a drastic measure not employed by previous workers. Such a disarrangement should be expected.

Plate 9, in portions depicting twisting, shows clearly the two major fibers of the axoneme in the flagellum of *Euglena*. Plate 3A indicates the helical structure in the sheath rather well. Such plates as 2, 5 and 7 also suggest this helical structure. Other plates, in which the sheath does not stand out perceptibly from the axoneme, may be of interest in that there occur at more or less regular intervals along the sides of the flagellum dark spots which may represent the helix closely appressed to the axial core. Plates 1B, 9, 11, and 12 show such indications. If this is the correct interpretation of such appearances, the pitch of the helix on the flagellum of *Ochromonas* is considerably greater than that of *Euglena* (i. e., the coil is less tightly wrapped, forming longer spirals). Further, if this be the correct interpretation, the spots shown in Plate 9 will be of especial interest. Many of them are lighter in the center, indicating that the coiled fibril may be hollow or tubular.

Plate 1A, which is an electron micrograph of a diatom, is included to show our method of computing sizes. Large numbers of this type of diatom, a species of *Gomphonema*, were collected several years ago, cleaned, and calibrated. Lateral transverse rows of pores may be seen extending from a median longitudinal solid
area like the barbs in the vane of a feather. The pinnate rows along one side of the median line are usually more nearly perpendicular to the axis of the median line than are the rows along the other side of the line. The average distance between consecutive rows is about one-third of a micron. We obtain measurements by averaging the distances between every fourth row. This average is about 1 μ. After taking a series of micrographs at a given magnification, a diatom is photographed at the same magnification, in order to provide a scale for measurement. This method and the initial calibration of the diatoms were worked out by Dr. Prebus.

MECHANICS OF THE FLAGELLUM

Observations on Living Organisms

Flagellar action is, in most cases, very difficult to observe in normally-moving or freely-swimming creatures. The flagellum is hard enough to see when still, and when in active motion is beyond the capabilities of the human eye. For this reason, most studies on living flagella have been made on organisms under abnormal conditions. They have been chilled, anesthetized, compressed, placed in viscous media, or simply observed in the latter stages of approaching death, when the water beneath the cover glass was drying up. Realizing that normal activity is hardly to be expected under such circumstances, yet assuming that certain basic phenomena should remain constant, I have made a few observations under some of the above-mentioned conditions.

The most convenient method I have found for rendering flagellar motion visible involves the use of methyl cellulose (Methocel, Dow Chemical Co.). A drop of 10% solution of this substance is mixed on a slide with a drop of culture, then a cover glass added (Marsland, 1943). The resulting mixture, of rather high viscosity, slows down the strokes of flagella or cilia, and also is of a very different refractive index from water, such organelles becoming much more easily visible than in water.

Among the structures observed by this method were the flagella of Peranema trichophorum, Euglena gracilis, and Trichonympha sp., the undulating membrane of a trichomonad from the gut of Reticulitermes flavipes, and the cilia of Paramecium sp. The optical system employed included a Spencer 4 mm. objective (N.A.—0.85) and a 20 X Planoscopic ocular, with a resulting magnification of about 880 X. In every case, the wave impulse traveled from the base toward the tip, in a spiral course, producing rotation of the tip. All of these observations directly confirmed certain conclusions of Lowndes (see historical review). In Euglena the flagellum was usually directed back more or less along the body. I was somewhat surprised to find this sort of movement in cilia, as I expected to see the paddle-stroke described by Gray and others. However, the cilia were observed to alter the direction of their strokes quite readily, beating forward, directly outward, or backward (and toward or away from the observer). The spiral, flagellum-like stroke or undulation was most conveniently observed when the cilia were beating directly outward, or away from the body surface. The cilia beating thus created a current away from the body. I have wondered whether this spiral undulatory stroke in cilia might be due to the greater density or viscosity of the medium employed in these experiments. Cinematic photography of such cilia under more nearly normal conditions should aid in clearing up the matter. Alverdes (1922) made an extensive study of ciliary movement in several species of Paramecium, Stentor, etc. He ascribed to the cilia considerable versatility of movement. He also described interesting experiments on the shedding and regeneration of cilia by Paramecium. He kept the organisms in a 0.1% solution of chloral hydrate for about 48 hours, then transferred them to fresh water, and, after 3 to 9 hours, observed the regeneration of the cilia. The cilia began beating when only stubs.
This technique, combined with good cinematic photomicrography, might produce very interesting results.

I have observed, without altering the medium in any way, flagella in colonies of *Volvox* which seemed capable of performing almost any movement possible for a filament attached at one end. Since the colony was probably suffering under adverse conditions, the movements probably were not normal, but they certainly served to emphasize the versatility of movement possessed by the flagellum. To quote again from Krijgsman's summary (translation): "... its movements at times can not be explained according to simple mechanical principles." It is too easy to agree with him.

**Experiments on Locomotor Mechanisms.**

"**ARTIFICIAL FLAGELLATES.**" In order to test the forces produced by rotating and gyrating objects, a device was worked out as shown in Fig. 18, whereby structures comparable both to bodies and to flagella of flagellate organisms could be studied in this connection. Originally it was devised for comparison with a flagellum, but when the significance of Lowndes's hypothesis became apparent to me (upon the receipt of his later papers), it was extended to a study of body gyration.

This hypothesis, it will be recalled (see historical review), suggests that the major component of force producing the forward locomotion of a monoflagellate results from the rotation and gyration of the body of the organism, and not directly from the action of the flagellum. Consequently, a model was constructed in the shape of a sample protozoan (e.g., see Fig. 22) in order to test the locomotor effect produced by the rotation and gyration of such a body. As figured in the accompanying diagram, *B* represents this body, with *a* representing its axis. The arrows encircling the axis of forward progression *A* serve to indicate the path of gyration of the body axis *a*. All of the rest of the diagram below the body *B* is simply included to show how the rotational and gyrational force is applied to *B*.

Thus in Fig. 18, *A* represents the axis of gyration and progression about which the cork body *B*, with axis *a*, is caused to rotate and gyrate. In the diagram, the body is gyrating clockwise as seen from the base or rear. (Throughout

![Fig. 18. Diagram of the "artificial flagellate." *B* represents a model of a monad body, which rotates and gyrates about its axis of progression, *A*. *a* represents the body axis. The lower part of the diagram merely indicates the apparatus employed to impart to the body *B* its rotational and gyrational force. See text for explanation and discussion.]
this paper, when the terms clockwise or counter-clockwise are employed, reference is made to the rotational path of the distal gyrating extremity as viewed from the apex of the gyration cone. A heavy rubber band, $F$, twisted in the desired direction, produces the rotation which, in turn, causes the part of the wire bent out of line to gyrate. It was often found desirable to use two rubber bands, in order to obtain greater speed and force. Beads below the bend in the wire served as bearings, and turned in the funnel-like flared end of a metal tube which was inserted in the cork stopper $C$. Since the rotation and gyration of $B$ caused the base ($C, D, E,$ etc.) to rotate and gyrate in the opposite direction, it was found necessary to reduce such rotation considerably; otherwise, the rubber band rapidly became untwisted. Two razor blades, $D$, inserted in the cork parallel to axis $A$, served as fins or keels in reducing rotation of this portion of the system. In order to balance the system to a specific gravity slightly above 1.0, water was added to test tube $E$ in the necessary amounts. Paraffin was found to be less satisfactory in achieving this balance. The angle ($\beta$) between $A$ and $a$ was altered simply by bending the wire. The body, $B$, may be replaced by other objects of diverse shapes and sizes.

1. Currents produced in fluids by gyrating structures. Experiments were performed with the "artificial flagellate" using smoke, in air, and minute suspended particles, in water, to observe the currents produced by the gyration of $B$.

![Diagram of currents produced by gyrating wires in water.](Redrawn from Metzner, 1920, Figs. 3, 4.)

The results of these experiments, which were not extensive, were in accord with those of Metzner (1920). In brief, if the path of gyration forms a simple cone, fluid is drawn into the cone primarily from its base, departing near the apex (Fig. 19). Thus a major current is produced toward the base of the cone in the surrounding medium. If the body were free to move, it would progress toward the base of the cone. Lowndes (1944 b) has also demonstrated rather fully the action of similar objects. For theoretical considerations as to the mode of function, see Metzner, (1920, pp. 53-58).

2. Locomotion of gyrating systems in water. A number of experiments were performed to determine the velocity achieved by the "artificial flagellate" in water, relative to the shape and position of the body $B$. In each case, both clockwise and counter-clockwise gyrations were tested, in order to eliminate such factors as the possibility that the razor blades ($D$) might be acting as screws or propellers. These experiments were carried out in a greenhouse tank (200 x 80 x 60 cm.).

It was found that when structures much larger than $B$ were placed on axis $a$ to gyrate, the rear end of the test tube $E$ was caused to gyrate considerably. Since this would exert a force in the opposite direction, it was objectionable. By placing
around the test tube a fairly close-fitting, rigid jacket, this gyration may be mini-
mized. A coarse screen or hardware cloth is perhaps best, although a test tube
slightly larger than \( R \) was first used for the purpose. In the latter case, there is
too much difficulty involved in the movement of water to fill the space vacated by
the progressing system. In the set-up as shown in the diagram, the gyration of
the test tube was sufficiently unimportant to be neglected in the gross observations
being made.

Velocity was measured horizontally and vertically. The latter measurements
represent much freer motion on the part of the “flagellate,” but are more difficult
to obtain under the conditions encountered. The “organism” is balanced so that
it sinks rather slowly. Then, wound up, it is held down, allowed to gyrate a few
times, and timed on its way up (its stable position is in the vertical axis). For
the most part, it was timed through a distance of 20 cm. A stop watch was used
in all cases. Horizontal runs were made by placing the test tube in some such
jacket as mentioned in the preceding paragraph and holding the jacket steady in
the horizontal plane. By the nature of this set-up, such runs were confined to a
distance of 5 or 6 cm. Since, in every case, \( B \) was buoyant, the gyration was
uneven in this plane. Another factor necessitating brief runs, both horizontally
and vertically, is that the force producing the gyration diminishes rapidly as the
rubber bands untwist. In consideration of these, and perhaps other conditions,
it is obvious that the measurements are necessarily inaccurate. To assume an
error of \( \pm 10\% \) would be optimistic. However, the figures are at least indicative,
and have some value thus.

Relative speed, as here employed, equals distance traversed by the organism in
one second divided by the length of the gyrating body. Thus, if the body \( B \) were
5 cm. long and the system moved at 10 cm./sec., the relative speed would be
10 ÷ 5, or 2. Among Protozoa which have been actually timed for rate of swimming,
the relative speeds vary from 0.25 (\( E\)ug\( e\)\( n\)\( a\) \( t\)\( e\)\( r\)\( i\)c\( o\)la, \( G\)un\( t\)h\( e\)r, 1928) or less, to
40.0 (\( M\)onas \( s\)t\( i\)g\( m\)\( a\)\( t\)i\( c\)a\( t\)i\( c\)a, \( L\)ow\( n\)d\( e\)s, 1944 b, 1945 a) or more.

With a long body (\( B \)), 14 x 1 x 1 cm., at 1 to 1.1 gyrations/second, traveling
horizontally out of a vial, with angle \( \beta \) at 23°, the maximum constant velocity
observed was 1 to 1.2 cm./sec., representing a relative speed of about 0.08. Using
the same set-up, but with angle \( \beta \) greater than 90° (see Fig. 19 b), the maximum
velocity was 0.33 cm./sec. This condition hardly corresponds to any natural one.

With a short body (\( B \)) as shown in the diagram, 4.7 x 3.7 x 2.4 cm., at an
estimated 6 gyrations/second, traveling vertically, with angle \( \beta \) at 29°, the max-
imum constant velocity was 25 cm./sec., representing a relative speed of about 5.
With angle \( \beta \) at 15.2°, a maximum velocity of 33 cm./sec. was observed. This
was under ideal conditions and was never quite duplicated. It represents a
relative speed of 7, the highest obtained in these experiments.

To test the forces produced by a flagellum undulating in a helix or spiral, the
body, \( B \), was made in such a shape by bending a wire and coating it with 1.5 to 2.0
mm. of paraffin. The form was approximately that shown in Fig. 16, but with a
less complete pitch. The complete pitch would be about 20 cm. The flagellum
rotated at about 18 turns/sec. With the flagellum rotating counter-clockwise, the
system moved forward at 34 to 35 cm./sec. With the flagellum rotating clockwise,
the system moved backward at 20 to 22 cm./sec. If a living flagellum beats from
the base outward, the latter is the only type of motion compatible with the system.
In order for an actual flagellum to execute a movement similar to the former, the
wave of contraction would have to begin at the tip of the flagellum. If this
occurred, it would constitute a “tractellum.” It probably does not occur in nature.

From our experiments with the “artificial flagellate,” we learn that the mere
rotation and gyration of a body in water can provide sufficient force to produce
rapid forward locomotion of the body. This greatly strengthens the hypothesis
advanced by Lowndes (1944 a).
UNDERWATER SWIMMING. These experiments test, in a fashion, the strength of the "pull" exerted by gyrating structures. In contrast with flagellate bodies, but like the flagella themselves, the gyrating objects do not necessarily rotate. The arms of the swimmer serve as the gyrating structures. The rate of gyration is approximately 1/sec. The gyrating portion is 60 cm. in length. Figures on velocity are computed from the distance traversed in about one-half minute. The body weight of the swimmer is about 130 lbs. or 59 kg. In each case, enough air has been expelled from the lungs to allow the body to sink to the bottom of the pool. All experiments were performed by the author. Timing was done by an observer with a stop watch.

(1) The body is horizontal, with one arm extended horizontally forward and gyrating, e.g. clockwise, in a relatively narrow cone. (Fig. 20 a) Result: the body moves horizontally forward, rotating counter-clockwise, in this case. Velocity, 10+ cm./sec. Relative speed, 0.16+. Total distance progressed, 10 ft.

(2) The body is vertical, with both arms extended horizontally forward, the right gyrating clockwise, the left counter-clockwise or vice versa. (Fig. 20 b) Result: the body moves horizontally forward, not rotating. Velocity, 10+ cm./sec. Relative speed, 0.16+. Total distance progressed, 10 ft.

(3) The body is horizontal, with both arms extended horizontally forward, the right gyrating clockwise, the left counter-clockwise. (Fig. 20 c) Result: the body moves horizontally forward, not rotating. Velocity, 33 cm./sec. Relative speed, 0.55. Total distance progressed, 32 ft. Reversing the gyrations, the velocity is considerably less, being at most about 23 cm./sec., with a relative speed of about 0.39. Total distance progressed, 22 ft. This may well be due to an unintentionally weaker stroke, as it is more difficult and tiring to the experimenter.

(4) The body is in the same position as (3), but both arms are gyrated in the same direction. Result: the body rotates in the opposite direction at approximately the same rate, little forward movement being accomplished.

(5) The body is in position (3), but the arms are pendulated or swung back and forth in one plane, instead of being gyrated. Result: no detectable forward component.

Minor components are omitted in this consideration, for the sake of simplicity. In many cases they are results of the musculature of the arm. Other experimental positions were tried, but contribute no additional significant data and are hence omitted.

The results of these experiments provide additional evidence in support of Lowndes's hypothesis. They serve further, however, to show that rotation of the gyrating object is not necessary to the production of a forward component. Mere gyration of an object (an arm or a flagellum) can produce an effective locomotor force.
DISCUSSION

From the historical review given in this paper, it is apparent that a considerable mass of knowledge has been accumulated on the subject of flagellar structure, but that, in large part, the data have not previously been assembled and organized. Certain phases of the subject, to be sure, have been well summarized. For instance, Vlk (1938) treats whip- and flimmer-flagella as such rather thoroughly, but neglects the internal structure. However, he is not to be criticized for this, as little new knowledge has appeared relative to the matter within several decades. The subject has awaited a new technique which could permit of more minute investigation. The electron microscope provides this new angle of attack through its much greater resolution and magnification. This paper presents the results of the first intensive study of the protozoan flagellum employing the electron microscope.

By means of this instrument, we find that the flagellum, at least in the forms studied, consists of a dense fibrillar axial region and a surrounding sheath of much less density. This much has been previously accepted, though never before so clearly demonstrated. Just how many fibrils there are comprising the axoneme, I cannot tell. The number probably varies to some extent. The sheath seems to contain a fiber which encircles the axoneme in the form of a helix. This sort of composition of axoneme and sheath appears to be in close agreement with the structure of mammalian sperm tails as described by Schmitt and others, but has never before been described in the protozoan flagellum, unless the brief remarks of Uleha (1911) be considered such. Unlike the sperm tails, certain flagella possess lateral or terminal external extensions of the sheath. These structures have been recorded since 1889, but are not yet universally accepted by protozoologists. Several electron micrographs of the lateral structures in question (the flimmer) are included in this paper (e.g., Plates 4, 9, 11, 12). Of course, such pictures constitute no conclusive proof of the nature, or even natural occurrence, of the flimmer. But they certainly contribute to the evidence in favor of their acceptance as normal structures, rather than as artifacts. If Vlk had had at his disposal the equipment of Lowndes (1935), Harvey and Loomis (1931), or Pipper (1940), perhaps the question might have been settled for the most skeptical by the actual photographing of flimmer on living flagella. However, the facts that they have been demonstrated by a number of basically different techniques, on dried or moist flagella, following various fixatives or none at all, and that they are amazingly constant in appearance, position, etc., place the overwhelming weight of evidence in favor of the view that they represent normally-occurring structures in many types of organisms. The presence and type of flimmer or whip comprise valuable taxonomic characters, and will doubtless be so considered when they are more readily demonstrable.

There are a number of devices as yet untried in the study of these structures, some of which should certainly contribute to our knowledge of their nature. I mention them in the hope that someone, to whom they may be available, may be interested in furthering this study. By means of the shadow technique of Williams and Wyckoff (1944, 1945 a, b), the flimmer should be made obvious. A study of living flagella might be made with the ultra-violet microscope (Lucas, 1930, 1934; Schmitt, 1939; Lavin and Hoagland, 1943), whereby twice the resolution of the ordinary microscope might be obtained. Phase difference microscopy (Richards, 1944) offers interesting possibilities in the study of living flagella, yielding the benefits of stains and fixing reagents, as does the ultraviolet microscope, without requiring the use of such deleterious agents. (That is, it serves to differentiate structures and substances which we commonly bring out only by staining.) Fluorescence microscopy (Ellinger, 1940; Metcalf and Patton, 1944) also gives promise, especially, perhaps, in the study of flagellar motion. If nat-
urally fluorescing flagella occur, these would be ideal for such a study, but a fluorescent vital stain such as Fluorescein might serve well.

It is interesting to view the results of Kunstler and Fischer in the light of details brought out in electron micrographs. If the reader will again refer to Fig. 1, he may notice certain similarities to some of the plates. The interpretation of the appearances, however, is rather different. The axial canal of Kunstler is probably the axoneme. Plate 7 shows, in places, an apparent breaking up or segmentation of the axoneme into pieces resembling the "vacuoles" of Kunstler, as seen in Fig. 1a. Figs. 1b and c are reminiscent of the helix of the sheath as seen in the electron micrographs, the corrugations depicted by Kunstler probably representing the surface appearance of the structures. It must be borne in mind that Kunstler had at his disposal nothing like the resolution and magnification of the electron microscope; his observations were remarkably good, in consideration of this fact. Segmentation of the flagellum may somehow be produced by the action of osmic acid, in combination with other factors. Kunstler had used this reagent in his preparations, as have I. Gelei (1926) showed that such appearances resulted from certain reagents, among them osmic acid in combination with other chemicals.

Fischer apparently found a continuous axoneme in many flagella, but took great pains to demonstrate or argue that it was but an artifact. He, too, did a good job with what he had, and presented a rather convincing argument. However, in the light of evidence since brought forth, including much in this paper, it is highly probable that he was mistaken in his interpretation of at least the nature of the axial fiber.

Regarding the structure of the flagellar filament, Plate 7 presents an interesting picture, which, as Dr. Prebus points out, may well represent at the denser end the natural appearance of the flagellum. Here the helical structure in the sheath is evident, but the inner axial fibers are obscured by the density of the surrounding matter within the sheath. In the less dense portions of this filament we see what is perhaps the result of a loss of material from the sheath or intermediate substance. This lighter portion corresponds to the appearance of most of the flagella shown, and brings out the internal structure, but may thus represent a state of disintegration of the flagellum. The ground substance which may have escaped is precisely that which is considered to be the contractile portion of the flagellum by many workers. For my part, I find it easier to attribute the major contractility to the fibrillar structures remaining.

In this connection, I might mention a possible mechanism for the progress of the contractive wave which has not heretofore been suggested, so far as I know. It has no basis in experimental evidence, and is simply proposed for consideration. Since the flagellum appears to have a fibrous core which is encircled by a helical fiber, as represented diagrammatically in Fig. 21, it is possible that the spiral fiber transmits the impulse. If along this fiber passed a wave of chemical change, comparable to that in a firecracker fuse or a nerve fiber, perhaps releasing H ions wherever it passed over the axial fibers, it could stimulate local contraction in such parts of the axial fibers as might be contiguous with it. In conjunction with twisted axial fibers (Dellinger, 1909), this might account for undulatory movements of a spiral nature. In such organisms as Peranema, the axial fibrils might be stiffened or less sensitive in the basal portion, thus accounting for the fact that normally only the distal part shows obvious activity, whereas, if stimulated to a

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A second possible explanation for the appearance shown in Plate VII is that a portion of the flagellum might have contracted, bringing the coils of the helix into closer approximation and producing the denser and thicker appearance seen in the upper portion of the flagellum shown. It is difficult to determine precisely what actually happened to produce this effect, and the interpretation of such an appearance must remain uncertain for the present.
greater extent, the entire flagellum is thrown into undulations. Also, if the axial fibrils along one side should be less reactive, the resultant stroke might be paddle-like. With a bit of imagination, this hypothetical mode of functioning can be fitted to most observed facts. At least, it fits the facts better than any other system proposed, so far as I have been able to discover. However, it remains purely speculative.

Likewise in the realm of speculation to an extent, but also of interest, in a different way, is the appearance of the axial core of the flagellum of *Euglena* (Plates 2 A; 5; 9). There seem to be two equal fibers running side by side, which are displayed especially well where the flagellum (or merely the fibers) is twisted. These fibers are probably composed of yet finer fibrils (see Dellinger, 1909), though such are not evident in the plates. Now it happens that *Euglena* has a bifurcate flagellar root. Two distinct branches, arising from two separate basal granules, unite in the gullet at about the level of the stigma (eyespot) to form the single flagellum (Wager, 1899, et al). Hartmann and Chagas (1910), in consideration of the bipartite root, considered *Euglena* as derived from a biflagellate. It seems probable to me that the two fibers within the flagellum definitely strengthen such a view, and might be regarded as the two ancestral axial fibers or axonemes in close association.

Another interesting source of speculation is the relationship between the bacterial flagellum and that of the protozoon. Electron studies have been made on many bacteria. Among authors reproducing numerous electron micrographs of bacterial flagella are Piekarski and Ruska (1939) and Mudd and Anderson (1944). According to these and other authors, the flagella average from 14 to 50 m in diameter, and often occur in clumps or tufts, as they do in the spirochete *Treponema* (Mudd, Polevitsky, and Anderson, 1943). Their precise nature still appears doubtful. The suggestion has been made (e. g., Polevitsky, 1941) that they may be hollow tubes, but Mudd and Anderson (1942, p. 106) find no support for this idea in cases which superficially present such an appearance. Mudd, Polevitsky, Anderson, and Chambers (1941) figure an electron micrograph of *Bacillus subtilis* with the protoplasm shrunken away from the cell wall. On the other hand, Mudd and Anderson (1944) figure *Vibrio cholerae*, showing a “single polar flagellum, which seems to traverse the cell wall to join the bacterial protoplasm.” At any rate, bacterial flagella, whatever may be their exact nature, and even though they may not arise in a tuft, often entwine to function as a unit and may even remain adherent, forming a filament composed of many fibrils and perhaps encased in a gelatinous (?) sheath. (See historical review for references.) Now recall the flagellar or sperm tail structure with its 9 to 12 fibrils, each 25 to 50 m in diameter, ordinarily encased in a sheath. The possibility certainly is suggested that the ancestral protozoan flagellum may have been derived from such a permanently united clump of fibrils as occurs in some of the bacterial forms. Should this prove
to be the actual phylogenetic origin of the protozoan flagellum, the concept of the fibrils as contractile units, rather than as mere elastic rods, would be definitely strengthened. However, the fibrils might well serve both as contractile and as supportive structures.

With regard to the mode of functioning of the locomotor flagellum, my observations and experiments confirm the conclusions reached by Lowndes. The living flagella observed beat in spiral undulations, the waves of contraction always progressing from base to tip. My experiments with an "artificial flagellate" and with underwater swimming demonstrated that sufficient force is produced by the simple conical gyration of a body to account for rapid locomotion toward the base of the cone. Fig. 22 and its legend describe the propulsive mechanism as conceived by Lowndes. I have never, to my knowledge, watched the swimming of this particular organism, but I have observed other similar ones. From my own experiments and the work of Mr. Lowndes, I have no doubt that the gyration of the organism could account for a relative speed of 40 (that given by Lowndes for Monas stigmatica), if the body gyrated with sufficient rapidity. But in the forms I have observed, it is difficult to imagine how the body gyration alone could produce such speed. At a rate of body gyration of only about 1/sec. the relative speed may be at least 5. I should think that the forward component produced by the flagellum would aid greatly in the attainment of high speed. But perhaps my imagination is not keyed to the physics of the micro-world of the flagellum. Apparently the mechanics and physical relationships (e.g., the relative viscosity of water, a major item in the protozoan world) to which we are accustomed are tremendously altered, practically non-existent, at this level of dimensions. For an interesting discussion of some of the differences see Bidder (1923, pp. 304-307).

Brief mention may be made of the possibility of studying living organisms with the electron microscope. It would obviously be desirable to study such a structure as the flagellum in the living, intact state. Several papers have appeared relative
to the possibility of observing living structures with the instrument, but, so far as I know, the only objects studied thus or even very practical for such study have been spores. Perhaps the most ambitious attempts have been made with the closed cell described by Abrams and McBain (1944). The obvious difficulties result from greatly reduced contrast and Brownian movement, which prevent photographing of the objects under observation. Another major factor is the killing effect (aside from heat production) due to "... the electrons bounding down through the specimen and breaking the chemical bonds of the specimen. Breaking the bonds is detrimental to all living organisms. Every bond in living organisms does not have to be intact, but the critical molecules in the living organisms must have a specific structure in the organism for it to grow and reproduce itself. When you consider that, in maximum intensity, the beam reaches a high value of 1,000 electrons per square angstrom per second, you can realize that it wouldn't take the scattering of many electrons to kill such an organism." (Anderson, 1943.)

In terms more familiar to the biologist, this electron bombardment would amount to 100,000,000,000 electrons striking a surface of 1 square micron each second. The average protozoan body would present a surface of a considerable number of square microns. Thus the prospects of studying a living flagellum, even if it were perfectly stationary and met the other requirements, would not be encouraging.

**SUMMARY**

1. The work of previous investigators in this field is reviewed and discussed.
2. The structure of the flagellum of several species of Protozoa has been investigated by means of the electron microscope. Micrographs are included which show that:
   
   (a) The flagella studied are of approximately uniform diameter throughout their entire length.
   
   (b) Each flagellum consists of a denser axial core (axoneme) and a less dense sheath surrounding the core.
   
   (c) In the flagella of *Euglena* and *Astasia* the axial core appears to consist of two closely approximated fibers of equal size.
   
   (d) The sheath appears to contain or to consist of a coiled fiber which encircles the axial core in a helix.
   
   (e) The flagella of *Euglena* and *Astasia* bear, along one side, what appears to be a single row of delicate filaments (flimmer) extending from the sheath. These flimmer have an average length of about 5–6 times the diameter of the flagellum proper.
   
   (f) The long flagellum of *Ochromonas* bears similar filaments along both (or all?) sides.
   
   (g) The flagella of *Chilomonas* bear no such lateral filaments.

The significance of certain details present in the electron micrographs is discussed.
3. The presence of two equal fibers in the flagellum of *Euglena*, together with the fact (demonstrated by previous investigators) that the flagellum of this form has two separate roots, indicates the ancestral formation of its flagellum by the fusion or union of two flagella. The hypothesis of a biflagellate ancestry of *Euglena* is strengthened.
4. The possibility of the origin of the primitive protozoan flagellum through the union within a common matrix of several simple fibrillar flagella (as are present in certain bacteria) is suggested. Evidence for such a theory is presented.
5. Observations were made on the activity of living flagella. The results were in direct confirmation of Lowndes's contentions that:

(a) The flagellum beats in spiral undulations.
(b) The waves of contraction progress from the base toward the tip of the flagellum.
(c) As they progress, the waves often increase in amplitude.
(d) The flagellum usually serves to push, rather than to pull the organism through the water, although it arises from the anterior end of the body.

6. Experiments were performed by means of actual underwater swimming by the author and with an "artificial flagellate" to determine the forces produced by gyrating bodies in water. In both cases, a simple conical gyratory movement was found sufficient to produce locomotion. Employing the artificial flagellate, it was found that rotation and gyration of a small body at about 6 gyrations per second was capable of producing a relative speed of as much as 7.0 (i.e., 7 lengths per second), in spite of the fact that the body had to pull along after it an object much larger than itself. This confirms the theory of Lowndes that the rotation and gyration of the body alone may adequately account for the locomotion of many flagellates, without any forward component produced directly by the flagellum.

7. A hypothesis is suggested concerning the mechanism of flagellar function: Perhaps the helical fiber of the sheath transmits an impulse which stimulates local contraction in the underlying fibrils. Possible ways by which such a mechanism might account for the various types of flagellar movement are mentioned.

8. Promising techniques, recently developed in other fields, are suggested for further research on flagella.

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A. Portion of a diatom used for measurements and estimation of magnification. EM 1703 D.
B. Astasia klebsii. Portion of flagellum showing unilateral flimmer and indications of helix of sheath as in Plate 9. EM 1712 D. Technique #1.

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*Euglena gracilis*. Extremities of bodies and portions of flagella. Fig. A, EM 1704 D. Fig. B, EM 1696 D. Prepared according to technique #1.
Euglena gracilis. Portions of flagella. In Fig. A (EM 1691 D), the helical nature of the sheath is apparent. Flimmer may be seen in Fig. B (EM 1687 D). Technique #1.
Euglena gracilis. Part of a flagellum showing unilateral flimmer especially well.
EM 1700, 1701 D. Technique #1.

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Euglena gracilis. The flagellum extends from the body, loops, and returns to lie alongside the body. EM 1710, 1711 D. Technique #1.
Euglena gracilis. In the upper section, the flagellum is seen to emerge from the body. It has been snapped by a rupture of the supporting film, and parts of the axoneme are exposed. The length of the flagellum is about 13 μm. EM 1697, 1698, 1699 D. Technique #1.
*Euglena gracilis.* Portion of the flagellum showing two very different appearances. The upper part perhaps represents the more nearly natural condition. See discussion in text. Note the transversely striated appearance of the upper, denser portion, produced by the closely wrapped helical fiber. EM 1692, 1693, 1694 D. Technique #1.
Euglena gracilis. Portion of flagellum showing a swollen appearance, perhaps due to the escape of flagellar plasm during drying. EM 1707, 1708 D. Technique #1.
*Euglena gracilis.* Portion of flagellum showing unilateral flimmer. In places where the flagellum is twisted, the two equal fibers of the axoneme are made rather obvious. The helix of the sheath appears to be closely appressed to the axoneme (note regular bumps along sides of flagellum. EM 1020, 1621 D. Technique #2.
Chilomonas paramecium. Anterior end of body and the two flagella. No flimmer are to be observed. EM 1723, 1724, 1725 D. Technique #1.
Ochromonas variabilis. Portion of body and flagellum showing flimmer on both (all ?) sides, not in a single lateral row as on the flagellum of Euglena. The dimensions of Ochromonas flagella vary with the size of the organism, but in those photographed they ranged for the most part between 4 and 6.5 μ in length, averaging about 0.1 μ in diameter. The flimmer seem to average about 0.5 μ in length and are less than 0.01 μ in diameter. Note apparent helix as in Plates 9 and 1 B. EM 1715 D. Technique #1.
Ochromonas variabilis. The entire body of the organism is shown, with most of the flagellum. The small second flagellum may be seen near the base of the long flagellum. No flimmer have been observed upon it. Apparent helix as in Plate 11. EM 1737, 1739 D. Technique #1.
THE EYE-BANK FOR SIGHT RESTORATION

More than half a hundred hospitals in nine states are already cooperating with The Eye-Bank for Sight Restoration, Inc., in a nation-wide effort to help restore or remedy the vision of America's estimated 15,000 persons blinded because of corneal affections, it was announced recently by Mrs. Henry Breckenridge, executive director of the Eye Bank, 210 East 64th Street, New York.

In Greater New York alone, 32 hospitals are associated in the movement to make available for distribution healthy corneal tissue for those whose sight may be restored through corneal graft operations by which ocular opacity is overcome, the announcement stated.

In addition, 8 hospitals in other New York state cities, together with 6 in New Jersey, 3 in Connecticut and one each in six other states have become actively affiliated with the movement.

Organized only last February, the Eye Bank was established for the collection, preservation and distribution of healthy corneas which may be obtained only from persons either living or immediately after death. Inasmuch as corneas may be preserved and utilized for transplanting to the eyes of others for only 72 hours, speedy collection and distribution is essential as soon as they are obtained. Whenever cooperating hospitals have eyes available, the Red Cross Motor Corps rushes them to the Eye Bank for distribution to persons requiring the corneal graft operation.

In addition to extending this activity to hospitals throughout the United States, the Eye Bank is presently engaged in a nation-wide effort to obtain support for its work through solicitations for memberships and donations of eyes after death.