

Annual Laboratory Reports of the Biology Society

February 1940 to and through January 1941.

Histology Section

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These reports are abstracted from the running notes, too voluminous to copy.

(N.B. - This is almost a thesis! J.L.)
4500 words.

To:
Mr. Schur

4 pages

fall 1940 - Leberberg

The major accomplishments and experiments of the Histology Group are exemplified in the slides that it has prepared. Since the preparation of these slides follows, generally, a basic preparatory process, it would be well to indicate this process. In the preparation of tissues for histological study, the tissue must be killed and hardened in the condition it was when alive, and preserved against future decay and deterioration. This is accomplished by the use of a 'fixative' solution, and the process is called fixation. The tissue must then be made ready for cutting very thin slices; in order to accomplish this, the fixed tissue, after having had its fixative thoroughly washed out, is dehydrated, the water being replaced by paraffin, through an intermediate stage of alcohol and a solution of ~~alcohol~~ paraffin in dioxane. The tissue is now said to have been embedded. The embedded tissue can now be sectioned; this is done by a microtome, and which delivers sections of about 10 to 12 μ ($1\mu = .0001 \text{ cm.} = \frac{1}{25,000} \text{ inch}$) as indicated. These sections are now mounted and attached to a slide, and the slide, holding a thin section of tissue embedded in paraffin, is ready to be stained. To do this, the paraffin is removed by xylol, the xylol is removed with dioxane and that with water. After the proper immersion in the desired staining fluid or fluids, the slide is washed thoroughly, put through dioxane and xylol and is finally covered with a drop of Canada Balsam dissolved in xylol, and a thin cover-glass. This is for mechanical protection and to make the section transparent. The tissue-section has had to be stained in order to bring out and differentiate the various elements and cells in the tissue.

Notes respecting the work of the for larger part of the first term of this last year were lost during revision, so that a good deal of chronological information is lacking; However, all information relative to the preparation of the tissues that is necessary for the interpretation of the results. Dates, unless otherwise stated, refer to the date of primary staining.

Site in February 1940, a fat body ^{was} freshly dissected out of a male frog, a small fragment, 2 x 5 mm. placed in 10% biological formal in water, for 2 or three days, washed one hour in 35% methyl alcohol, then dehydrated overnight in dioxane, infiltrated 5 hours in dioxane-paraffin, 3 hours in molten paraffin, and embedded. The block was then sectioned at 18 μ , stained in Delafield's hematoxylin and in 42% eosin in 50% dioxane, and mounted under balsam. In June, 1940, the slide was borrowed by Mr

Weinberger, and thence disappeared in some recess of the Biology Department.

From the same frog, a piece liver was dissected out and subjected to a similar process, with the sections at 12 μ , stained in Delafield's haematoxylin. Unfortunately, this section is somewhat crumpled. This was not sectioned or stained until April 16, 1940.

From the same frog, and fixation, etc., the cartilage and muscle of the episternum. Stained after 12 μ section in Delafield's. 4/16/40.

From the same frog, and fixation, etc., the spinal cord of the frog in the cervical region, and, also, the lower brain. Sectioned at 10 μ , stained for 3-4 seconds in Delafield's, washed, then restained in 1% Methylene Blue, differentiated in 99% KOH one, and finally mounted under balsam. A very poor photomicrograph from this slide, effective magnification about 450 X, ~~was~~ enlarged 2.5 X. The picture, which is the waist of a series, was taken by H. F. Smith. It shows the ciliated ependymal epithelium surrounding the central canal. A series of these sections were stained at about this time.

April 27, 1940, a smear of human blood, was stained about one minute in Wright's stain.

(Part of a more detailed notation follows: -)

(Mon., April 1 - "No work was allowed today in 521.")

April 2, 1940 - dissection of a rabbit in 419: "The maxillary turbinates are curious, unfamiliar structures. I am not sure whether the paired set of nerves found just subcutaneously above the rear maxillary teeth are the superior maxillary or one of the facial - facial, trigeminal - nerves. They pass just under the orbit and enter the brain near the optic nerve. Also there is a small white patch, medially at the roof of the mouth, between the soft and hard palate. What is it? Without even a temporary use of the incubator (which was at the time full of bacteriological tubes) it seems almost hopeless to consider most topics of histological work in the near future. Oh, for a freezing microtome! No dehydration. No difficult and messy infiltrations, no rehydrations to wrench a section from the slide. The block of the frog's central nervous system is well embedded. Everything is so disorganized." Quotation from my journal, 4/2/40.

4/3/40. "A cross-section of the frog spinal cord was stained because of examinations on April 11 and 12, there will be no work this week. Slide was apparently not clean. A unstained, dehydrated, restained. More experience in technique required."

4/14/40. Remained afternoon and evening till 6:50 P.M. with Teachers-in-Service Course. Spent one hour staining nerve slides. Both pyrene and aniline are unsatisfactory in arresting destaining in acid alcohol. Stain was hyptrial and error until right depth of stain was revealed. Eosin counterstain only confuses the field.

4/15/40. Till about 2 o'clock, aid Mr. Towne in cleaning Petri dishes. Thence, some further microtomy of blocks of frog nervous system.

4/16/40, same as above. In lecture before ^{Study} biology technique used in histology was demonstrated.

4/17/40. 12 slides, all nervous, were stained with methylene blue, in a deofane modification of a Nissl stain.

4/18/40. Till 6:45 P.M. at the ~~by~~ Service Course - see 4/4/40.

"All afternoon and evening was spent in assisting and in staining some more of the unstained sections thus far. Today I used an 85% deofane in water solution to destain, absolute deofane to arrest differentiation. The chief difficulty lay in obtaining sufficient intra-cellular detail simultaneously with inter-cellular contrast. In earlier afternoon fixation of a 10 mm chick embryo that had been kept about 4 hours in salt solution for macroscopic examination in Baum's.

4/19/40. "to 5 21 blue staining of the last nerve slides was completed. One in haematoxylin and one in safranin. The remainder in Nissl - Methylene blue. The Canada Balsam we have must be quite acid as the haematoxylin stained slide has turned reddish again.

4/22 - 4/23/40. Embryo dehydrated at home. In afternoon infiltrate. Later, in evening, at Sid Babacia's incubator; imbed the embryos on 4/24/40.

May 1, 1940. Sectioning of embryos, more anterior region. Cephalic region found not imbedded at all. Upper and abdominal sections poorly, much wrinkled. Some sections, more posterior, flattened out nicely. About 12 sections were made.

May 8, 1940. Stain many such sections, largely methylene blue; poor wrinkling, staining. (Slides: II/22-25, II/1).

May 9, 1940. "Upon request to Mr. Verbrugge, I was able to obtain today a number of frog embryos and larvae (tadpoles) for embryology. All were placed in Baum's.

May 13, 1940. Stain all embryo slides thus - (Slides: II/2, 3, 4, 7.) Prepare whole mount of hydra; unsuccessful attempt to kill a planaria intended

May 16, 1940. Section blocks of ~~tadpole~~ embryo frog, as begun on
May 9, 1940. Yolk causes poor sectioning.

May 20, 1940. Stain two slides from 5/16/40. Hematoxylin.

May 23-24. Biology Exhibit. Stain a number of slides, ~~etc~~: chick
embryo, and a block of head of snake embryo secured by John Jaquay.
Also stain a section of Cat cerebrum secured by him.

May 23, 1940 - Frog blood smears, stained with methylene blue and eosin.

May 27, 1940. - Some blood smears, stain with Wright's.

May 28, 1940. Stain a few sections of cat cerebellum - Methylene Blue, Wright's.

June 5-6, 1940. Preparation of about 200-250 sections of stomach,
lung, heart, intestine and tongue ^{of frog} prepared largely for Mr. Wemburg's
shop course. In month of June, 1940. a series of sections of stomach,
intestine, as above, were stained, a great deal in assistance to Mr.
Wemburg. Also, a ^{block} section of kidney was embedded.

Second Term 1940, September to January 1941.

1 School opened Monday, September 9, 1940.

September 13, 1940 - Blocks A 1-14 were sectioned. Among them, small intes-
tine and kidney of a male frog. (Slides VI 117, 22, VII 121, etc.) These
had been prepared in June of the previous term with J. Jaquay, Bourin's
fixation. The sections were not mounted, but floated in a pan of water till

September 16, when they were mounted. and, on

September 17, when the types of slides indicated above were stained at home
with hematoxylin and eosin.

September 18, 1940 - the group prepared 105 slides for the Bio Dept con-
sisting of different colored threads variously mounted. Study of stomach
and kidney.

September 19, 1940 - Examination of I/8. Note possible granulation or
ciliation of cells of frog marrow at 430X.

September 20, 1940 - Examination at 970X of many slides, particularly
I/8.

September 23, 1940 - A lecture meeting in the Histology Group,
dealing with the use of the microscope, demonstrating the oil-immersion
microscope, and lecturing on the structure of the elementary structures. Also,
preliminary examination of prepared blood smears at 970X. There
was also a discussion of Howell's Theory of Blood Coagulation.

September 25, 1940 - A lecture on the general structure of the cell and
the theories of histological fixation and staining. Then, the mechanism of
the microtome, and general basic histology & technique.

continued - 9/25/40. Some sections were taken of tissues the same as previously taken. Wright

- 9/27/40. Staining Technique, introductory. Group members stained sections previously returned.

- 9/30/40. Wright Stain Technique on blood smears; also examination thereof. General group work.

October 2, 1940. Various stains applied to Blood Smears to determine optimal staining: all were fixed in absolute Methyl alcohol after the smear dried. Then dried again. Then stained, washed and dried. Stains used were picric acid, methyl violet, picric acid followed by Methylene Blue; light green, Carbol-Fuchsin, methylene blue and alcoholic gentian violet. Nuclei of leucocytes seen best stained by methyl and gentian violet. These also stain the hemoglobin of the erythrocytes.

- October 9, 1940. Fixation Theory reviewed; Locking Cutie histological technique demonstrated. All Group work.

October 14, 1940. Blood Histology for entire group. Remounted and replein slides made therefore previously thereof. Also group examination on Wright staining technique. Preparations 1, 2, 3, 4, 5, 6, 7, and 02 outlined or attached shut were performed by the entire group. Slide I/8 on frog marrow was shown to demonstrate 01. Materials for 03 were not available. Results were highly successful, but detailed notes were taken more by the members of the group.

For two weeks, we tried to get a rabbit at a usable time, and were unsuccessful until October 30, 1940. Also, Jewish holidays entirely disrupted a few schedules: 10/3, 4; 10/17, 18; 10/24, 25; also, work was not allowed on 10/21/40

October 23 - Histology Theory review for group. Also theory on connective tissue.

October 28 - Lecture from 10/23/40 continued; examination of slides showing cartilage, fibrous, areolar and elastic tissue. No rabbit was available.

October 30 - Same as October 28, ^{outlined} preparations regarding the nails of Skat Two were discussed

October 30. A rabbit was finally gotten, so that preparations 1, 2 and 3, 4, and 6 on Skat 2 were attempted, with reasonable success, except in the case of 3; there was no emulgent; develop-

ment did not proceed; it was available with from external sources. Cartilage was studied indirectly from organ slides - rat embryos, late chick embryos, larynx. In preparation 1 and 2, a peculiar type of striated fibre, resembling algae, chassem, still unidentified. Van Heeren's had to be prepared at the time, but not hastily.

November 1 - Acute...

November 4 - Lecture and slides on cartilage, bone and bone development, with 11 of Sheet 2. Slides on rat and chick embryos, long ago prepared, were most helpful. A number of stomach slides were restained with Van Heeren's, Methylene Blue and Fast Green, and Wright's stain.

November 8, 1940 - Stain sections of kidney with hematoxylin-eosin. essay study.

November 13, 1940 - Due to general lack of knowledge of anatomy in group, a rabbit and a frog were rapidly dissected with explanation.
11/14, 15 - M. alternans.

November 18, 1940 - Lecture on muscle structure (summary due to lack of facilities) and also of the elementary canal in general.

11/20/40 - Lecture on structure of pharynx, esophagus and stomach, with examination of slides and discussion.

- in afternoon, at home, fix from a fox from fresh frog, in 10% formalin, the: liver, spleen, pyloric region of stomach, tongue, lung, kidney, oviduct, sciatic (spinal) nerve, episternum, fundal and cardiac regions of stomach. This is to be known as series B. Fixation proceeded to Monday, 11/25/40. Tissues whose sections do not appear in following accounts were discarded because of obvious decay before fixation.

Mon. 11/25 - series B transferred to 50% methyl alcohol for washing. Airt to Mr. Towne.

Tues, 11/26 - B series transferred to dioxane for dehydration overnight.

11/27 - placed in dioxane-paraffin for infiltration; later in afternoon, into molten paraffin over two nights, when, on

11/29 - The series, or a large part of it, was embedded and blocked off.

mount did not proceed.

December 2, 1940. ~~Stain~~ Section, blocks embedded 11/29/40. and mount on slides. Fix nucleus root tips in: Carnoy's, 10% formal, Zenker's, hemis, 79% CrO₃, Flemming's achromatic, Zenker's fixatives.

December 3, 1940. - Some sections were stained from blocks sectioned 12/2. The stomach was found to be somewhat autolyzed with a little detect remaining, however (hem-eosin.) Fixed root tips from yesterday washed in 70% Methyl Alcohol, with Li₂CO₃ or Lurie where indicated. Stained were - stomach, kidney and urovent. Cursory examination.

December 4, 1940 - root-tips in deoxane, overnight, for dehydration. Stain Frog Liver, series B, hem-eosin; Frog spinal cord, Series A with Methylene Blue. Section, nose stomach (A) liver, kidney, and spleen (B.) Oviduct was much ripped and wrinkled.

December 5, 1940; root-tips in deoxane - paraffin, then paraffin for imbed infiltration; Buy slide labels. King classification to date, clean, relabel slides.

December 6, 1940: Embed root-tips; Help Mr. Torne.

December 9: Group members stain stomach, liver, kidney cells; ^{stain} tissue still imperfect, but fair.

December 11 - help Mr. Torne.

December 12 - Section and stain root-tips (no. 4 - ~~as fixed~~ with Flemming's achromatic - ~~chrom~~ dichromate - chromic acid - acetic acid fixative. Results inconclusive. Also section Spleen B. Close group examination of slides ~~12/13~~ on Rab, chick and snake embryos respectively.

12/13. Examine stained, mounted sections of Spleen B - with hem-eosin. "Sinuses and blood vessels are very characteristic, but well-defined. Malpighian bodies are not at first recognized. Capsule traversed from trabeculae dense pulp. Striated venous walls. Also, stain smear of blood of turtle. Apparently, type of amphibia. Rearrangement and house-cleaning of group property.

12/16/40^m - Aid Mr. Torne - preparation of glass holders with nichrome wires for bacteriology.

12/18/40. - Duodenum, Ureter, peritoneum, liver, and esophagus-trachea were taken from a rabbit and all fixed in Bouin's. Subsequently washed through to 70% Methyl Alcohol, wherein they now repose.

12/18/40 - Dissection and fixation in Bouin's of skin, peritoneum, liver, appendix, spleen, duodenum, pylorus, stomach, oedodenum, pancreas, fat body (subcut), kidney, lung, heart, esophagus and trachea of a male white mouse. Subsequent washing; now in 70% Methyl Alcohol. Fix in 10% formal, subsequent washing - the brain and vertebral cord.

12/19/40 - ~~open~~ "subsequent operations" of yesterday. ↑

12/20/40 - Prepare slide of Parameria caudatum, hematoxylin stain. Fix in Schaudinn's, wash through iodine alcohol, water. Stain, dehydrate in diorane, clear in xylol, mount in xylol-balsam. Stain not too precise or clear. Also, stain stomach, pylorus B, in hem-rorin. Take inventory.

Christmas vacation.

January 2, 1941. Group buys a quantity of chemicals, with personal funds. 1/3/41. Fr. Stain a section of liver B with Iron-Hematoxylin. Mount in 4% Ferric Alum, 25 min.; wash one minute in tap water; stain 5 min. in 1/2% hematoxylin aq.; differentiate to suit with Ferric alum, wash, dehydrate, etc.

Stain some wet blood smears fixed in absolute MeOH. Proceed as above for iron-hematoxylin.

Stain a number of specimens from a culture mailed Spirostromum. From members Blepharisma. ^{protogoa.} Iron-hematoxylin. (See 12/20/40 for technique.)

1/6/41. Restain Spirostromum from 1/3 to satisfy. Mount in xylol-damar.

1/9/41. - Sutor, again, a number of blocks in series A, B, C (root tip) (date.)