## SPINAL CONDUCTION AND KINESTHETIC SENSI-TIVITY IN THE MAZE HABIT<sup>1</sup>

## K. S. LASHLEY AND JOSEPHINE BALL

The importance of kinesthetic and organic sensations has been much emphasized in recent psychological speculation, both behavioristic and otherwise. Our most widely taught theory of the emotions is based upon a doctrine of sensory reverberations from visceral activity, the doctrine of "current control" of the speed and accuracy of movement still has many adherents, and the theory of implicit movements as the organic basis of thinking extends the same concepts to the psychology of thinking. In general, these developments represent a tendency, fostered by recent studies of nerve conduction, to seek the immediate completion of the conception of chain reflexes for that of continued intraneural activity.

Toward the development of this point of view the analysis of the sensory control of the maze habit has contributed no small part. Watson ('07) successively and simultaneously eliminated all of the important distance receptors of the rat without serious interference with the animal's ability to learn and execute accurately the maze habit. This seemingly left kinesthetic and organic sensitivity as the sole remaining basis for the habit. As Watson pointed out, these first studies, by the method of sensory elimination, gave only negative evidence without indication of the nature of the intraorganic processes which control the maze running.

In their study of behavior in a maze with alleys adjustable in length, Carr and Watson ('08) obtained further evidence that the movement series is internally conditioned and that after distraction the animal regained orientation from the kinesthetic pattern

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aroused by running some segment of the maze. They then clearly enunciated the theory that, "the 'movement to come' is released at the proper time by the afferent (kinesthetic) impulses aroused by the movement which has just been made," and this interpretation has been followed in almost all succeeding discussions of the maze habit. One of the most definite recent statements of this chain-reflex hypothesis is that of Washburn ('16). "When one is playing a piece of music for the first or second time, each movement has to have the stimulus of the notes on the page; when a piece has been long practiced, each movement sets up the next one 'automatically.' This really means that, as one movement is performed, the sensory processes occasioned by the contraction of the muscles involved excite the motor paths for the next movement. The stimulus for one movement is the kinesthetic excitations received from the preceding movement."

Vincent's later work ('15) showed the occasional importance of visual, olfactory and tactile cues, but did not question the essential conclusions from Watson's studies.

We were first led to doubt the validity of this interpretation of the maze habit by observations on the maze-running of rats after cerebellar injuries (Lashley and McCarthy, '27). Marked changes in the motor pattern were seen to have no effect upon accuracy of orientation in the maze, even in blind animals under conditions where other than supposedly kinesthetic cues seemed to be eliminated. This raised the question as to whether the habit is controlled by kinesthetic sensitivity or by some wholly intraneural mechanism, once orientation has been obtained. Are automatized sequences of movement the result of a mechanism by which each movement arouses sensory cues to initiate the next, or of some mechanism in which a central organization, once aroused, discharges successive motor impulses constituting the series, with comparative independence of the sensory consequences of the movements?

The experiments with lesions to the cerebellum are inconclusive because we know neither the important components of the movement system nor the afferent impulses which may be essential to the habit. Perhaps the only crucial experiment for the question would be one with animals having section of all the somatic sensory fibers. This seems technically impossible and the question may therefore be one which can be answered only in terms of relative probabilities deduced from indirect evidence. Loss of ability to run the maze after partial destruction of kinesthetic sensitivity might serve to establish the chain reflex theory. Survival of the habit after even severe disturbance of kinesthetic sensitivity may always be interpreted as due to failure to eliminate the significant afferent impulses. Nevertheless, the probability of the chain-reflex hypothesis is reduced in proportion to the extent of anesthesia and the absence of effects of anesthesia upon the performance of the habit.

With these limitations, it seemed to us that some significant data might still be obtained by a study of the effects of extensive destruction of the spinal afferent tracts upon the accuracy of maze running. We have, therefore, carried out a series of experiments in which animals trained in the maze were subjected to spinal lesions and subsequently tested for retention of the habit. The senior author alone is responsible for the surgical and histological part of the work. The training and tests of performance have been carried out by both of us independently on different sets of animals, with essentially similar results.

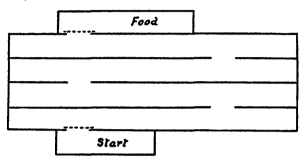
#### METHODS

Training. The animals were trained with food as incentive and five trials per day in a maze with 8 culs de sac (fig. 1) until 10 consecutive runs without error were made. This maze had been constructed to record errors automatically so that animals could be tested in it in total darkness. The recording platforms were arranged symmetrically on each side of the openings from alley to alley so that they gave no directive tactile cues. Two groups of rats were trained with somewhat different subsequent procedures. The first consisted of young rats purchased from a local dealer. They were rested for seven days after training, were then retrained until 10 consecutive errorless trials were obtained (preliminary retention tests) and subjected to operation. Seven days after operation they were again retrained to 10 successive

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errorless trials (post-operative retention tests). During all the work with them these animals were very unstable, easily distracted during the period in the maze and rarely eager for food. They are number 1, 2, 3, and 6 of the protocols. Their records were difficult to interpret because of their erratic behavior, so the experiment was repeated with another group.

These were older animals, reared in the laboratory. They were were given 50 trials of overtraining after reaching the criterion of 10 errorless trials. They were then rested for ten days, given preliminary retention tests and also ten trials with the maze in darkness. (This was accomplished by covering the maze with





Start, starting compartment; Food, food compartment. The broken lines represent swinging doors serving to prevent back-tracking from the maze and food compartment.

black cloth and several layers of heavy paper. Observation through a peephole failed to reveal light leakage after thirty seconds adaptation for our eyes.) The animals were then operated and from six to ten days later, depending upon the rate of recovery, were given postoperative retention tests.

Surgical. Under ether anesthesia the third cervical neural arch was removed and the cord exposed by a transverse incision of the dura. The point of a thin-bladed iridectomy knife was passed through the cord at the depth to which we wished the lesion to penetrate and the overlying fibers then cut through with the edge. This insured that all fibers peripheral to the line of penetration of the knife were cut and not merely pushed aside. The injuries involved either section of the dorsal funiculus, section of both lateral funiculi, or section of the ventral funiculus. The last injury involved lifting the cord from the canal with consequent compression and shock and only one animal, and that with a partial lesion, survived. After the injury to the cord the muscles and skin were united above the cord by interrupted sutures and the wound dressed with collodion.

Histological. At necropsy the brain and cord were removed and prepared in serial sections by the Marchi method. For most cases complete serial sections were preserved from below the level of the lesion to the superior colliculi. The figures of plates 1 and 2 are made from camera outlines. Solid lines are used to represent general regions and contours; stippling in all cases represents degenerated fibers; solid black areas represent unabsorbed clots. For the plates one section was sketched at the level of maximum injury (a) and one (b) at a sufficiently higher level to avoid the degeneration of the shorter ground bundles and present only tracts which could be traced into the medulla.

There are few data available on the spinal tracts of the rat so we have been compelled to work out their position as well as possible from our material. A certain fallacy may arise here since our knowledge of the tracts is limited to what we have destroyed and our estimates of the character of the lesions are based upon that knowledge. However, the lesions cover every part of the cord except the median ventral columns so we may be sure that in one or another animal we have severed every organized tract in the cord and that the demonstration of the tracts is limited only by the level of the lesions and the defects of the Marchi technique as a method for the rat.

#### THE SPINAL TRACTS OF THE RAT

The conduction paths of the spinal cord of the rat have not been described in detail. Interest has centered chiefly in the pyramidal tracts because of their deviation from the usual lateral position. They have been described and figured by Lenhossek ('89), Bechterew ('90), Goldstein ('04), Van Der Vloet ('06), King ('10), Ranson ('14), and Linowecki ('14). Ranson ('14) has described the position of Clarke's column and the tract of Lissauer. The rubro-spinal tract has been traced by Held ('90) and by Papez ('23), but briefly and without figures so that its exact position in the cord is not clear. We have not found descriptions of other tracts.

The series of cases studied in the present experiment has involved the interruption of all the tracts of the cord in one or another animal and we have been able to trace these in so far as they are revealed by the Marchi technique. The pyramidal tract does not stain readily by the Marchi method, as Ranson and others have pointed out, because of the thinness of its medullation. The same seems true of other long descending tracts, for in only a few of our specimens have we been able to trace degeneration below the lesions. In contrast, many ascending tracts show clearly, with well defined limits.

#### Descending tracts

*Cortico-spinal.* The descending fibers from the cortex occupy the median ventral position in the dorsal funiculus (plate 1, fig. C). Decussation of the pyramidal tracts to this position seems to be complete and Ranson ('13, '14) reported no other descending degeneration after hemidecerebration.

Rubro-spinal. This has been described in a preliminary note by Papez ('23). He does not figure it and states its position in the cord only as corresponding to that in other mammals. We find a large clearly limited bundle ventro-lateral to the dorsal horns (plate 1, figs. B and C) which is probably the rubro-spinal tract.

Schultz's comma. After section of the dorsal funiculus the comma appears as a small, well defined descending bundle lying between the fasciculus gracilis and fasciculus cuneatus on each side (plate 1, fig. B).

No other clearly defined descending tracts are visible in our preparations. After lesions to the lateral and ventral funculi many degenerated fibers may be traced caudad in these regions, but they appear rather uniformly scattered and as our preparations do not include lesions above the cervical cord it is impossible to distinguish the source of the various scattered fibers.

## Ascending tracts

Fasciculus gracilis. In degeneration of the dorsal funiculus this tract appears as a clearly defined bundle which can be traced to its complete termination in the nucleus gracilis (plate 1, fig. 4b).

Fasciculus cuneatus. This bundle lies lateral to the fasciculus gracilis (plate 1, fig. 4b). It gives off fibers to the gray matter throughout its length and receives additions from the dorsal cervical roots (plate 2, fig. 6b).

After destruction of the lateral funiculus, degeneration appears throughout an extensive zone in the ventro-lateral region of the cord (plate 2, fig. 8 and 9b). Traced into the medulla these ascending fibers are seen to break up into four principal groups (plate 1, fig. A). It has not been possible to determine with certainty the origin of these bundles in the cord, but their most probable positions in the cervical cord, judged by tracing downward from the medulla, are indicated below.

Fasciculus spino-cerebellaris dorsalis. The fibers lying in the more dorsal portions of the lateral funiculus may be traced forward to the medulla where they separate from the remainder of the degenerated mass to pass laterad to the posterior cerebellar peduncles (plate 1, fig. A, s.c.d.).

Fasciculus spino-cerebellaris ventralis. The more ventral fibers of the lateral funiculus may be distinguished in the medulla as a broad band of fibers which ultimately turn laterad to reach the anterior cerebellar peduncle (plate 1, fig. A, s.c.v.).

Spino-quadrigeminal system (?). After destruction of the lateral funiculus a large number of degenerated fibers appear scattered in the ventral region of the reticulated substance of the medulla (plate 1, fig. A, s.q.). Traced forward they ascend and are lost, in our material, in the region of the inferior colliculi. Traced caudad, they could be followed to the most ventral portions of the lateral funiculus and to scattered fibers of the ventral funiculus.

Fibers to the median longitudinal bundle. After lesions to the ventral portion of the lateral funiculus or to the ventral funiculus a few degenerated fibers appeared in the fasciculus longitudinalis

medialis of the medulla, (plate 1, fig. A). These could not be traced to their termination because of defective fixation of our material. They seemed to be derived chiefly from the scattered fibers of the ventral funiculus. Some of the fibers of this and the preceding systems may reach the thalamus.

In our preparations, which included section of every part of the cord, no other definite degenerated tracts could be detected. There is no assurance that other important ascending tracts may not have remained unstained owing to conditions of medullation similar to that of the pyramidal tract. This is, however, of little importance for our experiments since, whether stained or not, there is no question but that they were cut in some of our animals.

In general, the tracts of the rat's cord present no fundamental differences from those of higher mammals, to which we can ascribe the results obtained in our studies of behavior. The ascending tracts of the dorsal funiculus and the spino-cerebellar tracts are large and sharply defined. The pyramidal tract is restricted to definite area and this seems true also of the rubro-spinal. On anatomical grounds there seems no more reason to predict a diffuseness of function in the cord of the rat than in that of the monkey or man.

The path of the proprioceptive impulses is perhaps not vet certainly established, but the weight of evidence from clinical material indicates that sensitivity to movements and bodily posture is disturbed only by lesions within the posterior columns. Head ('20) states that, "... the impulses associated with passive position and movement and with tactile discrimination do not, within the limits of the spinal cord, reach the point where they are recombined, but continue uncrossed to pass along the fibers of primary afferent systems in the posterior columns. It is not until they reach the posterior column nuclei (nucleus gracilis and nucleus cuneatus) that they pass from a primary to a secondary sensory system" (p. 401), and again, "We therefore find, that the only definite consequence of destruction of the posterior columns is to produce loss of tactile discrimination (compass test) and of the sense of passive position and movement on the same side as the lesion" (p. 402).

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## GENERAL EFFECTS OF LESIONS

For each of the regions of the cord involved in the experimental destructions the animals showed definite syndromes. The picture after section of the dorsal funiculi seemed primarily one of reduced sensitivity to bodly position. In walking the animals had a sprawling gait, with feet rather widespread. They tended to walk on the dorsal surface of the fore feet, less frequently of the hind feet. There was occasional incoördination, especially a lack of syncronization of the fore and hind legs. In rapid walking or running, the hind legs tended to drag in recovery from the backward step, and in turning to one side the opposite fore leg was often dragged stiffly behind. In running rapidly in the maze all legs were recovered too slowly from the backward step so that the animals seemed to run ahead of their feet and fall forward. When tested at the edge of a table with one leg unsupported this leg was often allowed to hang flaccid until movements of progression were initiated (a condition noted in many animals after section of the dorsal roots of a limb). When the animals were placed on a board which was tipped at various angles their posture in maintaining balance seemed practically normal. Only crude tests for cutaneous sensitivity could be used. They gave no indication of abnormality, but were unreliable for any but the grossest changes.

Section of the lateral funiculi gave a picture suggestive of cerebellar ataxia. The gait was staggering with exaggerated balancing movements of the tail. The animals frequently fell to one side, especially in making the turns of the maze. Tested at the edge of the table, the unsupported limbs were recovered as promptly as by a normal animal. On a tipping board, adjustments to change of position were sometimes slow or defective, but often normal in all animals. There were suggestions of disturbances of cutaneous sensitivity, a slight analgesia and a tendency to bite at the fore feet during the first few days after operation.

In the one case with lesion in the ventral funiculus the behavior was almost normal, with perhaps a slight hyperextension of the legs for the first few days after operation.

#### PROTOCOLS

## Destruction of fasciculi gracilis and cuneatus and of the pyramidal tract

No. 1. Young male, somewhat unstable during training. Training record: Time, 1463 seconds; errors, 67; trials, 45. Preliminary retention tests: Trials, 10; time, 53 seconds; errors, 0.

Dorsal funiculus sectioned at third cervical segment. On the following days there was some incöordination in walking, with a tendency to walk on the dorsum of the fore feet. The hind limbs were dragged occasionally in walking and in turning toward either side the fore leg of the other side was dragged. There seemed to be some analgesia.

Retention was tested ten days after operation. The record for 15 trials was the following:

Time	Errore	Time	Errors
18	0	8	0
34	0	10	0
83	3	10	0
92	3	22	1
43	0	. 10	0
9	0	9	0
7	0	8	0
6	0		

Locomotion was slowed considerably in the maze, owing largely to the motor incoördination, but the first two trials were made without error and the total record gave clear evidence that the maze pattern was retained

Lesion: Section through the site of injury showed that practically all of the cord above the level of the central canal had been destroyed (plate 1, fig. 1a). The f. gracilis, cuneatus and the pyramidal tract together with the dorsal horns of the gray matter were destroyed. Section at the level of the first cervical segment showed complete degeneration of the f. gracilis and cuneatus except for the lateral areas representing the fibers of the first and second cervical nerves (plate 1, fig. 1b). Scattered degenerate fibers in the lateral and ventral funiculi probably originate from injury to the third pair of nerves and to the dorsal horns of the gray matter. No. 2. Young male. Training marked by periods of refusal to run the maze. Training record: Time, 5409 seconds; errors, 153; trials, 20. Preliminary retention tests: Time, 63 seconds; errors, 0; trials, 10.

On the first day of preliminary retention tests he would not run in the maze and did not eat when placed in the food compartment.

Dorsal funiculi sectioned at third cervical segment. On the following days he tended to drag his legs and walk on the dorsum of the feet, like number 1. Retention was tested ten days after operation. On the first day he showed little evidence of hunger and behaved much as on the first day of the preliminary retention tests, making the following record:

Time	Errora
330	8
360	8
142	4
63	1

He was not fed after these tests and on the next day made the following record:

Time	Errors	Time	Errors
21	0	17	0
10	0	15	0.
17	0	15	0
27	0	9	0
13	0	6	0
14	0		

The record does not show perfect retention but is far better than that for initial learning. In view of his earlier records, we are justified in concluding that this case does not give conclusive evidence for loss of the habit after spinal lesion.

Lesion: Section at the third cervical segment (plate 1, fig. 2a) shows destruction of the greater part of the cord above the level of the central canal. The f. gracilis, cuneatus, and practically all of the pyramidal tract were interrupted. A section below the first cervical roots (2b) shows complete degeneration of the f. gracilis and cuneatus, except for the portions supplied by the second cervical nerve, and extensive degeneration in the right spino-cerebellar tracts corresponding to the degeneration in the right lateral funiculus, shown at the level of injury.

No. 3. Young male. Behavior normal during initial training. Training record: Time, 1669 seconds; errors, 97; trials,

20. No preliminary retention tests given.

The dorsal funiculus was sectioned deeply. On the following days there was marked incoördination in walking. The fore feet were usually flexed with the dorsum on the ground. Hind feet were frequently dragged in walking and were not drawn up when placed over the edge of a support. When making coördinated steps the feet were widespread, so that the belly touched the ground. There was probably some positive disturbance of sensitivity, as on the eighth day he amputated his left fore paw. Retention was tested fourteen days after operation. On the first day he would not eat in the maze and his behavior was erratic. The record for the first day was the following:

Time	Errors
24	2
<b>4</b> 6	4
38	2
689	20

He was not fed and on the next day made the following record:

Time	Errors	Time	Errors
17	0	27	0
60	0	26	4
14	2	9	0
7	0	8	0
8	1	7	0
7	1	8	0
8	0	12	1

The behavior on the first day did not resemble that of an untrained animal and there was indifference to food. On the following day, when the incentive to run was increased, errorless trials were obtained from the first. Like the record of number 2, this is somewhat ambiguous. It indicates some retention, but is impossible to interpret.

Lesion: At the third cervical level the position of the lesion was clearly marked by scar tissue and clots (plate 1, fig. 3a). It involved all of the dorsal funiculus, the greater part of the gray matter on the right, and the dorsal third of the lateral funiculus, including the rubrospinal tract. At the second cervical segment (plate 1, fig. 3b) there was complete degeneration of the f. gracilis and cuneatus and considerable degeneration in both lateral columns. No. 4. Large adult male. Given 50 trials overtraining. Training record: Time, 1676 seconds; errors, 49; trials, 29. Overtraining: Time, 279 seconds; errors, 1; trials, 50. Preliminary retention tests: Time, 106 seconds; errors, 3; trials, 16.

The dorsal funiculus was deeply incised at the third cervical segment. On the following days he walked with a sprawling gait, falling occasionally in making turns and frequently stepping on the dorsum of the feet. Retention was tested five days after operation. The record was the following:

Time	Errors	Time	<b>Errors</b>
7	0	6	0
6	0	8	0
12	0	6	0
7	0	8	0
5	0	8	0

The record shows perfect retention and a speed in traversing the maze only slightly inferior to the preoperative record. The maze was covered and performance tested in darkness. The record was the following:

Time	Errors	Time	Errors
28	2	7	0
22	0	7	0
8	0	8	0
8	0	6	0
10	0	8	0

After the exploration of the covered top of the maze in the first trial no further errors were made when moving in total darkness.

Lesion: Section at the third cervical level showed complete interruption of the dorsal funiculus, destruction of the posterior horns and a considerable amount of degeneration in the right lateral funiculus (plate 1, fig. 4a). Section above the first cervical level (plate 1, fig. 4b) showed complete degeneration of the f. gracilis and cuneatus with considerable degeneration in the region of the dorsal spino-cerebellar tract.

These four cases all show complete interruption of the fasciculus gracilis, fasciculus cuneatus and pyramidal tracts, with more or less degeneration in the regions of the rubro-spinal tracts and of the dorsal spino-cerebellar bundles. Two cases, numbers 2 and 3, showed somewhat erratic behavior in preoperative tests and similar behavior after operation so that their records are not clear cut. They do, however, show some indication of retention. The other two, with equally extensive destruction give unequivocal evidence for perfect retention of the maze habit. Destruction of the dorsal funiculus did not in the least affect their ability to make the correct turns of the maze.

# Destruction of fasciculus gracilis and fasciculus cuneatus without extensive injury to the pyramidal tracts

No. 5. Large male, more than usually stable. Initial training: Time, 957 seconds; errors, 32; trials, 14. Preliminary retention tests: Time, 35 seconds; errors, 0; trials, 10. Overtraining: Time, 182 seconds; errors, 1; trials, 50.

Ten trials with the maze darkened were given before operation.

Time	Errora	Time	Errors
25	2	4	0
18	3	7	0
5	0	6	0
4	0	5	0
4	0	4	0

The dorsal funiculus was sectioned at the third cervical segment. For the first few days the animal tended to circle to the right in walking. This condition cleared up quickly, leaving the usual picture of reduced sensitivity to posture. Retention was tested eight days after operation, with the following record.

Time	Errors	Time	Errors
8	0	13	0
5	0	4	0
16	0	8	1
10	0	6	0
7	0	12	0

On the following day, with the maze darkened, he made the following record:

Time	Errora	Time	Errors
38	2	8	0
36	2	7	0
10	0	6	0
8	0 '	6	0
18	1	12	0

The post operative tests show some slowing in locomotion but no loss of ability to traverse the maze correctly. With vision eliminated the record was as good as that made under similar conditions before the spinal lesion. Lesion: Section above the third cervical level showed complete destruction of the f. gracilis and f. cuneatus with little if any involvement of the pyramidal tracts (plate 1, fig. 5a). Section at the first cervical level showed degeneration restricted almost entirely to the two ascending fasciculi (plate 1, fig. 5b).

## No. 6. Young male, very unstable throughout the experiment. Initial training: Time, 710 seconds; errors, 53; trials,

50. Preliminary retention tests were not given.

The dorsal funiculus was transsected at the third cervical segment. On the following days little abnormality of behavior was noted beyond a tendency to walk on the dorsum of the paws and drag them slightly. Retention was tested seven days after operation, with the following record, made in three consecutive days:

Time	Errora	Time	Errors	Time	<b>Errors</b>
322	12	23	2	8	0
201	11	15	0	13	2
80	3	15	0	12	0
33	1	10	0	7	0
34	1	8	0	11	1
16	1	8	0	8	0
9	0	11	1	5	0
12	1	11	0	5	0
7	0	22	2	6	0
15	2	92	6	5	0

This record is not significantly better than the initial training record. The behavior during retention tests was characterized throughout by prolonged periods of quiescense in the starting compartment and by failure to eat after the food had been reached.

Lesion: There was active infection of the wound, inflammation of the cord with necrosis of the dorsal horns at the third cervical level (plate 2, fig. 6a). Section at a higher level showed extensive degeneration of the f. gracilis and cuneatus with some degenerated fibers in the spinocerebellar tracts, chiefly on the left (plate 2, fig. 6b).

Of these two cases the first alone can be considered. The infection rules out the negative evidence of the second and leaves only as significant the fact that, in spite of the infection and destruction of tissue, he was able ultimately to traverse the maze without error. Number 6, with greater actual destruction showed perfect retention of the maze habit and ability to traverse the maze in darkness as well after as before the spinal injury.

## Partial destruction of the dorsal and of one lateral funiculus

No. 7. Large male, about 200 days old. Initial training: Time, 1077 seconds; errors, 38; trials, 21. Overtraining: Time, 247 seconds; errors, 0; trials, 50. Preliminary retention tests: Time, 41 seconds; errors, 0; trials, 10.

Preliminary tests with the maze darkened gave the following:

Time	Errors	Time	Errors
85	5	7	1
8	1	4	0
5	0	4	0
5	0	5	0
7	1	5	0

An attempt was made to divide the dorsal funiculus, but the knife was driven deeply into the left side of the cord. General behavior after operation was not markedly different from that of animals with only the dorsal funiculus cut, except that adjustments to tipping the substratum were made rather slowly and inaccurately. Retention was tested six days after operation with the following record.

Time	Errors	Time	Errors
11	0	10	0
22	1	10	0
11	0	7	0
12	0	8	0
10	0	8	0

Tests with the maze darkened gave the following record:

Time	Errors	Time	Errors
35	0	33	0
25	0	14	0
38	0	27	0
55	0	43	0
22	0	30	0

The prolonged time in these tests is due to the fact that the animal was disturbed by the cover of the maze and repeatedly stopped during every trial to push against it with his nose.

Lesion: The cut extended diagonally from the outer margin of the right f. cuneatus through the left dorsal horn to the lower margin of the left spino-cerebellar tract, interrupting all the fibers above this line (plate 2, fig. 7a). Section at a higher level shows almost complete degeneration of the f. gracilis and cuneatus of both sides, degeneration of the left spino-cerebellar tracts and many degenerate fibers in the region of the spino-tectile bundles (plate 2, fig. 7b).

This case, with nearly complete section of the f. gracilis and cuneatus and interruption of the spino-cerebellar tracts of one side gave clear evidence of ability to run the maze even in darkness.

Section of the lateral funiculi

No. 8. Large male, about 200 days old. Initial learning: Time, 5861 seconds; errors, 83; trials, 20. Overtraining: Time, 431 seconds; errors, 0; trials, 50. Preliminary retention tests: Time, 55 seconds; errors, 0; trials, 10.

Preliminary tests with the maze darkened gave the following results:

Time	Errors	Time	Errors
125	9	7	0
16	1	5	0
15	1	3	0
8	0	4	0
10	0	3	0

The lateral funiculus was divided on each side at the level of the third cervical segment. On the following days he showed tremor and hyperextension of the legs, staggering gait with a tendency to fall to the side, especially in turning, and constant balancing movements of the tail. There was some difficulty in adapting to inclination of the substratum. Retention was tested ten days after operation with the following record:

Time	Errors	Time	Errors
10	0	10	0
11	0	· 12	0
8	0	11	0
11	0	8	0
11	0	7	0

With the maze darkened, the record was the following:

Time	Errors	Time	Errors
115	4	22	0
46	1	18	0
15	0	20	0
15	0	100	2
18	1	13	0

This record is somewhat inferior to the preoperative one, but gives conclusive evidence of ability to traverse the maze in darkness.

Lesion: At the third cervical level the lesions of the two sides were separated longitudinally by about one millimeter, that on the right being the more cephalad (plate 2, fig. 8). On the left the knife had passed though the fasciculus cuneatus and diagonally downward to interrupt the entire lateral funiculus, involving the dorsal horn of the gray matter as well. On the right the lesion extended vertically from the lateral margin of the fasciculus cuneatus through the dorsal and ventral horns and involved the entire lateral funiculus (plate 2, fig. 8).

No. 9. Large male, about 200 days old. Initial training: Time, 1297 seconds; errors, 57; trials, 18. Overtraining: Time, 338 seconds; errors, 0; trials, 50. Preliminary retention tests: Time, 47 seconds; errors, 0; trials, 10.

Preliminary tests with the maze darkened gave the following record:

Time	Errors	Time	Errora
175	11	5	0
6	0	4	0
5	0	5	0
5	0	5	0
5	0	5	0

The lateral funiculi of both sides were cut at the third cervical level. The subsequent behavior of the animal was much like that of number 8, but with a less steady gait and some indication of analgesia. In retention tests he fell almost every time in making a turn in the maze. Retention tests were given six days after operation, with the following record:

Time	Errors	Time	Errors
34	1	8	0
10	0	9	0
8	0	10	0
10	0	9	0
8	0	9	0

Tests with the maze darkened gave the following:

Time	Errors	Time	Errors
37	2	9	0
27	1	24	1
13	0	11	0
12	0	9	0
9	0	10	0

Lesion: Section at the level of the injury showed total destruction of both lateral funiculi with little or no injury to the dorsal or ventral funiculi. Section at the first cervical roots showed degeneration of the spino-cerebellar tracts of both sides and ascending degeneration of fibers from the third dorsal root in the right f. cuneatus. (Plate 2, figs. 9a and 9b.)

No. 10. Male, about 200 days old. Initial training: Time, 2022 seconds; errors, 80; trials, 45. Overtraining: Time, 322 seconds; errors, 3; trials, 50. Preliminary retention tests: Time, 85 seconds; errors, 1; trials, 14.

The lateral funiculus of each side was divided at the third cervical segment. On the following days he showed hyperextension of the legs, staggering gait with tendency to fall to the side, constant balancing movements of the tail and some tremor of the legs.

Retention was tested five days after operation with the following record:

Time	Errora	Time	Errors
14	0	8	0
12	0	10	0
45	2	29	0
12	0	10	0
9	0	8	θ

Tests with the maze darkened gave the following:

Time	Brrors	Time	Errors
56	1	18	0
82	2	11	0
29	0	17	0
52	1	10	0
20	0	10	0

In these tests the additional time was consumed in pushing at the cover of the maze.

Lesion: Section at the third cervical level showed lesions restricted to the dorsal halves of the lateral funiculi without invasion of the gray matter (plate 2, fig. 10a). At the first cervical level there was partial degeneration of the dorsal spino-cerebellar tract of each side with no other involvement.

These three cases, involving interruption of all the paths of the lateral

funiculi and considerable motor disturbance, resembling a cerebellar ataxia, showed perfect retention of the maze habit with ability to traverse the maze in the absence of visual cues.

## Injury to the ventral funiculus

No. 11. Large male, about 200 days old. Initial training: Time, 2491 seconds; errors, 71; trials, 23. Overtraining: Time, 647 seconds; errors, 7; trials, 50. Preliminary retention tests: Time, 83 seconds; errors, 1; trials, 11.

Preliminary tests with the maze darkened gave the following:

Time	Berors	Time	Errors
86	8	16	1
18	1	6	0
11	1	15	1
10	0	7	0
8	0	6	1

The cord was exposed, raised, and the knife passed through the ventral columns. Following the operation there was little disturbanace of behavior beyond a slight tendency to drag the hind feet in walking. Retention tests were given six days after operation.

Time	Errors	Time	Errors
9	0	13	0
15	0	25	0
15	0	14	0
13	0	12	0
10	0	11	0
58	1		

Tests with the maze darkened gave the following:

Time	Errors	Time	Errors
54	4	28	0
17	0	18	0
20	0	13	0
15	Ö	35	1
14	0	28	0

Lesion: In dividing the cord for fixation the segment containing the lesion was destroyed. A section immediately above the lesion shows extensive degeneration throughout the ventro-lateral column of the left side with little involvement of the right. It seems certain that the knife

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passed diagonally downward to destroy all of the funiculus below the ventral horn on the left side, without serious injury to the right (plate 2, fig. 11a). Ascending the cord the degenerated fibers disappear rapidly into the gray matter so that only a few are left at the level of the decussation of the pyramids (plate 2, fig. 11b). These lie chiefly in the ventral spino-cerebellar bundle and in the spino-tectile bundles.

This case, with extensive destruction in the ventral funiculus, showed quite undisturbed ability to traverse the maze.

#### RESULTS OF THE EXPERIMENTS

The records of the animals in the various tests are summarized The animals of the first group tested are marked with in table 1. Their records in postoperative retention tests are an asterisk. much worse than those of the others. This is unquestionably due largely to the fact that they were not rendered sufficiently hungry on the first day of retention tests. No. 6 alone showed retardation sufficient to produce anything like a normal learning curve in postoperative tests and he showed evidence of infection in the cord. Numbers 2 and 3 made errors only on the first day of retention tests and on later days, when eager for food, made no errors. Number 1 gave evidence of retention from the first. All these cases ran the maze without error at some time after operation so that they at least show the possibility of acquisition of the habit after section of the dorsal funiculi. However, we are justified in discounting completely their failures after operation in view of the records of the second group.

Number 4 with interruption of the entire dorsal funiculus including the pyramidal tract and number 5 with section of the f. gracilis and cuneatus only made no errors in postoperative tests under the conditions of training. In tests with the darkened maze they made fewer errors after operation than the average for all animals under the same conditions before operation. From them we may conclude that accuracy in running the maze is not disturbed by complete destruction of the afferent and efferent paths of the dorsal funiculus. Number 7 showed a similar ability to traverse the maze after almost complete destruction of the f. gracilis and cuneatus and partial destruction of one lateral funiculus.

Numbers 8, 9 and 10 made practically perfect records in postoperative tests after section of the lateral funiculi. Their motor incoördination was severe, they fell at almost every turn in the maze yet recovered and turned correctly without getting into the *culs de sac*. The lesion in number 8 certainly involved all of the tracts of the lateral funiculus with a considerable part of the left f. cuneatus as well. The records of these cases show that none of the tracts of the lateral funiculus is essential for maze running, either under ordinary conditions or in darkness.

TABLE 1					
Summary	of	results	from	protocols	

The errors made in the various tests are given as probably the most reliable of the three criteria.

NUMBER	ERRORS LEARNING	ERRORS PRELIMI- NARY RETENTION	ERRORS PRELIMI- NA RY DARKNESS	ERRORS POST- OPERATIVE	ERRORS POST- OPERATIVE DARKNESS	LESION
1*	67	0		7		Entire dorsal funic- ulus
2*	153	0		21		Entire dorsal f.
3*	97			37		Entire dorsal f.
4	49	3		0	2	Entire dorsal f.
5	32	0	5	0	5	Dorsal f.
6*	53			44		Dorsal f.
7	38	0	8	1	0	Dorsal and lateral f.
8	83	0	11	0	8	Lateral f.
9	57	0	11	1	4	Lateral f.
10	80	1		2	4	Lateral f.
11	71	1	12	1	5	Ventral f.

Number 11, with severe lesion on the left in the ventral funiculus, gave no evidence of disturbance of the maze habit. The lesion did not involve that portion of the funiculus lying between the ventral horns but shows that the other parts of the funiculus are not essential.

The only significant difference between the preliminary and postoperative tests is in the time required for traversing the maze. The average time per trial for trials without error in the preliminary retention tests (cases 4, 5, 7, 8, 9, 10, and 11) is 5.4 seconds. The corresponding average for the postoperative tests is 9.8 seconds. However, animals with lesions in each region occasionally traversed the maze as quickly after as before operation so that this retardation can not be taken as evidence for a necessity to "feel their way" with greater care than before operation. As a matter of fact all the animals showed ample motor disturbance to account for all the excess time of the postoperative tests and their behavior gave not the slightest indication of an increased dependence upon tactile or olfactory cues.

## MOTOR CONTROL OF THE MAZE HABIT

The most surprising result of these experiments was the capacity of the animals to orient in the maze after section of the descending tracts. The inclusion of these tracts in the lesions was inadvertant and the lack of effects wholly confounding. After the first few days the motor capacities of the animals seemed but little disturbed. Their incoördinations of movement resembled the syndromes of sensory disturbances in man rather than any direct paralytic symptoms. Coördinations between sensory and motor systems on opposite sides of the lesions appeared in all cases shortly after the operations. These included such activities as postural coördinations of the head and trunk in sitting erect, turning the head toward the locus of protopathic stimulation on the limbs and trunk, and washing the face with the forepaws, as well as orientation in walking.

The lack of specific symptoms following section of the pyramidal tract is to be expected from the similar absence of motor disturbances following ablation of the entire stimulable cortex. Possibly some of the disturbances of gait after section of the lateral funiculi are referable to the destruction of the rubrospinal paths rather than of the spino-cerebellar, but there would be little reason to assign the control of habitual movements to these paths. This leaves only the less definite thalamico-spinal ponto-spinal and similar fibers as potential conductors of the orienting movements and since, in one case or another, the lesions interrupted practically all of the cross-sectional areas of the cord, we are forced to the conclusion that the motor impulses essential to the performance of the acts described above are not restricted

to any of the definite tracts, but descend, either diffusely or over alternative paths in the different funiculi.

Similar conditions in the motor functions of the cord have been described by other investigators. Porter's work ('95) on the path of the respiratory impulses is well known. Weiss ('79) found after hemisection of the cord in dogs a rapid restitution of motor functions and Mott ('92) reported a similar recovery in the monkey. Osawa ('82) and Borgherini ('86)<sup>2</sup> found that a double hemisection, when the lesions were separated by six or seven spinal segments, might be followed by restoration of motility of the hind legs in the dog. All these observations point to a great plasticity or diffuseness of motor conduction in the cord.

In contrast to this condition of the motor paths, the sensory paths seem much more definite and less plastic. All of these investigators report a much slower recovery of sensory functions after spinal lesion with never a complete restitution. Why this difference between afferent and efferent conduction should exist is still wholly obscure, but the difference seems to be clearly established, both by earlier and by our own observations and is of considerable importance for the interpretation of our data.

## SENSORY COMPONENTS OF THE HABIT

The lesions to the dorsal funiculi produced serious sensory disturbances, if we may judge from the defects of posture shown by the animals. Lesions to other parts of the cord, although followed by few signs of sensory defect, were complete enough to assure that all ascending paths were interrupted. It is clear that the ability of the animals to traverse the maze was unimpaired by the observed sensory defects and further, that no particular afferent path of the cervical cord is essential for the performance of the maze habit. There seem to be four possible alternative interpretations of the data:

1. The kinesthetic impulses were eliminated by the lesions and the animals shifted to other sensory cues for direction in the maze. The following controls of such sensory cues seem to rule

<sup>2</sup> Cited from Sherrington ('98).

out this interpretation. Olfaction. With the older group at the termination of the preliminary retention tests the maze was rinsed out and allowed to stand open and unused for six days before the postoperative retention tests. There is no evidence to indicate that under such conditions directive odors would remain. Moreover, the rat which is following an olfactory trail shows an unmistakable pattern of movements (Vincent, '15) and none of the animals in our experiments showed any significant sniffing at the critical points in the maze during the postoperative tests. Audition. In passing from one alley of the maze to the next the animal is confronted by two passages of unequal length and must always turn toward the longer. It is conceivable that differences in echoes from the two ends of the passage might give directive cues, but evidence which we have of the enormous difficulty of such a discrimination of localization of sounds for the rat makes such a speculation preposterous. Differences in air pressure in the two passages are positively excluded by the fact that the alleys were covered only by coarse netting. Vision. The maze was covered in a way to exclude all light. The animals were disturbed by the presence of the cover and spent much time in pushing against it through the wire netting but their behavior in this respect did not differ before and after operation and their accuracy after operation was in general better than before. Cutaneous sensitivity. The sides of the maze and the angles of the doorways between the alleys offer tactile stimuli. The form of the maze requires that the animal pass through each doorway as he reaches it. Traversing the alleys and turning through the doors may therefore be controlled by tactile stimuli. But as the animal passes through each doorway he is confronted by the alternatives of a right or left turn with identical tactile situations at each side. These are the critical points in the maze, the most difficult for the normal animal to learn (Hubbert and Lashley, '17) and they offer no differential cutaneous cues.

2. The important proprioceptive systems for the maze habit are the vestibule and the muscles of the neck, whose innervation through the first and second cervical nerves was spared in the cases studied. The great importance of these systems in postural

adjustment has been shown by Magnus and Leeuwen ('14) but our observations on the behavior of the rats in the maze do not bear out the assumption of especial importance of these systems for the maze habit. The tests in darkness seem especially significant here. When placed in the covered maze all the animals were markedly disturbed. They pushed against the cover, sniffed at the cloth, moved forward a few inches and repeated the exploration. In the first trials in the darkened maze the progress of all the animals could be traced by their continuous pushing at the flexible cover. This behavior does not indicate a disturbance of orientation, for it may be induced by placing a strip of cloth across the top of the maze without significantly modifying visibility of the path. It is an expression, rather, of the tendency of the rat to explore thoroughly any changed condition in a familiar situation. The lack of real disturbances in orientation is illustrated by the record of No. 7 who required thirty-five to fifty-five seconds for each of the first four trials in the darkened maze, whose progress could be traced by a continuous jerking of the cover of the maze, yet without a single entry into a cul de sac. On the other hand, these exploratory movements of the head involve a continuous alteration of the pattern of stimulation from the vestibules and neck muscles which does not provide an adequate pattern for a proprioceptive motor chain. For, even if some of the original elements of the chain are retained, such as stereotyped movements of the head at the critical turns of the maze, there is required some mechanism by which these significant movements may be distinguished from the others with which they make up a continuous series. To say that the habit is carried out by utilization of these movements seems thus to beg the whole question, since the significant movements can be elicited and effective for subsequent behavior only in case the animal is already oriented and this orientation can not be ascribed to the previous series of irregular movements of the head.

3. The proprioceptive paths of the cord are scattered in such a way that a sufficient number of afferent impulses to control orientation may ascend through any part of the cord which happens to remain intact. It may be argued that, if the motor impulses can pass caudad over any intact area of the cord, the sensory impulses may likewise pass cephalad. Opposed to this interpretation are the very obvious disturbances in sensitivity to posture after section of the dorsal funiculus and the ataxic symptoms after section of the lateral funiculi; facts which suggest as definite a localization of afferent paths in the rat's cord as in that of higher forms. Evidence from the dog and monkey (page 94) also suggests that sensory conduction in the cord is more definitely restricted than motor. Finally we have seen that, anatomically, the tracts are perfectly definite and the corresponding appearance of functional disturbances suggests as distinct a localization of afferent conduction paths as occurs in higher forms.

Thus, none of these three hypotheses seems admitted by the conditions of our experiments. The theory of chain-reflex arcs demands a continuity of adequate stimuli and a constancy of motor response which do not correspond to the observed facts. Marked reduction of proprioceptive sensitivity does not reduce accuracy of maze running although it results in motor disturbances sufficient to alter completely the pattern of motor activity. The hypothesis most in harmony with the facts seems to be one assuming that the successive activities of the maze habit are largely determined by central nervous activities. We are completely ignorant of the details of the mechanism implied here but may hazzard some such guess as the following.

4. The engram of the maze habit consists of some central organization in which the general direction and succession of turns are so recorded that, once the series is initiated, the essential sequence of movements may be performed in the absence of sensory control and with considerable variation in the actual movements produced. This is, of course, a very vague statement and implies conceptions of plasticity in nervous functioning which run counter to the whole doctrine of conditioned reflexes, yet the problem seems at present to admit of no greater simplification.

The maze habit presents many complexities. There is clear evidence that the animals acquire some general orientation in

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addition to the habits of making particular turns and at present this orientation can only be described as an abstraction of a general direction from the diverse directions of the successive The variations which may occur in the motor pattern in allevs. traversing the maze, best illustrated by the disturbances following cerebellar lesions, point to a functional equivalence of motor activities similar to the transfer of motor habits to an unpracticed limb reported earlier (Lashley, '24a, '24b). It is not within the scope of the present paper to enter into a discussion of the problems suggested here. It must suffice that the maze habit involves intricacies which are not accounted for by any of the theories hitherto proposed, that it has common elements with the problems of rational learning and purposive acts, and that its analysis must await a better understanding of such complexities of behavior. Our present experiments serve only to emphasize the inadequacy of the chain-reflex theory of maze learning. Earlier work has shown the difficulty of interpreting the habit as a succession of reactions to exteroceptive cues. Our data indicate an exactly similar difficulty in basing the habit on proprioceptive ones.

## DEFECTS OF THE GENERAL THEORY OF CHAIN-REFLEXES

Attempts to explain behavior in terms of simple reflexes elaborated only by combination in chain-reflex arcs have proved of little value for an understanding of the more intricate problems of psychology and, where they have been tested by experiment, have received little factual support. Perhaps the simplest phase of the problem is that involved in the control of accuracy of movement. Are movements of definite extent or duration limited by the sensory components of the movements or by some preliminary central set which determines the limit of movement independently of sensory processes? In many cases the sensory processes obviously play a rôle in the control of accuracy, but it seems certain that such "current control" is not essential to accurate movement. Observations of Bowditch and Southard ('81), on the comparative accuracy of visual and kinesthetic control, of Loeb ('97) on the bilateral symmetry of movement, and of Woodworth ('99) on the relation of speed to accuracy all indicate a relative independence of sensory control. Lashley ('17) has shown that complete anesthesia to movement after lesion to the dorsal roots or spinal funiculus may not result in any disturbance in the accuracy of extent or duration of movements which are not opposed by external resistance.

The results of these studies seem explicable only on the assumption that the speed and extent of movement (intensity and duration of muscular innervation) may be determined in the initial set before any overt movement has actually occurred, and that "current control" enters in only when there is some departure from the conditions for which the initial adjustment was made.

Many swift sequences of movement, such as the execution of rapid musical passages, demand the same sort of explanation of control through the initial set. In sight reading of music it is often impossible to take in the individual notes and the player must learn to respond to groups of notes as does the reader to groups of letters or words. The effective stimulus is momentary yet determines the release of a series of movements in definite order and at such a rate that the initiation of each can not be ascribed to the completion of the preceding.

The theory of maintenance of emotional status by proprioceptive excitations is not in better case. In the large number of studies of bodily changes in emotion there is not one established invariable correlation between muscular pattern and emotional state. On the contrary, the picture of motor changes during emotion is highly variable, in contrast to the relatively stable motivating effects of the emotions and there is at least as much reason to interpret facial and bodily expression in emotion as an actual "expression," determined in part as a social gesture and in part by chance irradiation of intense neural excitation (Landis '24), as to accept the hypotheses of James and Lange.

Finally, such experimental evidence as we have on the question of implicit movements in thinking (Thorsen, '25) directly opposes the chain-reflex theory and suggests that, when implicit movements do occur, it is as a result of irradiation and not a part of the normal process of thinking.

The chief attractiveness of the chain-reflex theories lay in their promise that psychology might progress by simple objective methods and escape the need for indirect inference concerning neural complexities which could not be directly observed. The accumulation of evidence against the existence of observable muscular contractions which can serve as a basis for the chain reflexes forces us either to seek the completion of the reflex in action currents without observable muscular contraction (Watson '24-25), or to turn to the central nervous system for explanation of maintenance as well as coördination of activity. There is no direct evidence supporting the excitatory effects of such action currents and it will not be less difficult to disentangle the complex electrical phenomena of tonus and subliminal movement than to measure cerebral changes directly, so that the advantage of greater objectivity can not be claimed for the motor theory.

On the other hand, for the two simplest types of activity where the theory of chain reflexes might be expected to apply, the current control of movement and maze running, the evidence seems definitely to oppose the motor theory and to favor the existence of some wholly central mechanism as the determiner of the motor sequences.

#### SUMMARY

Rats were trained in a maze with eight *culs de sac* and subjected to partial section of the spinal cord in the upper cervical region. A series of cases with complete interruption of the dorsal or lateral funiculi and one animal with partial section of the ventral funiculus were obtained. The ability of the animals to traverse the maze was then tested. In spite of serious sensory disturbances, some animals with each type of lesion traversed the maze without significant errors after each type of lesion. Control experiments indicated that they did not fall back upon exteroceptive cues for orientation and that the persistence of the habit can not be referred to the remaining proprioceptive sensitivity. From this it is argued that the maze habit can not be interpreted as a series of kinesthetic-motor reflexes but must be referred to some intraneural mechanism capable of producing an integrated sequence of movements in the absence of directive sensory cues.

#### REFERENCES

- BECHTEREW, W. VON. Ueber die versecheidenen Lagen und Dimensionen der Pyramidenbahnen, usw. Neurol. Zentralb., 1890, 738-741.
- BOWDITCH, H. P., AND SOUTHARD, W. F. A comparison of sight and touch. Jour. Physiol., 1881, iii, 232-254.
- CARR, H., AND WATSON, J. B. Orientation in the white rat. Jour. Comp. Neurol. and Psychol., 1908, xviii, 27–44.
- GOLDSTEIN, K. Zur vergleichenden Anatomie der Pyramidenbahn. Anat. Anz., 1904, xxiv, 451-454.
- HEAD, H. Studies in Neurology, London, 1920.
- HELD, H. Der Ursprung des tiefen Markes der Vierhügelregion. Neurol. Centralb., 1890, ix, 481-483.
- HUBBERT, H. B., AND LASHLEY, K. S. Retroactive association and the elimination of errors in the maze. Jour. Animal Behav., 1917, vii, 130–138.
- KING, J. L. The cortico-spinal tract of the rat. Anat. Record, 1910, iv, 245-252.
- LANDIS, C. Studies of emotional reactions: II. General behavior and facial expression. Jour. Comp. Psychol., 1924, iv, 447-511.
- LASHLEY, K. S. The accuracy of movement in the absence of excitation from the moving organ. Amer. Jour. Physiol., 1917, xliii, 169–194.
- LASHLEY, K. S. Studies of cerebral function in learning. V. The retention of motor habits after the destruction of the so-called motor areas in primates. Arch. Neurol. and Psychiat., 1924a, xii, 249-276.
- LASHLEY, K. S. Studies, etc. VI. The theory that synaptic resistance is reduced by the passage of the nerve impulse. Psychol. Rev., 1924b, xxxi, 369-375.
- LASHLEY, K. S., AND MCCARTHY, D. A. The survival of the maze habit after cerebellar injuries. Jour. Comp. Psychol., 1926, vi, 423-433.
- LENHOSSEK, M. VON. Ueber die Pyramidenbahnen im Rückenmarke einiger Säugetiere. Anat. Anz., 1889, iv, 208-219.
- LINOWECKI, A. J. The comparative anatomy of the pyramidal tract. Jour. Comp. Neurol., 1914, xxiv, 509-530.
- LOEB, J. Untersuchungen über den Fühlraum der Hand. Arch. f. d. ges. Physiol., 1897, xli, 107-127.
- MAGNUS, R., U. LEEUWEN, W. S. V. Die akuten und die dauernden Folgen des Ausfalles der tonischen Hals- und Labyrinthreflexe. Arch. f. d. ges. Physiol., 1914, clix, 157-217, 224-249.
- Morr, F. W. Results of hemisecting the spinal cord in monkeys. Philos. Trans. Roy. Soc. London, 1892, clxxxiii (B), 1-59.
- PAPEZ, J. W. The rubro-spinal tract, Marchi method. Anat. Record, 1923, xxv, 147.
- PORTER, W. T. The path of the respiratory impulse from the bulb to the phrenic nuclei. Jour. Physiol., 1895, xvii, 455–485.
- RANSON, S. W. The fasciculus cerebrospinalis in the albino rat. Amer. Jour. Anat., 1913, xiv, 411-424.

RANSON, S. W. The tract of Lissauer and the substantia gelatinosa rolandi. Amer. Jour. Anat., 1914, xvi, 97-126.

RANSON, S. W. A note on the degeneration of the fasiculus cerebrospinalis in the albino rat. Jour. Comp. Neurol., 1914, xxiv, 503-507.

SHERRINGTON, C. S. The Spinal Cord. In Schafer's Text-book of Physiology, London, 1898-1900, ii, 783-883.

- THORSEN, A. M. The relation of tongue movements to internal speech. Jour. Exp. Psychol., 1925, viii, 1-32.
- VAN DER VLOET. Ueber den Verlauf der Pyramidenbahn bei niederen Saugetieren. Anat. Anz., 1906, xxix, 113-132.
- VINCENT, S. B. The white rat and the maze problem. Jour. Animal Behav., 1915, v, 1-24; 140-157; 175-184; 367-374.
- WATSON, J. B. Kinaesthetic and organic sensations; their rôle in the reactions of the white rat to the maze. Psychol. Rev. Monogr. Suppl., 1907, viii, 1-100.
- WATSON, J. B. The place of kinaesthetic, visceral and laryngeal organization in thinking. Psychol. Rev., 1924, xxxi, 339-348.

WATSON, J. B. Behaviorism. New York, 1924-25.

- WEISS, N. Untersuchungen über die Leitungsbahnen in Rückenmarke des Hundes. Sitzungsb. d. k. Akad. d. Wissensch., Wien, 1879, lxxx, (Abt. 3), 340-356.
- WOODWORTH, R. S. The accuracy of voluntary movement. Psychol. Rev. Mongr. Supple., 1899, iii, (No. 13), 1-114.

## PLATE 1

FIG. A. Composite diagram showing degeneration at the level of the trapezoid body after section of the lateral and ventral funiculi.

FIG. B. Descending degeneration after section of the dorsal funiculus.

FIG. C. Descending degeneration after section of both lateral funiculi.

c.t., trepezoid body; f.c., fasciculus cuneatus; f.g., fasciculus gracilis; m., median longitudinal bundle; n.c.V., nucleus of fifth nerve; p., cortico-spinal tract; r.s., rubro-spinal tract; s.c.d., dorsal spino-cerebellar tract; s.c.v., ventral spinocerebellar tract; s.q., spino-quadrigeminal system (?); S.c., Schultze's comma.

FIGS. 1 TO 5. Cases with lesions in the dorsal funiculus. The figures marked a represent sections through the level of maximum extent of the lesion (third cervical), those marked b, sections at about the first cervical level. The numbers of the figures correspond to the numbers of the animals in the text.

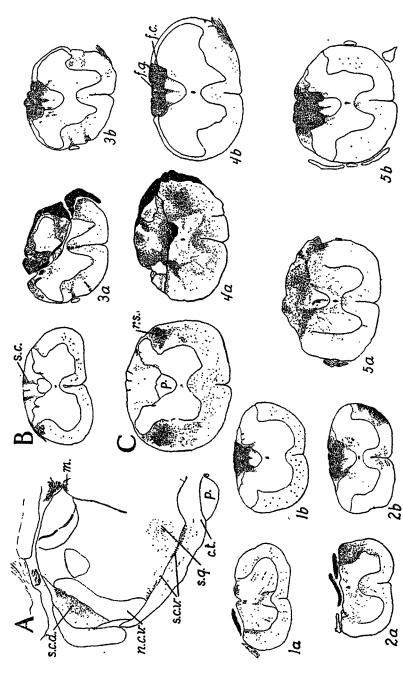
The stippling represents degenerated fibers, solid black unabsorbed clots.

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SPINAL CONDUCTION AND KINESTHETIC SENSITIVITY X. 5. LABELLEY AND JOSEPHINE BALL



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## PLATE 2

FIGS. 6 TO 11. Sections from cases bearing corresponding numbers in the text. a, level of maximum lesion; b, first cervical level or level of decussation of the pyramids.

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PLATE 2

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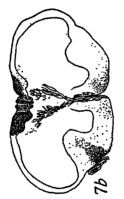
SPINAL CONDUCTION AND KINESTHETIC SENSITIVITY K. S. LASHLEY AND JOSEPHINE BALL







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