BILATERAL ASYMMETRY AND ITS RELATION TO CERTAIN
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INHERITED AND NON-INHERITED DEVIATIONS FROM BILATERAL SYMMETRY

Of the three axes of symmetry which characterize bilateral animals, two
may be defined in terms of function or of the incidence of external forces.
The terms anterior and posterior relate to the direction of locomotion.
The former end is, by definition, the end which moves forward, and it is
here, accordingly, that we find the mouth, the principal sense organs and
various organs of offense and defense. Again, the dorsal and ventral sides
of the body are differentiated with reference to the action of gravity, light,
contact with environment, etc. The expressions right and left, on the
contrary, correspond to no such simple relations to incident forces. Some
degree of functional and structural differentiation may occur, to be sure,
between these two sides of the body, but such differentiation could, in every
case, undergo complete reversal without affecting the welfare of the organism. It is nevertheless an interesting fact that many of these "normal"
deviations from bilateral symmetry are, within a given species, constantly
dextral or sinistral, as the case may be. To consider a few familiar instances
among the vertebrates alone, the birds have but a single functional ovary, the left, while in the snakes the left lung is greatly reduced. The flounders lie upon one side, which may be either the right or the left, but in most cases this is the same in all members of the species. Extensive structural differentiation has arisen among these fishes in connection with this habit. Certain cetaceans have conspicuous asymmetry in the structure of the skull. In all of the true vertebrates, including man, there are marked asymmetries of a constant character, affecting the internal organs, particularly the digestive and circulatory systems. Finally, in man, we have the familiar functional differentiation of the right and left fore-limbs, accompanied by a no less marked functional differentiation of the two sides of the brain. Less evident, though quite measurable, structural differentiations are to be found in the bones of the two sides of the skull, of the fore-limbs, and perhaps also of the hind-limbs.

That all these cases of normal asymmetry are hereditary in a real sense hardly needs argument. The sole alternative explanation, namely, that they are due to environmental influences which affect the two sides unequally, and which are repeated generation after generation, needs only to be stated in order to be refuted. The absurdity of such an explanation is most evident in those cases, e.g., the bony fishes—in which no intra-uterine development occurs. It may be true that much of this asymmetry arises during development as a result of the mutual pressure of the various parts, but it must, at the outset, have a germinal basis. As to the "mode of inheritance," in the Mendelian sense, of these cases of fundamental structural asymmetry we know practically nothing. Indeed it is not at all clear at present how far the Mendelian principles of segregation, dominance, etc., apply to the normal ground-plan of organization. We shall discuss this phase of the subject later.

But besides these cases of undoubtedly heritable asymmetry of structure, which is common to an entire species, genus, or even phylum, we have a vast array of individual variations in symmetry, whose heritability, in the great majority of cases, is entirely unproved. Most of these are comprised in two main classes: (1) The appearance of asymmetry in structures which are normally symmetrical. Here belong the loss of one of a pair of organs, the acquirement of a supernumerary member on one side only, and the unequal development of paired structures which normally exhibit a fairly exact balance. (2) Reversal of symmetry, in normally asymmetrical structures or entire organisms. Here belong sinistral specimens in an ordinarily dextral species, or vice versa.
In investigating the inheritance of such cases of asymmetry, three possible alternatives should be borne in mind. (1) We might have the specific transmission either of the same unilateral character on the same side, or at least of corresponding inequalities in size in the case of paired structures. (2) We might find the asymmetry or lack of balance alone to be inherited, without there being any agreement between parents and offspring as to which side was better developed. (3) The structure or abnormality in question might itself be inherited, even as to varying grades in size or weight, yet there might be no inheritance in respect to its relations to the axis of symmetry.

After a considerable search for previously recorded data and a careful study of our own, it seems evident to us that each of these three modes of inheritance is realized in different cases, and that no general law can be formulated to cover all characters of all organisms. In certain cases, there seems to be an undoubted specificity in the transmission both of the character and of its position as left or right; in certain others, the mere asymmetry or lack of balance is transmitted, without reference to the side of the body; in others yet, the structure or structures themselves may be transmitted, though departures from symmetry either in size or position seem to be strictly non-hereditary. We shall proceed to give a few illustrations of each of these conditions.

Specificity of transmission in respect to the side affected

The available data are derived mainly from incidental mention of such cases by various writers. It is surprising that so few geneticists have recognized a distinct problem here.

DARWIN (1868, p. 456), after remarking that "it might have been anticipated that deviations from the law of symmetry would not have been inherited," cites several cases, on the authority of other observers, of the inheritance of one-sided abnormalities. In two of these instances, it is stated that the abnormality recurred upon the same side of the body. One relates to the appearance of stags in a certain forest in Germany, of which "many were observed with only one horn on the right side of the head." But this is an old record, dating back to 1788. The other case cited is that of the cow which "lost a horn by suppuration, and she produced three calves which had on the same side of the head, instead of a horn, a small bony lump attached merely to the skin." WEISMANN (1891, p. 82) (following HENSEN) later remarked of this case that the loss of the horn in the mother was probably due to some local weakness of a congenital nature. He offers the same explanation (here following RICHTER) of another case.
cited by him (loc. cit., p. 451), of a father "who lost his left eye by inflamma-
tion fifteen years before he was married, and who had two sons with
left eyes malformed (microphthalmic)." The point of present interest
is the statement that the defect in each of these cases was on the same
side among all the related individuals.

WEISMANN (1893, p. 55) also states: "I know of a family in which a de-
pression of the size of a pin's head in the skin in front of the left ear has
been transmitted through three generations."

A similar and much more recent case is that reported by SCHOFIELD
(1917, p. 517), of "a family in which a bilobed ear has been transmitted
through four generations,1 and in which "only the right ear shows the
characteristic in question. . . . Its appearance is not constant,
for it may skip one or even two generations and then be present." Further-
more, there is no indication of sex-linkage. SCHOFIELD's chart represents
eight affected individuals out of 22, all of whom he has seen "except the
female shown in the first generation." The chance that this abnormality
should occur upon the same side in eight cases out of eight is only one in
128 (i.e., $2 \times \frac{1}{2}$), according to the laws of probability. It is likely,
therefore, that we have to do here with a mode of transmission which
insures a specificity in the position as well as in the nature of the defect.
The chances of coincidence are much greater in the cases cited by DARWIN
and WEISMAN, inasmuch as the numbers of related individuals are smaller.

An interesting case, which unfortunately has not been as carefully studied
as we might wish, is furnished us by a prominent business and professional
man, X, who takes considerable interest in larger biological problems,
particularly those bearing on heredity. This man has a left eye which
shows a decided outward squint, and which is defective in visual power,
though not wholly blind.2 The defect has appeared, with varying degrees
of intensity in a considerable number of X's relatives, and so far as he
knows, is always on the left side. X's maternal grandmother and the
latter's father both had "the eye," though it is not definitely known which
one was affected. In X's mother the left eye was defective to such an
extent that she did not use it. Three of his children and at least two of
his grandchildren have likewise shown muscular defects of a similar nature,
which may or may not be accompanied by somewhat impaired vision. In

1 This abnormality seems to be of a nature closely similar to one described and figured by
WEISMANN (1891, pp. 453-455), and discussed by him in connection with the alleged inheritance
of mutilations.

2 A prominent oculist, who has examined X, writes that his "left eye is amblyomyopic as the
result of a high astigmatic error of fraction."
all cases, it is the left eye which is said to be affected, though the direction of the "squint" does not agree in all cases. The same defect has occurred in some of X’s more distant relatives, and the character was said by his mother to have appeared so frequently that it was called the "B— eye" after his mother’s family. Unfortunately, however, the evidence is not sufficient as to the identity of this defect, in its nature and incidence, among all the various members of this family. We, ourselves, have not been able to make any personal observations in the matter. But even as it stands, the case seems sufficiently striking to be recorded, and it gives evidence of a specificity of transmission, in relation to the side affected.

Dr. C. B. Bridges has kindly consented to our including here an account of a curious asymmetrical mutant which appeared in his Drosophila cultures several years ago, and of which he has not yet published a description. This mutant, “rotated abdomen,” writes Dr. Bridges, appeared among the offspring of “a star pale dichaete male” which was “crossed to a female showing none of these characters, but taken from a stock related to the pale. Rather less than a quarter of the offspring showed a new mutant character,—the abdomens were twisted at the end. This mutation was in all cases to the left, and was through from 60 to 90 degrees, with no apparent overlap between the rotate and the normal conditions. It was equally frequent among females and males. Neither of the parents had shown this character, which was evidently a simple recessive.” Linkage relations showed that “the gene for rotate . . . is in the third chromosome. . . . The viability of the rotates was not good, as evidenced by their low percentage. . . . Both sexes proved sterile, probably because of difficulty in copulation. . . . The race was continued for some generations, by breeding together heterozygous brothers and sisters. . . . Aside from the rotation no structural changes were observed.” (Bridges, MS.).

Equally definite evidence of a specificity of transmission in relation to the right and left sides of the body is to be drawn from various cases of reversed asymmetry. In exceptional individuals among normally asymmetrical species we meet with a lateral inversion, which may affect either the entire body structure, or certain organs only. The case of left-handedness in man is a familiar illustration of this phenomenon. This condition has long been known to be hereditary, in the sense that left-handed parents are far more likely to have left-handed offspring than are normal, right-handed parents. Jordan (1911, 1914) has compiled a large number of family histories of left-handed persons, and the tendency of this trait to “run in families” seems beyond question. On the whole, too, the trait ap-
pears to behave as an alternative character in inheritance, though ambidexterity and the lower grades of right- and left-handedness may be regarded as intermediate conditions. Far less certain is the possibility of classifying left-handedness as a dominant or recessive character. In many pedigrees it seems to behave as a simple recessive, though in at least three pedigrees the children of two left-handed parents were in part right-handed. In some cases the pedigree seems more intelligible on the assumption that left-handedness is dominant. "A strict Mendelian interpretation of this group of charts," says Jordan (1914, p. 68), "involves the further assumptions of degrees of bias to sinisterity and variation in relative hereditary prepotency."

It is interesting that this lateral inversion of the functions of the forelimbs (and apparently also of the brain) seems to be quite independent of that more radical transposition of internal organs known as "situs inversus." The latter condition (Bardeleben 1909) occurs only about once in 10,000 individuals, while left-handedness occurs in two to four percent of individuals. Again, the reversed position of the organs does not appear to be more common in left-handed persons.

Concerning the inheritance of dextrality and sinistrality in mollusks, the evidence is somewhat contradictory. That the conditions normal to entire species or genera are transmitted, in the same sense that other bodily characters are transmitted, can hardly be questioned. But the case of exceptional reversed individuals of a species is somewhat different. Crampton (1916), in his study of Tahiti land-snails of the genus Partula, examined many broods of young taken from the brood pouches of the females. Several subspecies, foremost among these being P. otaheitana sinistrorsa, and P. otaheitana sinistralis, differ from most of the other varieties of their species in being predominantly sinistral. There is, however, in each case, a considerable minority of dextral specimens. Crampton's tables make it plain that there is a strong correlation between mothers and offspring in respect to the dextrality and sinistrality of the shell. Despite the relatively small proportion of dextral individuals in the general population, an overwhelming majority of their offspring were likewise dextral. The correlation here shown is between offspring and female parents only, since the condition of the father was in no case known. Either inheritance, in respect to this character, is in some way strongly "matroclinous" or there must be a high degree of assortative mating.

It must here be added, however, that Lang "had no success in raising

*Possibly cytoplasmic; see below.
sinistral snails by inbreeding two sinistral individuals for two generations” (Przibram 1911); but Lang was dealing with a very different species.

The inheritance of asymmetry per se, with no agreement between corresponding sides of the body in related individuals

Some species of mollusks are indifferently right- or left-handed, within a given species, and Lang’s experiments, just cited, make it probable that in such cases the direction of growth is not inherited. Likewise certain species of flatfishes (Jordan and Evermann 1896, pp. 2623, 2625) are indifferently right- and left-handed, and in certain fiddler crabs (Yerkes 1901) the large cheliped of the male may occur indifferently on the right or left side. Concerning the last two cases we have no genetic evidence, but we are perhaps justified in asserting the probability that in these cases also the condition of asymmetry is inherited, but that dextrality or sinistrality is not.

In some of the teleost fishes, the relative positions (dorsal or ventral) of the right and left optic nerves in the optic chiasma are quite inconstant, and Larrabee (1906) has shown that there is no tendency for these relations to be inherited.

Newman (1917) offers evidence, for the armadillo, that unsymmetrically placed abnormalities in the scutes may be inherited, either in the same specific position as in the parent, or in a position corresponding to a mirror image of the latter. He likewise describes some less simple cases of inherited asymmetry of position, for which he devises special interpretations, and finally there are many cases in which the only thing inherited seems to be the abnormality itself, without regard to its position, whether dextral or sinistral, symmetrical or unsymmetrical.

Wilder (1904, p. 452) who studied the resemblances in finger-print pattern between duplicate twins found in some cases that there was a simple agreement in the pattern, without reversal; in other cases, a “mirror-image” relation between corresponding fingers of the two twins, while in still others there was no close correspondence.

In the experiments of Przibram (1907), with cats having right and left eyes of dissimilar color, the offspring, in some cases, showed the same condition as the affected parent, in one case, at least, the relation of the eyes was reversed, in still others, the offspring of affected parents had both eyes colored alike, this color corresponding to one of the parental colors. Various other abnormalities seem to be inherited indifferently on either or both sides, and all such cases perhaps belong more appropriately in the next subdivision.
The non-inheritance of asymmetry in the case of a single or paired structure or abnormality, the latter in itself being inherited

Illustrations of this class of phenomena could probably be multiplied indefinitely. From an admittedly very incomplete survey of the literature, we gather the impression that such human abnormalities as harelip, cleft palate, polydactyly, congenital cataract, etc., while in a high degree hereditary, have no tendency in related individuals, to have a corresponding position in relation to the main axis of symmetry. The defects may be either symmetrical or unsymmetrical, and if the latter, they may occur indifferently on one side of the body or the other. In fact, the various observers have been so little impressed by any such tendency toward agreement in position that they frequently fail to state which side of the body is affected (see Pearson et al. 1912).

Castle (1906), starting with a single male guinea-pig having a small supernumerary digit on the left hind foot, built up, by selection, a strain having four toes on each hind foot. Although, from the first, a slight majority of the animals were sinistral in this regard,¹ there was no specificity in transmission, and an endeavor to increase the sinistrality through selection was unsuccessful. Castle and others have likewise found that the position of the asymmetrical color patches in the hair-coat of the guinea-pig could not be fixed by heredity, though the spotted condition itself, as well as the colors, were inherited (Wright 1917).

From the data offered by Sollas (1909) we may conclude that there is no obvious inheritance of asymmetry in the case of supernumerary mammae in guinea-pigs.

Another case of non-inheritable disturbance of symmetry, regarding which we are able to offer rather extensive data, is derived from our own studies of the skeleton of the deer-mouse, Peromyscus. We offer no apologies for giving a rather detailed account of this case, despite the fact that the results are so completely negative. Indeed we introduce it at this point in our discussion just because it is so conclusively and satisfactorily negative. We regard it as not altogether fortunate that so many writers on genetics have an aversion to offering negative results, or to dealing with material which is likely to yield negative results. As we shall point out in a later section, the entire significance of certain of our findings depends upon the fact that the characters under consideration are demonstrably non-hereditary.

The case to be considered at this point is that of certain abnormalities

¹ See also Bond (1920) in regard to supernumerary digits in hybrid fowls.
of the sacrum. It has long been known that in man and certain other vertebrates (Bateson 1894, p. 110; Bumpus 1897) the attachments of the two sides of the pelvis may be unsymmetrical, the ilium on one side being fastened to the vertebral column, one vertebra in advance of that on the opposite side. Related to this (perhaps its direct consequence) is the fact that the vertebrae concerned are likewise unsymmetrical. The anterior of these has, on one side, the characteristically modified transverse process of a sacral vertebra, on the other the type of process proper to the lumbar series (figure 1).

An examination of the skeletons of 2288 specimens of *Peromyscus maniculatus*, of several subspecies, brought to light 99 specimens showing an asymmetrical sacrum. The normal sacrum of this species consists of four vertebrae, though five are met with in rare cases. The main attachment of the ilium, on each side, is to the anterior one of these vertebrae though in a large proportion of cases there is a lesser attachment to the second one of the series. Various grades of asymmetry exist. In the most marked cases, the first sacral vertebra furnishes the sole support for the ilium of one side, while the second furnishes the sole support for the ilium of the opposite side (fig. 1, B and C). In less extreme cases, each ilium may be attached to both of these vertebrae, though the chief attachment on one side is to the first, that on the opposite side to the second. This likewise results in a more or less marked asymmetry of the two vertebrae concerned. In 52 of the cases, the anterior attachment was greater on the right side,
being smaller or lacking on the left; in 47 cases the situation was reversed. It is curious that very few border-line cases were found. With two or three exceptions, the sacrum was either almost perfectly symmetrical or quite obviously unsymmetrical. Thus the abnormality was of a sort which one might reasonably have believed to arise through mutation, and which we should expect to be inherited in alternative fashion. Our evidence seems to be conclusive that it is not inherited at all.

In addition to the 99 cases of asymmetry above mentioned, 6 cases were found in which the sacrum was perfectly symmetrical, though the specimens were abnormal in having the chief attachment, on both sides, to the second vertebra, with only a minor attachment to the first (figure 1, D). In the unsymmetrical specimens the chief attachment of the ilium may be said to have been shifted back on one side, while in these last-named specimens the attachments have been shifted back on both sides.

When pedigree charts were plotted, it seemed probable at the outset that the abnormal individuals tended in a high degree to be related to one another. In one case, indeed, four affected specimens occurred among the offspring of a single father and two mothers (all three parents being sibs). But a more thorough study of the data showed quite conclusively that the occurrence of the abnormality was fortuitous, and that the appearance of the charts as originally plotted was illusory. When all members of a family, normal as well as abnormal, were included, this appearance of concentration in particular descent lines vanished.

A brief synopsis of these pedigree studies is worth while. Let us first consider the proportion of abnormal individuals among the offspring and more remote descendants of abnormal ones. Of the 105 abnormal individuals (99 + 6) only 13 were parents of specimens whose skeletons were preserved. These 13 had 49 offspring, of which only 1 (= 2 percent) was abnormal. The percentage of abnormal ones in the general population was 4.6 percent.

Considering more remote descendants, as well as offspring, 15* abnormal ancestors had 136 descendants, of which 4 (= 2.9 percent) were abnormal. Here again, the proportion of abnormal individuals was smaller among the descendants of abnormal individuals than it was in the population at large.

Unfortunately, there was no case in which two abnormal individuals were mated, or at least had offspring.

Again, let us consider the proportion of abnormal individuals among

*Of course all the “ancestors” are also to be ranked as “parents.” The difference in number is due to the fact that the skeletons of certain members of intermediate generations were not preserved.
the parents and ancestors of abnormals. The total number of parents and ancestors was 161 (eliminating duplications), of which 5 ( = 3.1 percent) were abnormal. We once more meet with the unexpected result that the proportion of abnormal individuals was smaller among the ancestors of abnormals than among the general population!

Lastly, we present figures showing the proportion of abnormality among the sibs of abnormal individuals. Sixty-seven abnormal individuals had 880 sibs (including half-brothers and half-sisters)\* of which 52 ( = 5.9 percent) were abnormal.

In this last instance, the percentage is slightly higher than among the population at large. We do not believe, however, that either here or in the cases previously cited, the slight differences among these percentages have any significance. On the contrary, the figures as a whole, seem to furnish definite proof that the character in question is strictly non-hereditary.

Another question which was tested in connection with these sibs, was whether related individuals having the abnormal sacrum tended to agree in respect to the type of abnormality. Those cases in which the attachment of the pelvis to the first sacral vertebra was stronger on the right side were termed "right-handed," the opposite condition being termed "left-handed." As has just been stated, there were 52 cases in which an abnormal individual had an abnormal sib. In 20 of these cases, both related individuals were right-handed or both were left-handed; in 26 one was right-handed and the other left-handed; while in the remaining six, one was asymmetrical, while the other displayed the symmetrical type of abnormality shown in figure 1. Thus there were actually more cases of disagreement than of agreement.

As a corollary of the doctrine that somatic modifications are non-hereditary, it is frequently assumed that all non-hereditary characters are of somatic origin, and indeed the terms somatic and non-hereditary have come to be used interchangeably. If, by calling a character "somatic," we mean that it has arisen during ontogeny through some specific influence of the physical environment, it seems probable that the term is inapplicable in the present case. It is not at all obvious how any feature of the environment (even of the intra-uterine environment) could exercise such a localized influence on the developing fetus as to cause unilateral displacements in the attachment of the pelvis. It is more likely, perhaps, that this condition has arisen through one of those chromosomal mishaps which are commonly held to be responsible for the phenomena known as "bud

\* Duplications are not eliminated in this case.
variation" and "somatic segregation." That such an unstable condition of the chromosomes, in these cases, arises de novo during ontogeny has not, so far as we know, been proven. May there not be characters of strictly germinal origin which are, none the less, not transmitted to further generations?

Certain further examples of non-hereditary asymmetry in hereditary structures will form the chief subject-matter of the second half of this paper. These are slight deviations from strict bilateral symmetry occurring among certain paired bones of the body. As already stated, the significance of certain of these results for genetics depends upon the very fact that the characters in question are not hereditary.

Discussion

We have thus far found that departures from symmetry in normally symmetrical parts, as well as reversals of asymmetry in normally unsymmetrical parts, may be inherited, in the same sense that any other structure or character of an organism is inherited. We have also found that quite similar conditions may be strictly non-hereditary. Various theoretical questions, of course, crowd themselves upon us. Why, for example, should there be such a disagreement in the hereditary behavior of these characters in different cases? And how is it possible, according to current ideas of the mechanism of heredity, that there should be any specific transmission of right- and left-handed characters as such? A satisfactory solution of such questions is probably impossible in the present state of our knowledge, but certain reflections may not be unprofitable at this point.

In the first place, to repeat certain former statements, there seem to be at least three more or less independent factors concerned in the hereditary transmission of bilateral asymmetry. (The word factor is not here used in the Mendelian sense.) One of these is the inheritance of a character, structure, or abnormality, single or paired, regardless of symmetry. A second is the inheritance of a tendency toward balance or lack of balance, regardless of whether related individuals agree in respect to the side of the body (right or left) on which the character in question is better developed. A third is the specific transmission of dextrality or sinistrality, as affecting a particular part or the general organization of the body.

Needless to say, neither of the last two processes can occur without the first,—there can be no transmission of asymmetry, unless the asymmetry is inherent in some part or parts which are themselves transmitted. On the other hand, we have found that disturbances in the symmetry of hereditary parts are not always hereditary.
Again, it would seem that, in some cases, at least, the third process is something superadded to the second. Within a single group, e.g., the gastropod mollusks, we have a hereditary tendency toward asymmetry, which is commonly specific,—related individuals being constantly dextral or sinistral,—but which may be quite neutral, related individuals being indifferently dextral or sinistral. And similarly, in certain unsymmetrical crustaceans, the larger cheliped of the male is normally on the same side of the body in some species, indifferently on the right or left in others.

The asymmetries of a gastropod are due primarily to a more rapid growth of one side of the body, just as the various contortions in the alimentary canal of a higher vertebrate arise, during development, from the fact that the originally straight inner tube grows faster than the inclosing body. Teleologically speaking, it is necessary that the digestive system should be asymmetrical in order that it should be sufficiently long. And a similar statement would probably apply to the molluscan body. The question is why the excess of growth should occur predominantly on the same side in related individuals.

Crampton (1894) was the first to show that in the development of a sinistral gastropod *Physa heterostropha* there occurs a reversal in the direction of the oblique cell-divisions which give rise to the 4-cell and later cleavage states. This principle may be one of very general applicability. According to Conklin (1916, p. 177):

"In these sinistral snails, and probably in all animals showing inverse symmetry, the embryo is inversely symmetrical and every cleavage of the egg from the first to the last is the inverse of that which occurs in dextral snails. There is a good reason to believe that in such cases the unsegmented egg is also inversely symmetrical as compared with the more usual type. In all of these cases there is a direct correspondence between the polarity and symmetry of the oösperm and the polarity and symmetry of the developed animal."

Some single, very slight change in the spatial relation of parts in the egg may thus be responsible for the entire reversal of symmetry in the adult.

Przibram (1911) found that in certain unequal-limbed ("heterоchelate") crustacea, which normally have a heavier and differently shaped pincer on the right side, removal of the right pincer would cause the weaker, left-handed one to develop the structure proper to the right side, without, however, losing the left-hand type of symmetry. Again, in the lobster, supernumerary pincers may appear, after injury to a chela, on either side of the body, taking the form of lateral excrescences upon the limb concerned. These accessory pincers frequently have a reversed symmetry, i.e., a symmetry proper to the limb of the opposite side. Yet they never-
theless retain the same type of structure ("crusher" or "nipper") as the
main pincer of the appendage to which they are attached. Przibram
concludes, very reasonably, that these cases of inverted symmetry cannot
be "due to the presence there of latent determinants of the opposite body-
side." But he goes much further than this and draws "the conclusion
that there are no distinct determinants for right and left body-sides; but
that the bilaterality depends entirely upon the distribution of anlagen in
opposite directions with reference to the dorso-ventral and antero-posterior
axes." He admits that such a conclusion seems inconsistent with the
existence of asymmetries which "are manifested in the structural plan of
the entire organism" and which "are transmitted unchanged from one
generation to the other." He sets aside such objections as inconclusive,
however, "because, even though rarely, we do nevertheless find a reversal
of the fundamental asymmetry," and because the two instances of such
reversal which he cites (situs inversus in man and sinistrality in Helix) happen
to be, so far as he knows, non-hereditary. Przibram concludes with the
admission that we know nothing "of the causes of these asymmetries and
why in certain species one side of the body is particularly predisposed to
modification."

We are not arguing for the existence of "distinct determinants for right
and left body-sides," but we do regard the evidence which we have pre-
sented as showing conclusively that there may be, and frequently is, a
specific transmission of deviations from bilateral symmetry. To say
that the normal dextrality of a crab’s claws is not inherited because we
can reverse this relation by removing one of them is like saying that the
eye of the same animal is not inherited because its removal under certain
conditions will result in its replacement by an antenna.

If all unilateral, or otherwise asymmetrical, inheritable structures were
correlated in such a way that any reversal of symmetry affected all of these
parts alike, the mechanism of their transmission would be more readily
conceivable. Some slight variation in the relative position of substances
in the egg might suffice to account for the initial bias which determined
whether the organization as a whole should become dextral or sinistral.
But no such constant relation seems to exist. Right- and left-handedness,
in man, seem to bear no relation to situs inversus, and while the latter
condition may not be inherited (of this we have no knowledge) the normal
position of the viscera certainly is hereditary, in the same sense that any
other major feature of our organization is hereditary.

Weismann (1893), after mentioning the inherited unilateral peculiarity
referred to above (page 448) expresses his belief that "we are logically com-
pelled to assume a particular element of the germ-plasm for each peculiarity of this sort, not because heredity may be manifested in details so minute, but because the transmission of such details may be independent." The modern chromosome school of geneticists regards every such independently inherited character as conditioned primarily by a single genetic factor or "gene" having a particular "locus" in one or another chromosome. These two views, which, for present purposes do not differ very widely, have one common weakness, namely, that they fail to render any account of the spatial relations of the fully developed parts. According to the investigations of Morgan and his school, there is a very precise spatial arrangement within each of the chromosomes, of the genes which condition the various inheritable characters of the adult. But there seems to be no possible relation between the spatial arrangement of these genes and the arrangement of the adult parts. Genes which affect the development of widely separated parts of the body may occur alongside of one another, while genes which produce closely similar modifications of the same organ may be parcelled out among different chromosomes. If there is any bilaterality, either of chromosomes or genes, it certainly bears no relation to the bilaterality of the adult organism.

Such a mechanism, while accounting admirably for the segregation, linkage, etc., of the various Mendelian factors with which these authors are chiefly concerned, seems ill-adapted to account for spatial arrangement among the inherited parts. Various writers have seen in the cytoplasm a more fit medium for carrying such fundamental characters as symmetry, polarity, and the general ground-plan of organization. Since the cytoplasm available for the purposes of heredity is nearly or quite restricted to the egg, it has been contended that the ground-plan of organization is of purely maternal origin. Says Conklin (1916, p. 184):

"We are vertebrates because our mothers were vertebrates and produced eggs of the vertebrate pattern; but the color of our skin and hair and eyes, our sex, stature, and mental peculiarities were determined by the sperm as well as by the egg from which we came. There is evidence that the chromosomes of the egg and sperm are the seat of the differential factors or determiners for Mendelian characters, while the general polarity, symmetry and pattern of the embryo are derived from the cytoplasm of the egg."

It is true that Conklin (1917) later receded somewhat from this position, conceding that while "there is cytoplasmic inheritance through the female only . . . . . . . these cytoplasmic characters are themselves of biparental origin." This, he added "is Mendelian inheritance though somewhat com-

* Of course they are fundamentally different in some other respects.
licated by the fact that every ontogeny has its beginnings in the preceding generation." Still later, however, CONKLIN (1920) has definitely committed himself to a non-Mendelian origin of "a few orienting differentiations such as polarity and symmetry." If the genes or Mendelian factors of the nucleus were the only differential factors of development, as is sometimes assumed, "these genes would of necessity have to undergo differential division and distribution to the cleavage cells; since this is not true, it must be that some of the differential factors of development lie outside of the nucleus and if they are inherited, as most of these early orientations are, they must lie in the cytoplasm." (p. 403).

LOEB (1916, pp. 246, 247) takes a similar and yet more radical position. He offers evidence in support of "the view, according to which the determiners in the chromosomes only tend to give special characters to the embryo or to the adult while the cytoplasm of the egg may be considered the real embryo." Expressed in other words, this author regards as most tenable "the idea that the Mendelian factors of heredity must have the embryo to work on and that the organism is not to be considered a mere mosaic of Mendelian factors."

In apparent contradiction to the viewpoint just outlined is the fact that certain of the disturbances of symmetry discussed in the preceding pages are known to be transmitted by the male parent as well as by the female. This is true of left-handedness in man,8 of the ear defect described by SCHOFIELD, and the eye defect of the "X" family. It is also true of the mutant "rotated abdomen" discovered by BRIDGES. In the latter case, indeed, the character is definitely known to be a Mendelian recessive, and its linkage relations have been partially determined. As regards its mode of transmission, therefore, this character seems to be quite comparable with many other mutant characters which have been described for Drosophila.

How then can we reconcile the facts referred to in the last paragraph with the viewpoint set forth in the immediately preceding ones? Only, it would seem, by bearing in mind the distinction, already insisted upon, between the inheritance of a structure or abnormality itself and the inheritance of its position in relation to the axis of symmetry. The former may be supposed to be transmitted alike by the male and female parents, the latter by the female alone. This statement may appear paradoxical in more than one respect, but our meaning will become clearer, we trust, in the ensuing discussion.

8 In a number of the family histories given by JORDAN, 1914, left-handedness appears to have been transmitted through the male alone for three and even four generations.
It is a well-known fact that the ovarian eggs of certain animals already show a marked bilateral symmetry, and that this symmetry corresponds to the bilateral symmetry of the adult organism. It is well within the range of possibility that the symmetry of the unfertilized egg, even when this is unrecognizable, dates back to the earlier stages of oogenesis, and indeed that there is just as much a “continuity of symmetry” for the egg cytoplasm as there is a “continuity of germ-plasm,” for the constituents of the chromatin of both sexes.

It may also be assumed that this bilateral symmetry of the egg implies a correlative asymmetry, in the case of organisms displaying hereditary asymmetries in their adult structure. What else, for example, can account for the reversal in the direction of cleavage in the case of sinistral gastropods? And does it seem improbable that more specific and definitely localized departures from bilateral symmetry may be accounted for in the same way? May there not be, even in the egg, qualitative differentiations of the future right and left halves of the body, similar in nature, though less in degree, to those obvious differentiations of the cytoplasm which foreshadow the anterior and posterior, dorsal and ventral, regions of the body in the eggs of certain animals?

Thus far, such an explanation would seem to leave out of account the possibility of inheritance through the male parent. But we think that a conceivable mechanism for such inheritance can be discerned here. The sperm nucleus, and for that matter the egg nucleus, is believed to play its part in ontogeny through its influence upon the cytoplasm. Suppose the case of a unilateral abnormality, say an eye defect, which is inherited in a specific manner, and through the male as well as the female descent lines. This means that something in the sperm nucleus—be it “determinant,” “determiner,” “factor,” or “gene”—is so constituted as to impair the normal growth of the tissues of the human eye in a certain specific manner. But it does this only in collaboration with certain materials of the egg, including those of the egg cytoplasm. If we grant qualitative differentiations in the cytoplasm of the two bilateral halves of the egg, is it not conceivable that a “gene” of a certain constitution from one sperm-nucleus would combine more readily with components of the cytoplasm of the right side of the egg, whereas a similar “gene” from another sperm-nucleus (a “stereo-isomer,” perhaps, of the first) would combine more readily with the cytoplasm of the left side? In this way, the male parent could

*Such a suggestion does not necessarily imply any thorough-going preformation view, i.e., the assumption of a part-for-part correspondence between elements of the fertilized egg and elements of the adult structure.
determine a condition of asymmetry in the offspring, without this condition being present, in any sense, in the paternal chromosomes.\(^\text{10}\)

It is not, of course, necessary to assume that such a combination takes place in the undivided egg. Since division products of the original sperm-nucleus are contained in every cell of the adult organism, it is quite possible that these effects could take place at any stage of ontogeny. It has been contended by some writers that the left eye, along with some other parts, is normally inferior in right-handed persons (Jordan 1911, pp. 20, 21); and such a difference might be due to an inferiority in the fetal blood-supply on that side. It is thus plain that the left-handed incidence of an inherited eye defect does not necessarily imply any specific pre-localization of eye-determining parts in the cytoplasm of the egg. Only a pedigree combining left-handedness and some inherited unilateral eye-defect could furnish critical evidence on this point. We are badly in need of evidence upon the simultaneous inheritance of independent unilateral variations.

The authors are not in a position to elaborate this hypothesis in detail; nor do we pretend that it is a wholly original one. But it harmonizes, we believe, with various speculations which have been made in the fields of stereochemistry, immuno-chemistry, etc.\(^\text{11}\)

Those cases in which the heritable abnormality in question consists in nothing else than a departure from a normal condition of symmetry (e.g., Bridges's flies), or in the reversal of a normal condition of asymmetry (e.g., left-handedness), could be explained in a quite similar manner. Some hereditary factor from the left-handed male might be supposed to have the effect of stimulating (or retarding) the growth of tissues having a certain chemical constitution, irrespective of their position in the body. Owing to the preexisting differentiation of the egg cytoplasm into qualitatively unlike right and left halves, this paternal factor might be supposed to favor the growth of the left side (or the right, if the brain is to be regarded as taking the initiative in the process). Such a differential result would be due to the selective affinity of the chemical substances constituting this factor for substances in the cytoplasm of one or the other side of the body. In the case of a right-handed parent, the affinities involved would be of a reverse nature.

\(^{10}\) Conklin (1920, p. 403) asks the question: "How can identical factors give rise to different products in different cells," and answers: "This is evidently due to the fact that while the division of chromosomes is non-differential, that of the cell body is often differential and the same chromosomes and genes acting upon different kinds of cytoplasm will produce different results."

\(^{11}\) This appears to be the type of explanation which Dr. Bridges would apply to the inheritance of "rotated abdomen," if the senior author correctly understands the views expressed during a conversation with him.
We may suppose the existence of eggs in which the materials responsible for more rapid growth passed indifferently to the right or the left side of the body, at (or before) the time of the first cleavage. This would result in an inherited asymmetry, indifferent as to whether dextral or sinistral. Finally, the cytoplasmic substances with which the paternal "gene" combined might be disposed without any relation to the main axis of symmetry. The last two conditions would give rise to the second and third alternatives in respect to transmission which have been discussed above (page 447).

NON-INHERITED DEVIATIONS FROM SYMMETRY AND THE THEORY OF MULTIPLE FACTORS

The "multiple-factor" hypothesis of the inheritance of quantitative differences which display no obvious tendency to segregate is now too well known to need any explanation in this paper. Those readers who are not familiar with this highly interesting outgrowth of Mendelism are referred to the excellent discussions of Castle (1916, chapter XXI), and of Babcock and Clausen (1918, chapter X).

As is well known, the chief evidence for the theory that quantitative genetic differences depend upon the cumulative action of independently segregating Mendelian factors consists in the fact that the second and subsequent generations of hybrids frequently show increased variability over the first hybrid generation, when races presenting such differences are crossed. This hypothesis is logical and consistent, and, formally at least, it saves the theory that all inheritance belongs to the Mendelian category. One element of its strength is the surprising facility with which its exponents are able to meet classes of facts which, on first glance, seem quite irreconcilable.

Thus, various authors have found that the range of variation in the second and subsequent hybrid generations may exceed that of the parent races themselves. Such results would, on the face of things, seem to prove too much. But an answer lies ready at hand. Punnett and Bailey (1914), for example, found that in crossing races of fowls of widely different size the later hybrid generations included some individuals which were larger and smaller, respectively, than either of the parent extremes. These authors conclude that the larger race had the constitution $AABBCCdd$, in a previous paper (Sumner 1920) the senior author presented data showing that in subspecific crosses in Peromyscus there was a slightly preponderating tendency toward an increase of variability in the $F_2$ generation. Further data, not yet published, emphasize the reality of this increased variability, and show, indeed, that it holds for some characters in which the parent races do not differ from one another.
the smaller being \( aabbccDD \). As a consequence of the independent segregation of these size factors, individuals having the constitution \( AABBCCDD \) and \( aabbccdd \) were obtained.

Other cases have been described in which there was an increase in variability in the \( F_2 \) generation, in respect to characters in which the parent races did not differ at all. When the senior author first encountered cases of this sort in his own material, he devised the following interpretation as being possible, though he believed far from probable.\(^3\) The two parental races, though not differing \textit{phenotypically}, in respect to the character in question, none the less might be supposed to differ \textit{genotypically}. The quantitative value of the character in the two races might be supposed to be due to different factor combinations which merely chanced to bring about the same resultant condition. Thus, to take an imaginary example, the value 20, which is common to the two races, might be due, in one case, to the cumulative action of two pairs of factors, having the potential values 14 and 6 respectively; while their allelomorphs in the other race might be supposed to have the values 8 and 12. Segregation and recombination would give us the values 14, 17, 20, 23, and 26, with 20 as the modal condition.\(^4\) This explanation was so obvious and plausible (albeit wholly unconvincing) that it gave us no surprise to learn that practically identical interpretations had been offered by other writers some time previously (see, for example, East and Jones 1919, pp. 174, 175.)

In the present paper we are setting forth data which would seem to be much more baffling to the radical Mendelian. We are presenting cases in which the \( F_2 \) variability shows a marked increase, \textit{in respect to characters which are not inherited at all}. We think, too, that the numbers are sufficient to establish this fact statistically, with reasonable certainty, and that other causes than hybridization are not likely to be involved. If this be true, the current explanation of the increased variability of second-generation hybrids, as being due wholly to gametic segregation of the Mendelian type, is, to say the least, weakened. We may have to fall back upon the unanalyzed fact that variability does increase in later generations of hybrids, a fact which has been recognized by breeders for over a century, and for which Mendelian segregation only partially accounts.

For a number of years the senior author has been accumulating osteological material derived from the wild races of one species of deer-mouse (\textit{Peromyscus maniculatus}) as well as from various pure and mixed strains

\(^3\) These results have not thus far been published.

\(^4\) This, of course, is on the assumption that the duplex condition of each factor calls forth twice as great an effect as the simplex.
used in the course of genetic experiments. The bones constituting this collection have been cleaned by trained preparators, according to a fairly constant method. The number of these partial skeletons now exceeds 2000, of which some 800 are derived from individuals so related that their bones can be used in studies of parent-offspring correlation. At first, the bones which were prepared comprized the skull (including the mandible), certain regions of the vertebral column, the left femur and left innominate bone. For these earlier series, therefore, the two halves of the mandible are the only paired structures available. Later, both femurs and both innominate bones were saved, expressly for use in connection with these studies of asymmetry. The numbers of the various bones which have been available for measurement differ in different cases and are stated below in the separate discussion of each.

The chief question which it was sought to answer by means of these rather laborious studies is this: Are the slight differences between the right and left members of a bilateral pair of bones inherited? The method employed was as follows: The two members of a pair were weighed or measured, as the case might be. The value obtained for the left side was divided by that obtained for the right side, and the resulting "sinistrodextral" ratio was expressed as a percentage. These percentages were then treated like any other values derived from the measurement of a series of organisms. The means, standard deviations and probable errors were computed, and finally the coefficients of parental-filial correlation were determined.

In computing these correlation coefficients, the values for parents and offspring were lined up in neighboring columns, the parental value, in each case, being taken as many times as the number of offspring of that particular parent. Thus the "n" of the formula is the number of offspring in a given series, not the number of parents. As is doubtless true of most other investigators who have computed parent-offspring correlation coefficients in this way, we are confronted with the question of the proper determination of the probable error of this coefficient. What shall be the "n" employed in this latter formula? Shall it be the number of parents, or the number of offspring, or some intermediate value? It seems to be commonly believed that the last of these alternatives is the correct one. Surprising as it may seem, we learn on the high authority of Professor Karl Pearson that no satisfactory method of determining the proper value of "n" has

13 Any slight inequalities of treatment due to the strength of the fluids, temperature, duration of maceration, etc., might conceivably lead to differences between two individuals, but they should not lead to differences between the right and left members of the same individual.

15 In a letter to the senior author.
been formulated for such cases. This being true, we have adopted, as a makeshift, the use of the arithmetical mean between the number of parents and number of offspring. This seems quite as legitimate as the use of the more elaborate formula proposed by MacDowell (1920, p. 428).

Studies of the mandible

The two halves of the mandible were carefully separated with a scalpel, in those specimens (a small minority) in which they had not already fallen apart in the process of maceration or of subsequent cleaning. It was found possible to effect a fairly clean fracture in practically all cases, and we are convinced that no appreciable part of the differences found are due to this possible source of error. Specimens which were damaged in any way—whether through the breakage or loss of a tooth, or the breakage of the bony substance—were rejected from the series, and each half mandible was carefully searched for such defects before being weighed. This led to the rejection of a considerable percentage of the entire collection.

In weighing, the procedure was as follows. The weight of the left half of the mandible was first determined to the nearest milligram. The two halves were then placed in opposite scale-pans, and the lighter side was compensated by the movement of the rider until an exact balance was established. This gave the difference between the two halves, which was recorded in tenths of a milligram. As a check on the first reading of this difference, the two halves were in every case transposed, and a second reading made, the mean of the two values being employed in the subsequent computations. The absolute difference (L—R) was then divided by the (approximate) weight of the left half of the jaw, the quotient being the percentage difference between the two sides. The latter was added (algebraically) to 100 percent, thus giving the sinistro-dextral ratio. A ratio of 100 denoted, of course, perfect equality. Figures lower than this denoted a preponderance of the right side, higher ones a preponderance of the left side. Fractional percentages were not used. Differences of less than half a percent were discarded, fractions greater than this being counted as one percent.

The mean ratio was 100.027 ± .027. In other words, the mean indicates a very slight and quite non-significant sinistrality. The purely accidental character of this departure from symmetry is further evidenced by the fact that of the four means included in figure 2, two indicate a condition of dextrality, two of sinistrality.

The limit of sensitiveness of the balance employed was actually about two-tenths of a milligram, but this was sufficiently accurate for present purposes.
Two main problems have been investigated in relation to these jaw weights, as well as to the other measurements of paired bones: (1) the heredity or non-heredity of the individual deviations from symmetry; (2) the variability of first- and second-generation racial hybrids in respect to this character.

(1) Are individual deviations from symmetry in the weight of the two halves of the mandible hereditary? Under this head would be included either the specific inheritance of dextrality or sinistrality in respect to jaw weight, or the inheritance of a mere lack of balance, without regard to which side preponderated in two related individuals.

Considering in order the two aspects of this question, coefficients of parental-filial correlation were first worked out in respect to the sinistro-dextral ratio. According to the mode of treatment just described, it is obvious that sinistral individuals would give plus deviations, dextral individuals giving minus ones. A positive correlation coefficient would imply a tendency toward correspondence, either as to dextrality or sinistrality.

To consider the father-offspring correlation first, the available material comprizes 87 fathers and 779 offspring. A small proportion of those which figure as "offspring" likewise figure as "fathers" since three successive generations are included in some cases. The coefficient resulting from this entire lot, treated collectively, is \(-0.072 \pm 0.032\). The relative size of the coefficient and its probable error would render it improbable that the former was significant, even though it had not been negative. Separate coefficients were computed for the 11 different groups of individuals into which the population may be divided, and for the sons and daughters separately, within each of these groups. The various groups differed among themselves in respect to either the subspecies (or hybrid form) comprised, or to differences in the history of the stock which need not be detailed here. Of these 22 single coefficients, 16 chanced to be negative and only 6 positive, the mean being \(-0.082\).

For the computation of the mother-offspring coefficient, 173 mothers and 766 offspring were available. The coefficient for this lot, taken collectively, was \(+0.057 \pm 0.031\). Here again, the relative size of the probable error would in itself render the coefficient of very doubtful significance. One's doubts are greatly strengthened when the fact is considered that the evidence of a positive correlation for the mothers is even less than that of a negative one for the fathers!

Considering, as above, the figures for minor groups, we have 15 positive coefficients, 6 negative ones and one of 0. The mean of these 22 is identical with that for the entire series, i.e., \(+0.057\).
Our experiences in the computation of these results compel us to call attention to the danger of attributing any considerable significance to mere coincidences of sign in a series of coefficients or of differences. There was more than one point, in the course of these studies, when the probability seemed to us high that we had significant positive coefficients of mother-offspring correlation, and negligible ones for the father-offspring relation. For example, the first 8 group-coefficients which were computed for the mothers were all positive, while the first 16 of these comprised 13 positive, two negative and one zero coefficient. These sixteen groups included, indeed, all the individuals that it was originally intended to deal with in these studies. The remaining 6 groups considerably reduced this preponderance of plus signs; and the significance of this preponderance is entirely destroyed by the fact that the father-offspring series shows an even greater preponderance of negative coefficients.

Finally, there is no evidence, either for the mothers or fathers, of a sex-linked transmission of this character, which might be obscured by the throwing together of sons and daughters in computing these general coefficients.

As regards the inheritance of dextrality and sinistrality in jaw weight, accordingly, the verdict for both sexes seems to be that there is none. At least, this seems to be true of those small deviations which form the basis of our coefficients. Whether single, heritable "mutations," large or small, ever occur in respect to this character, we do not know. We can only say that we have found no evidence of such in our tables.

The second question regarding inheritance relates to the possible transmission of asymmetry *per se*, regardless of sign. For this purpose, the departure of each individual ratio from 100 percent, *in either direction*, was set down; and these differences (or grades of asymmetry) were used as the variables, instead of the ratios themselves. Averages were computed for these differences, and the plus and minus deviations of the individual variants from their mean were entered, as in any other case of correlation.18

Thus proceeding, a father-offspring correlation of \(-0.026 \pm 0.032\) and a mother-offspring correlation of \(+0.007 \pm 0.031\) were obtained. There is, accordingly, as little tendency for parents and offspring to resemble one another in respect to the degree of asymmetry in their jaw-bones as there

18 To illustrate: Animals having ratios of either 99 or 101 would be credited with the difference 1, those having ratios of 98 or 102 with the difference 2, etc. The mean of these differences, was not far from 1 in each case. Accordingly, individuals having a 1 percent difference between right and left would have a deviation of 0, those with a 2 percent difference would have the deviation \(+1\), those with a zero difference (100 percent) would have the deviation \(-1\), etc.
is for them to resemble one another in respect to which side of this member is the heavier.

(2) Differences between the first and second generations of race-hybrids in respect to variability in bilateral asymmetry. The standard deviation for the 1202 sinistro-dextral ratios is 1.386 ± 0.19. The variability differs somewhat in the four major groups into which the material has been divided. Thus we have:

Wild (174 specimens) .......................................................... 1.44 ± 0.052
Pure races, cage-bred, first generation (438) .......................... 1.18 ± 0.027
Subspecies hybrids, F₁ (326) ................................................. 1.35 ± 0.036
Subspecies hybrids, F₂ (264) .................................................. 1.71 ± 0.058

The frequency distributions for the various lots are shown in figure 2.

Figure 2.—Histograms showing frequency distributions of the sinistro-dextral ratios of the two halves of the mandible. The abscissas are the ratios, the ordinates being the frequencies of each. The frequencies are also indicated by the numbers at the tops of the various columns. The number in parenthesis under each of the four histograms is the total number of individuals on which it is based.

The difference between the figure for the F₁ and F₂ generation (.36) is nearly six times its probable error (.062) and is therefore of highly prob-

19 Skeletons of cage-bred mice of pure race, later than the first generation, were not prepared.
able statistical significance. The histograms show that this difference of variability cannot be due wholly, or even mainly, to the influence of a few extreme variants. An obvious feature of the $F_2$ histogram is the decrease in the central column, as compared with the lateral ones.

But there are other differences among these four standard deviations which are of more than possible significance, notably that between the wild and the cage-bred mice, belonging to pure races (i.e., subspecies). The difference in this case is more than four times its probable error.

We shall postpone a discussion of these differences until the various other cases of asymmetry have been considered. It is well to state here, however, that this increase of variability in passing from the $F_1$ to the $F_2$ generation holds true for each of three different crosses when treated separately. The figures are as follows:

<table>
<thead>
<tr>
<th>Cross</th>
<th>$F_1$ (std.dev.)</th>
<th>$F_2$ (std.dev.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calistoga-Carlotta</td>
<td>$1.25 \pm .049$</td>
<td>$1.66 \pm .091$</td>
</tr>
<tr>
<td>Carlotta-Victorville</td>
<td>$1.31 \pm .065$</td>
<td>$1.63 \pm .076$</td>
</tr>
<tr>
<td>Eureka-Victorville</td>
<td>$1.54 \pm .080$</td>
<td>$1.84 \pm .096$</td>
</tr>
</tbody>
</table>

The probable errors are of course larger here, owing to the smaller numbers comprised in these separate groups. The differences are 4.0, 3.3 and 2.4 times their probable errors, respectively.

The femur

Femur measurements of 960 individuals are available for these studies. The measurements made were of two sorts, linear measurements and weights. These two have been treated as quite independent characters. It was early found that there was no constant correspondence in individual cases between preponderance in weight and preponderance in length. The longer bone of a pair was frequently the lighter and vice versa. In fact, the correlation between the sinistro-dextral ratios for femur length and femur weight was only $+ .285$. More surprising yet is the fact that when we consider the grand averages for the entire series, the femur is preponderantly dextral in respect to length and preponderantly sinistral in respect to weight. The figures are $99.882 \pm .027$ and $100.226 \pm .055$, respectively. As already stated, these figures are based upon nearly a thousand indi-

20 The omission of the only really extreme variant (89 percent) reduces the standard deviation to 1.58, which is still far removed from that of the $F_1$ group.

21 The names employed are those of towns near which the specimens were trapped. The Calistoga mice belong to the subspecies gambeli, the Carlotta and Eureka mice to the subspecies rubidus, while those from Victorville belong to the subspecies sonoriensis.
viduals. It would seem not only that the difference between these two figures is statistically significant, but that the departure of each from 100 percent is of very probable significance.

Various writers have reported significant differences in length between the right and left femurs in man and even in some other animals. But these results have not been wholly consistent, and as a rule have been based upon insufficient numbers. The most extensive biometric study of this bone is doubtless that of Pearson and Bell (1919). This is based upon some 900 separate human femurs (not paired), derived from disinterred remains. These writers find that "the differences in right and left members of none of the chief lengths . . . . appear to have any significance" (p. 130). Certain other characters, on the other hand, were found to display marked dextro-sinistral differentiations. For example, "the torsion . . . . is always markedly greater in the left than the right bone. Further the obliquity is most definitely and markedly greater in the left bone than in the right" (p. 132).

Donaldson and Conrow (1919, p. 304), conclude that "the post-natal asymmetries which appear to arise largely from use in man (Gaupp 1909) are not evident in the rat, and there is nothing systematic in the slight deviation which we have observed for the values of symmetrical bones."

Femur length

The measurement employed by us was the "maximum length from top of head to bottom of internal condyle" (Pearson and Bell 1919, p. 4). A dial-caliper was employed for this purpose, graduated to tenths of a millimeter. Each femur was measured twice, the two being measured in alternation. Differences less than 0.1 millimeter were not included. Differences of one-tenth or more could be relied upon as due, in the great majority of cases, to actual differences in the length of the femur. Instrumental errors and differences due to the presence of foreign matter rarely reached this magnitude. Roughly speaking, the femur length of these mice averages about 15 millimeters.

The very close approach to equality in the length of the right and left femurs of most specimens was a source of surprise. The great majority of differences, when such were recorded at all, did not exceed one-tenth of a millimeter. The histograms (figure 3) show the low variability in the sinistro-dextral ratios. Analysis of these histograms reveals the fact that nearly 50 percent of the cases are placed in the 100-percent columns. Since differences of one-half percent or less were counted as zero, this means that in nearly half of the specimens the right and left femurs did
Figure 3.—Showing sinistro-dextral ratios for femur length. (See legend for figure 2).
not differ from one another by more than half of one percent. Furthermore, all but 5 percent of the specimens belonged in the three central columns of these histograms; i.e., 95 percent of them showed a difference not exceeding 1.5 percent. On the other hand, there was a small minority of variants which departed rather widely from the condition of perfect symmetry.

As regards heredity, the same procedure was adopted for femur length as for jaw weight. The father-offspring coefficient, based upon the entire lot of 64 fathers and 732 offspring, is \(+.023 \pm .034\). Of the sixteen separate group-coefficients, 8 are positive, 7 negative and one 0. The mean of these is \(+.010\).

The mother-offspring coefficient gives one a first impression of furnishing proof of a genuine correlation. The figure for the entire series of 136 mothers and 716 offspring is \(+.186 \pm .031\). This coefficient, accordingly, is just six times its probable error. A comparison of these maternal and paternal coefficients, taken by themselves, would seem to indicate a probability that we had to do with a case of inheritance through the female parent only.

An analysis of this case showed us, however, that we were on very treacherous ground. Reference to the histograms (figure 3) reveals the presence of a single extremely aberrant wild specimen, having a ratio of 115, or in other words giving a deviation from the average of \(+15\), the largest in the entire series. This specimen was the mother of seven offspring, of which four agreed in giving plus deviations, while two gave minus ones, and one a zero deviation. Now it happened that one of these offspring showed a deviation of \(+10\) percent, this being the second-largest deviation for femur length. The product of these two \((= +150)\) is more than half of the algebraic sum of all the deviation products (i.e., of the numerator, in the formula for the correlation coefficient). That this coincidence was due to chance and not to heredity, we think probable for several reasons. Not only was it true that only four out of seven children of this mother showed positive deviations, but only three out of seven grandchildren showed positive ones, and none of these was extreme. Two were negative and two zero. Again, when femur weight is considered, we have the following seven deviations for this fraternity: \(+6, +5, -5, -1, 0, 0, 0\).

For the foregoing reasons, the coefficient of mother-offspring correlation was computed again, excluding this mother and her seven offspring. This time the figure is \(+0.105 \pm 0.033\). Even here, however, the coefficient is more than three times its probable error and, taken by itself, would be regarded as indicating the probability of a correlation. This
figure cannot, however, be considered apart from the coefficients for the three other characters. The figure in question is the only one found by us which approached a condition of probability in the statistical sense. It is certainly surprising, too, that if the correlation in respect to femur length is a real one, there is no correlation shown in the weight of these same bones.

Considering the sixteen separate group-coefficients of mother-offspring correlation, we find that 13 are positive and 3 negative, the mean of the lot giving us $+0.129$. (This is when the single aberrant female and her seven offspring are included. Omission of these last greatly reduces the figure.)

The coefficients of correlation in respect to degree of asymmetry (regardless of sign) are: father-offspring, $+0.059 \pm 0.034$; mother-offspring (excluding the "aberrant” family), $-0.008 \pm 0.033$.22

Passing to the differences between the first and second generations of race hybrids, the following are the standard deviations for the four groups of animals previously considered:

- Wild (96) ...................................................... $1.72 \pm 0.083$
- Pure races, cage-bred (211) .................................. $0.69 \pm 0.034$
- Subspecies hybrids, $F_1$ (552) ............................... $0.88 \pm 0.029$
- Subspecies hybrids, $F_2$ (300) ............................... $1.11 \pm 0.028$
- Subspecies hybrids, $F_2$ (300) ............................... $1.47 \pm 0.040$

The figure for the wild mice differs greatly, according to whether or not the single extremely aberrant individual is included. In any case, the number of animals is not great, so that the standard deviation for this group has not the same significance as those for the three other groups. As in the case of jaw weight, the variability is seen to be much greater for the $F_2$ than for the $F_1$ generation, the difference being more than 7 times its probable error. As in the previous case, also, the figure for the $F_1$ hybrids is significantly greater than that for the cage-bred mice of pure race, though the difference is not so great as that between the two hybrid generations. Comparison between the wild mice and any of the other lots, in this respect, is difficult for the reasons already indicated. The histograms (figure 3) should also be referred to in this connection. Analysis of those for the $F_1$ and $F_2$ series is particularly instructive.

Tabulating, as before, the figures for the separate lots of hybrids, we have:

22 The figure, with this family included, was not computed.
The differences are 7.0, 1.3 and 4.0 times their probable errors respectively.

**Femur weight**

As already stated, the mean value of the sinistro-dextral ratio for this character is $100.226 \pm 0.055$. Some interest attaches to the probable correlation between the ratios for femur weight and for jaw weight. There is, we find, a slight tendency for animals which are dextral or sinistral in respect to one of these pairs of bones to show a corresponding condition as regards the other. The correlation coefficient is $+0.090 \pm 0.023$, the former thus being nearly 4 times its probable error. These figures are based upon 869 individuals.

The question of the possible inheritance of a dextral or sinistral bias in femur weight was tested in the same manner as was done for the characters already considered.

The father-offspring coefficient was found to be $+0.071 \pm 0.034$, for the entire series. Of the 16 separate group-coefficients 8 were positive, 7 negative, and one 0, the mean of these being $+0.039$.

The mother-offspring coefficient was $+0.049 \pm 0.032$, for the entire series. There were 9 positive and 7 negative coefficients for the separate groups, the mean of these being $+0.027$.

Coefficients were not calculated for degrees of asymmetry, regardless of sign.

As regards variability in the four main sections of the mice, we have:

- Wild (96) .......................................................... $3.19 \pm 0.155$
- Pure races, cage-bred (212) ....................................... $2.37 \pm 0.116$
- Subspecies hybrids, F₁ (353) ...................................... $2.10 \pm 0.069$
- Subspecies hybrids, F₂ (299) ...................................... $2.28 \pm 0.058$

As in the two previous cases, there is a considerable and significant difference between the F₁ and F₂ generations, here amounting to about 6.5 times its probable error. There is also a much smaller and only possibly
FIGURE 4.—Showing sinistro-dextral ratios for femur weight. (See legend for figure 2).
significant difference between the \( F_1 \) mice and the cage-bred mice of pure race, the latter, as in the two previous cases, showing a lower variability. (See, also, figure 4.) The difficulties in the way of comparing the wild mice with any of the other lots are the same as were mentioned in the discussion of femur length. When the single extreme variant (the same as in the last case) is excluded, the variability is pretty close to that for the \( F_1 \) animals.

**Pelvis**

The maximum length of the innominate bone was taken, from the anterior end of the ilium to the most posterior point in the pubis. This series of measurements is considerably smaller than any of the others, because of the number of bones which it was necessary to reject. Bones were rejected which were obviously deformed. Some of these were rather badly contorted. Again, a large number were excluded owing to breakage during the cleaning process. Others still—and many such—were excluded because of scarcely perceptible damage which was inflicted during the process of making the caliper measurements, and which sometimes led to considerable

![Figure 5](image_url)

**Figure 5.**—Showing sinistro-dextral ratios for innominate (pelvic) bones. (See legend for figure 2.)
errors. Very slight pressure from the instrument was in some cases sufficient to inflict such damage, owing to the brittle character of the pelvic bones. In fact, it was found necessary to examine each specimen with a high-power hand-lens at the time of measurement.

The mean ratio for the 747 pairs of innominate bones was $99.901 \pm .018$. Thus, as in the case of the femur, we find this pair of bones to be predominantly dextral, at least as regards length. This preponderance is of high statistical probability as is evident from the relatively small probable error.

The histograms (figure 5), as well as the standard deviation ($0.736$ for the lot) show that the variability of this character is far lower than that of any character previously considered.

The father-offspring correlation for the pelvic ratio was $+ .055 \pm .040$, based upon 58 fathers and 498 offspring. The mother-offspring correlation was $+ .007 \pm .039$ for 113 mothers and 486 offspring. There is thus no evidence of heredity here. No separate group-correlations were computed, nor were the correlations in respect to degree of asymmetry.

The variability of the pelvic ratio for the four chief groups of mice is shown by the following standard deviations:

<table>
<thead>
<tr>
<th>Group</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild (78 individuals)</td>
<td>$0.82 \pm 0.044$</td>
</tr>
<tr>
<td>Pure races, cage-bred (125)</td>
<td>$0.77 \pm 0.033$</td>
</tr>
<tr>
<td>Subspecies hybrids, $F_1$ (279)</td>
<td>$0.75 \pm 0.021$</td>
</tr>
<tr>
<td>Subspecies hybrids, $F_2$ (265)</td>
<td>$0.73 \pm 0.021$</td>
</tr>
</tbody>
</table>

It is probable that none of these slight differences have any statistical significance.

**Discussion**

Two main problems have been dealt with in the second section of this paper: the problem of the inheritability of the individual tendencies toward dextrality and sinistrality in respect to these paired bones, and that of the relative variability of these departures from symmetry in the first and second hybrid generations. For this purpose, sinistro-dextral ratios were computed for three paired bones, and for two measurements (length and weight) in the case of one of these bones (the femur).

Parent-offspring correlation coefficients were computed, in order to determine the possibility of heredity in respect to each of these four series of ratios. Since the father-offspring and mother-offspring coefficients were computed separately, eight such coefficients were obtained. With a single exception, none of these coefficients was of such a magnitude that it would be regarded as having any statistical significance, when judged by customary standards. In two cases they were smaller than their own prob-
able errors, and in all but three cases (one of these being negative) they were less than two times their probable errors. The single apparently significant figure (the mother-offspring coefficient for femur length) was shown to be reduced to a scarcely significant magnitude by excluding one very aberrant mother and her seven offspring. Indeed the exclusion of a particular one among these offspring would have had nearly as great an effect. An added reason for doubting the significance of this single coefficient for femur length is the fact that the corresponding coefficient for femur weight is trivial. It would be hard to believe that dextrality and sinistrality in respect to femur length is hereditary, while that in respect to femur weight is not.

Of the eight coefficients referred to, it is true that seven carry the positive sign. This can hardly be regarded as anything more than accidental, when the facts just mentioned are considered, and when it is also borne in mind that the single negative figure is second-largest among the eight, being about two and one-fourth times its probable error.

For jaw weight and for femur length we also tested the question of the inheritance of degrees of asymmetry, regardless of whether this was predominantly dextral or sinistral. Here again, the results are negative. Of the four coefficients obtained, two bear the plus sign and two the minus, none of them being large enough to be significant.

This lack of inheritance in respect to the proportional size of the right and left members of a pair of bones stands in contrast to what we should have had a right to expect if we had tested the question of the inheritance of the absolute size of any one of these bones. On the analogy of numerous other cases in which parental-filial correlation has been determined for various characters, we should have been likely to find such a correlation in respect to either the absolute size of any of these bones, or the relative size (ratio to body size). Such determinations have not been made by us.

Passing to the relative variability of the two hybrid generations, a rather detailed analysis of our results is scarcely avoidable, since we regard our findings here to be among the most important results contained in the present paper. It has been shown that the sinistro-dextral ratios for jaw weight, femur weight and femur length show considerably higher variability in the F₂, as compared with the F₁ lots. The differences in the magnitudes of the standard deviations were of undoubted statistical significance, and moreover the same differences held without exception when the three main crosses were considered separately. As regards the ratios for the innominate bone, on the contrary, there were no such differences in variability, but it must be recalled that the variability of this character was very
low throughout the series. It should also be pointed out that exact measurements of this bone were more difficult than in the case of the femur.

The interpretation of the increased variability in the $F_2$ generation is somewhat complicated by the fact that certain other differences, apparently likewise significant, were found between the standard deviations of other lots besides the two hybrid generations. For example, the $F_1$ mice consistently showed a higher variability (for the first-named three characters) than did the cage-bred animals belonging to natural subspecies. This difference was smaller than that between the two hybrid generations themselves, but in at least two cases it was probably a real one. The case of the wild mice is peculiar. Owing to the small numbers and the presence of a single highly aberrant individual, it is perhaps safer to throw out of consideration the standard deviations for the two femur characters. As regards jaw weight, on the other hand, the figure for the wild mice significantly exceeds that for the pure cage-bred ones and about equals that for the $F_1$ hybrids.

To say that a difference is "statistically significant" merely means that it is not likely to be due to random sampling. Nothing is implied as to the actual cause of the difference. The differences between the two hybrid generations, discussed above, cannot, without further consideration, be attributed to the fact that we are dealing with subspecies hybrids. The occurrence of differences between some of the other lots of animals just referred to, shows that the possibility of other causes cannot be overlooked. One possible cause of such differences will be considered in some detail.

For reasons already stated, it seems safer to restrict our comparisons chiefly to the three cage-bred lots of mice. This procedure seems further justified by the fact that we are here dealing with individuals which have developed under closely similar environmental conditions. Whether, or to what extent, the conditions of life have any influence upon the degree of skeletal asymmetry is a question which we shall next consider. In the absence of any definite information on this point, it would seem wiser not to throw together individuals having such widely different histories as wild and captive mice.

In the three cage-bred lots we find a graded series in respect to variability as follows: (1) Pure races, (2) $F_1$ hybrids, (3) $F_2$ hybrids. As already mentioned, the step between 2 and 3 is considerably greater than that between 1 and 2.

Considerable reference has been made, in earlier reports upon these Peromyscus studies (Sumner 1915, 1918, 1920), to the occurrence of abnormalities of growth and even of marked deformities among the cage-bred
mice. It may be objected that all of these differences in variability (which are based upon differences of asymmetry) may be due to differing degrees of abnormality, resulting from the artificial conditions of life in captivity. The F₃ generation is more variable than the F₁, it may be contended, owing to its having been subjected to these conditions for one additional generation. We have given much attention to this point, and we think that such an explanation can be eliminated in the present case for several reasons.

1. There has been no obvious tendency for stunted individuals, even those showing an abnormal shortening of the long-bones, to be unsymmetrical. The most extreme case of asymmetry in the femurs was found in a wild mouse. On the other hand, it was a source of surprise to what an extent the normal growth of the femurs could be arrested without disturbing the exact balance between them.

2. If the asymmetry were due, in any measure, to the abnormal effects of captivity, we should expect to find the cage-bred mice, of pure race, to be more variable than the wild, which is the reverse of the truth. In respect to the jaw-bone, these cage-bred mice are considerably more symmetrical than the wild. Again, the F₁ hybrids (or about three-fourths of them) were derived from the mating of wild mice. Why, then, this constant increase in variability over the other first-generation, cage-bred animals, i.e., those of pure race?

3. The abnormal mice are commonly stunted in size, and the degree of abnormality is closely associated with the degree of stunting. Accordingly, we divided the F₂ mice into two equal groups, comprizing the smaller and larger individuals, respectively, and computed standard deviations for each half. Of the four characters considered in the present paper, two showed a higher variability for the smaller individuals, two for the larger, the mean of the four being slightly greater for the larger ones. It is not likely, therefore, that stunting has been a factor in increasing the variability.

4. In two of the three sets of hybrids, and for both sexes of these, the mean relative length of the femur (ratio to body) was actually less in the F₁ than in the F₂ generation. One of these was the “Calistoga-Carlotta” cross, in which the greatest increase in F₂ variability was manifest. The third set of hybrids, in which the femurs were relatively shorter in the F₂ generation, showed the least increase in the later generation. The inference is the same as that drawn in the preceding paragraph.

*The senior author was at first disposed to regard this as a probable explanation of the increase in F₃ variability shown in certain other characters (Sumner 1920, p. 397).*

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For these reasons, we believe that we are warranted in rejecting the supposition that these differences in variability are due to differences in the degree of abnormality, induced by the unnatural conditions of captivity. The pure races, F₁ hybrids, and F₂ hybrids probably differ among themselves in the variability of their sinistro-dextral ratios for the very reason that they do belong to "pure" and to first- and second-generation-hybrid stocks, respectively. It is not, it is true, commonly contended that first-generation hybrids are more variable than the parent races, at least when the latter are of pure stock. But it must be remembered that the wild "races" used in these experiments are far from being "pure" in the sense of being genetically uniform. Each of them comprises a mixture, displaying a considerable range of variations, some of which breed true. The F₁ generation from such a cross is not, therefore, wholly comparable with that derived from the crossing of pure strains.

Finally, the reader may be disposed to object that the "variability" which we have dealt with in this entire discussion is of a quite different sort from that ordinarily considered in genetic studies. It may be said that an increase in "variability," in our series, means merely an increase in the extent of asymmetry in the animals concerned. In a certain sense this is true, though it must be pointed out that an increase in asymmetry would not necessarily result in an increase in variability. If the increase in asymmetry were constantly in the same direction (e.g., toward a higher degree of dextrality in a given pair of bones), we might find no increase whatever in our standard deviations. It is the fact that we have a greater variety in the kind and degree of asymmetry that results in our obtaining larger standard deviations. We do not believe that there is any essential difference between variability of this sort and that resulting from differences in the absolute size or number of parts.

On the whole, therefore, we feel justified in insisitng that this constant and considerable increase in variability in the F₂ generation of hybrids, in respect to certain characters which are demonstrably non-hereditary, greatly weakens the force of the evidence which is commonly offered in favor of the "multiple factor" hypothesis.

SUMMARY

(1) Cases are discussed in which there was an undoubted specificity in the transmission both of an asymmetrical character itself and of its position as left or right. In others, the mere asymmetry or lack of balance appeared to be transmitted, without reference to the side of the body. In others yet, the structure or structures alone were found to be transmitted,
though departures from symmetry, either in size or position, were strictly non-hereditary.

(2) It is pointed out that the chromosome mechanism of heredity, as revealed through the observations and speculations of recent years, is ill-adapted to account for the transmission of definite spatial relationships, such as are involved in symmetry, etc. The egg-cytoplasm seems to be a more fit medium to account for such transmission. The fact that certain of the more striking cases of inherited asymmetry are known to be inherited through the male, as well as through the female, appears, on first thought, to be at variance with the latter suggestion. But a reconciliation with this view is shown to be possible, if we assume that specific substances in the sperm nucleus may have a selective affinity for one or the other half of the egg-cytoplasm, in consequence of qualitative differentiations in the latter. Thus the male parent could determine a condition either of symmetry or asymmetry in the offspring, without this condition being present, in any sense, in the paternal chromosomes.

(3) Considerable attention is devoted to an asymmetrical abnormality of the sacrum, which was found in about four and one-half percent of all specimens of Peromyscus. This proved to be strictly non-hereditary.

(4) The most extended presentation of data relates to the small differences between the right and left members of certain paired bones. The bones employed were the mandible, femur and innominate bones of Peromyscus. The two halves of the mandible were weighed, the femurs were both weighed and measured, and the innominate bones were measured. The first showed no constant asymmetry; the second were predominantly dextral in length, and predominantly sinistral in weight; the last were predominantly dextral in length. These departures from symmetry were slight, though of probable significance statistically, the number of specimens being great.

(5) No heredity was manifested in relation to individual deviations from symmetry in these bones. This question was tested by obtaining the “sinistro-dextral ratios” for all these bones, and then computing coefficients of parental-filial correlation. The evidence as a whole is decidedly negative. We also tested the question whether parents and offspring tend to agree in respect to grades of asymmetry (regardless of whether dextral or sinistral). No such tendency was revealed.

(6) Perhaps the most significant feature of our results is the fact that the $F_2$ generation of subspecific hybrids showed a considerably higher variability than the $F_1$ generation, in respect to this non-inheritable character, bilateral asymmetry. This may be fairly held to weaken the force of much of the evidence offered in favor of the “multiple factor” hypothesis.
LITERATURE CITED


