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Normal and abnormal bilirubin metabolism

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NORMAL AND ABNORMAL BILIRUBIN METABOLISM

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NORMAL AND ABNORMAL BILIRUBIN METABOLISM

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The term bilirubin immediately calls forth in the mind of the physiologist the normal processes of biliary pigment formation and the normal hepatic functions. Bilirubin to the pathologist usually brings up pictures either of liver damage or of some hemolytic process with deposition of this pigment in the tissues of the body. In the mind of the chemist the extensive chemical analyses of hemoglobin, bilirubin and the related pyrroles done by Fischer are recalled. To the mind of the medical clinician, the term bilirubin integrates the above pictures as a single working unit portrayed to him by his patient.

It is the purpose of this paper to present the normal and abnormal aspects of bilirubin metabolism giving some of the major contributions that have taken this subject out of the realm of conjecture and have placed it upon a scientific basis. Some points are still not explained, and these points of dispute will be pointed out. It is not the purpose of this paper to discuss the various pathological entities that may cause disruption in the normal bilirubin metabolism, but there will be an attempt to show that these entities may be classed into groups fundamentally the same as far as the principle involving the disrupted bilirubin metabolism is concerned; and, that through simplified classification of these processes, bilirubin disorders are more readily understood. Finally, an attempt will be made to show the various practical
applications that can be made upon the basis of the known facts of bilirubin metabolism.

C. Robert Hankins

3/24/39
PART I

NORMAL BILIRUBIN METABOLISM
Berzelius is credited with the first scientific studies of the biliary constituents, or bile. He studied bile as a whole, and gave a fair analysis of the various constituents as to their percentage composition. To an orange-red pigment found in the bile he attached the term cholepyrrhin. This work was done in 1840. (21)

Virchow in 1847, (13,39) made the first observations of extrahepatic formation of bile pigment. He found a substance which he called hematoidin occurring in tissues around old hemorrhages discovered at autopsy. In his report he stressed the close relation of hemoglobin to this substance, and also stated its close relation to the biliary pigment termed cholepyrrhin by Berzelius. However, Virchow could not prove conclusively that the pigment formed under such circumstances is identical with cholepyrrhin (or bilirubin) and he therefore gave it the name hematoidin.

Frerichs and Stadler, 1850, (39) worked with bile and renamed the orange-red pigment bilirubin. The name is derived from "bilis" meaning bile, and "ruber" meaning red. They found that a pigment resembling bilirubin could be produced in vitro by the action of sulphuric acid upon bile salts, and more important, that the injection of bile acids into the blood stream of an animal would be followed by the appearance of undoubted bilirubin in the urine. Their conclusion was that the body could transform bile acids into bile pigment.
Kuhne, 1858, (39) repeated and confirmed a forgotten or unnoticed observation of von Dusch that bile acids are powerful hemolytic agents; he insisted that the experiments of Frerichs and Stadler did not prove the origin of bilirubin from bile acids, for those investigators had not taken into account the fact that a large amount of hemoglobin is set free in the plasma by the injection of bile acids. Kuhne was unable to satisfy himself that the injection of hemoglobin alone, in the absence of bile acids, would be followed by bilirubinuria, and he was forced to hold to the idea that the bile acids were necessary in some way for the formation of bile pigment.

Herrmann, 1859, (39) was able to produce bilirubinuria at will by inducing intravascular hemolysis with injections of distilled water. This was the first clear demonstration that simple liberation of hemoglobin into the blood stream may be followed by an increased output of bile pigment in the urine. Neither Naunyn, 1868, nor Steiner, 1873, could confirm Herrman's results, a fact that is now easily explained, since the absence of bilirubinuria is by no means a proof that there has been no increased formation of the pigment. Tarchanoff, 1874, on the other hand not only confirmed Herrman's work, but, with the use of bile fistula animals, carried the proof of the relation of hemoglobin to bile pigment still further by demonstrating, for the first time, that the introduction of pure hemoglobin into the circulation is followed by a marked increase in the amount of
bile pigment excreted by the liver.

As early as 1769, Morgagni taught that the liver excreted the bile which was brought to it, preformed, by the blood. An altogether different view was held for many years following the experiments of Minkowski and Naunyn, 1886, (13) who found that jaundice was produced in geese by poisoning with arseniuretted hydrogen, whereas jaundice could not be so produced if the liver was removed immediately after the administration of the substance. From this they concluded that the liver was necessary for the production of jaundice, and the belief arose that bile pigment was formed by the hepatic cell.

McNee, 1913-14, (30) repeated the experiments of Minkowski and Naunyn, and found small traces of bile in the urine and the tissues of hepatectomized geese following the administration of the arseniuretted hydrogen. McNee observed, in histologic studies of the normal goose liver, that the von Kupffer cells contained erythrocytes and gave a marked iron reaction. In the liver of jaundiced geese, the Kupffer cells were seen to contain, besides many erythrocytes, large quantities of yellowish green pigment, which resembled biliverdin, although chemical proof was lacking. McNee concluded that in geese bile pigment was formed by the endothelial (Kupffer) cells of the liver and by the small number of endothelial cells found elsewhere (spleen and bone-marrow). In geese, hepatectomy accomplishes removal of the greater part of the reticulo-endothelial system, but in mammals it does not. Thus
the conception of bile pigment formation returns to the concept of Morgagni, 1769, that bile constituents are produced in the body and carried to the liver via the blood stream, at least in relation to bilirubin.

Before reviewing further physiological experimentation, one should consider the type of cell of which the Kupffer cell is an example. During the last half century, certain fixed cells and certain wandering cells having phagocytic properties have been studied by many workers. Ribbert, 1904, (13) showed the interrelation of all these different cells in their common ability to engulf particles of lithium carmine injected into the circulation. To Aschoff belongs the credit for the conception that these reticular and endothelial cells of the various tissues possess common properties and functions, and may be designated together as the reticulo-endothelial system.

As these physiological principles were being studied, the biochemists were doing much work in determining the composition of the bile pigments. The discoveries of Virchow and Herrman were the guiding lights in their studies. It was up to them to show the chemical similarity of bilirubin to hemoglobin, or if possible definitely prove the derivation of bilirubin from hemoglobin. Fischer has probably been the greatest student of this subject, and to him is attributed most of the advancement made in this field. Without going through the long processes involved in determining this technically difficult problem, it
is the purpose of this paper to present the modern concept of this problem.

As stated before, the chemists started working with hemoglobin and attempted to prove the derivation of bilirubin from this source. Hemoglobin is classified as a hematoprotein. The hematoproteins are present in all protoplasm, but are found more accessible in the blood. Their purpose is to carry oxygen, and they are associated with the processes of oxidation in every tissue of the body. Hemoglobin is insoluble in itself, and so must be carried about physically in the red blood cells rather than in solution. Hemoglobin is a sub-group of the conjugated proteins. Conjugated proteins are made up of two radicals, one being the protein radical and the other being termed the prosthetic group. In the case of hemoglobin, the protein fraction is made up of a globin and the prosthetic group is a porphyrin. The prosthetic group is a relatively small part of the molecule, roughly making up about 4% of the molecule. Porphyrins, chemically, are constructed on the porphine nucleus; this nucleus is made up of 4 pyrrole rings arranged in rosettes and called tetra-pyrroles. The porphyrins are found widely distributed throughout nature in both animal and plant kingdoms, their main characteristic being their ability to produce color in reflected light, or, in other words, they are pigments.

When hemoglobin is treated, under appropriate conditions, with glacial acetic acid and sodium chloride, and the mixture
warmed gently, a substance is obtained which crystallizes readily as brown crystals--this substance has been called hemin. (See the plates at the end of this paper for formula and reactions mentioned in this chemical discussion of bilirubin) The other product of this cleavage is globin. Hemin has been synthesized by Fischer. (6) It contains four methyl-pyrrol radicals and an atom of iron, and may be represented as shown in the aforementioned plates. If the crystals of hemin are treated with sodium hydroxide, the corresponding base, heme, is liberated; this is the original prosthetic radical of hemoglobin.

The porphyrin nucleus related to hemin is called protoporphyrin. (see plate) With the introduction of an iron atom into the molecule, neutral heme is formed, which may be isolated as hemin, indistinguishable from the product derived from hemoglobin. This is indirect evidence to the fact that hemoglobin is built up from preformed protoporphyrin. It is assumed by the chemist that the body is capable of forming this nucleus as a necessity to the making of hemoglobin. (Evidence will be given later to substantiate this fact from the physiological viewpoint.)

Bilirubin has been studied extensively by Fischer and he has given the substance a definite formula. (See plate) As yet, he has not been able to synthesize this substance, so the formula rests merely upon analysis of the substance. As shown by his formula, Fischer believes that bilirubin arises by oxidative scission of the porphyrin nucleus of hemoglobin, the iron
being lost in the process. The method by which the iron is lost, and the nature of the compound in which it is lost is still a vague subject. This iron containing radical is called hemosiderin; the material can be demonstrated to contain iron; and beyond that, there is little information. Lemberg (22,23) believes that the process leads to biliverdin first, and then to bilirubin. Biliverdin is an oxidized product of bilirubin, containing two more oxygen atoms, but otherwise the same empirically. In his experiments, Lemberg has been able to obtain bilirubin from hemin as outlined in the plates previously mentioned, and has identified the intermediary products. He suggests that the formation of bile pigments in the animal body proceeds in a similar fashion yielding biliverdin which is afterwards reduced to bilirubin. He admits that from a physiological point of view much remains to be done, but he believes that this model will provide a good guide to further research.

Andrews (1) and Rich (41) have proven that the hematoidin found by Virchow is the same substance as bilirubin.

In summarizing the modern chemical viewpoint, we must admit that definite proof of the formation of bilirubin from hemoglobin is still lacking. An important step in clearing up this matter would be the synthesis of bilirubin, and the demonstration that the formula as given is correct. But in all fairness, we must say, the only chemical evidence of derivation of bilirubin from hemoglobin is the chemical similarity of the bilirubin to the porphine
nucleus. Lemberg's work seems to be the closest approach to
definite proof of this derivation.

Knowing these chemical relationships of bilirubin to hemo-
globin, we can now proceed more intelligently to follow the
physiological developments that have been made in the last few
years.

Interesting observations were made by Peabody and Brown (13)
on the phagocytosis of erythrocytes in the bone marrow. These
workers employing the Berlin blue iron reaction, observed in the
bone marrow the presence of erythrocytes and granules of iron in
the same phagocytes. They concluded that the phagocytes of the
bone marrow ingest erythrocytes and that the hemosiderin (an iron
containing substance derived from the breaking down of hemoglobin)
is formed within the phagocytes from hemoglobin.

P. Corr (8) recently studied material from 71 necropsies
with respect to the distribution of hemosiderin and bilirubin.
Sections of liver and spleen and occasionally of bone-marrow,
kidney, lymph node, lung, suprarenal gland and pancreas were
stained for iron by the Berlin Blue method. Hemosiderin was
found in varying amounts in the splenic phagocytes, in the hepatic
and Kupffer cells of the liver and in the reticulo-endothelial
cells of the bone marrow. Corr concluded that the reticulo-
endothelial cells of the spleen and the bone marrow were most
active in breaking down the hemoglobin. He believed that the
Kupffer cells and large wandering phagocytes served as a
reserve of the reticulo-endothelial system.

Haldeman (13) anesthetized a group of dogs and removed pieces of liver, spleen, bone marrow, lymph nodes and various other tissues. The hemolyzed corpuscles of 100 cc's of dog's blood were injected into the juglar vein. At intervals of 15 minutes to 3 hours thereafter, additional pieces of the same organs were excised. Samples of blood from the juglar vein, corresponding in time to the various sets of tissues removed were studied by spectrophotometric methods to determine the quantity of bilirubin circulating in the blood at the various stages of the experiment. Since it is known that iron is liberated in the breaking down of hemoglobin into bilirubin, it is reasonable to conclude that the cells that contain iron are those concerned in the conversion of hemoglobin into bilirubin. Furthermore, when samples of various tissues are removed before the intravenous injection of hemoglobin and at regular intervals of time thereafter, it may be justifiably assumed that the tissues showing the striking increase in iron content are those actively concerned in the process of converting hemoglobin to bilirubin, while those tissues containing a constant amount of iron (as the spleen) probably have to do with the storage of the iron until it is again built up into hemoglobin. Haldeman concludes that the bone marrow plays the major role in the formation of bilirubin, with the liver and spleen and possibly the lymph nodes having lesser parts in the process.
Mann and Bollman (31) to afford conclusive proof of this extrahepatic formation of bile pigment made a series of studies on hepatectomized dogs. These demonstrated clearly that bilirubin formation continued in the absence of the liver. By measuring the arterial and venous flow to the various organs and then comparing the bilirubin content indicated that the spleen and the bone marrow, particularly the latter, are the chief sites of formation of bilirubin. They also found bilirubin in the splenic vein and in the veins returning from the bone marrow after injections of hemoglobin solution into the arteries afferent to these structures.

Rich (40) stated it was obvious that if red blood cells could be kept under continuous observation after being ingested by clasmatocytes, or while disintegrating in the tissue fluids outside of living cells, a much clearer idea of the process might be obtained. This Rich has tried to do by the use of the Lewis tissue culture. Fresh red blood cells were added to cover glass cultures containing wandering phagocytic cells of mesodermal origin, and the fate of the red cells was followed under the microscope. It was found that the phagocytes wandered among and ingested the red cells in large numbers, and that, as the hemoglobin was broken down within the phagocytic cells, beautiful rhomboid or needle-shaped crystals typical of bilirubin, and often diffuse, bright green biliverdin appeared within the cytoplasm of the living phagocytes. These pigments gave a perfect Gmelin test for
bilirubin under the microscope and possessed every other chemical and physical property of bile pigment for which they were tested. The Prussian blue reaction often showed the presence of an iron-containing residue within the phagocytic cells by the side of the bile pigment. Up to the stage in which crystals of bile pigment first appeared within the phagocytes, no crystals were ever found outside of these cells, although there were often great numbers of red cells disintegrating extracellularly. Later, as the clasmatocytes themselves began to die and to disintegrate, the intracellular crystals and soluble pigment were, of course, set free; so that in older cultures there might be numerous crystals lying outside of cells some of which apparently formed extracellularly from soluble bile pigment. It was certain, however, from the study of hundreds of such cultures that the hematoidin crystals can make their first appearance within phagocytic cells, and that they were not phagocytized preformed. The experiments are not yet entirely free from the possibility that bile pigment may have been present invisibly in soluble form in the tissue fluids, and until this possibility of phagocytosis of the pigment in soluble form from the surrounding media can be controlled, it cannot be stated with any certainty that hemoglobin can be converted into bilirubin within phagocytic cells of the reticuloendothelial variety. However, this is one of the best proofs we have of the extrahepatic formation of bilirubin in the reticuloendothelial system.
Despite the extensive evidence of the formation of bilirubin extrahepatically, some men still cling to the idea that the hepatic cell in itself is able to produce bilirubin. The main evidence to date that the hepatic cell makes bilirubin is that the jaundice produced by certain substances, notably toluylene-diamine, is prevented by the removal of the liver. Toluylene-diamine produces both increase in formation of bilirubin and retardation of its excretion, together with demonstrable injury to some of the hepatic cells. In order to explain the jaundice produced by this substance, on the basis that the retained bilirubin is formed by the hepatic cells, it is necessary to postulate that toluylene-diamine injures the hepatic cells in such a manner that they form more bilirubin than normal and excrete less than normal. Mann (33) believes that this consideration must remain as a possibility until the mechanism of the jaundice caused by the drug is conclusively determined, although it appears rather unique that injury to a cell should cause it to increase its activity in one respect and decrease it in another.

Among numerous data which have been assembled both in support of and against hepatocellular bilirubin formation, one observation of Lubarsch, 1921, (13) deserves mention. This is the appearance of bilirubin in distant metastases from primary carcinomas of the liver. Lubarsch states that since these lesions do not contain von Kupffer cells, the presence of bile pigment is evidence of formation by hepatic cells. On the other hand, Aschoff, 1922,
has pointed out that these cells might be serving simply in an excretory capacity, removing bilirubin from the blood, the only function which he attributes to the normal liver cell insofar as bilirubin metabolism is concerned.

Lepehne (9) felt that the question of bile pigment formation could best be explained as a dualistic role, one in which both hepatic and reticulo-endothelial cells participate. This view of Lepehne's is difficult to accept since it seems unlikely that glandular epithelial cells, (the hepatic cells) and cells of mesenchymal origin (the v. Kupffer cells) would both possess the rather specific function of converting hemoglobin into bile pigment. In the following it will be seen that epithelial cells of ectodermal or entodermal origin have not been shown to have this ability, while the evidence already discussed makes it clear that the reticulo-endothelial cells generally are able to make this conversion. In connection with this argument, van den Bergh and Papendieck (9) have seen marked variation in the relative amount of bilirubin taken from hemorrhagic fluids of different sources. In an extensive series of tests, it was found that bilirubin was regularly present in those fluids within mesothelial-lined cavities, but was present only twice in epithelial-lined cysts. It also showed that epithelial specialization of cells which were originally mesodermal is associated with a diminished production of the ferment necessary to bilirubin formation. In this connection it is noteworthy that bleeding into the gastro-intestinal canal does not result in bilirubin formation. Cases of gastro-intestinal
cancer are not infrequently seen, having blood in the feces and at the same time exclusion of bile from the duodenum because of neoplastic disease. Hemoglobin or hematin is readily demonstrable in the feces of these patients, but bilirubin or its derivatives are absent.

The evidence for the extracellular formation of bilirubin is rather convincing; in fact, Leschke's study of bilirubin formation in the spinal fluid has proved that bilirubin may be formed extracellularly, probably by means of a ferment. (9) This work was based upon Ober-Mayer and Popper's demonstration, 1908, that the yellow color of spinal fluid, as in the Froin syndrome and in other blood containing fluids, was due to the presence of bilirubin. Leschke injected washed erythrocytes in the human lumbar canal and noted bilirubin formation which did not occur with control erythrocytes kept sterile in the incubator. After centrifuging, the supernatant fluid was placed in the incubator with more washed erythrocytes while the other half was incubated without addition of cells. An increase of bilirubin in the half containing erythrocytes was observed. This pointed strongly to a fermentative mechanism as the basis for the transition of hemoglobin to bile pigment, which was also suggested by the studies of van Czike, the results of which indicated that small amounts of bilirubin may form in vitro, when blood alone is allowed to stand.

Besides the possibilities of bilirubin being formed intracellularly and extracellularly from hemoglobin, the possibility of the body
making this pigment independently of hemoglobin must be considered. As mentioned under the chemical discussion of this subject, it has been suggested that the body is capable of making the pyrrole nucleus, and thus might be capable of manufacturing bilirubin. Hawkins (20) found that anemic dogs formed new hemoglobin from the entire amount of foreign hemoglobin administered, but at the same time formed additional bilirubin equivalent to the entire amount of hemoglobin given. These observations indicated that new hemoglobin molecules cannot be utilized in the formation of new hemoglobin, although other portions of the hemoglobin molecule may be incorporated in the new hemoglobin. Whipple (49) also agrees that the body is capable of producing pyrrole rings. Whether the body produces some of the pyrrole rings directly for bile pigment formation is a question, but it does produce them for the formation of hemoglobin, using the old globin and iron fractions in the process.

Rich (39) in summarizing the evidence that hemoglobin furnishes a source of bile pigment, states: "1. that hematin and bilirubin are closely related chemically; 2. that hemoglobin spilled into the tissues almost anywhere in the body may be transformed, locally into bile pigment; and 3. that the presence of an excess of hemoglobin in the circulation, whether introduced in pure form experimentally or liberated during hemolysis in experimental or pathological conditions, is regularly followed by an increased formation of bile pigment." Mann and Bollman (32)
have shown that bile pigment continues to be formed in animals after complete removal of the liver and there can be no doubt that hemoglobin is the material from which it is derived.

Further summary to this point on the evidence presented, one may conclude that the major site of bilirubin formation is located within the reticulo-endothelial system, particularly within the bone marrow and spleen. It is also reasonable to conclude that the bilirubin that is formed within the liver is formed within the von Kupffer cells and not the hepatic cells.

It is of some interest at this point to consider briefly the methods used to determine bilirubin concentrations in the blood. These methods can be grouped into four main groups: oxidation method; icteric index; methods based on Ehrlich's reaction; and spectrophotometric methods.

The first method, that of oxidation of bilirubin has proven to be rather inaccurate and has been replaced by other methods. The principle involved was that the bilirubin of the sera was oxidized to biliverdin, the resulting color being compared with previously made color charts. The results were obtained directly from the color chart with which it compared.

The icteric index is a method of direct comparison of the blood sera with a known standard. Assuming the yellow color of blood sera is principally due to bilirubin, methods have been
evolved to determine the amount of bilirubin by comparing the color of the serum with artificial standards of approximately the same shade of yellow color—usually a solution of potassium dichromate. If in crystalloidal state, a given value of bilirubin would impart a greater color intensity to a given volume of sera than that same amount of bilirubin if it were in a colloidal state. The icterus index, then, may be regarded as a functional as well of the physical state of bilirubin as of its total quantity. (Barron, 2) Elton (10) in correlations from over 1700 sera examined found: "1. The icterus index fails to conform consistently with any constant proportion of total bilirubin; 2. For a given icterus index the bilirubin content of the serum is higher when it exists in the colloidal state than when it is crystalloidal; 3. Colloidal bilirubin accumulates in the bloodstream at the icterus index 16.6, but fails to impart a higher color intensity to the serum in which it is suspended until it undergoes a physical change, expressed by the development of a direct type of van den Bergh." In further criticism of this method, Barron (2) states, "As the chemistry, the amount and the daily variation of the lypoohromes (lutein, carotin, etc.) of the blood is unknown, the writer is of the opinion that these methods in which bilirubin as well as all other yellow substances in the blood serum are estimated, ought to be discarded. It is well known that the yellow color of sera increases with a diet rich in vegetables, producing a pseudo-jaundice." In summary of the icteric index
we may say that it is useful in showing an increased bilirubin content in the blood, but it must not be relied upon to give a quantitative estimation of the bilirubin present.

The van den Bergh reaction is probably the best known method of testing for bilirubin. Erhlich, 1883, first made the observation that when a diazonium salt is added to a solution of bilirubin, a new compound, azo-bilirubin, is formed, the solution becomes colored, and the color depends upon the reaction of the solution—red when neutral and blue when the solution is acid. Van den Bergh applied it first to clinical work in 1913, thus transferring the outlook on jaundice from the color of the skin to quantitative changes in the blood. (McNee, 29)

The van den Bergh test consists in overlaying a small amount of clear serum with the diazo reagent and observing for a purplish or bluish red ring at the line of contact. The diazo reagent consists of two parts, A and B, mixed together just before making the test. Solution A is a 0.5% solution of sulphanilic acid in distilled water. Solution B is a 0.5% solution of sodium nitrite in distilled water. If a reddish violet color reaction occurs immediately on adding the reagent to the serum and reaches a maximum within about 30 seconds, the reaction is known as "immediate direct." When with the same technique, a reddish color is observed to develop only after some time has elapsed (1-10 minutes) and deepens to a violet hue, the reaction is known as "delayed direct." When a reddish color appears immediately and, only after some time deepens to a violet hue, the reaction is known
as a "biphasic direct." When no color reaction whatever occurs after addition of the reagent to the blood directly, but does occur after the blood proteins have been precipitated with alcohol and the reagent has been applied to the clear alcoholic filtrate, the reaction is known as the "indirect." The van den Bergh is delicate enough to determine the presence of bilirubin in dilutions as great as 1:1,500,000 in alcohol. The significance of the various reactions obtained in this test will be pointed out later.

The spectrophotometric method for determination of bilirubin calls for a costly instrument and careful technique, and is not suitable for most laboratories. There is no doubt that it is by far the most accurate and sensitive way of measuring bilirubin. It is possible to determine the character and the shape of the spectrophotometric curve of bilirubin for a dilution as low as one-fiftieth of the smallest amount measurable by the van den Bergh technique.

By use of the above procedures a number of facts have been elicited. Normal blood serum contains between 0.1 and 0.25 mg of bilirubin per 100 cc; or in other words, it is present in dilutions of 1:1,000,000 to 1:400,000. (37) As the reaction is frequently spoken of in "units" of bilirubin, it may be here observed that one unit corresponds to a dilution of 1:200,000. In other words, normally, blood serum contains 0.2 to 0.5 units.

Barron (2) believes the normal bilirubin content to vary
between 0.1 and 0.5 mg%. It is possible some of the higher normal figures include some individuals classified among the group of latent familial jaundice or those affected with slight liver insufficiency. However, daily variations occur. The highest reading is found in the morning when fasting; it falls after breakfast, the lowest figure being obtained generally 3 to 6 hours after meals. A prolonged fast seems to increase the bilirubin content of the blood. Sex seemingly plays no difference in the bilirubin level.

Van Berghman (10) estimates that in the average 70 kilogram human adult the daily bilirubin production is about 500 mg, which, in terms of 100 cc units of blood serum volume, may be computed as approximately 14.3 mg. The normal bilirubin level does not exceed 0.2 mg%. Visible jaundice may appear in the white race when this level rises to 1.5 or 2.0 mg and may often be detectable in the sclerae at much lower levels. So, the body must have mechanisms for adequate excretion of bilirubin.

It is almost unnecessary to state the fact that the liver is the main organ of excretion of bilirubin. The bilirubin is removed from the blood in the liver sinusoids and is excreted in the bile. The normally functioning liver keeps the bilirubin level down to figures just previously mentioned. The exact threshold value of the liver in relation to bilirubin has not been
shown; but if it does exist, it is very low.

It appears that, as for sugar and a number of other blood constituents, there is a renal threshold for bilirubin; the normal threshold is about 2.0 mg% or 4 units. It is regularly found in the urine when the concentration is 3.5 to 4 units. (Rabinowitch, 37) A curious exception must be made to this statement; bilirubin giving the indirect van den Bergh reaction is not excreted by the normal kidney regardless of how high the bilirubin sera concentration is. This fact will be discussed later. So we can say that the kidney plays a part in the excretion of bilirubin only under pathological conditions.

The fate of bilirubin in the intestinal tract after leaving the common duct brings up some rather interesting factors. Sackey (43) has proven that bilirubin is not reabsorbed in the small intestine. He placed bilirubin in isolated sterile jejunal loops for a period of two hours and was able to quantitatively obtain the bilirubin back from the loop contents. He has also shown that no change takes place in the nature of the bilirubin while in the small intestine. He has placed bilirubin in intestinal juice and incubated it for 2 hours with no change in the bilirubin.

Blankenhorn (3) devised an ingenious means of obtaining blood from the portal vein of a dog without opening the abdomen. He made corresponding studies of the juglar and portal blood, and the lymph from the thoracic duct. He found that "bilirubin is
not reabsorbed from the intestine by way of the portal vein in healthy animals. Bilirubin may be absorbed from the intestine by the lymphatics but only in minute amounts."

With this evidence, it is reasonable to say that the bilirubin passes through the small bowel with little quantitative loss due to absorption, and with no chemical alteration. But in the large bowel a large proportion of the bilirubin is changed into another substance called urobilin, or into urobilinogen. These substances are reduced bilirubin. Their relation to bilirubin may be seen on the plate showing chemical properties and formulae of the bile pigments. This change in the bilirubin is supposedly due to bacterial action in the colon. The urobilin formed in the intestines is in part excreted in the stools, in part destroyed, and in part resorbed by the portal blood stream and carried back to the liver. Normally the liver removes this urobilin and excretes it in the bile. If the liver does not excrete this urobilin, it is then excreted by the kidney, the renal threshold being practically nil for urobilin. Normally the liver removes this urobilin from the blood to such an extent that urobilin does not appear in the urine in readily demonstrable quantities.

To prove this transformation of bilirubin, Friedrich Muller (45) obtained urobilinogen by bringing bilirubin in contact with peptone solution and putrefactive bacteria obtained from fecal material. (1892)

Methods of estimating urobilinogen and urobilin are:
1. Fluorescence in the presence of zinc salts; 2. Spectroscopic absorption bands; 3. Production of red color with Ehrlich's aldehyde.

The fluorescence in solutions of zinc salts was one of the first methods of determining the quantity of urobilin present. Acriflavine is used as a standard, being standardized against a zinc acetate filtrate from a standard known solution of urobilin with regulated pH (equivalent to a 0.05 N HCl solution). By determining standards in this way, a measurement of the urobilin content is effected by comparing its fluorescence at a great dilution with a standard containing acriflavine, calibrated in turn against pure urobilin. (24)

The spectroscopic method is the most accurate, but again requires expensive apparatus and special technique.

For a simple test, we are again indebted to Ehrlich (1886). The reagent in the original form consisted of a 3 per cent paradimethylamidobenzaldehyde in a 50% solution of HCl. Neubauer, 1903, was the first to use it clinically. When added to urobilinogen solutions a red color develops. The test is not specific for urobilinogen, being common to a variety of chemically related substances, but, for practical purposes as far as urine is concerned, it may be assumed to be so. (37)

Wallace and Diamond, 1925, (47) devised a quantitative test. The test is, essentially, one of serial dilution of urine and the addition of a definite amount of reagent to a definite quantity
of diluted urine. The reagent for this purpose consists of a 2% solution of the aldehyde compound in 20% HCl. The amount of pigment is judged by the greatest dilution of the urine in which the faintest pink color is discernable on the addition of the reagent. The normal figure obtained in this manner ranges from a 1:10 to 1:20 dilution.

Urobilinogen is a normal constituent of urine in man. It is present in smallest amount in the morning urine, or may be absent altogether--rising somewhat in the afternoon and evening. In constipated individuals with increased putrefactive changes in the large bowel, the quantity of urobilogen rises above normal. (47) Urobilin never appears in freshly voided urine, but appears upon oxidation of the urobilogen.

McMaster (24) has shown that the normal presence of urobilin in the bile and feces of dogs depends upon the passage of bile pigment to the intestine, either through the normal channels, or by abnormal ones, as when it is fed by mouth. He devised a method whereby animals can be totally, partially or intermittently deprived of their bile, without infection of this secretion or of the duct system. By this means he was able to study urobilin physiology and to make the above observations. He has shown that complete loss of bile from the body resulted in the total disappearance of urobilin and urobilinogen from the bile, feces and urine. Partial loss of the bile resulted in a corresponding reduction in the urobilin of the dejecta.
Feedings of sterile urobilin-free dog bile to intubated dogs losing all of their bile and having no urobilin in it, or in the feces or urine, were followed by the appearance of the pigment in the hepatic bile secreted shortly thereafter. When the feedings were stopped, the urobilin soon disappeared.

Total obstruction of the bile flow caused disappearance of the urobilin of the bile and stool. Later as the animals became heavily jaundiced, the pigment appeared again in very small quantity in the feces. Autopsy at this time showed that the intestinal mucosa was deeply tinted with bilirubin, some of which undoubtedly had passed into the lumen of the bowel and had there been changed to urobilin.

McMaster (24) also employed intermittent diversion of the bile stream of animals from the intestine to a collecting apparatus, and showed that while bile pigment still reached the intestine urobilin was present in the bile secreted by the liver, but that almost at once after the bile had been diverted from the gut urobilin disappeared from it. He has shown that the bacillus putrificus and other intestinal flora were capable of changing bilirubin to urobilin.

So in conclusion we can say that urobilin is an end product of the action of bacterial flora of the large intestine on bilirubin; that bilirubin in itself is not reabsorbed from the gut in any quantitative manner, but this end-product is rather readily absorbed; that for the production of urobilin, bilirubin must reach
the large intestine; that final excretion of the urobilin is via the feces in the normal individual, only a small proportion being eliminated by the kidneys.

The following is a graphic summary of the changes from hemoglobin to urobilin with the intermediary steps; reading downward:

| Hemoglobin | Hematin | Bilirubin | Biliverdin | Urobilinogen | Urobilin |

The following chart was taken from Hewlett (15) after Willis and Addis. This gives in summary the fate of the bile pigments in the various organs of the body.

<table>
<thead>
<tr>
<th>Reticulo-endo. System</th>
<th>Bile Passages</th>
<th>Intestine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin</td>
<td>Bilirubin</td>
<td>Urobilinogen</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Intestine</th>
<th>Liver</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urobilinogen</td>
<td>(Part excreted in bile) (Part returned to form Hemoglobin?)</td>
</tr>
<tr>
<td>(Part absorbed) (Part destroyed)</td>
<td>(Part enters general circulation and excreted by kidney).</td>
</tr>
</tbody>
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PART II

ABNORMAL BILIRUBIN METABOLISM
Having considered the normal physiology of bilirubin, one can now view the pathological disturbances of its metabolism with clearer insight. Knowing the normal physiology, one can point to the process of the bilirubin metabolism in which the fault lies in the pathological process.

The most objective sign of abnormal or pathological bilirubin metabolism is jaundice or icterus. This phenomenon is defined by Dorland as "a syndrome characterized by hyperbilirubinemia and deposition of bile pigment in the skin and mucous membranes with resulting yellow appearance of the patient." Jaundice comes from the French word "jaune", meaning yellow.

Jaundice, being such an evident thing, has been described in connection with disease from the times of the early Greeks. The Greeks called the disease icterus. This word is derived from the Greek word applied to an animal resembling our cat. The eyes of this animal were said to be of the yellow color characteristic of jaundice. Others believe that the word is derived from the name of a bird called icterus; the sight of this bird was a cure to the jaundiced patient, but death to the bird, as the belief goes. (21)

The Latins called jaundice "morbus regius", from the yellow color of gold, the "Rex metallorum". For the same reason, another Latin synonym is "aurigo" from "aurum, meaning gold."
Jaundice could hardly fail to be noticed early in the
history of mankind, as the change of complexion and color would
be a striking appearance. Accordingly there is abundant mention
of it in the genuine Hippocratic writings, which most often speak
of it as a complication of other diseases rather than as a
disease in itself. In the "Aphorisms" it is said to be a bad
sign if the liver becomes hard during jaundice; and in the third
book of the "Epidemics", that an habitual pain in the neighborhood
of the liver, with other symptoms, preceded a jaundice.
It is disputed if Hippocrates conceived any relation between the
bile secreted by the liver and the symptom of jaundice. (21)

Galen asserts that "the yellow bile, when it is carried all
over the body, still keeping its own nature, causes a disease
called jaundice". He believed that "weakness of the gall bladder,
which does not draw to itself the bile out of the liver as it is
accustomed, leaving the blood impure, is the cause for jaundice".
Galen also states that jaundice may be found without any associated
liver damage. (21) In the light of our present day knowledge,
it is remarkable that such accurate conclusions could be drawn
from pure observation.

Returning to the present-day conception of jaundice, Rice (38)
in view of the known physiology of bilirubin, states that jaundice
will occur if any of the four following qualifications are met:

1. "If the threshold of the liver for bilirubin
excretion became greatly raised."
2. "If for any reason, bilirubin were produced faster than the normal liver cells could excrete it."

3. "If the mechanism of the liver were so disturbed that the amount of bilirubin normally produced could not be satisfactorily removed from the blood."

4. "If there occurred any combination of the above."

According to Rice, the first is hypothetical, with no evidence of its existence, and which it is unnecessary to involve in order to explain any form of jaundice.

The second classification, an uncomplicated excessive production of bilirubin, is rarely a cause of jaundice for two reasons, according to Rice: "First, because the normal liver is able to excrete much more bilirubin than is ordinarily delivered to it; and second, because an excessive production of bilirubin is, as we shall see, practically always associated with other conditions which tend to impair the excretory power of the liver." There is a tremendous reserve power in the liver in excreting bilirubin. The excretory power of more than 95% of the liver substance of the dog or monkey can be abolished, and yet the remaining 5% of the liver tissue will suffice to prevent the development of jaundice. In the human, atrophy of over half of the liver has been shown not to produce any noticeable jaundice. The fact that the liver possesses a much greater capacity for excreting bilirubin than it is called upon to use normally makes it clear that up to a certain limit an over-production of bilirubin will be taken care of without
the development of cutaneous jaundice. Just what this limit is in the human being is not accurately known, but the numerous pathological conditions in which the amount of pigment in the stools is greatly increased above the normal in absence of jaundice stand as evidence that marked increases in the amount of bilirubin produced can be excreted by the normal liver well enough to prevent visible jaundice. It is imaginable that with excessive production over a long period of time the excretory power of the liver may be impaired due to the strain put upon it. There is no evidence that this occurs; on the contrary one would believe that an uncomplicated demand for increased work would, through the process of work hypertrophy, increase the liver's efficiency, as in the case of the other excretory organs, rather than depress it. Overproduction of the pigment is ordinarily associated with conditions other than mere overstrain which tend to depress the excretory power of the liver (e.g. anoxemia, febrile disease, immaturity of liver cells); thus it is believed that jaundice rarely, if ever, results from simple overproduction of bilirubin.

The third possibility (liver damage to the extent that the normal bilirubin production can not be excreted) undoubtedly causes certain cases of jaundice. No jaundice is known to result from a functional depression of the excretory power of the liver without actual loss of cells through necrosis. There is no pathological condition which is known to impair the function of the
living hepatic cells to such a degree that they are unable to prevent jaundice by excreting the amount of bilirubin normally found. If, however, the total amount of functioning liver substance is reduced by necrosis and cellular damage, jaundice will result wherever the loss of liver tissue reaches a point at which the activity of the cells remaining alive is insufficient for the proper removal of the pigment from the blood (e.g. acute yellow atrophy, yellow fever, chloroform poisoning). (38)

The fourth possibility (a combination of any of the preceding possibilities) is the most common. A liver with reduced capacity for excreting bilirubin may be able to rid the blood of the amount of pigment formed normally; if bilirubin is produced in excess, such a liver may be quite unable to excrete it at all, and jaundice will result from the retention of the excess pigment. (38)

In line with the preceding discussion, the clinician classifies jaundice as being hepatogenous or hemotogenous in origin. The hepatogenous jaundice is caused by the inability of the liver to excrete via the bile any or all of the bilirubin brought to it by the blood stream. The pathology of this process is found entirely within the liver. Rice (38) has termed this regurgitation jaundice. The cause of this jaundice is blockage of the bile passages so that bile cannot be excreted (calculus, inflammatory processes, or neoplastic disease). The commonly accepted view on the manner of origin of the jaundice is that after occlusion of the common bile duct the bile continues to be excreted by the liver
parenchyma, and since it is unable to follow its normal course to the duodenum, it is damned back into the delicate bile capillaries. The increase in tension in these very delicate structures causes them to rupture; the extravasated bile is absorbed either by the blood capillaries or by the lymphatics and is then distributed throughout the body, being in part deposited in the tissues and in part excreted by the urine. Bloom (4) has proven that the removal of bilirubin early in obstructive jaundice is via the lymphatics, then later by the blood. That is, before the bile canaliculi rupture, the bilirubin is resorbed by the lymph in the space immediately about the canaliculi and returned to the systemic system before rupture has taken place.

The hemotogenous jaundice has been termed retention jaundice by Rice (38). This implies merely an abnormal retention of the bilirubin within the systemic circulation. Basically it is concerned with a minimal amount of liver damage and a pronounced tendency for peripheral blood destruction (e.g. hemolytic icterus, any poison causing intravascular hemolysis).

Besides hepatic dysfunction due to hepatic disease, there is a type of constitutional nature. The constitutional type shows a light yellow or sallow tint to the skin, however, the conjunctiva is not colored. There is no demonstrable hepatic or splenic damage. The urine contains appreciable amounts of bilirubin and considerable amounts of urobilin. This condition has not been adequately explained, but it has been found in the
people falling within this group a tendency to hepatic disease, especially cholecystitis. (42)

Before continuing further in the field of jaundice it is necessary to reconsider some of the factors in the van den Bergh test for bilirubin. As will be remembered, van den Bergh got four different types of reactions with his reagent, namely the immediate direct reaction, the delayed direct reaction, the biphasic direct reaction, and lastly the indirect reaction. Van den Bergh was unable to discover the factor which underlies these different reactions, nor has this problem been solved up to the present time. A voluminous literature has accumulated which relates to this subject.

Van den Bergh (45), in 1918, found a direct reaction occurred with serum of a patient having obstructive jaundice (regurgitation type). He found that an indirect reaction resulted in a patient having hemolytic icterus (retention type of jaundice). Because of this and other work, van den Bergh suggested that there was some chemical difference between the two forms of pigment, or perhaps the pigment was modified by its attachment to the protein of the serum; the form present in the serum of the obstructive jaundice was probably circulating in a free state, while that giving the indirect reaction was bound with serum protein and, in order to detect its presence, had to be liberated from the proteins by the alcohol. Thus, van den Bergh believed that in the passage through the hepatic cells the bilirubin was changed chemically, thus
accounting for the difference in reaction in the obstructive and
hemolytic types of jaundice.

Blankenhorn (10) found that there was present in bile and in
many icteric sera a form of bilirubin which would pass through an
animal membrane, showing that since it acted as a diffusate on
dialysis, it must be in the crystalloid state. He found, however,
that this was a relatively unstable form, for on repeated dialysis
of the diffusates less and less would pass through the membrane,
indicating that it readily reverted to a non-dialyzable or colloidal
form. Collinson and Fowweather (11) applied this observation to
the van den Bergh reaction, and found that the crystalloid form
alone gave the direct positive reaction, while the non-dialyzable,
or colloidal, form was direct negative, reacting with the diazonium
salt only after the addition of alcohol and giving the indirect
reaction. They conclude that the difference is largely in physical
state; the direct negative being free bilirubin, a suspensoid
colloid; the direct positive being a pure crystalloid.

Vaughan and Hubbard (46) after studying xanthochromic spinal
fluid, concluded that the type of reaction depends upon the
quantitative relationship between the pigment and protein. They
found with material of low protein concentration that when the
bilirubin exceeded 0.3 mg% the van den Bergh reaction became
direct. They believe, therefore, it is not necessary to consider
that passage through the liver cells is the determining factor
that changes the indirect to the direct type of bilirubin. Gregory
and Andersch (12) have shown through clinical studies that as high as 26 mg% of bilirubin can be found in spinal fluid with an indirect van den Bergh; therefore they do not believe that the quantity of bilirubin present in the sera determines whether the van den Bergh will be direct or indirect in contradistinction to Vaughan and Hubbard's work.

Fowweather (11) has reviewed the literature well on the van den Bergh reaction, and the following are only a few of the many theories put forward.

Davies and Dodds, 1926, believed that bilirubin was responsible for the direct reaction, whereas an oxidation product of bilirubin was responsible for the indirect reaction.

Roberts, 1928, put forward the view that bilirubin in the free colloidal condition was the pigment of indirect sera, while in direct sera the bilirubin was in combination with some substance the nature of which is as yet undetermined.

Newman, 1928, opposed the view that the pigment of indirect sera was free bilirubin and claimed that the pigment of direct sera was a sodium hydrogen bilirubinate.

McGowan, 1930, considered that the pigments of direct and indirect sera were identical, the types of van den Bergh reaction being dependent on the alkali reserve of the blood; indirect reactions were obtained when the alkali reserve was diminished.

Snider and Reinhold, 1930, claimed that the type of
van den Bergh reaction was dependent on the concentration of the pigment which is present, direct reactions being associated with high concentrations of bilirubin, indirect reactions with low concentrations.

Kuster believes a keto-enol modification of bilirubin exists explaining the varied reaction.

Fowweather, 1926 and 1931, put forth the idea that the pigment of indirect sera was free bilirubin as an acid, while the pigment of direct sera was a salt of bilirubin, probably the ammonium salt.

Mann and Bollman (32) have shown that the van den Bergh reaction for serum bilirubin is indirect in animals that have become jaundiced following complete removal of the liver. The amount of bile pigment that accumulates in the blood for the first 24 hours after the removal of the liver is comparable in amount to that which is found after an equal interval of time following ligation of the common bile duct and extirpation of the gall bladder. In the latter case the bilirubin of the blood gives a direct van den Bergh reaction. Removal of the liver when obstructive jaundice is present produces serum bilirubin with a true biphasic reaction obtained by the van den Bergh method. If the liver is removed several hours after ligation of the common duct and removal of the gall bladder, at a time when a definite direct van den Bergh reaction is present in the serum, the amount of bilirubin reacting directly will be subsequently unchanged.
Additional bilirubin will accumulate in the blood after removal of the liver, so that the total amount of bile pigment in the blood will progressively increase at the same rate that it was increasing following ligation of the common bile duct, but the bilirubin that is added to the blood after removal of the liver gives only the indirect reaction. Thus Mann and Bollman's experiments show the relative importance of liver passage of bilirubin in relation to the way it will react to the van den Bergh reaction, but it also shows a possible mechanism for explaining the delayed direct and the biphasic reactions. McNee, 1922, (29) believes blood from obstructive jaundice would yield an immediate direct reaction, hemolytic jaundice would account for the indirect reaction, and jaundice due to a toxic agent and which, furthermore, may be due to a combination of obstructive and hemolytic types would account for a biphasic reaction.

Probably one of the most ingenious of the theories put out thus far is that of Barron's. (2) Because of its apparent rationality and the usefulness in explaining the various van den Bergh phenomena it is given in more detail than the other theories. It must be remembered, however, that this theory is not accepted by all. Barron, in contradistinction to van den Bergh, does not believe that the passage of bilirubin through the hepatic cells changes the bilirubin in any manner; but to him it has seemed more probable that there is some substance present in the bile which is responsible for the difference in the reaction. In other words, he believes
that the type of reaction depends upon the medium in which the pigment is contained rather than upon any alteration of the pigment itself. Barron has shown that, while pure bilirubin at the pH of blood gives the direct or prompt action, if it be added to normal plasma it then gives the indirect reaction. This Barron believes to be the result of the adsorption of bilirubin by the plasma proteins, the pigment being in that way prevented from reacting promptly with the reagent. If, however, substances which have the property of lowering surface tension are first added to the plasma and then bilirubin is introduced, or if such substances are added to the plasma at the same time as the bilirubin, the reaction of the pigment remains direct. This occurs apparently for the reason that substances which lower surface tension seem to be adsorbed by proteins more readily than is bilirubin by the plasma proteins, and it is significant that these substances are present in whole bile and occur in increased amounts in the plasma in various forms of jaundice in which the blood bilirubin gives the direct reaction. Bile salts and acids have the faculty of lowering surface tension and are the active principles in this role in the case of bile. In obstructive or regurgitation jaundice, whole bile, containing both bilirubin and bile salts, is regurgitated into the systemic circulation. The bile salts collect at the protein interface leaving the bilirubin free to react with the diazo compound, giving a direct reaction.

When sodium bilirubinate solution giving a direct reaction is
added to normal serum in increasing amounts up to 12 mg%, a
typical indirect reaction takes place. When the concentration
is increased to 16 mg%, the reaction becomes of the biphasic type.
When the concentration is higher than 16 mg%, a direct reaction
is obtained. Barron believes that up to 12 mg% the bilirubin is
being adsorbed by the protein; from 12 to 16 mg%, small enough
amounts of bilirubin are left free from the protein interface, or
relatively free of this interface so that a biphasic reaction may
take place; from 16 mg% on upwards the bilirubin remains free in
the plasma and thus gives the direct reaction. The direct reaction
can be obtained from the first by adding bile acids which have
the surface tension lowering effect in the case of bile.

Whatever be the ultimate explanation of the different forms
of the reaction, a great amount of experimental and clinical in-
vestigation has made it perfectly clear that when the plasma
bilirubin in a case of jaundice is subjected to the van den Bergh
test, the direct reaction indicates that whole bile, containing
bile acids and cholestrin as well as bilirubin, has been regurgi-
tated into the blood stream, and therefore that one is dealing
either with obstruction of the ducts or with necrosis of the liver
cells, for these are the two conditions which permit bile to
escaped from the canaliculi into the blood; the indirect reaction,
on the other hand, informs us that the bilirubin in the plasma
under examination has not been regurgitated into the blood from
the canaliculi, but that it represents pigment which the liver has
not yet been able to remove from the blood stream. Bloom (4) brings out a little recognized fact concerning obstructive jaundice that gives very good proof of the above summarizing statement. He found that bilirubin in early obstructive jaundice is removed via the lymphatics surrounding the canaliculi before rupture of these canaliculi ever takes place. During this period when the bile pigment is thus being resorbed, the van den Bergh shows an indirect reaction. He has shown histologically that ruptured bile canaliculi are not found in the early stages of obstructive jaundice. But as soon as the bile canaliculi have ruptured, pouring whole bile into the lymphatic and blood streams, the van den Bergh becomes direct positive.

The biphasic reaction is due to a small amount of direct reacting pigment in plasma containing a large amount of indirect reacting pigment. According to Barron's view, a biphasic reaction would occur, for example, if bilirubin continued to be added to the plasma, the proteins of which had already adsorbed all of the pigment they were capable of holding. This form of reaction does not yield the clear cut information which the pure direct or indirect reaction yields, for it can occur as a result of a high degree of pigment retention in the absence of regurgitation, or it may occur in certain stages of biliary regurgitation caused by obstruction or by necrosis of liver cells. However considered in relation to findings in the stools and urine in a given case, it is frequently possible to determine whether the biphasic
reaction results from the retention or from the regurgitation of bilirubin.

In contradistinction to the condition of jaundice, there may exist a condition of hypobilirubinemia. This condition is found in all secondary anemias where blood lost by hemorrhage is not promptly replaced. It can also be associated with alterations in the hematopoietic system (diminution of the number of red blood cells thrown into the general circulation) which have as a consequence a diminished production of bilirubin. Hypobilirubinemia is found in chronic nephritis with cardiac complication, secondary anemias, aplastic anemias, and chlorosis. Murphy, in 190 cases of secondary anemia due to different conditions finds a persistant hypobilirubinemia. This condition is due, then, to a general lack of blood destruction in the body. This fact may be taken as further proof of the derivation of bilirubin from hemogoblin.

The classical work on urobilin has been done by McMasters and his co-workers already mentioned before. Besides studying the normal physiology involved in urobilin, they have also included pathological study that explains very nicely the abnormal physiology occurring in the human patient.

He has shown (24) that urobilin is not formed normally unless bilirubin reaches the large intestine. In further experimentation, (26) urobilinuria has never found after liver damage except when bile pigment was present in the intestine. Thus, for example,
it appeared during the first days after ligation of the common duct, but disappeared as the stools became acholic. When this had happened, a small amount of urobilin-free bile, given by mouth, precipitated a prompt urobilinurea. After obstruction of the duct from one-third of the liver, mild urobilinurea was found, but no bilirubinurea. In animals intubated for the collection of a part of the bile only, while the rest flowed to the duodenum through the ordinary channels, liver injury caused urobilinurea, unless indeed it was so severe as to lead to bile suppression, when almost at once the urobilinurea ceased, though the organism became jaundiced. He concludes that "urobilinurea is an expression of the inability of the liver cells to remove from the circulation the urobilin brought by the portal stream with the result that the pigment passes on to the kidney and urine. Urobilinurea occurs with far less degree of liver injury than does bilirubinurea."

McMasters (27) also found that animals rendered urobilin-free by the collection of all the bile from the intubated, uninfected common duct, remain urobilin free during and after extensive blood destruction due to intravenous injections of hemolyzing agents and also reinjections of the animals own hemolyzed blood. On the other hand, when bile flow into the intestine is uninterrupted, urobilinurea occurs during blood destruction caused in any of the ways mentioned, and it parallels, both in severity and duration, the destructive process. He concludes, "Urobilinurea, occurring during blood destruction, is primarily the result of an increased
excretion of bilirubin from which, in turn, an unusually large quantity of urobilin is formed within the intestine. The liver fails to remove from the portal blood all of the latter pigment which is resorbed and consequently some of it reaches the kidneys and urine."

And again, McMasters (28) shows that experimental infection of the intubated and previously sterile biliary tract of the dog with particles of the stools leads to a formation of urobilin from the bilirubin of the bile as it flows through the ducts. No urobilinurea occurs, however, unless temporary biliary obstruction is produced, or the liver parenchyma is injured. Then urobilinurea develops, despite the fact that no bile is reaching the intestine and, by corollary, no urobilin being formed there. "Cholangitic urobilinurea, as one may term the phenomenon just described, to distinguish it from the urobilinurea having origin in the pigment absorbed from within the intestine is far more pronounced in animals possessing a healthy gall bladder than in those with a pathological gall bladder or with one prevented from functioning by severance of the cystic duct. These facts suggest that there may be an active absorption of urobilin from the normal gall bladder. There can be no doubt that the pigment is absorbed from within the bile ducts." He further states that there is no justification for the belief that urobilin is ever formed through the action of the liver parenchyma.

In summarizing this work on urobilinurea, we may say;
urobilinurea occurs in excessive hemolysis and whenever there is an excessive excretion of bile pigments by the liver (as when a stone suddenly becomes dislodged from the common duct allowing a large amount of bile to empty into the gut at once); no urobilinurea occurs with obstruction of the common bile duct since the bilirubin must reach the intestine to be changed into urobilin, but when hepatic cells are affected by bacteria or toxins, urobilin may be formed in the bile ducts themselves resulting in urobilin being formed within the liver itself. The excretion of urobilin from the portal blood stream is one of the most delicate functions of the liver, and is one of the first functions to become impaired in a pathological process; whereas, the liver may still be function­ing well in regards to bilirubin excretion, yet some increased quantity of urobilin may be escaping the liver and be showing in the urine. Thus, the test for urobilin in the urine is a good functional test of the liver and rather delicate in nature, too, and may be one of the first signs of liver dysfunction.

An attempt will now be made to correlate those factors brought out in the normal and abnormal metabolism of bilirubin and show their application in clinical medicine.

Being presented with a clinical case of jaundice, one may usually come to an accurate diagnosis by studying the type and amount of the biliary pigments within the blood, the urine, and the stools. By studying these factors, one can determine whether
or not the jaundice is of the regurgitation type or of the retention type. A summary of the characteristics of these two types of jaundice will be given at this point.

Retention jaundice, in its pure form, whatever the etiology, is recognizable clinically by the following characteristics:

1) The bilirubin of the plasma, never having passed through the liver cells and being, therefore, unmixed with the constituents of the bile, gives the indirect van den Bergh reaction; 2) The stools contain an increased amount of urobilin because, in the face of the increased production of bilirubin the liver is excreting more of the pigment than normally, even though it cannot excrete enough of the excess to prevent jaundice; and 3) The urine contains an increased amount of urobilin but no bilirubin; (it is not yet clearly understood why only the direct reacting bilirubin can escape readily from the blood to the urine; but if it be true that the indirect reacting bilirubin is bound to plasma protein while the direct reacting bilirubin is not, it is clear that the latter form could be excreted more readily by the kidneys than the former); and again, the urine will be free from bile salts because bile salts are formed in the liver and are absent from the urine unless they have been poured into the blood by the escape of whole bile from the canaliculi into the blood stream. Pathologically, the liver may show cloudy swelling, or atrophy, according to the etiological reason for their mal-function, but there is no widespread necrosis; and the bile ducts are patent.
Regurgitation jaundice, on the other hand has the following characteristics: 1) Plasma bilirubin gives the direct van den Bergh because the bile canaliculi are altered in such a way that whole bile escapes into the blood stream; 2) The stools contain less urobilin than normally since no bile or very little bile is getting into the intestine—the acholic stool; and 3) The urine contains bilirubin, because the direct plasma reacting bilirubin escapes into the urine. Many clinicians have noted a bradycardia and a pruritis accompanying regurgitation jaundice. At first these symptoms were attributed to the hyperbilirubinurea. But recent work of Horrall and Carlson and of Still (19), in which pure bilirubin spectro-photometrically identical with circulating serum bilirubin was injected intravenously, has found bilirubin to have no toxic effects. The toxic effects of jaundice were found to be due to bile salts mainly. The bradycardia (according to Stewart and King) is due to the bile salts which act by increasing the tonus of the vagus nerve.

The following chart was taken from Rich (38) and summarizes well the differentiating points of the two types of jaundice:

<table>
<thead>
<tr>
<th>TYPE</th>
<th>VAN DEN BERGH (Bilirubin)</th>
<th>URINE (Bile Salts)</th>
<th>FECES (Urobilin)</th>
<th>Urobilin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Retention jaundice</td>
<td>Indirect</td>
<td>0</td>
<td>0</td>
<td>/</td>
</tr>
<tr>
<td>Regurgitation jaundice</td>
<td>Direct</td>
<td>/</td>
<td>/</td>
<td>0 or /</td>
</tr>
</tbody>
</table>

Increased
Decreased
"Into these two main groups all various forms of jaundice can be rationally and helpfully placed from the standpoint of pathogenesis, for the results of laboratory findings, as listed above, will enable one to determine the fundamental mechanism responsible for the jaundice in a given case and will permit one to form a judgement regarding the general type of anatomical alteration present in the liver." (Rice, 38)

We may now mention some of the other clinical applications that are concerned with bilirubin metabolism.

Perkin (36) believes that a routine check of the van den Bergh of patients taking arsenicals affords the earliest possible indication of damage to the liver. If a record of the blood bilirubin is made at the commencement of the treatment and repeated at regular intervals during the course, one may proceed with safety as long as the reading remains normal. A rise of a few mgs. is significant and the patient should be treated accordingly. Any considerable rise calls for immediate cessation of treatment. Once an arsenical hepatitis is established, determination of the bilirubin offers a reliable evidence concerning the severity and progress of the lesion.

Bochner-Mortensen (5), from his results and comparison with the results of other liver function tests, concludes that in its limited field of application, the bilirubin tolerance test is a sensitive test of the liver function. While inapplicable in hepatic disorders with bilirubinurea, it is useful in the diagnosis
of cirrhosis and in evaluation of the condition of the liver in chronic hepatitis, cholangitis, and in examination of possible injury of the liver in infectious diseases and different disorders such as diabetis, exophthalmic goiter and certain skin and brain disturbances. It may prove to be especially suitable for the investigation of possible lasting functional disorders of the liver following acute hepatitis.

Rozendal (42) states that the mechanism for secretion of bilirubin is a sensitive one. It is influenced by emotion and by disturbances of the sympathetic nervous system, as well as by very minor degrees of hepatic injury. An increased concentration of bilirubin may be the one laboratory evidence of dysfunction and it should be used more widely. It will in all probability disclose many unsuspected cases of mild functional and organic disturbances of the liver as well as a group of individuals of a constitutional type especially susceptible to disease of the gall bladder.

The diagnostic value of hypobilirubinemia in the differentiation of pernicious anemia from aplastic or secondary anemia is now a well known procedure. (2) The increase of urobilogen in pernicious anemia associated with an icteric tint of the sclera and skin and often a palpable liver presents difficulty at times in the differentiation between hepatic disease and primary anemia. Aside from blood morphology, the examination of the feces indicates an increased pigmentation to a dark brown color in contradistinction to the light
or acholic feces accompanying liver disease. In the anemias, bile is never found in the urine, while in hepatic diseases associated with icterus, bile is always present.

The constant presence of urobilogen in catarrhal jaundice, and its total absence in mechanical icterus, such as encountered in carcinoma of the head of the pancreas and of the biliary tract, forms a marked distinguishing diagnostic feature between this benign form of jaundice and those due to malignant causes. The presence of urobilogen, even in traces, during the state of jaundice promptly rules out an obstruction or mechanical icterus.

Bilirubin metabolism has been studied more carefully than any other function concerned with the liver to the present date. Knowledge of this has been a great aid in diagnosis of liver dysfunction, but can not be relied upon too much. Quite a bit of confusion is present regardless of the nice theories that have been concocted to explain the biphasic and delayed direct van den Bergh tests. These results occur quite frequently in cases of jaundice and are of little diagnostic value in single tests, but when repeated over some time, they often change to a direct or true indirect reaction. (47) However, great aid is obtained by studying the constituents of the feces and urine. The tendency has been to shy away from the use of the van den Bergh test and to rely upon the lactose tolerance tests and such related tests. These newer tests certainly have their place since they test an altogether different function of the liver. But, in the light
of our present knowledge, the tests for bilirubin and urobilin in the blood, urine and feces are still important steps in diagnosing a given case of jaundice.

Recent experiments with bilirubin have offered what may be a truly great advance in our knowledge of the purpose of bilirubin. To date it has been considered only as a waste product of hemoglobin metabolism. Najib-Farah (35) working on pneumococcal infections in rabbits noted that the rabbit sera contained no bilirubin. Knowing the lytic action of bile on the pneumococcus, he added bilirubin to the rabbit sera. Whereas before the pneumococci grew rapidly in the sera, the growth of the virulent pneumococci was inhibited by the addition of bilirubin. The organisms undergo agglutination and disintegration (partial and complete lysis), the capsules being denuded of their contents. The possibility of bilirubin being an aid to body defense in certain infections is indeed a new concept—the possibility that bilirubin may be an agglutinin and a lysin in vivo is interesting, and certainly is a line of thought that is worthy of further study.
HEMOGLOBIN

HEMIN
Protoporphyrin

Bilirubin

Urobilinogen
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