On Bilirubin-metal Complex Compounds in Relation to Black Pigments of Gallstones

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Gallstones occasionally contain black pigments which consist mostly of polymers of bilirubin derivatives. In order to study whether any bile pigment-metal complex is also concerned with such black pigments of the gallstone, attempts were made at synthesis of bilirubin-Cu complexes. When free bilirubin and cupric chloride were mixed in a solvent mixture of chloroform and ethanol, a dark blue or black substance was formed. This pigment was soluble in ethanol and showed a characteristic absorption at 350 m\(\mu\) (Soret band) and two visible absorption bands at 595 m\(\mu\) and 645 m\(\mu\). Infrared spectroscopy has revealed that this compound is a complex salt of bilirubin and copper having an analogous structure to metalloporphyrins, in which bilirubin has a ring structure of tetrapyrroles and copper is located in the center of nitrogen atoms of pyrroles. On the other hand, addition of cupric chloride to bilirubin in sodium hydroxide solution resulted in sedimentation of a black substance. This pigment has proved spectrometrically to be a bilirubin-Na-Cu complex formed by coordination of copper to sodium bilirubinate. Thus, it seems to be possible that the black pigments of the gallstone include some metal-complexes of bile pigments.

There is an unusual variety of the gallstone which is characterized by black appearance of the surface and transections. My previous experiments revealed that black pigments were contained in such gallstones which differed from melamins\(^1\) and consisted mainly of polymers of bilirubin derivatives.\(^2\) However, considering the presence of copper\(^3\) and other metal elements in bile at significant concentrations, it is also possible that metal complexes of bile pigments, e.g., bilirubin-copper complexes, are produced in vivo and incorporated into the gallstone, and take part in its black coloration. To evaluate this possibility on experimental basis, a bilirubin-copper complex and a bilirubin-sodium-copper complex, being dark blue or black in color, were prepared in vitro and their structures and general properties studied by spectrometric methods.
Outline of methods for preparation of bilirubin complexes

**Bilirubin-Cu complex:** This complex compound was produced when a chloroform solution of free dibasic acid bilirubin (I) and an ethanol solution of cupric chloride (II) were mixed and the mixture (III) allowed to stand at room temperature. The color of the mixture III ranged between green and blue according to the mole ratio of bilirubin vs. Cu and also to that of chloroform vs. ethanol. In order to prevent oxidative denaturation of the complex compound by the effect of excess Cu++, the solution III had to be freed from unreacting Cu++. For this purpose, the solution III was transferred into a separation funnel and washed with distilled water repeatedly, supplementing a small portion of ethanol to the solution before every step of the washing. The potassium ethylxanthogenate test was utilized to detect Cu++ in the washings. The washed solution (IV) freed from Cu++ was evaporated up and dried in nitrogen flow. The residual solid material (V) was extracted with chloroform and then the residue (VI) extracted with ethanol. Finally, the ethanol extract (VII) was evaporated and dried up in nitrogen flow, and the objective complex compound was recovered as a solid substance (VIII).

**Bilirubin-Na-Cu complex:** Free dibasic acid bilirubin dissolved in a 10^{-2}M aqueous solution of sodium hydroxide was added to an aqueous solution of cupric chloride. A brownish-black precipitate was separated by centrifuging, and washed with distilled water, ethanol and chloroform, and then dried in vacuo into a solid substance (IX).

**Measurements**

In order to clarify the reaction of the bilirubin-Cu complex and to determine the optimum conditions for its production, spectrometric observation was performed with a Hitachi 139 type spectrophotometer and a Hitachi EPI-S_{2} type infrared spectrophotometer.

**Results**

1. **Conditions for the production of bilirubin-Cu complex**

   **Formation of bilirubin-Cu complex:** In the first series of the experiment the visible spectrum of the solution III was investigated at various mole ratios of Cu to bilirubin in an equivolume mixture of ethanol and chloroform. Fig. 1 shows the spectra of the solution containing excess of Cu over bilirubin after standing for one hour at room temperature. The spectra a, b and c of this figure, corresponding to small excess of Cu (bilirubin: Cu=1:1-6), show two portions of absorption band on either side of an isosbestic point at about 500 m\mu. Of these two absorptions, the one at 450 m\mu is a characteristic absorption of free bilirubin.
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Fig. 1. Visible spectra of the solution III at various mole ratios of bilirubin vs. cupric chloride. Solvent, ethanol : chloroform = 1:1. One hour after mixture of the solutions I and II.

Therefore, the other one which is located at about 645 mμ and associated with shoulder absorptions on both sides represents a newly formed complex compound. On the other hand, the spectra d and e, corresponding to large excess of Cu (bilirubin: Cu = 1:9–12), are devoid of the bilirubin absorption but show three distinct absorptions at 495, 595 and 645 mμ. This multiplicity of absorption suggests a possibility of secondary change of the complex in the presence of large excess Cu++. The next experiment, that is follow-up of the spectrum of the solution III, was carried out to evaluate this possibility.

Fig. 2 shows the changes of the visible spectrum of the solution III of bilirubin: Cu = 1:3 when it was left standing at room temperature. An isosbestic point is seen at about 660 mμ. As time elapsed, the absorption in the range of longer wave length than this was weakened, and an absorption at 595 mμ, which was a mere shoulder of the 645 mμ absorption in an initial phase of the reaction, gradually became a distinct absorption maximum. This change is interpreted as indicating maturation of the complex compound, i.e., transformation of a primary product into a more stable form. The result of a similar study at a larger excess of Cu (bilirubin: Cu = 1:9) is shown in Fig. 3. In this case, the 645 mμ absorption decreased and the 595 mμ absorption increased in the intensity with the lapse of time, and a new absorption peak appeared at 495 mμ one hour or more
Fig. 2. Change of the spectrum of the solution III by time. Bilirubin : Cu=1:3. Solvent, ethanol : chloroform=1:1.

Fig. 3. Change of the spectrum of the solution III by time. Bilirubin : Cu=1:9. Solvent, ethanol : chloroform=1:1.

after beginning of the reaction. It is concluded from these results that the stable bilirubin-Cu complex produced by the above-mentioned method is represented by absorptions at 595 μm and 645 μm, and the absorption at 495 μm is due to oxidation product of this compound.

Fig. 4 is the result of another series of experiments in which the ratio of bilirubin to Cu was fixed at 1:6 and the composition of the solvent altered variously. The result suggests that ethanol accelerates the formation reaction of the complex.
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Fig. 4. Visible spectra of the solution III for various compositions of the solvent. Bilirubin : Cu=1:6. One hour after mixture of the solutions I and II.

Isolation of bilirubin-Cu complex: The above results show that the formation of the bilirubin-Cu complex proceeds effectively in the presence of a large excess of Cu++. But, excess Cu++ causes a change of the product into the substance which shows a maximum absorption at 495 m\(\mu\). Therefore, in order to obtain the bilirubin-Cu complex, the excess Cu++ has to be removed by washing with distilled

Fig. 5. The effect of well-timed washing of the solution III on its visible spectrum. Bilirubin : Cu=1:40. Solvent, ethanol : chloroform=1:1.
water after an appropriate period. The following experiments were performed to
determine the optimum timing of the washing. Forty or 45 times excess of Cu++
was added to bilirubin in an equivolume mixture of ethanol and chloroform, and
the mixture was followed up spectrometrically. When the solution III of the
above composition was thoroughly washed with distilled water after one hour's
standing at room temperature, the absorption peak at 492 mµ did not appear in
the resulting solution IV (Fig. 5). But, when the washing was carried out after
one and a half hours, that is after the absorption peak at 495 mµ already became
prominent, a new absorption peak at 525 mµ appeared in the solution IV (Fig. 6).

![Fig. 6. The effect of delayed washing of the solution III on its visible spectrum.
Bilirubin : Cu = 1: 40. Solvent, ethanol : chloroform = 1: 1.]

Accordingly, it was found that the best result was obtained when the solution III
was washed after one hour's standing at room temperature. On each occasion,
the washing of the solution III had a tint of purple with weak fluorescence, and
was positive for the potassium ethylxanthogenate test.

The solution IV, freed from excess Cu++, was evaporated up in nitrogen flow,
and the residue V was extracted with chloroform. Small absorption bands of
500–550 mµ range disappeared with this procedure. The residue VI
treated with chloroform was finally extracted with ethanol. The ultraviolet-
visible spectrum of the ethanol extract VII is shown in Fig. 7. In the visible
region, this extract exhibited two distinct absorptions at 595 mµ and 645 mµ,
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Fig. 7. Ultraviolet-visible spectrum of the sample VII.

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exactly the same positions as the stable bilirubin-Cu complex identified in the crude solution III (Fig. 2). In the ultraviolet region, a very characteristic band was identifiable at 350 μm.

2. Infrared spectra of the products
   a) Bilirubin-Cu complex

The bilirubin-Cu complex recovered from the ethanol extract VII as the solid material (VIII) was lustrous powder of dark blue or black in color. The infrared spectrum of this substance measured by the KBr-disk method is shown in Fig. 8 with that of free bilirubin for comparison. There were a number of differences between these two spectra, among which the following were noteworthy: (1) In free bilirubin, the absorption peak at 1,692 cm⁻¹ was due to the stretching vibration of C=O of carboxylic -COOH (νC=O), but in the bilirubin-Cu complex this shifted toward higher wave numbers, around 1,710 cm⁻¹. (2) Free bilirubin did not show the 1,100 cm⁻¹ band which was remarkable in the bilirubin-Cu complex. (3) The absorption band due to the stretching vibration of N-H (νNH) at 3,400 cm⁻¹ was much weaker in the bilirubin-Cu complex than in free bilirubin.
b) Bilirubin-Na-Cu complex

The bilirubin complex produced in aqueous solution in the presence of Na\(^+\) and Cu\(^{++}\) (Sample IX) was non-lustrous black powder which was insoluble in water and organic solvents. The infrared spectrum of this substance was characterized by the absence of \(\nu_{C=O}\) of carboxylic acid and by the presence of strong and broad absorption bands at 1,560 cm\(^{-1}\) and 1,400 cm\(^{-1}\) which are attributable to \(\nu_{C=O}\) of carboxylate -COO-. However, the absorption due to \(\nu_{C=O}\) of free carboxylic acid appeared at 1,710 cm\(^{-1}\) when this substance was treated with dilute hydrochloric acid in a solvent mixture of ethanol and chloroform.

**DISCUSSION**

It is known that macrocyclic tetrapyrroles, porphyrins and their analogous compounds have an intense absorption band at about 400 m\(\mu\) (the Soret band\(^{4,5}\)), while bile pigments in which the conjugation of tetrapyrroles is interrupted lack this characteristic absorption. It is also known that the squareplanar divalent metal complexes of porphyrins generally have a Soret band and two visible bands, usually referred to as the \(\alpha\) and \(\beta\) bands.\(^6\) On the basis of such facts, the three absorption bands identified in the sample VII (350, 595 and 645 m\(\mu\)) are presumed to correspond to the Soret band and to the \(\alpha\) and \(\beta\) bands of metalloporphyrins, respectively. This suggests that this substance has a metalloporphyrin-like structure in which the macrocycle may still be interrupted but a kind of ring configuration has been attained by coordination with Cu. On the other hand, the infrared spectrum of this compound shows that the propionic acid groups attached...
to the pyrrole nuclei remain as free acid forms unlike in the case of the complex of bilirubin with calcium. Therefore, Cu atom must be located in nitrogen atoms of two pyrroles as N-Cu-N. Moreover, the appearance of a sharp absorption at 1,100 cm⁻¹ and decrease of the ν\textsubscript{NH} absorption (3,430 cm⁻¹) occurred in complex formation of bilirubin with Cu. Since these phenomena are known to be specific to porphyrin derivatives, it is suggested that the sample VIII is a metalloporphyrin-like bilirubin-Cu complex. From these results it is clear that bilirubin, produced by oxidative cleavage of the α-methine bridge (-CH=) of the porphyrin macrocycle, forms a ring structure by complex formation.

The present result disagrees with the opinion of Bentley that bilirubin, unlike biliverdin and mesobiliviolin, does not form a metal complex in spite of the presence of replaceable hydrogen atoms and nitrogen atoms available for coordination. He proposes the structure of a biliverdin-metal complex as shown in Fig. 9 in which one of the two terminal pyrrole nuclei is converted into the lactam form and a stable hydrogen bond forms between the lactam carbonyl and the hydroxyl group of hydroxypyrroline. As an evidence for such a structure, it has been claimed that a methoxyl derivative of biliverdin (terminal hydroxyl groups substituted by -OCH₃) does not form a metal complex. But, Fischer and his coworkers synthesized the Zn-complexes of bilirubin and biliverdin derivatives in which hydroxyl groups had been replaced by methoxyl and the propionic acid groups esterified. Since the prototropy takes place in bilirubin (Fig. 10), it seems that this compound is able to form metalloporphyrin-like complexes.

In the synthesis of bilirubin-Cu complex in the presence of a great excess of Cu++, it was observed that the product (λ\textsubscript{max}=595 mM and 645 mM) gradually changed to a purple substance (λ\textsubscript{max}=495 mM). This is apparently due to oxidative effect of Cu++. However, the stability of the bilirubin-Cu complex was satisfactory in solutions containing a small excess of Cu++ or after removal of excess Cu++ by washing with water. Lemberg reported a similar phenomenon that a mesobiliverdin-Cu complex was oxidized into a mesobilivioilin-Cu complex.

![Fig. 9. Structure of a Zn-complex of bilirubin (Bentley).](image-url)
in the presence of excess Cu++. The infrared spectrum of the other bilirubin complex, the sample IX, which was produced by reaction of cupric chloride with bilirubin dissolving in an aqueous solution containing sodium hydroxide, was different from that of the above-mentioned bilirubin-Cu complex. From the infrared spectrum of the sample IX, it is observed that unlike the latter complex or free bilirubin, this substance has carboxylate groups instead of free carboxylic groups. This suggests that Cu coordinates to nitrogen atoms of sodium bilirubinate.

The bilirubin-Cu complex and the bilirubin-Na-Cu complex prepared in vitro in this experiment are dark blue or black in color and quite resemble the pigments of genuine black colored gallstones. Of course, until identification of these complexes in the bile or gallstone, it remains obscure whether such bilirubin-metal complexes are produced in vivo and contribute toward black coloration of the gallstone. However, it is known that copper is a common component of bile and bile also includes other metal elements such as iron, magnesium and manganese, and these facts suggest the possibility of formation in vivo of metal complexes of bile pigments. Such possibilities are also supported by a number of reports which described the presence of bile pigments containing metal elements in vivo: for instance, a “green pigment” that is an iron containing tetrapyrrrole compound as an intermediate product during transformation of hemin.
to bilirubin,\textsuperscript{12} and an unusual derivative of bilirubin associated with iron.\textsuperscript{15}

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