A New Method of Blood Galactose Estimation for Mass Screening of Galactosemia

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FUJIMURA, Y., KAWAMURA, M. and NARUSE, H. A New Method of Blood Galactose Estimation for Mass Screening of Galactosemia. Tohoku J. exp. Med., 1981, 133 (4), 371-378 — A new method for quantitative determination of galactose in blood by fluorescence of NADH was described. The assay system consisted of β-galactose dehydrogenase, NAD, buffer and a denatured blood disc (3 mm diameter), and the reaction was carried out for 1 hr at 37°C. Denaturation of hemoglobin was accomplished by exposing the blood disc to a vapor of formic acid in an air-tight container; this procedure completely eliminated false positive cases of galactosemia. This method can be applied in a wide range of galactose concentration from low (0 mg%) to high levels (200-1,000 mg%) with the accuracy of 8.0±0.3 mg% from a coefficient of variation of 3.5%. Semi-quantitative assay was also possible by using a spot test like Beutler’s method. The galactose content in one disc paper (3 mm diameter) of blood containing 5 mg% galactose is approximately 0.1 μg. The newly developed method is satisfactorily applicable in neonatal mass screening and clinical cases. — galactose; mass screening of galactosemia; β-galactose dehydrogenase; fluorometry

Gal occupies an important place in carbohydrate metabolism, and it serves as an efficient metabolite in various hepatic diseases, galactosemia and galactokinase deficiency. Notably, Rommel’s method (Rommel et al. 1968) has thus far been accepted as a workable technique in Gal determination. The technique, however, has proved inadequate for mass screening in that it requires perchloric acid for deproteinization and a spectrophotometer. Moreover, it requires relatively large quantities of blood specimen and is time-consuming. In the field of galactosemia screening, Beutler’s method (Beutler and Baluda 1966) and Paigen’s method modified by Guthrie (personal communication) are widely used currently, but both of them suffer from defects. In the former, a high incidence of false positive of galactosemia occurs which results from inactivation of the enzyme during mailing

Received for publication, April 12, 1980.
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The following abbreviations were used: GADH, β-galactose dehydrogenase; NAD, nicotine adenine dinucleotide; NADH, reduced nicotine adenine dinucleotide; Gal, galactose; Gal-1-P, galactose-1-phosphate; UDP-Gal uridine diphosphate galactose.
and preservation of test materials under a high temperature and humidity.

On the other hand, we found that Paigen's method determines not only Gal but the related compounds such as UDP-Gal, Gal-1-P and lactose (unpublished). Thus, development of a new technique to determine Gal exclusively has been awaited. Previously we reported (Fujimura et al. 1977a, b) a semi-quantitative spot test of galactose which is a modification of the method of Rommel et al. and uses approximately one-hundredth the volume of blood sample required in the original procedure. The minimum levels of Gal which could be determined by our method ranged 10 to 20 mg%. When combined with the denaturation of hemoglobin, the range of the newly developed assay expanded to cover the range from 6–8 mg% to 1,000 mg%. Then, this technique was further improved so as to be applicable to mass screening for galactosemia using only a single 3-mm Guthrie test filter paper which was impregnated with a minute amount of blood specimen from the neonate or adult. Moreover, the assay system, which is based on fluorospectrophotometry, quantitatively eliminates false positive reactions frequently encountered by Beutler's method and the Paigen's method modified by Guthrie.

The present paper describes the new method devised by the authors and some results obtained by the application of this technique to neonatal mass screening and clinical cases.

**Materials and Methods**

*Specimen.* Standard paper discs for blood galactose assay: Fresh blood samples collected at fasting before meal were left in a refrigerator (0–4°C) overnight, and hematocrit was adjusted to 50–60%. Next, given amounts of Gal were added into the blood, and each of the following blood specimens containing no Gal added (nearly 0 mg% by galactose kit, Boehringer Mannheim), 2, 4, 6, 8, 10, 12, 20, 30, 50, 75, 100, 200, 300, 500, 700 and 1,000 mg% of Gal were dripped onto Guthrie's test paper (made by Daiichi Kagaku Co.). The samples were then dried in air to prepare the standard samples.

General test materials: Neonatal blood filter papers (blood samples should not be autoclaved) sent in for Guthrie's test were used.

False positive cases by Beutler's test: Thirty blood samples selected from false positive cases by mass screening tests for galactosemia, with the procedure of Beutler and Baluda (1966), were studied.

Positive and false positive cases by Paigen's test: Thirty-five blood samples positive or false positive for Gal by Paigen's test were studied. The Gal levels by Paigen's test were over 20 mg% in 3 blood samples, 12 mg% in 3, 12 to 8 mg% in 5, 8 mg% in 4, and 6 to 8 mg% in 20.

**Hemoglobin denaturation treatment.** Each sample filter paper obtained as above was cut into discs of 3 mm diameter to be placed in the hemagglutination tray for TPHA (non-fluorescent, made by Tominaga Co.) or the microtiter trays (either Cook's V type or disposable tray V type, 125 mm × 14 mm, 96 holes). The discs were kept free for 1 hr in an air-tight glass container saturated with formic acid vapor. The trays were taken out, and placed in a desiccator so that acid could be removed. Then, the desiccator was evacuated by suction and placed in an incubator kept at 37°C for a few hr to overnight for removal of acid. Samples thus obtained did not liberate hemoglobin into solution at all.

**Principle of the test.** The test is a modification of the method of Rommel et al. (1968), utilizing GADH which reacts specifically with Gal, as previously described (Fujimura et al. 1977a, b).
Quantitative assay. Reaction system. Reagent system 1: This represents the complete reagent system and consists of 10 µl of 13 mM NAD (10 mg/ml from Boehringer), 10 µl of 0.35 M phosphate buffer (pH 8.0), 10 µl of a 1:100 dilution of 5 mg/ml GADH (Boehringer, code No. 104981) solution and one blood-impregnated 3 mm paper disc which has been treated for hemoglobin denaturation. Reagent system 2: Similar to Reagent system 1 except that 10 µl of distilled water were used in place of GADH. Reagent system 3: Similar to Reagent system 1 except that 10 µl of distilled water were used in place of NAD. Reagent system 4: Similar to Reagent system 1 except that the blood-impregnated paper disc (specimen) was omitted. These reagent systems were placed in small test tubes, respectively, stoppered, and incubated at 37°C for 1 hr.

Galactose assay by fluorospectrometry. After 1 hr of incubation at 37°C, 3 ml of distilled water were added to each tube to terminate the reaction and fluorescent transmittance of each mixture at excitation wavelength of 340 nm and emission wavelength of 450 nm was recorded by a Shimazu fluorospectrometer model 502 or a Hitachi fluorospectrometer model 204, with the reference reading for 1.17 µg quinine sulfate (Katayama Chemicals; 8.4 mg in 10,000 ml of 0.01 M phosphate buffer, pH 7.4) adjusted to 100% transmittance. For estimation of low concentration of Gal (0-30 mg%), the fluorometry was made using a quinine sulfate reference of lower concentration (0.29 µM or 8.4 mg in 40,000 ml of 0.01 M phosphate buffer, pH 7.4) adjusted to 100% transmittance.

Concentration and activity of GADH. GADH stock solution was diluted to 1:10, 1:100, 1:200, 1:500, 1:1,000, 1:2,000, 1:5,000, and 1:10,000 with 0.35 M phosphate buffer. The Gal assay was performed by using 10 µl of each of the dilution enzyme solutions.

Time course of reaction. Blood Gal assays were carried out in the complete reagent system with a 3-mm 100 mg% Gal blood discs, and fluorescence was read at 10, 20, 30, 45, 60, 90 and 120 min of incubation. 95% of maximal activity was obtained at 10 min and 100% at 20 min of incubation.

Error in Gal assay by the present method. Eight pieces of discs (3 mm diameter) were, at random, punched out from a standard blood paper containing 5 mM Gal, and treated for denaturation of hemoglobin. Gal assays were performed with these discs and fluorometric transmittance of the mean Gal concentration X, standard deviation s.D. and coefficient of variation C.V. (%) = s.D./X were calculated.

Semiquantitative spot test. A total of 30 µl of the reaction mixture composed of 10 µl of 13 mM NAD, 10 µl of buffer (pH 6.0-8.0) and 10 µl of 300- to 500-fold diluent of 5 mg/ml stock solution of GADH, was placed in a microtiter tray where a blood disc (denatured) had been placed. The top of the tray was sealed with tape to prevent drying, and the mixture was allowed to react for 1 hr at 37°C. Then, the mixture was spotted on the Watmann No. 1 filter paper as in Beutler's method. Following air dry, the fluorescence strength was determined under irradiation of excitation light at 365 nm or 253 nm or both, using a fluorescence detector, Superlight SL-DI (from Nikko Sekiei K.K.) and using the standard fluorescence references consisting of the various ratios of NADH/NAD mixture (Fujimura et al. 1976).

RESULTS

Hemoglobin denaturation by formic acid or acetic acid

Since hemoglobin interferes with the determination of NADH fluorescence, factors leading to denaturation of hemoglobin were studied. Hemoglobin on the disc was exposed to vapor of formic acid at various saturations and found to be completely denatured when exposed for 1 hr to vapors at higher than 33%. Furthermore, when the discs with denatured hemoglobin as processed as described in Methods section were incubated with the assay mixture at 37°C, the final pH remained to be 7.6-8.0, showing no appreciable decline. Acetic acid was not so good as formic acid since it was not easy to completely eliminate remaining acid.
Preparation of calibration curves with standard blood discs for Gal assay

The fluorometric estimation of blood galactose was conducted with Reagent systems 1 to 4, using hemoglobin-denatured discs containing 0, 2, 4, 6, 8, 10, 12, 20, 30, 50, 75 and 100 mg% Gal. Reagent systems 2, 3 and 4 served as the controls. As can be seen in Fig. 1-A, a linear plot of fluorescent transmittance was obtained against the Gal concentrations from 0 to 100 mg%. It was possible to scale up the difference of system 1 minus system 2 in 0–30 mg% of Gal by adjusting the reference with 0.29 μM quinine sulfate to 100% transmittance (Fig. 1-B).

Concentration and activity of GADH

In Fig. 2 are shown the results obtained with various concentrations of GADH at a fixed concentration (75 mg%) of Gal on the blood disc. The enzymatic activity fell drastically at dilutions beyond 1:100, and the values at 1:100 and 1:1000 dilutions of the enzyme were somewhat higher than the activity obtained for the undiluted enzyme stock solution. When the assay was carried out with blood-impregnated paper discs containing 20 mg% Gal, however, the enzyme activity of GADH was not appreciably decreased even at the 1:1000 dilution. The activity of GADH was not significantly lost when the disc preparation was left at 0–4°C for one week (Fig. 2).
Error in assay

The extent of error in the measurement of blood Gal content by the present assay method was examined by repeating the assay with the same specimens (Table 1). A mean fluorescent transmittance \( \bar{X} \) of 59.5 was obtained for eight discs impregnated with blood containing 100 mg% Gal. The standard deviation (s.d.) was calculated to be 2.1 and the coefficient of variation \( C.V. = \frac{s.d.}{\bar{X}} \) was 3.5%; hence the measured values being within a 5 per cent range of error. This method was in the accuracy and the sensitivity of 8.0±0.3 mg% of Gal. The results indicate usefulness of the assay method.

Blood Gal assay in false positive cases by Beutler's test

Blood samples from 30 patients found to be false positive for galactosemia by...
Beutler's test were assayed by the present method. As shown in Fig. 3, the blood Gal level was less than 4 mg%, being by far lower than the lower limit of the positive range, in all the 30 cases studied.

In the Paigen-Guthrie method, determination involved not only Gal but Gal-1-P, UDP-Gal and lactose in some cases. With this in mind, some of the cases proved as positive or false-positive by Paigen's test were studied by our method. In three cases of liver diseases in which the blood Gal value was over 20 mg% by Paigen's test, the assay by our method revealed concentrations of 10.1, 25.7 and 50.4 mg%, respectively, as shown in Table 2. Three patients showing blood Gal levels of 12 mg% and five patients showing 8 to 12 mg% by Paigen's test were all shown to be entirely normal with values between 2 and 4 mg% by the present test. It was found that the blood Gal content returned to normal levels in all these cases sent in after a few weeks by re-examination of the Paigen's method. In 4 cases displaying 8 mg% of Gal in the blood and in 20 cases showing 6 to 8 mg%, the tests by the author's method were all normal.
DISCUSSION

The new method of Gal quantification which we have developed is characterized by simplicity and expediency of the procedure that is comparable to Beutler's method (Beulter and Baluda 1966); the method uses Guthrie's test filter paper so that blood can be sucked and the blood test material can be sent by mail for investigation.

Another advantage inherent in authors' technique is that the technique is free from a high incidence of false positive cases of galactosemia due to enzymatic inactivation which might occur during mailing or preservation of blood. Still another merit may come from expediency which allows direct measurement of Gal itself. The new technique is suitable and valuable as the mass screening method because the amount of blood required is so small; about 1/100 that needed in Rommel's technique (Rommel et al. 1968). Tengström and Wranne (1968) and Tengström (1969) introduced an assay system with galactose oxidase (de Verdier and Hjelm, 1962) using paper discs as the carrier for blood specimen for estimation of blood Gal by an autoanalyzer. However, their method requires a disc diameter of not less than 14 mm which is practically unsuitable for mass screening by the Guthrie test. Grenier and Laberge (1973) reported a mass screening method of fluorometric determination of NADH for estimation of Gal levels with GADH using blood-impregnated paper discs of 7.94 mm in diameter. This size of the disc seems to be still oversized for mass screening. It has been pointed out that assays by the procedure of Grenier and Laberge (1973) are unreliable unless the test is repeated at least twice on the same specimen; this is probably because the measurement is interfered with by a high degree of nonspecific fluorescence arising from hemoglobin and reagents. We have experienced a similar phenomenon in the initial state of development of the present assay system.

Statistical data analysis revealed a remarkably low coefficient variation, 3.5%, in the estimation of error in the assay by the present method, providing evidence for potential usefulness of this assay method.

The enzyme GADH exhibited sufficiently high activities at dilutions up to approximately 1:1,000 in phosphate buffer when assessed using a 5 mg/ml enzyme stock solution. Therefore, we decided to use a 1:100 dilution of the stock solution in the assay system in anticipation for adequate margins of safety. When the stock solution was tested without dilution, the activity obtained was significantly lower than those obtained with the 1:10 and 1:100 dilutions (Fig. 2). Probably this is due to inhibition caused by high contents of ammonium sulfate in the stock solution.

By using our method, several blood samples diagnosed as false positive by Beutler's test were shown to be normal with Gal values of less than 4 mg%. The majority of cases in which Paigen's tests were false positive for galactosemia were also found to be normal by both the present and clinical methods. In the Gal tolerance test, too, Paigen's method appears to produce a higher value than the real value quantified by Rommel's method, whereas the test by the authors'
method yields value which is close to that obtained by the Rommel's. Thus, our method may be valuable for eliminating such frequently encountered false positive reactions in mass screening. The incidence of "false positive" in the assay by the present method may probably be at a level lower than 1/6000, this is reported by Grenier and Laberge (1973).

References