Annual Summary of External Quality Assurance Survey for G6PD Blood Quantitative Test in Philippines (2024)

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1. Introduction

Preventive Medicine Foundation Quality Assurance Program Center (PMF QAP Center) has been providing "EQA Program for Glucose-6-Phosphate Dehydrogenase (G6PD) Blood Quantitative Test "for G6PD confirmatory laboratories in Taiwan since 1988. In cooperation with PMF QAP Center, the Newborn Screening Reference Center (NSRC) Manila, has adopted this EQA program for the newborn screening referral hospitals in the Philippines since 2009. This EQA program has been officially accredited by Taiwan Accreditation Foundation (TAF, a member of ILAC Mutual Recognition Arrangement Signatories) for conformity to international standard ISO/IEC 17043:2010 since 2017 (Accreditation No.: P016) and was extended the certification on December 30, 2022.



Figure 1. Certification Accreditation (Certificate No: P016-221230)

2. Participants

Thirty-two G6PD confirmatory laboratories have participated in the EQA program in 2024. (Fig. 2 and 3) There is one more participant than last year.



Fig. 2. Distribution of participating laboratories in Philippines.

(• Participating laboratory, n=32)

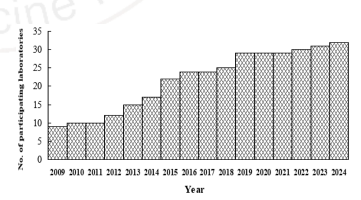


Fig. 3. Number of participating laboratories in Philippines (2009 ~ 2024)

- 3. Quality Control Sample (QC Sample)
 - 3.1 Three QC samples were used in each survey.
 - 3.2 The QC samples were lyophilized hemolysate prepared from human red blood cells with no extra G6PD added. (Taiwan IVD Register. No.: MOHW-MD-(I)-No.004851)
 - 3.3 The homogeneity and stability of QC samples conform to the requirements of the international standard ISO/IEC 17043:2010.

4. Surveys

4.1 There were three EQA surveys performed in 2024. (Table 1).

Table 1. 2024 EQA survey schedule

No.	Survey No	Shipping Date*	Reporting Deadline*	Survey Result Released*
1	RH2024-01	02/26	03/04	03/07
2	RH2024-02	05/27	06/03	06/11
3	RH2024-03	09/23	09/30	10/09

^{*} Date: MM/DD

- 4.2 In 2024, 93 sets of QC samples were sent to participants, 93 (100%) reports were returned.
- 4.3 Most laboratories received the QC samples within $1 \sim 3$ days (median = 1 days) after samples were sent out when the survey started.
- 4.4 More than 32.3% of the participants, which were less than previous years, reported that dry ice was sublime completely when they received the QC samples. However, there is no unsatisfactory report was found due to the shipping temperature.
- 4.5 The reports returned time was between 1 and 7 days (median = 6 days) after the survey started. All reports were returned within the target time (7 calendar days after the survey started).
- 4.6 The survey summary reports were released on the website between 3 and 6 working days after reporting deadline, which conformed to the target time (7 working days after reporting deadline).

5. Evaluation Criteria

- 5.1 The assigned value (Xa) = the median of all the results reported for this QC sample.
- 5.2 SD for proficiency assessment (σ_p) = 7% × Xa ; but when Xa < 2.9 U/g Hb , σ_p = 0.2 U/g Hb.

- 5.3 Adjusted SD for proficiency assessment (σ_p') = ($\sigma_p^2 + u_{Xa}^2$)^{1/2}. σ_p' is used for proficiency assessment when uncertainty of the assigned value $u_{Xa} \ge 0.3\sigma_p$.
- 5.4 z score = D / σ_p ; D = X Xa, $\sigma_p = SD$ for proficiency assessment. When adjusted SD (σ_p) is used for proficiency assessment, z score = D / σ_p .
- 5.5 The evaluation criteria for measurement result of "each QC sample ":
 - a) Acceptable : $|z| \le 2$;
 - b) Caution : 2 < |z| < 3;
 - c) Unsatisfactory : $|z| \ge 3$.
- 5.6 The performance evaluation criteria for participant survey report:
 - a) Acceptable : all results |z| < 3 and more than one result $|z| \le 2$;
 - b) Acceptable with Caution: only one result $|z| \ge 3$ (other results |z| < 3) or more than one result 2 < |z| < 3;
 - c) Unsatisfactory: more than one result $|z| \ge 3$.

6. Result of EQA surveys

- 6.1 Three EQA surveys for G6PD quantitative test were performed in 2024.
 - a) Ninety-one (97.8%) reports were "Acceptable";
 - b) One (1.1%) reports was "Acceptable with Caution";
 - c) One (1.1%) reports was "Unsatisfactory";
 - d) The "Acceptable with caution" rate and "Unsatisfactory" rate of the report were similar to recent years (Fig. 4).

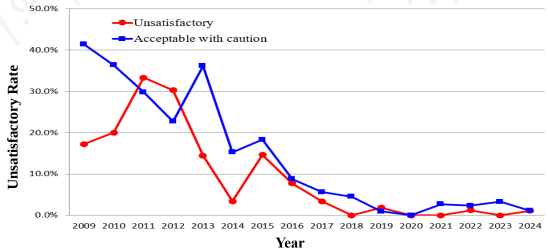


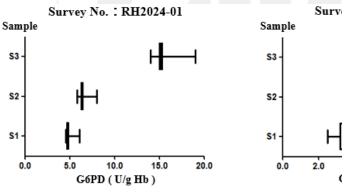
Fig. 4. "Unsatisfactory" and "acceptable with caution" rates of the survey reports ($2009 \sim 2024$).

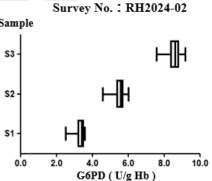
6.2 The G6PD activity (assigned value; Xa) of 9 QC samples used in 3 surveys (RH2024-01, RH2024-02 and RH2024-03) were between 3.4 and 16.2 U/g Hb (Table 2). The distributions of G6PD test results reported in each survey were shown in Fig. 5.

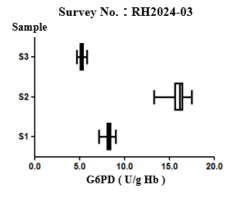
Table 2. Summary	y of the survey	v results of each	OC san	nnles in 2024
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Survey	Sample	N	Median* (Xa)	Mean*	SD*	CV%	Min*	Max*
	S1	31	4.8	4.8	0.17	3.5%	4.6	6.1
RH2024-01	S2	31	6.4	6.4	0.21	3.3%	5.8	8.0
	S3	31	15.1	15.2	0.45	3.0%	14.0	19.0
	S1	30	3.4	3.4	0.18	5.3%	2.5	3.6
RH2024-02	S2	30	5.6	5.6	0.28	5.0%	4.6	6.0
	S3	30	8.6	8.6	0.32	3.7%	7.6	9.2
1	S1	32	8.3	8.3	0.29	3.5%	7.2	9.0
RH2024-03	S2	32	16.2	16.0	0.62	3.9%	13.3	17.5
1	S3	32	5.3	5.3	0.22	4.2%	4.7	5.8

^{*}U/g Hb







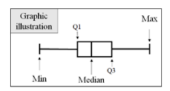


Fig. 5. Distribution of G6PD test results of each survey in 2024

6.3 Between Laboratory Variations

All the interlaboratory C.V. of QC samples in 2024 were all lower than 10% ($3.0 \sim 5.3\%$; Table 2), which were better than 2023 ($3.4 \sim 7.0\%$).

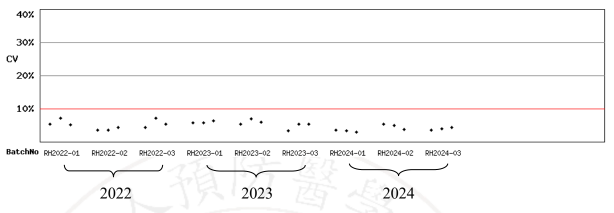


Fig. 5. Inter laboratory C.V. vs. surveys ($2022 \sim 2024$)

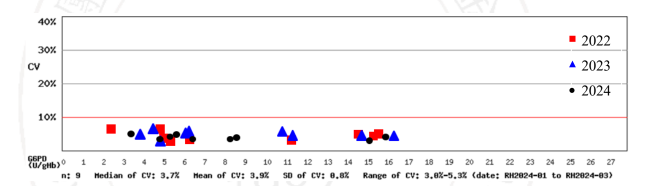


Fig. 6. Inter laboratory C.V. vs. G6PD activities ($2022 \sim 2024$)

6.4 All the results of EQA surveys for G6PD blood quantitative test in 2024 were posted on website:

< https://g6pd.qap.tw/113rep-phi.htm >

The content of the website including following parts:

- a) Summary report of G6PD and Hemoglobin (Hb) quantitative test results of each survey;
- b) Long-term observation of EQA survey results for G6PD quantitative test;
- c) Distribution of G6PD test results of each survey;
- d) Distribution of Hb test results of each survey;
- e) Deviation graphs (z score, D%, SDI) for individual laboratory;
- f) Intermediate measurement precision of G6PD quantitative test;
- g) Repeatability of G6PD Quantitative Test.

7. Conclusion of the customer satisfaction survey

In the 2024 customer satisfaction survey, the return rate was 55% (17/32). Among the returned questionnaires, 82% of the participants rate "Excellent" performance and 18% of the participants rate "Great" performance in overall satisfaction.

