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Development of Arduino Uno-Based TCS3200 Color Sensor and Its Application on the Determination of Rhodamine B Level in Syrup

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Abstract: The use of the notorious synthetic dye, rhodamine B, in food and beverage products has been widely reported. This application urges the need to develop an analytical method that can provide reliable rhodamine B data with an easy operational technique. Therefore, this research is aimed to develop an Arduino Uno-based TCS3200 color sensor and study its application to determine rhodamine B levels in syrup. The design of the analytical instrument included TCS3200, an Arduino Uno microcomputer, an Integrated Development Environment (IDE) software, a black box container, and a 24×2 matrix display screen, where samples were prepared via absorption using wool thread. With a linear range of 1–20 mg/L, our proposed colorimetric sensor had recoveries of 96.25–110.3%, which was better compared to that was obtained from the UV-vis (81.8– 100.6%) method. The detection and quantification limits of the sensor were 2.766 and 8.383 mg/L, respectively. The syrup samples used in this study were purchased from the local stores in Banda Aceh. Based on the proposed TCS3200 color sensor, the highest rhodamine B concentration from the syrup sample was 16.74 mg/L. The t-test analysis in this study revealed that the Rhodamine B levels quantified using the newly developed TCS3200 color sensor were not statistically or significantly different from the UV-Vis spectrophotometer method.

Keywords: color sensor; TCS3200; rhodamine B; Arduino Uno; Zn(CNS)₂; IDE software

INTRODUCTION

As a form of consumer protection efforts, sensor technology for food or beverage products has been developed intensely [1-3]. For example, Fourier Transform Infrared (FTIR) spectroscopy has been employed to separate halal and non-halal meatballs [4]. The color spectroscopy method has also reported the detection and analysis of synthetic dye rhodamine B in chili powder [5]. Dyes used in food products are derived from natural and artificial chemicals. Synthetic dyes are widely used because it is more cost-efficient and available. Additionally, the synthetic dye may give a brighter color to the food or beverage product [6].

As one of 30 synthetic dyes available, Rhodamine B is considered as a dangerous dye, where its use in food or beverage products has been prohibited. Nevertheless, Rhodamine B is often used in processing industries, papers, and fabrics [7-8]. Moreover, it could be employed as a ligand to bind metal ions [9]. However, in Indonesia, rhodamine B is still very popular as a food coloring agent, including in iced syrup. The syrup is intentionally added with rhodamine B, so the products obtain a more attractive appearance [10]. Methods that have been previously developed to identify rhodamine B in food ingredients include thinlayer chromatography (TLC) [11], voltammetry [12], and the standard method using UV-Vis spectrophotometer (the best option for identifying compounds with color). However, UV-Vis spectrophotometry has several drawbacks; not portable, complicated, and expensive [13]. Hence, this study tried to overcome the stated drawbacks by developing a simple measurement method using a portable sensor.

The sensor system developed in the present work was based on the TCS3200 color sensor constructed by the console to overcome external noise and program library modification [14]. The TCS3200 color sensor has been widely reported for different applications, including measuring levels of cyanide [14], nitrogen [15], and heavy metals [16]. In the case of colorimetric sensors, analytes should first be reacted with a complexing agent to cause a color change [17-18]. In this study, the sensor detects color degradation from tissue paper that has been spiked with reagents, so its sensitivity is specifically improved for rhodamine B analysis. The reagent used was $Zn(CNS)_2$, which can cause a color change from red to purple due to the formation of the rhodamine B-Zn-thiocyanate ((RhB) complex $2Zn(CNS)_4$) [19].

The color intensity contributed by the presence of rhodamine B was converted through the sensor output pin in the form of a square signal in which its frequency depends on the concentration. The box's signal with varied frequency was then processed using a microcontroller on Arduino Uno. In this processing, four filters were used, namely green, blue, red, and *no filter*. In this case, *no filter* was excluded because the three parameters were sufficient to represent the color degradation of rhodamine B in the sample [20]. Filter settings were performed by providing low and high logic in the Arduino IDE program, following the reported study [21].

The distance between the sample and the 8×8 diode array was set at 3 cm, following the sensor system's geometry. The console's color was made black so the color could be absorbed fully, and influence from the degradation of colors that enter the diode array could be avoided. After obtaining the concentration of rhodamine B using the Arduino Uno-based TCS3200 color sensor, the results were compared with the standard UV-Vis spectrophotometric method. Finally, the analysis results were compared to obtain the data on sensitivity and accuracy of the newly developed sensor [22].

EXPERIMENTAL SECTION

Materials

UV-Vis The materials used were а spectrophotometer (Thermo Fisher Scientific, Selangor Malaysia), a color sensor TCS3200 (ICTAOS/AMS), a console, and an Arduino Uno (wavgat). Syrup samples tested for rhodamine B levels were procured from local stores in Banda Aceh. The standard rhodamine B was purchased from The National Agency of Drug and Food Control of Indonesia (BPOM RI). All other chemicals used, i.e. NH4OH, NaOH, HCl, C2H5OH, CH3CO2Na, ZnCl₂, CH₃CO₂H, and KCNS, were obtained from Merck (Selangor, Malaysia) in analytical grade.

Hardware Design

The hardware design was initiated by developing a console for the TCS3200 color sensor, then connecting the output port of the color sensor via a jumper cable to the Arduino Uno microcomputer to process frequency data and convert it into 8-bit RGB digital data. There were 256 color digit variations for each RGB color component that was sortable and distinguishable by the processing. These color digit variations were displayed on the computer screen and converted to reduce color variations. These color variations were also recorded in .xls format (MS Excel) (Fig. 1).

Development of the TCS3200 Sensor Console

The TCS3200 console sensor was designed in black to absorb all color wavelengths. The distance between the diode array and the color object was 3 cm. The console was arranged in such a way that light from outside could not enter. The TCS3200 sensor was positioned opposite the color sample, which was absorbed into a filter paper. Four LED units with white



Fig 1. Schematic diagram of hardware design

wavelengths would hit the filter paper, and the intensity light reflected the diode array following the color intensity of the sample.

Software Design

Construction of the software design was initiated with a blink test on the Arduino Uno system to determine the response and performance of the microcomputer. The software used was Arduino IDE with available opensource libraries – C programming language. The program library was modified to enable the required color filters, Arduino Uno pins, the required display format, and data storage mode (Fig. 2).

Rhodamine Analysis Using TCS3200 Color Sensor

Construction of the calibration curve for rhodamine B

Briefly, the rhodamine B solution was added with 3.0 mL of Zn-thiocyanate. Then, the standard solution $(RhB)_2$ -Zn $(CNS)_4$ with different concentrations measured the RGB value with the TCS3200 color sensor and absorbance with UV-Vis at the maximum wavelength obtained. The solution was prepared with 100 mg/L rhodamine B as the stock solution, which was then diluted using distilled water into standard solutions with varying concentrations ranging from 1 to 20 mg/L. These solutions were prepared to determine the maximum wavelength of rhodamine B and as a database for the TCS3200 color sensor. Following that, a solution of 1 mL ZnCl₂ 2 M and 2 mL KCNS 2 M as a reagent was made to detect the presence of rhodamine B, as suggested by a previous report [23].



Fig 2. Display of the Arduino IDE Software main menu

Determination of rhodamine B level using the TCS3200 color sensor

The standard curve of $(RhB)_2$ -Zn $(CNS)_4$ was obtained by measuring the RGB values of the standard solution $(RhB)_2$ -Zn $(CNS)_4$ using the TCS3200 sensor. The concentration of rhodamine B used was 1 to 20 mg/L, which were priorly reacted with reagents. Measurements were carried out three times, and the concentration was averaged. Thereafter, RGB values were converted into a color index, namely Hue, Intensity, and Saturation (HIS). Conversion of RGB values to HIS values was carried out using the following Equations.

Red color index
$$(I_R) = \frac{R}{R+G+B}$$
 (1)

(3)

Green color index (I_G) =
$$\frac{G}{R+G+B}$$
 (2)

Blue color index $(I_B) = \frac{B}{R+G+B}$

The HIS color model was designed to resemble the perception of human vision, while the RGB values resembled the image of the display system [20]. The results of the calculation of the HIS value were then plotted as the dependent variable (*y*-axis) to the variation of concentration (RhB)₂-Zn(CNS)₄ (*x*-axis).

TCS3200 color sensor method validation

Method validation included accuracy, precision, sensitivity, and linearity, which were conducted based on the suggestion from a previous report [22].

Syrup sample preparation

Samples of commercial red syrup were purchased from local stores in Banda Aceh. Each sample (10 mL) was taken and put into an Erlenmeyer which was subsequently mixed in 20 mL of 25% ammonia solution (dissolved in 70% ethanol) for 24 h and evaporated on a hot plate. The evaporation residue was dissolved in 10 mL distilled water containing acid (10 mL distilled water and 5 mL acetic acid 10%). Wool thread (15 cm) was dipped into the acid solution and simmered for 10 min until the dye colors appeared on the wool thread, then lifted. The wool thread was then washed with distilled water, and the wool thread was dissolved in ethanol 70% and heated to a boil (Fig. 3). This solution was used as the sample, per suggestion by a published work [24]. The wool thread was used to extract



Fig 3. The extraction of rhodamine B from commercial red syrups using wool thread. Wool thread was dipped into the dissolved syrup residue for 10 min (a). Rhodamine B-containing wool thread before re-immersed to ethanol 70% and boiled (b)

rhodamine B-containing samples in an acidic environment. A comparative study has reported that wool thread has the highest dye adsorption as compared with silk and nylon [25]. Adsorption of dye analyte in wool thread is determined by its O- and N-containing functional groups, which has been reported in many published papers [26-28]. The dyed wool was then immersed in ethanol 70% and boiled until its original color returned. The obtained solution was analyzed for its rhodamine B levels using the TCS3200 color sensor and a reference method – UV-Vis spectrophotometry.

Quantitative Analysis

The prepared sample was added with Znthiocyanate and then dipped in filter paper. Rhodamine B levels were measured using the TCS3200 color sensor [29]. The concentration was obtained based on the linear equation obtained from the calibration curve.

Method Comparison using Two-Way t-Test

Results of the samples between the TCS3200 color sensor and the UV-Vis spectrophotometry method were compared [30]. In addition, a two-way *t*-test was carried out to see the significance between the newly studied TCS3200 color sensor method and the reference method using UV-Vis spectroscopy by calculating the *t* value for each method and then comparing it with the t_{theoretical}.

RESULTS AND DISCUSSION

Maximum λ of Rhodamine B Complex

The complex $(RhB)_2$ -Zn $(CNS)_4$ was produced to give rhodamine B a specific color, allowing easier analysis. The solution of rhodamine B, which was initially red, turned to purple and was then measured using a UV-Vis spectrophotometer at a wavelength ranging from 574 to 600 nm. The UV-vis absorbance corresponding to the $(RhB)_2$ -Zn $(CNS)_4$ complex scanned from 574 to 600 nm is presented in Fig. 4.

Based on the measurement results, the UV-Vis spectrometer spectrum of $(RhB)_2$ -Zn $(CNS)_4$ showed a maximum absorption (0.442 au) at a wavelength of 590 nm. The difference in wavelength between rhodamine B and $(RhB)_2$ -Zn $(CNS)_4$ is due to a shift in



Fig 4. UV-Vis spectrometer spectrum of (RhB)₂-Zn(CNS)₄ showing a maximum wavelength at 590 nm

wavelength towards the bathochromic direction caused by substitution, solvent effects, and the influence of the chromophore group [31]. The successful formation of the (RhB)₂-Zn(CNS)₄ complex was indicated by a color change from red to purple and a shift in wavelength. The equation for the reaction between rhodamine B and Zn(CNS)₂ can be seen in Fig. 5.

Based on the graph, we can see three regression equations obtained from each RGB index value, namely I_R y = 0.0028x + 0.3411; I_G y = 0.0032x + 0.3513 and I_B y = -0.0058x + 0.3059. The values of the determination coefficient (R^2) were 0.9792,0.9700, and 0.9729 respectively. The R index had the best determination coefficient (R^2) of 0.9792. Therefore, the regression

equation for the R index was used to determine the concentration of rhodamine B in the sample.

Measurement using UV-Vis Spectrophotometer

The standard curve of $(RhB)_2$ -Zn(CNS)₄ was measured at a wavelength of 590 nm by a UV-Vis spectrophotometer. The concentration of rhodamine B that was used ranged from 1 to 20 mg/L, which was priorly reacted with reagents. Measurements were carried out three times and averaged for each concentration. The absorbance measurements can be seen in Fig. 6. The regression equation y = 0.0023x +0.0773 had a determination coefficient (R²) of 0.9927. Hence, it can be concluded that the concentration was



directly proportional to the absorbance, meaning that the absorbance for the complex $(RhB)_2$ -Zn $(CNS)_4$ was dependent on rhodamine B concentration.

Method Validation

Accuracy

The accuracy of the proposed sensor method was based on the recovery (%), representing the value proximity of the standard concentration solution to the actual concentration. The concentrations of $(RhB)_{2}$ -Zn(CNS)₄ used were 1, 10, and 20 mg/L for the analysis with TCS3200 and UV-Vis color sensors. The actual concentration values and the percent recovery values from each method can be seen in Table 1. The recovery % calculation for the TCS3200 color sensor was still within the allowable error range of 90–110% [32]. However, at a concentration of 1 mg/L, UV-Vis had a recovery value below the permissible range (81.8%). Therefore, our proposed method was suggested to have better accuracy for determining rhodamine B levels at a low concentration (1 mg/L).

Precision

The precision was determined to see the proximity of the value changes in the repetition process. The precision value was derived from the standard curve with a respective concentration of $(RhB)_2$ -Zn(CNS)₄ (1, 10, and 20 mg/L), expressed by the variation coefficient (VC). The precision values for both methods based on intra-day and inter-day repetition are presented in Table 2. The variation coefficient value obtained by the two measurements increased with the decrement in the concentration of the standard solution. The method is accurate if it provides a variation coefficient value of less than 2% [32]. Nonetheless, inter-day repetition yielded higher variation coefficient, especially when rhodamine B concentration was 1 mg/L.

Linearity

Linearity is the functional area of sample measurement. The linearity of measurements using the TCS3200 color sensor and UV-Vis spectrophotometer for a concentration range of 1–20 mg/L is depicted in Fig. 5 and 6, respectively. Several studies used a



Table 1. Recovery percentages of TCS3200 sensor and UV-vis spectrophotometer

Concentration (mg/L)	Actual cor (mg	ncentration g/L)	Recovery (%)		
	TCS3200	UV-Vis	TCS3200	UV-Vis	
1	1.030	0.818	103.5	81.80	
10	11.03	10.06	110.3	100.6	
20	19.25	19.03	96.25	95.15	

[Phodomino B] (mg/I)	Intra-day variation coefficient (%)		Inter-day variation coefficient (%)	
[Kilodaninie D] (ilig/L)	TCS3200	UV-Vis	TCS3200	UV-Vis
1	0.291	0.721	7.966	8.563
10	0.268	0.521	1.294	1.664
20	0.253	0.357	0.509	0.851

 Table 2. VC values of TCS3200 sensor and UV-vis spectrophotometer obtained from intra-day and inter-day repetition

non-linear calibration curve because the sensor system formed an exponential response [33]. However, in the present study, the quantitative analysis was conducted based on linear regression.

Sensitivity

The sensitivity value is shown from the slope of the complex standard curve of (RhB)₂-Zn(CNS)₄ for each method. Based on the linear regression standard curve equation, the slope value for the TCS3200 color sensor measurement method was obtained from the regression equation y = 0.0028x + 0.3411, which was 0.0028. While the slope value for the UV-Vis spectrophotometer measurement method was obtained from the regression equation y = 0.0253x + 0.0773 is 0.0253. Based on the constructed standard curve, we calculated the limit of detection (LOD) by multiplying the standard deviation of response by 3.3 and dividing with the slope. Meanwhile, the limit of quantification (LOQ) could be obtained by multiplying the standard deviation of response by 10 and dividing it with the slope. The LOD obtained for the TCS3200 color sensor and UV-Vis spectrophotometer was 2.766 and 1.715 mg/L, respectively. These values explain why the inter-day precision for the 1 mg/L rhodamine B sample obtained for both methods exceeded the acceptable maximum variation coefficient (< 2%). As for the LOQ, the values reached 8.383 and 5.196 mg/L for the TCS3200 sensor and UV-Vis spectrophotometer, respectively. Lower LOD and LOQ in UV-Vis spectrophotometer suggest its superiority in comparison to the TCS3200 color sensor, in terms of sensitivity.

Quantitative Analysis Using the TCS3200 Color Sensor

Samples were measured using a series of tools that had been readily connected to the TCS3200 color sensor. The measurement was carried out by dipping the filter paper into the sample solution to which 3 mL of Zn(CNS)₂ reagent had been added, then dried and measured using the TCS3200 color sensor in dark conditions. Measurements were carried out three times on each sample with 3 cm-long distance between the sensor and the sample. Such distance was given to allow even distribution of the emitted light from four Light Emitting Diodes (LEDs) to the sample and the photodiode, in which the sample could emit a current proportional to the basic color of received light.

Table 3 shows that the RGB value obtained from each sample is a code to indicate a specific color. The HIS value in the table was obtained using Eq. (1-3). The I_R value was used to determine the concentration of rhodamine B in the sample because it had the best R² (0.9792) among the others (Fig. 6). The total concentration of rhodamine B obtained from the measurement using the TCS3200 color sensor based on the I_R value can be observed in Table 4, showing the concentration of each sample with five repetitions. The red index value obtained from Eq. (1) has the same function as the absorbance value, the dependent variable in determining the concentration. Therefore, the concentration of rhodamine B in the sample was calculated by substituting the red color index value of the sample into the standard curve regression equation $(RhB)_2$ -Zn $(CNS)_4$ R index.

Following the analysis, we found that samples A, B, and C contained rhodamine B with an average of 1.74, 16.74, and 5.10 mg/L, respectively. However, sample A had a rhodamine B concentration lower than the LOD of both the TCS3200 and UV-Vis spectrophotometer (2.766 and 1.715 mg/L, respectively). In this case, the response generated from sample A could not be differentiated from that of the blank standard. Hence, the presence of rhodamine B in sample A could not be

					1			
Repetition		RGB	Measurem	nent	HIS Va	alue Measu	rement	Color
		R	G	В	I _R	I_{G}	I_B	Color
	1	232	213	224	0.346	0.317	0.334	
	2	233	213	224	0.347	0.317	0.334	
le ,	3	233	212	224	0.348	0.317	0.334	
fun	4	231	211	222	0.343	0.312	0.330	
Š	5	230	210	223	0.346	0.316	0.336	
	Ā	231.8	211.8	223.4	0.345	0.316	0.334	
	1	206	108	215	0.389	0.203	0.406	
umple B	2	206	107	216	0.388	0.202	0.407	
	3	206	108	216	0.386	0.203	0.407	
	4	205	106	215	0.389	0.201	0.408	
Š	5	206	107	215	0.388	0.203	0.407	
	$\overline{\mathrm{D}}$	205.2	107.2	215.4	0.388	0.203	0.407	
	1	222	173	228	0.356	0.278	0.366	
()	2	221	173	227	0.355	0.278	0.365	
ole (3	221	172	226	0.357	0.276	0.366	
aml	4	220	171	227	0.355	0.276	0.367	
Š	5	220	172	228	0.354	0.277	0.367	
	$\overline{\mathbf{F}}$	220.8	172.2	227.2	0.355	0.276	0.366	

Table 3. RGB value samples

Table 4. Sample concentration value of TCS3200 color sensor

Samula (V)	Repetition (mg/L)				$\overline{\mathbf{V}}$ (m - /I)	
Sample (X)	1	2	3	4	5	A (mg/L)
А	1.75	2.10	2.46	0.67	1.75	1.74
В	17.10	16.75	16.03	17.10	16.75	16.74
С	5.32	4.96	5.67	4.96	4.61	5.10

confirmed by either method. As for sample C, the calculated concentration was lower than the LOQ of the TCS3200. Although its presence was confirmed, its quantitative concentration value was not reliable. Therefore, for the following analysis of comparing TCS3200 with UV-Vis spectrophotometer, samples A and C were excluded.

Comparing Methods Between the TCS3200 Color Sensor with UV-Vis Spectrophotometry Using the Two-Way t-Test

Method comparisons were carried out to see whether the TCS3200 color sensor had similar results to a UV-Vis spectrophotometer. The prepared samples were measured for five repetitions with UV-Vis at a wavelength of 590 nm and TCS 3200. Concentrations of rhodamine B in sample B were 16.74 and 17.26 mg/L for measurements using TCS3200 and UV-Vis spectrophotometer, respectively. *T*-test (a = 8.95%) performed on the obtained data revealed that the t_{experimental} and t_{theoretical} values were 1.21 and 2.31, respectively. Therefore, H₀ is accepted because the value of t_{experimental} < t_{theoretical}. H₀ states that differences of data obtained from TCS3200 and the UV-Vis spectrophotometer are not meaningful or significant. This analysis validates the high concentration of rhodamine B in sample B, calculated using TCS3200. It is worth mentioning that high concentrations of rhodamine B exposed to the human body could cause adverse health effects [34].

CONCLUSION

The analytical performance of the newly developed TCS3200 color sensor was satisfactory, considering that

the analysis could be run *in situ* and available at an affordable cost, and the components were free to access. The results showed that the color gradation only occurred in the R (Red) component, while the other components, G (Green) and B (Blue), were not concentration-depended. The *t*-test results showed that $t_{experimental} < t_{theoretical}$ suggesting the absence of statistical significance between the results obtained from the TCS3200 color sensor and the UV-Vis spectrophotometric method. The syrup samples procured from the local stores in Banda Aceh were tested qualitatively and quantitatively and was found to contain rhodamine B with high concentrations.

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Announcements

What have we done to improve the quality of our journals?

As a consequence of being indexed on Scopus and ESCI, the manuscripts we received increased during the 2010-2019 period. Details were given in Fig. 1. This increase is a formidable challenge. On the one hand, we are under pressure from authors who demand short review times and short waiting times for publication. On the other hand, we must strictly maintain the quality of the articles we publish (Fig. 2).

Number of Manuscript Received vs Accepted (2010-2019)



Figure 1. Number of Manuscript Received vs Accepted (2010-2019)



Figure 2. Rate of Article Acceptance (%) (2010-2019)

The following are the steps we have taken to guarantee and improve the quality of the published article:

- 1. Since 2017 we have used the Open Journal System (OJS) on all article management functions and facilities to manage articles more conveniently. The editors can supervise each other through the system. Also, editors can quickly monitor the performance of reviewers and authors.
- 2. We implement a more stringent filter at an early stage to reject inappropriate manuscripts to proceed to the review process. We have determined a minimum standard for articles to continue the review process.
- 3. We conduct strict plagiarism checks on manuscripts that we deem appropriate to proceed to the review process using professional software.
- 4. We hold editor training and discussion forums to improve editors' ability to handle manuscripts and synchronize mindsets in decision-making.
- 5. We invite editors with proven expertise from various countries and they are willing to help us voluntarily
- 6. We invite scientists from various countries as reviewers. Therefore, we really thank you for their commitment to voluntarily reviewing the articles.
- 7. We evaluate the review results from reviewers. As a result, our review time is relatively long; the average for the 2015-2019 period is 4-5 months (Fig. 3).
- 8. We do the copyediting stage carefully for articles that have been accepted. This step is needed to prevent substantial errors that are missed in the review process. Copyediting is also very useful to improve the readability of articles, the feasibility of illustrations, the suitability of citations and references, etc. This process takes an average of 3-4 weeks.
- 9. We improve the quality of the layout of articles with stringent standards so that the appearance of each article in each volume becomes uniform. We always try to shorten the time for article layout while maintaining quality.
- 10. We carry out the proofreading stage by the author and editor as a final check uploading the article in an online system —the time required at this stage is, on average, 2-3 weeks.
- 11. Finally, we post it in the "articles in press" section. Author and editor still have a chance to make revisions if they find unnecessary errors.

12. We are taking into account the waiting time between articles accepted to

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publish, which is sharply increased in the 2015-2019 period (from 4.3 to 8.8 months), see Fig 2. Through careful calculation, we manage the number of articles for each year's publication. Therefore, we increased the number of articles gradually from 43, 51, 68, 100, and 120. In 2020, we plan to publish 150 articles, which is distributed into 6 issues. The distribution of countries of origin of the authors for the 2015-2019 period is shown in Fig. 5



Figure 3. Average review time in the period of 2015-2019 (in weeks)







Figure 5. Distribution of Authors' Countries of Origin (2015-2019)

Posted: 2020-07-08

Publication Frequency and call for paper

After publishing four issues since volume 18 (2018), the number of submitted papers increases significantly. Therefore, to speed up qualified articles to be published internationally, Indonesian Journal of Chemistry publishes six issues (numbers) annually (February, April, June, August, October, and December) since 2020 (Volume 20). Therefore, we invite all authors to submit your qualified

manuscripts of original research articles, reviews, short communication in our Journal. Within two months (longest) from submission, the decision of acceptance or rejection has been made. Submission is only via online.

Posted: 2019-12-17

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by Muhammad S S, M Farhan, Zakaria Z, Muhammad Isa, Elly S, Sagir A, Elin Y, Leni Heliawati, Dkk

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Development of Arduino Uno-Based TCS3200 Color Sensor and Its Application on the Determination of Rhodamine B Level in Syrup

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Abstract: The use of the notorious synthetic dye, rhodamine B in food and beverage products has been widely reported. This application urges the need to develop an analytical method that can provide reliable rhodomine B data with an easy operational technique. Therefore, this research is aimed to develop an Arthrino Uno-based TCS3200 color sensor and study its application to determine rhodomine II levels in synup. The design of the analytical instrument included TCS3200, an Arduino Uno microcomputer, an Integrated Development Environment (IDE) software, a black box container, and a 24 × 2 matrix display server, where samples were prepared via absorption using wool thread. With a linear range of 1-20 mg/L, our proposed colorimetric sensor had recoveries of 96.25-110.3%, which was better compared to that was obtained from the UV-vis (81.8-100.6%) method. The detection and quantification limits of the sensor were 2.766 and 8.383 mg/L, respectively. The syrup samples used in this study were purchased from the local stores in Banda Aceh. Based on the proposed TCS3200 color sensor, the highest rhodamine B concentration from the syrup sample was 16.74 mg/L. The t-test analysis in this study revealed that the Rhodamine B levels quantified using the newly developed TCS3200 color sensor were not statistically or significantly different from the UV-Vir spectrophotometer method.

Keywords: color sensor; TCS3200; rhodomine B: Arduino Uno; Zn(CNS)/ IDE software

INTRODUCTION

As a form of consumer protection efforts, sensor technology for food or beverage products has been developed intensely [1-3]. For example, Fourier Transform Infrared (FTIR) spectroscopy has been employed to separate halal and non-halal meatballs [4]. The color spectroscopy method has also reported the detection and analysis of synthetic dye rhodamine B in chili powder [5]. Dyes used in food products are derived from natural and artificial chemicals. Synthetic dyes are widely used because it is more cost-efficient and available. Additionally, the synthetic dye may give a brighter color to the food or beverage product [6].

As one of 30 synthetic dyes available, Rhodamine B is considered as a dangerous dye, where its use in food or beverage products has been prohibited. Nevertheless, Rhodamine B is often used in processing industries, papers, and fabrics [7-8]. Moreover, it could be employed as a ligand to bind metal ions [9]. However, in Indonesia, rhodamine B is still very popular as a food coloring agent, including in iced syrup. The syrup is intentionally added with rhodamine B, so the products obtain a more attractive appearance [10].

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Methods that have been previously developed to identify rhodamine B in food ingredients include thinlayer chromatography (TLC) [11], voltammetry [12], and the standard method using UV-Vis spectrophotometer (the best option for identifying compounds with color). However, UV-Vis spectrophotometry has several drawbacks; not portable, complicated, and expensive [13]. Hence, this study tried to overcome the stated drawbacks by developing a simple measurement method using a portable sensor.

The sensor system developed in the present work was based on the TCS3200 color sensor constructed by the console to overcome external noise and program library modification [14]. The TCS3200 color sensor has been widely reported for different applications, including measuring levels of cyanide [14], nitrogen [15], and heavy metals [16]. In the case of colorimetric sensors, analytes should first be reacted with a complexing agent to cause a color change [17-18]. In this study, the sensor detects color degradation from tissue paper that has been spiked with reagents, so its sensitivity is specifically improved for rhodamine B analysis. The reagent used was Zn(CNS)₂, which can cause a color change from red to purple due to the formation of the rhodamine B-Zn-thiocyanate ((RhB) complex 2Zn(CNS)₄) [19].

The color intensity contributed by the presence of rhodamine B was converted through the sensor output pin in the form of a square signal in which its frequency depends on the concentration. The box's signal with varied frequency was then processed using a microcontroller on Arduino Uno. In this processing, four filters were used, namely green, blue, red, and no filter. In this case, no filter was excluded because the three parameters were sufficient to represent the color degradation of rhodamine B in the sample [20]. Filter settings were performed by providing low and high logic in the Arduino IDE program, following the reported study [21].

The distance between the sample and the 8 × 8 diode array was set at 3 cm, following the sensor system's geometry. The console's color was made black so the color could be absorbed fully, and influence from the degradation of colors that enter the diode array could be avoided. After obtaining the concentration of rhodamine B using the Arduino Eno-based TCS3200 color sensor; the results were compared with the standard UV-Vis spectrophotometric method. Finally, the analysis results were compared to obtain the data on sensitivity and accuracy of the newly developed sensor [22].

EXPERIMENTAL SECTION

Materials

The materials used were a UV-Vis spectrophotometer (Thermo Fisher Scientific, Selangor Malaysia), a color sensor TCS3200 (ICTAOS/AMS), a console, and an Arduino Uno (wavgat). Syrup samples tested for rhodamine B levels were procured from local stores in Banda Aceh. The standard rhodamine B was purchased from The National Agency of Drug and Food Control of Indonesia (BPOM RI). All other chemicals used, i.e. NH₄OH, NaOH, HCI, C₂H₃OH, CH₃CO₂Na, ZnCl₂, CH₃CO₂H, and KCNS, were obtained from Merck (Selangor, Malaysia) in analytical grade.

Hardware Design

The hardware design was initiated by developing a console for the TCS3200 color sensor, then connecting the output port of the color sensor via a jumper cable to the Arduino Uno microcomputer to process frequency data and convert it into 8-bit RGB digital data. There were 256 color digit variations for each RGB color component that was sortable and distinguishable by the processing. These color digit variations were displayed on the computer screen and converted to reduce color variations. These color variations were also recorded in .xls format (MS Excel) (Fig. 1).

Development of the TCS3200 Sensor Console

The TCS3200 console sensor was designed in black to absorb all color wavelengths. The distance between the diode array and the color object was 3 cm. The console was arranged in such a way that light from outside could not enter. The TCS3200 sensor was positioned opposite the color sample, which was absorbed into a filter paper. Four LED units with white

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Fig 1. Schematic diagram of hardware design

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wavelengths would hit the filter paper, and the intensity light reflected the diode array following the color intensity of the sample.

Software Design

Construction of the software design was initiated with a blink test on the Arduino Uno system to determine the response and performance of the microcomputer. The software used was Arduino IDE with available opensource libraries – C programming language. The program library was modified to enable the required color filters, Arduino Uno pins, the required display format, and data storage mode (Fig. 2).

Rhodamine Analysis Using TCS3200 Color Sensor

Construction of the calibration curve for rhodamine B

Briefly, the rhodamine B solution was added with 3.0 mL of Zn-thiocyanate. Then, the standard solution (RhB)₁-Zn(CNS), with different concentrations measured the RGB value with the TCS3200 color sensor and absorbance with UV-Vis at the maximum wavelength obtained. The solution was prepared with 100 mg/L rhodamine B as the stock solution, which was then diluted using distilled water into standard solutions with varying concentrations ranging from 1 to 20 mg/L. These solutions were prepared to determine the maximum wavelength of rhodamine B and as a database for the TCS3200 color sensor. Following that, a solution of 1 mL ZnCl₂ 2 M and 2 mL KCNS 2 M as a reagent was made to detect the presence of rhodamine B, as suggested by a previous report [23].



Fig 2. Display of the Arduino IDE Software main menu

Determination of rhodamine B level using the TCS3200 color sensor

The standard curve of (RhB)₂-Zn(CNS)₄ was obtained by measuring the RGB values of the standard solution (RhB)₂-Zn(CNS)₄ using the TCS3200 sensor. The concentration of rhodamine B used was 1 to 20 mg/L, which were priorly reacted with reagents. Measurements were carried out three times, and the concentration was averaged. Thereafter, RGB values were converted into a color index, namely Hue, Intensity, and Saturation (HIS). Conversion of RGB values to HIS values was carried out using the following Equations.

Red color index
$$(I_B) = \frac{R}{R+G+B}$$
 (1

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(2)

(3)

Green color index
$$(I_G) = \frac{G}{R + G + R}$$

the color index
$$(I_B) = \frac{1}{R+G+B}$$

The HIS color model was designed to resemble the perception of human vision, while the RGB values resembled the image of the display system [20]. The results of the calculation of the HIS value were then plotted as the dependent variable (y-axis) to the variation of concentration (RhB)=-Zn(CNS), (x-axis).

TCS3200 color sensor method validation

Method validation included accuracy, precision, sensitivity, and linearity, which were conducted based on the suggestion from a previous report [22].

Syrup sample preparation

Samples of commercial red syrup were purchased from local stores in Banda Aceh. Each sample (10 mL) was taken and purpto an Erlenmeyer which was subsequently mixed in 20 mL of 25% argunonia solution (dissolved in 70% ethanol) for 24 h and evaporated on a hot plate. The evaporation residue was dissolved in 10 mL distilled water containing acid (10 mL distilled water and 5 mL acetic acid 10%). Wool thread (15 cm) was dipped into the acid solution and simmered for 10 min until the dye colors appeared on the wool thread, then lifted. The wool thread was then washed with distilled water, and the wool thread was dissolved in ethanol 70% and heated to a boil (Fig. 3). This solution was used as the sample, per suggestion by a published work [24]. The wool thread was used to extract



Fig 3. The extraction of rhodamine B from commercial red syrups using wool thread. Wool thread was dipped into the dissolved syrup residue for 10 min (a). Rhodamine B-containing wool thread before reimmersed to ethanol 70% and boiled (b)

rhodamine B-containing samples in an acidic environment. A comparative study has reported that wool thread has the highest dye adsorption as compared with silk and nylon [25]. Adsorption of dye analyte in wool thread is determined by its O- and N-containing functional groups, which has been reported in many published papers [26-28]. The dyed wool was then immersed in ethanol 70% and boiled until its original color returned. The obtained solution was analyzed for its rhodamine B levels using the TCS3200 color sensor and a reference method – UV-Vis spectrophotometry.

Quantitative Analysis

The prepared sample was added with Znthiocyanate and then dipped in filter paper. Rhodamine B levels were measured using the TCS3200 color sensor [29]. The concentration was obtained based on the linear equation obtained from the calibration curve.

Method Comparison using Two-Way t-Test

Results of the samples between the TCS3200 color sensor and the UV-Vis spectrophotometry method were compared [30]. In addition, a two-way *t*-test was carried out to see the significance between the newly studied TCS3200 color sensor method and the reference method using UV-Vis spectroscopy by calculating the *t* value for each method and then comparing it with the toperate

RESULTS AND DISCUSSION

Maximum λ of Rhodamine B Complex

The complex (RhB)₂-Zn(CNS), was produced to give rhodamine B a specific color, allowing easier analysis. The solution of rhodamine B, which was initially red, turned to purple and was then measured using a UV-Vis spectrophotometer at a wavelength ranging from 574 to 600 nm. The UV-vis absorbance corresponding to the (RhB)₂-Zn(CNS), complex scanned from 574 to 600 nm is presented in Fig. 4.

Based on the measurement results, the UV-Vis spectrometer spectrum of (RhB)₂- (CNS)₄ showed a maximum absorption (0.442 au) at a wavelength of 590 nm. The difference in wavelength between rhodamine B and (RhB)₂-Zn(CNS)₄ is due to a shift in

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Fig 4. UV-Vis spectrometer spectrum of (RhB)2-Zn(CNS), showing a maximum wavelength at 590 nm

wavelength towards the bathochromic direction caused by substitution, solvent effects, and the influence of the chromophore group [31]. The successful formation of the (RhB)₂-Zn(CNS)₄ complex was indicated by a color change from red to purple and a shift in wavelength. The equation for the reaction between rhodamine B and Zn(CNS)₂ can be seen in Fig. 5.

Based on the graph, we can see three regression equations obtained from each RGB index value, namely $l_{\rm B}$ y = 0.0028x + 0.3411; $l_{\rm G}$ y = 0.0032x + 0.3513 and $l_{\rm E}$ y = -0.0058x + 0.3059. The values of the determination coefficient (R¹) were 0.9792,0.9700, and 0.9729 respectively. The R index had the best determination coefficient (R³) of 0.9792. Therefore, the regression equation for the R index was used to determine the concentration of rhodamine B in the sample,

Measurement using UV-Vis Spectrophotometer

The standard curve of (RhB)₂-Zn(CNS)₈ was measured at a wavelength of 590 nm by a UV-Vis spectrophotometer. The concentration of rhodamine B that was used ranged from 1 to 20 mg/L, which was priorly reacted with reagents. Measurements were carried out three times and averaged for each concentration. The absorbance measurements can be seen in Fig. 6. The regression equation y = 0.0023x =0.0773 had a determination coefficient (R⁷) of 0.9927. Hence, it can be concluded that the concentration was



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directly proportional to the absorbance, meaning that the absorbance for the complex (RhB)₂-Zn(CNS)₄ was dependent on rhodamine B concentration.

Method Validation

Accuracy

The accuracy of the proposed sensor method was based on the recovery (%), representing the value proximity of the standard concentration solution to the actual concentration. The concentrations of (RhB)₁-Zn(CNS), used were 1, 10, and 20 mg/L for the analysis with TCS3200 and UV-Vis color sensors. The actual concentration values and the percent recovery values from each method can be seen in Table 1. The recovery % calculation for the TCS3200 color sensor was still within the allowable error range of 90–110% [32]. However, at a concentration of 1 mg/L, UV-Vis had a recovery value below the permissible range (81.8%). Therefore, our proposed method was suggested to have better accuracy for determining thodamine B levels at a low concentration (1 mg/L).

Precision

The precision was determined to see the proximity of the value changes in the repetition process. The precision value was derived from the standard curve with a respective concentration of (RhB)₂-Zn(CNS), (1, 10, and 20 mg/L), expressed by the variation coefficient (VC). The precision values for both methods based on intra-day and inter-day repetition are presented in Table 2. The variation coefficient value obtained by the two measurements increased with the decrement in the concentration of the standard solution. The method is accurate if it provides a variation coefficient value of less than 2% [32]. Nonetheless, inter-day repetition yielded higher variation coefficient, especially when rhodamine B concentration was 1 mg/L.

Linearity

Linearity is the functional area of sample measurement. The linearity of measurements using the TCS3200 color sensor and UV-Vis spectrophotometer for a concentration range of 1–20 mg/L is depicted in Fig. 5 and 6, respectively. Several studies used a





Concentration (mg/L)	Actual cos (m	g/1,)	Recovery (%)	
	TCS3200	UV-Vis	TCS3200	UV-Vis
1	1.030	0.818	103.5	81.80
10	11.03	10.06	110.3	100.6
20	19.25	19.03	95.25	95,15

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Table 2. VC values of TCS3200 sensor and UV-vis spectrophotometer obtained from intra-day and inter-day repetition

1941 Frank (0) Country	Intra-day variatio	n coefficient (%)	Inter-day variation coefficient (%)		
(knodamine b) (mg/L)	TCS3200	UV-Vis	TCS3200	UV-Vis	
1	0.291	0.721	7.966	8.563	
10	0.268	0.521	1.294	1.664	
20	0.253	0.357	0.509	0.851	

non-linear calibration curve because the sensor system formed an exponential response [33]. However, in the present study, the quantitative analysis was conducted based on linear regression.

Sensitivity

The sensitivity value is shown from the slope of the complex standard curve of (RhB);-Zn(CNS); for each method. Based on the linear regression standard curve equation, the slope value for the TCS3200 color sensor measurement method was obtained from the regression equation y = 0.0028x + 0.3411, which was 0.0028. While the slope value for the UV-Vis spectrophotometer measurement method was obtained from the regression equation y = 0.0253x + 0.0773 is 0.0253. Based on the constructed standard curve, we calculated the limit of detection (LOD) by multiplying the standard deviation of response by 3.3 and dividing with the slope. Meanwhile, the limit of quantification (LOQ) could be obtained by multiplying the standard deviation of response by 10 and dividing it with the slope. The LOD obtained for the TCS3200 color sensor and UV-Vis spectrophotometer was 2.766 and 1.715 mg/L, respectively. These values explain why the inter-day precision for the 1 mg/L rhodamine B sample obtained for both methods exceeded the acceptable maximum variation coefficient (< 2%). As for the LOQ, the values reached 8.383 and 5.196 mg/L for the TCS3200 sensor and UV-Vis spectrophotometer, respectively. Lower LOD and LOQ in UV-Vis spectrophotometer suggest its superiority in comparison to the TCS3200 color sensor, in terms of sensitivity.

Quantitative Analysis Using the TCS3200 Color Sensor

Samples were measured using a series of tools that had been readily connected to the TCS3200 color sensor. The measurement was carried out by dipping the filter paper into the sample solution to which 3 mL of Zn(CNS)₂ reagent had been added, then dried and measured using the TCS3200 color sensor in dark conditions. Measurements were carried out three times on each sample with 3 cm-long distance between the sensor and the sample. Such distance was given to allow even distribution of the emitted light from four Light Emitting Diodes (LEDs) to the sample and the photodiode, in which the sample could emit a current proportional to the basic color of received light.

Table 3 shows that the RGB value obtained from each sample is a code to indicate a specific color. The HIS value in the table was obtained using Eq. (1-3). The It value was used to determine the concentration of rhodamine B in the sample because it had the best R² (9.9792) among the others (Fig. 6). The total concentration of rhodamine B obtained from the measurement using the TCS3200 color sensor based on the Is value can be observed in Table 4, showing the concentration of each sample with five repetitions. The red index value obtained from Eq. (1) has the same function as the absorbance value, the dependent variable in determining the concentration. Therefore, the concentration of rhodamine B in the semple was calculated by substituting the red color index value of the sample into the standard curve regression equation (RhB)+-Zn(CNS), R index.

Following the analysis, we found that samples A, B, and C contained rhodamine B with an average of 1.74, 16.74, and 5.10 mg/L, respectively. However, sample A had a rhodamine B concentration lower than the LOD of both the TCS3200 and UV-Vis spectrophotometer (2.766 and 1.715 mg/L, respectively). In this case, the response generated from sample A could not be differentiated from that of the blank standard. Hence, the presence of rhodamine B in sample A could not be

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			Table 3.	RGB valu	e samples	6		
Repetition		RGB	RGB Measurement			HIS Value Measurement		
		R	G	В	Iĸ	L	I.	- Color
	1	23.2	213	224	0.346	0.317	0.334	
mple A	2	23.3	213	224	0.347	0.317	0.334	
	3	233	212	224	0.348	0.317	0.334	
	4	231	211	222	0.343	0.312	0.330	
50	5	230	210	223	0.346	0.316	0.336	
	Ā	231.8	211.8	223.4	0.345	0.316	0.334	
2	1	206	108	215	0.389	0.203	0.406	T.
	2	206	107	216	0.388	0.262	0.407	
2	3	206	108	216	0.386	0.203	0.407	
E.	4	20.5	106	215	0.389	0.201	0.408	
20	5	206	107	215	0.388	0.203	0.407	
	Ð	205.2	107.2	215.4	0.388	0.203	0.407	
	1	22.2	173	228	0.356	0.278	0.366	1
le C	2	221	173	227	0.355	0.278	0.365	
	3	221	172	226	0.357	0.276	0.366	
E.	4	.Z20	171	227	0.355	0.276	0.367	
-25	5	220	172	228	0.354	0.277	0.367	
	F	220.8	172.2	227.2	0.355	0.276	0.366	

Table 4. Sample concentration value of TCS3200 color sensor

Camerala (M)		×				
Sample (A)	1	2	3	+	5	A (mg/L)
Λ	1.75	2.10	2.46	0.67	1.75	1.74
B	17.10	16.75	16.03	17,10	16.75	16.74
C	5.32	4.96	5.67	4.96	4,61	5.10

confirmed by either method. As for sample C, the calculated concentration was lower than the LOQ of the TCS3200. Although its presence was confirmed, its quantitative concentration value was not reliable. Therefore, for the following analysis of comparing TCS3200 with UV-Vis spectrophotometer, samples A and C were excluded.

Comparing Methods Between the TC53200 Color Sensor with UV-Vis Spectrophotometry Using the Two-Way t-Test

Method comparisons were carried out to see whether the TCS3200 color sensor had similar results to a UV-Vis spectrophotometer. The prepared samples were measured for five repetitions with UV-Vis at a wavelength of 590 nm and TCS 3200. Concentrations of rhodamine B in sample B were 16.74 and 17.26 mg/L for measurements using TCS3200 and UV-Vis spectrophotometer, respectively. T-test (a = 8.95%) performed on the obtained data revealed that the taxonana and taxonaa values were 1.21 and 2.31, respectively. Therefore, Ho is accepted because the value of terrenerated < thereicted. Has states that differences of data obtained from TCS3200 and the UV-Vis spectrophotometer are not meaningful or significant. This analysis validates the high concentration of rhodamine B in sample B, calculated using TCS3200. It is worth mentioning that high concentrations of rhodamine B exposed to the human body could cause adverse health effects [34].

CONCLUSION

The analytical performance of the newly developed TCS3200 color sensor was satisfactory, considering that

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the analysis could be run in situ and available at an affordable cost, and the components were free to access. The results showed that the color gradation only occurred in the R (Red) component, while the other components, G (Green) and B (Blue), were not concentration-depended. The *t*-test results showed that $t_{\rm opermost} < t_{\rm hereofcl}$ suggesting the absence of statistical significance between the results obtained from the TCS3200 color sensor and the UV-Vis spectrophotometric method. The syrup samples procured from the local stores in Banda Aceh were tested qualitatively and quantitatively and was found to contain rhodamine B with high concentrations.

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Development of Arduino Uno-Based TCS3200 Color Sensor and Its Application on the Determination of Rhodamine B Level in Syrup

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