WATER QUALITY ASSESSMENT OF TEKALA RIVER, SELANGOR, MALAYSIA


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Abstract. A study on water quality status of Tekala River, Selangor, Malaysia was conducted at two sampling times. A total of six sampling stations were selected along the river representing the upstream (S1 and S2), the middle stream (S3 and S4) and the downstream (S5 and S6) of the Tekala River. In this study, in-situ and ex-situ analyses were conducted to determine the quality of Tekala River. Physical, chemical and biological parameters included biochemical oxygen demand (BOD), chemical oxygen demand (SOD), ammoniacal nitrogen, total suspended solids (TSS), dissolved oxygen (DO), pH, temperature, total dissolved solid (TDS), salinity, conductivity, Escherichia coli, coliform and macroinvertebrate. According to the result obtained from this study, Tekala River is classified under Class I and Class II based on water quality index and National water quality standard. Two-way ANOVA showed a significant difference between parameters (ammoniacal nitrogen, BOD, pH, temperature, conductivity, DO and TDS) of sampling station. Significant difference was found between ammoniacal nitrogen, pH and temperature with sampling time. The correlation test revealed that there is relationship between Escherichia coli with ammoniacal nitrogen, temperature and DO. There is also relationship found between coliform with pH and BOD.

Keywords: water quality, biomonitoring, water pollution, water management, biological parameters

Introduction

Over the centuries, river has been very important to the human society. River has also provided water for irrigation, industrial and domestic uses. Additionally, river plays an important role in assimilating municipal and industrial effluents as well as runoffs from agricultural land and the surrounding area in a watershed (Al-Badaii et al., 2013). The importance of water quality has become a serious matter especially when it involves the assessment of producing the biologically-accepted water system as a whole (Ashraf and Hanafiah, 2017; Muhammad Mansoor et al., 2018). The demand for safe and clean drinking water has increased by leaps and bounds in developing countries that have deteriorated environment (Gelover et al., 2006). The target of the Millennium Development Goal (MDG) which is the ‘access to safe drinking water’ has set its track globally with about 6.1 billion people in 2010 using improved drinking water sources, an increase of over 2 billion since 1990 (MDG, 2012).

Selangor state can be divided into several districts which are Klang, Kuala Langat, Gombak, Kuala Selangor, Hulu Langat, Hulu Selangor, Petaling, Sabak Bernam and Sepang. Selangor is one of the most populated and industrialized state in Peninsular Malaysia. As the most populated state, rivers in Selangor are facing serious
contamination due to anthropogenic activities. On the other hand, study by Al-Badaii et al. (2013) found that Semenyih River, Selangor was contaminated with NH$_3$-N, TSS, COD and NO$_3$ and highly polluted with PO$_4$ and faecal coliform. The sources of contamination were originated from industrial, agricultural, livestock farming and erosion. Mohamad Ali (2010) stated that the main pollution in Selangor River is from poultry farms, wet market activities and industrial wastewater.

Tekala River is a recreational river that has potential to be contaminated with faecal pathogens (such as bacteria and viruses) from human sewage and animal manure. Contaminated recreational water can cause diseases. Therefore, this study aims to measure the water quality status of Tekala River based on the physico-chemical and biological parameters. In view of the above facts, Tekala River has been chosen due to its importance as recreational area. If this river is classified as not safe then further action should be taken to maintain and improve its quality.

Materials and methods

Study area

This study was conducted at Tekala River, Selangor, Malaysia (Fig. 1a) within Tekala River Recreational Park (TRRP) that was established since 1982. The highest level of the river leads up to a sparkling waterfall which cascades into a natural rock pool. TRRP is located 13 km from Semenyih via Jalan Semenyih – Hulu Langat and 50 km from Kuala Lumpur.

![Figure 1. Map of Tekala River and locations of six sampling points](image-url)
Measurement of ex-situ parameters

The water quality index was determined based on the physico-chemical and biological parameters. The parameters include biochemical oxygen demand (BOD), chemical oxygen demand (COD), ammoniacal-nitrogen (NH₃-N), total suspended solids (TSS), dissolved oxygen (DO), pH, water temperature, total dissolved solid (TDS), salinity, conductivity, Escherichia coli (E. coli), coliform and macroinvertebrate.

Samples collection and preservation

Water samples were collected from the selected river and have been collected for two visits in May 2017 and October 2017. A total of 18 water samples were collected from 6 sampling stations and the samples were kept in polyethylene bottles. For BOD sample, another 18 water samples were collected in glass bottle wrapped with aluminium foil. This is to avoid sunlight from penetrating the bottles and react with sample and cause changes in the BOD. During sampling for BOD, no air bubble was observed as it can influence the oxygen content of the sample. All bottles were preserved at 4–10 °C in an ice box before transported to the laboratory. Preservation of samples need to be done to maintain the actual composition of water samples and can minimize changes in chemical composition during transportation to the laboratory.

DO, water temperature, TDS, salinity, conductivity and pH values were measured in-situ. The COD, TSS, BOD, ammoniacal-nitrogen, E. coli and coliform in the water were analysed in the laboratory using standard procedure of HACH. For macroinvertebrate, samples were collected by using Surber net at 6 stations based on different habitats. All collected samples were transferred into plastic. Then, samples were preserved by using 70% ethanol in the laboratory for further identification.

Water was kept in dark condition and in temperature around 4–5 °C to reduce biological activity, to avoid changes and to preserve the actual condition of sample as it is before conducting any analysis. BOD samples were kept in glass bottle preserved at 30 °C for about 5 days prior to measurement. The use of glass bottle can avoid any changes in activity and microorganism metabolism. Samples for E. coli and coliform test were kept at 4–5 °C and analysis was conducted at least 4–8 h after sampling activity.

Biological Oxygen Demand (BOD) was determined by using conventional method by incubating the water samples for a standard time period in incubator at 20 °C and the oxygen consumption was determined at the end of the incubation period. The first measurement of DO values (DO₁) was carried out within 24 h after sampling by using YSI model 5000. The reading was recorded on a data sheet. The second reading of DO values was determined after 5 days of incubation (DO₅). BOD values were obtained from the difference in DO values of the first reading (DO₁) and the second reading (DO₅). The BOD value is expressed in milligrams per liter using Equation 1:

\[
\text{BOD (mg/l)} = \text{DO}_{\text{initial}} - \text{DO}_{\text{final}} \quad \text{(Eq.1)}
\]

Chemical oxygen demand

COD was measured by using Reactor Digestion Method approved by USEPA for reporting wastewater analysis. Samples were first homogenized to dissolve containing solids for better representative samples. The homogenized water samples were then added into glass vials containing strong oxidizing agent, potassium dichromate. This mixture was heated for 2 h at 150 °C to allow them to react. Colorimetric determination of COD
with high range using HACH DR/2010 was used to measure COD value in the water samples.

**Total suspended solid**

Whatmann filter paper with pore size of 0.45 µm was put in petri dish each and dried in the drying oven at the temperature of 105 °C for 1 h. After an hour, the petri dish was taken out and cooled for 30 min in the desiccator. Before performing filtration of the samples, the weighed filter paper was obtained. This was the initial weight of the filter paper (A). This filter paper then was used in filtration process. For the filtration process, the filter paper was inserted between the Buchner funnel and the suction flask. The water sample in the bottle was shaken and about 200 ml of sample was used in the filtration. After that, the filter paper was put back into petri dish and was dried in the drying oven at 105 °C and the final weight of the filter paper was recorded (B).

For the calculation (Eq. 2),

\[ x \text{mg/L} = \frac{(B-A) \times 1000}{\text{Volume of sample used (ml)}} \]  

where,

- A = initial weight of the filter paper (mg)
- B = final weight of the filter paper + soluble solid (mg)

**Ammoniacal-nitrogen**

Ammoniacal-nitrogen was measured by using HACH DR 2700 based on Nessler Method. 25 ml of sample and 25 ml of deionized water (blank) were transferred into glass cell each and then three drops of mineral stabilizer, three drops of polyvinyl alcohol and 1.0 ml of Nessler reagent were added to each transferred sample. The mixture was measured by using spectrometer at 425 nm. Deionized water was used as blank and the reading was recorded for each sample.

**Measurement for biological parameter**

*Escherichia coli* and *coliform*

100 ml of water sample was used for determination of *E. coli* and *coliform*. Water samples were filtered in cellulose membrane filter. After filtration, Eosin Methylene Blue (EMB) agar was added with no air bubbles observed on the agar. The agar was preserved for 22 ± 2 h at 35 °C. Organisms that produce a colony with golden-green metallic within 24 h of incubation on the agar were considered members of *E. coli* group. While organisms that produce colony with light purple were classified as *coliform* group.

Quantitative data was obtained by counting the number of each colony formed by assumption that each colony represents one bacterium. Colony of *coliform* and *E. coli* that grow was counted and expressed in unit of cfu (colony forming unit) (APHA, 1980) and calculated as below (Eq. 3):

\[ E. coli/100 \text{ml} = \frac{100 \times \text{Number of colony}}{\text{Volume of sample filtered (ml)}} \]
Macroinvertebrate

All the samples from the river were taken to the laboratory for identification process. Sample from different stations were put in different bottles. Each sample was analyzed by identifying specific characteristics for each species such as head shape, body segment and type of wings. Samples that had already been identified were preserved in 70% ethanol.

Water quality classification

Classification in water quality index (WQI) was determined based on the water quality index as below (Eq. 4):

\[
WQI = 0.22 \times \text{SIDO} + 0.19 \times \text{SIBOD} + 0.16 \times \text{SICOD} + 0.15 \times \text{SIAN} + 0.16 \times \text{SITSS} + 0.12 \times \text{SipH}
\]

(Eq.4)

where:
- SIDO = Subindex DO (% saturation)
- SIBOD = Subindex BOD
- SICOD = Subindex COD
- SIAN = Subindex NH\textsubscript{3}-N
- SITSS = Subindex SS
- SipH = Subindex pH

Statistical analysis was conducted by using IBM SPSS Statistics 20. Two-way ANOVA was conducted for comparing water quality parameter at different stations and sampling times. Correlation analysis was also conducted to determine the relationship between physico-chemical and biological parameters.

Results

Physico-chemical parameters

Mean value for total suspended solid (TSS) for two sampling times is shown in the Figure 2a. TSS ranged from 1.67 mg/L to 8.17 mg/L. The average value of TSS is 4.15 ± 0.48 mg/L and this value is categorized under Class I of NWQS. Based on the Two-way analysis of variance (ANOVA) result, there is no significantly different between TSS versus station and sampling time (P > 0.05, P = 0.746).

Figure 2b presents the mean value for ammoniacal nitrogen (NH\textsubscript{3}-N) for both sampling times. It was found that NH\textsubscript{3}-N were ranged from 0.03 mg/L to 0.31 mg/L with an average value of 0.18 ± 0.04 mg/L and falls under Class I of NWQS. Two-way analysis of variance (ANOVA) shows that there is significantly different between NH\textsubscript{3}-N and station (P < 0.05, P = 0.000). There is also significant difference between NH\textsubscript{3}-N and sampling times (P < 0.05, P = 0.038). Significantly different was also found between the effect of different stations and different sampling times on NH\textsubscript{3}-N (P < 0.05, P = 0.037).

Mean value for biological oxygen demand (BOD) for two sampling times is shown in Figure 3a is ranged from 0.04 mg/L to 1.42 mg/L. The average value of BOD is 0.78 ± 0.14 mg/L, categorized under Class I of NWQS. There is no significantly
different between BOD and sampling times (\( P > 0.05, P = 0.189 \)). However, there is significant difference between BOD and station (\( P < 0.05, P = 0.000 \)).

Figure 3b shows the mean value for chemical oxygen demand (COD) that were ranged from 0.33 mg/L to 1.67 mg/L, with an average value of \( 0.64 \pm 0.22 \) mg/L (Class I). Two-way analysis of variance (ANOVA) shows that there is no significant difference between COD with station and sampling time (\( P > 0.05, P = 0.656 \)).

\[ \text{Figure 2.} \ (a) \text{TSS values for six sampling stations;} \ (b) \text{Ammoniacal nitrogen values for six sampling stations, where; the upstream (S1 and S2), the middle stream (S3 and S4) and the downstream (S5 and S6)} \]

\[ \text{Figure 3.} \ (a) \text{Biochemical oxygen demand for six sampling stations;} \ (b) \text{Chemical oxygen demand for six sampling stations, where; the upstream (S1 and S2), the middle stream (S3 and S4) and the downstream (S5 and S6)} \]

\( \text{pH} \) was ranged between 5.00 to 6.8 with an average of \( 5.60 \pm 0.67 \), classified under Class III and IV for the first and second samplings, respectively (Fig. 4a). A significantly different was observed between \( \text{pH} \) and sampling times and station (\( P < 0.05, P = 0.025 \)) and (\( P < 0.05, P = 0.000 \)), respectively. Mean value for temperature for two sampling times is shown in Figure 4b ranged from 20.76 to 21.49 °C. There is significant difference between temperature and sampling times (\( P < 0.05, P = 0.000 \)) as well as between temperature and station (\( P < 0.05, P = 0.000 \)).

It was found that the conductivity for both sampling times were ranged from 0.013 mS/cm to 0.017 mS/cm (Fig. 5a). The average conductivity is \( 0.0014 \pm 0.0003 \) mS/cm and classified in Class I and IIA. Based on the result, there is no significant difference
between conductivity and sampling times (P > 0.05, P = 0.292), whereas there is significant difference between conductivity and station (P < 0.05, P = 0.006). For dissolved oxygen, the mean value for both sampling times were ranged from 5.32 mg/L to 7.86 mg/L as shown in Figure 5b. The average DO is 6.60 ± 0.29 mg/L (Class IIA and IIB). There is no significantly different between DO and sampling times (P > 0.05, P = 0.078). However, there is significant difference between DO and station (P < 0.05, P = 0.000).

Figure 4. (a) pH values for six sampling stations; (b) Temperature values for six sampling stations, where; the upstream (S1 and S2), the middle stream (S3 and S4) and the downstream (S5 and S6)

Figure 5. (a) Conductivity values for six sampling stations; (b) Dissolved oxygen values for six sampling stations, where; the upstream (S1 and S2), the middle stream (S3 and S4) and the downstream (S5 and S6)

Figure 6(a) shows the mean value for salinity for two times sampling ranged from 0.003 to 0.01 with an average of 0.007 ± 0.001 (Class I). No significantly different between salinity with station and sampling time was observed (P > 0.05, P = 0.813). The mean value for total dissolved solid ranged from 0.009 g/l to 0.011 g/l (Fig. 6b). The average of TDS is 0.009 ± 0.0001 and classified under Class I. There is no significant difference between TDS and sampling times (P > 0.05, P = 0.423), while significantly different was found between TDS and station (P < 0.05, P = 0.032).

Water Quality Index (WQI) for all stations times is shown in Figure 7. WQI has been considered as one criterion for surface water classifications, based on the use of
standard parameters for water characterization. This index is a numeric expression used to transform large quantities of water characterization data into a single number, which represents the water quality level (Mohamad Ali, 2010). The WQI values recorded were ranged from 85.45 to 94.86 with the highest WQI recorded at station 2, while the lowest WQI was measured at station 4. The results show that WQI obtained from this study fall under Class I (clean category). Overall, the WQI value of Tekala River was classified under Class II which indicated as clean.

Figure 6. (a) Salinity values for six sampling stations; (b) Total dissolved solid values for six sampling stations, where; the upstream (S1 and S2), the middle stream (S3 and S4) and the downstream (S5 and S6)

Figure 7. WQI value for six sampling stations

Biological parameter

Mean value for E. coli and total coliform for two sampling times are shown in Figures 8a and 8b. E. coli and total coliform values were ranged from 1 to 16 cfu/ml and 3 to 48 cfu/ml, respectively. The average value for E. coli and total coliform are 8 ± 2 cfu/ml and 21 ± 4 cfu/ml, respectively and both are classified under Class I. Pearson's
correlation was conducted to examine relationship between \textit{E. coli} and coliform with the physico-chemical parameters (\textit{Table 1}). Based on the test, there is relationship between \textit{E. coli} with \textit{NH}_3\textit{N}, temperature and DO. For coliform, there is relationship between coliform with pH and BOD.

\textbf{Macroinvertebrate}

Total value for macroinvertebrate for two sampling times is shown in Figure 9. The abundance of macroinvertebrate was ranged from 10 to 68 (Fig. 9). Pearson’s correlation was conducted to examine the relationship between abundance of macroinvertebrate with the physico-chemical parameters (\textit{Table 2}). Based on the test, there is relationship between abundance of macroinvertebrate with temperature of the river.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure8.png}
\caption{(a) \textit{E. coli} values for six sampling stations; (b) Coliform values for six sampling stations, where; the upstream (S1 and S2), the middle stream (S3 and S4) and the downstream (S5 and S6).}
\end{figure}

\textbf{Table 1.} Pearson’s correlation between \textit{E. coli} and coliform with physico-chemical parameters

\begin{table}
\centering
\begin{tabular}{lcccccccccc}
\hline
PARAMETER & \textit{E. coli} & Coliform & TSS & BOD & \textit{NH}_3\textit{N} & COD & pH & Temperature & Conductivity & DO & Salinity & TDS \\
\hline
\textit{E. coli} & 1 & & & & & & & & & & & \\
Coliform & -0.19 & 1 & & & & & & & & & & \\
TSS & 0.15 & -0.12 & 1 & & & & & & & & & \\
BOD & -0.03 & -0.47** & -0.094 & 1 & & & & & & & & \\
\textit{NH}_3\textit{N} & -0.43** & -0.102 & 0.111 & 0.256 & 1 & & & & & & & \\
COD & -0.25 & 0.206 & -0.229 & 0.089 & 0.118 & 1 & & & & & & \\
pH & 0.29 & 0.477** & -0.019 & -0.56** & -0.287 & -0.06 & 1 & & & & & \\
Temperature & -0.67** & -0.297 & -0.022 & 0.377* & 0.591** & 0.139 & -0.75** & 1 & & & & \\
Conductivity & 0.02 & 0.021 & -0.206 & 0.179 & 0.111 & -0.03 & -0.083 & 0.108 & 1 & & & \\
DO & 0.53** & 0.222 & -0.053 & -0.157 & -0.49** & -0.05 & 0.371* & -0.550** & -0.230 & 1 & & \\
Salinity & -0.21 & 0.086 & -0.045** & 0.112 & -0.003 & 0.040 & -0.231 & 0.190 & 0.014 & -0.08 & 1 & \\
TDS & 0.02 & -0.134 & -0.114 & 0.111 & 0.067 & 0.110 & -0.164 & 0.149 & 0.748** & -0.31 & 0.022 & 1 \\
\hline

\end{tabular}

\textit{**}Correlation is significant at 0.01 level (2-tailed)

\textit{*}Correlation is significant at 0.05 level (2-tailed)

\end{table}
Figure 9. Abundance of macroinvertebrate for six sampling stations, where: the upstream (S1 and S2), the middle stream (S3 and S4) and the downstream (S5 and S6)

Table 2. Pearson’s correlation between abundance of macroinvertebrate with physico-chemical parameters

<table>
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<tr>
<th>PARAMETER</th>
<th>X</th>
<th>TSS</th>
<th>NH₃N</th>
<th>COD</th>
<th>pH</th>
<th>Temperature</th>
<th>Conductivity</th>
<th>DO</th>
<th>Salinity</th>
<th>TDS</th>
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**Correlation is significant at 0.01 level (2-tailed)
*Correlation is significant at 0.05 level (2-tailed)

Discussion

Physico-chemical parameters

Total Suspended Solids (TSS) are solids in water include a wide variety of material, such as silt, decaying plant and animal matter, industrial wastes, and sewage. High concentrations of suspended solids can cause many problems for stream health and
aquatic life (Bilotta and Brazier, 2008). According to Wood and Armitage (1997), suspended solid in the river may be due to bank erosion, sediment load and anthropogenic activities or due to runoff from surrounding areas. In second sampling, station 2 surrounded by dense vegetation, thus recorded the highest TSS. Vegetation can reduce the erosion and increase sediment entrapment. However, there is no significant different between TSS with station and sampling time. The maximum threshold limit of TSS for Malaysian rivers which support aquatic life is 150 mg/l (Al-Badaii et al., 2013). TSS value recorded in this study are within the range of the limit.

Ammoniacal nitrogen (NH\textsubscript{3}-N) is a common pollutant in freshwater ecosystem and it is frequently found associated with organic compound or sometimes from industrial effluents (Alabaster and Lloyd, 1982; Magdalena et al., 2015). Abdel-Raouf (2012) stated that the presence of NH\textsubscript{3}-N in the water is mainly originated from the domestic sewage and waste water from certain types of industries. Ammoniacal nitrogen indicates nutrient status, organic enrichment and health of water body (Radojevic et al., 2007). Higher NH\textsubscript{3}-N values can be toxic to fish, but in small concentrations, it could serve as nutrients for excessive growth of algae (Al-Badaii et al., 2013). There is significant different between NH\textsubscript{3}-N with station and sampling times implying NH\textsubscript{3}-N is influenced by sampling location and time of sampling. The maximum threshold level of NH\textsubscript{3}-N for Malaysian rivers which support aquatic life is 0.9 mg/l. The value of NH\textsubscript{3}-N obtained from this study is still within the range.

Biochemical Oxygen Demand (BOD) is a measurement of the amount of dissolved oxygen used by aerobic microorganisms when decomposing organic matter in water. It is a measure of the quantity of oxygen used by microorganisms in the oxidation of organic matter. Microorganisms such as bacteria are responsible for decomposing organic waste. When organic matter such as dead plants, leaves, grass clippings, manure, sewage or even food waste is present in a water supply, the bacteria will begin the process of breaking down this waste. When this happens, much of the available dissolved oxygen is consumed by aerobic bacteria, affecting oxygen level needed for other aquatic organisms (USGS, 2016).

If there is a large quantity of organic waste in the water supply, there will also be a lot of bacteria present working to decompose this waste. In this case, the demand for oxygen will be high so the BOD level will be higher. When BOD levels are high, dissolved oxygen levels decrease because the oxygen that is available in the water is being consumed by the bacteria (Al-Badaii et al., 2013). Since less dissolved oxygen is available in the water, fish and other aquatic organisms may not survive. If there is no organic waste present in the water, there will be fewer bacteria present to decompose it and thus the BOD will tend to be lower and the DO level will be higher.

Chemical Oxygen Demand (COD) is a measurement of the oxygen required to oxidize soluble and particulate organic matter in water (Boyd, 1973; Kunlasak et al., 2013). It is used to measure the total quantity of oxygen-consuming substances in the complete chemical breakdown of organic substances in water. It does not differentiate between biologically available and inert organic matter. COD measurements can be made in a few hours while BOD measurements take five days (Othman et al., 2012). Generally, the lower COD level indicates a low level of pollution, while the high level of COD points out the high level of pollution of water in the study area (Al-Badaii et al., 2013). The COD value obtained in this study is low indicating this river has low level of pollution.
pH of a water sample is a measure of the concentration of hydrogen ions (Tank and Chippa, 2013). If the pH of water is too high or too low, most of the aquatic organisms will not be able to adapt to the extreme condition. pH can also affect the solubility and toxicity of chemicals and heavy metals in the water. Various concentration of solution, solid or gaseous that enters into river can influence pH of the river due to reaction of dissociation which may produce H+ or OH- ions (Chang et al., 1983; Lelis et al., 2016). Based on the result, pH ranged from 5.00 to 6.88. There is significance different between pH with station and sampling times indicating pH is influenced by sampling location and time of sampling. pH slightly decreased from station 1 to station 6. pH of water body can be affected by several factors. One of the factors is the amount of plant growth and organic material within a body of water. Carbon dioxide is released when this material is decomposed. The carbon dioxide combines with water to form carbonic acid. Although this is a weak acid, large amounts of it will lower the pH. Decaying vegetation produces organic acids, thus change the pH of the river (Gasim et al., 2007). Station 1 to station 6 were surrounded by vegetation. Different station may have different rate of decaying vegetation. This might explain the decreased of pH from station 1 to station 6. Rainfall could also affect pH of water body due to the interaction with carbon dioxide molecules in the atmosphere. This creates H$_2$CO$_3$ in the raindrops, lowering the pH value of rainwater. A pH level of 5.65, though acidic, is not considered acid rain. Acid rain requires a pH below 5.0. If rain falls on a poorly buffered water source, it can decrease the pH of nearby water through runoff.

Water temperature is one of the most important characteristics of an aquatic system. It affects dissolved oxygen levels, species composition, chemical and biological processes (Tank and Chippa, 2013). A change in water temperature can affect the general health of the aquatic organisms, thus changing the quality of the stream. There is significant difference between temperature with sampling times and station. The temperature of surface water is usually between 0 ºC and 30 ºC. The lowest temperature was recorded at station 1 and the highest at station 6 for both samplings. There is slightly increased of temperatures from station 1 to station 6. Usually there is an increase in temperature from the first station to the last station (Bordalo et al., 2001). Water temperature varies along the length of a river with latitude and elevation. The geographical location of sampling stations are differ on elevation and environment. Station 1 and station 2 were located in upstream, station 3 and station 4 in the middle meanwhile stations 5 and 6 in the downstream. Vegetation cover may influence water surface temperature. Area from station 1 to station 6 were covered by vegetation. However, the vegetation in the upstream was denser than middle and downstream. The vegetation acts as protection to surface water from directly being heated by solar radiation. The forest canopies will shield the water surface from the emission of long wave radiation from the sun. Denser vegetation gives more protection to the surface water thus this might explain the reason why station 1 had low temperature compared to other stations. Human activities in the surrounding area might also influence the water temperature. The middle and downstream had a lot of human contact as the river is a recreational area. Downstream area vegetation had some disruption as a lot of facilities being built in this area. Water temperature can also be influenced by variation of precipitation. According to Shuhaimi-Othman et al. (2008) large volume of water inputs and higher flow rate were responsible for cooling down the river temperature. There is only little difference of temperature between first and second samplings.
Conductivity can be defined as an ability of the aquatic system to transmit electric current based on the level of dissolved ions in a water body. Conductivity in water is affected by the presence of inorganic dissolved solids such as chloride, nitrate, sulfate and phosphate anions or sodium, magnesium, calcium, iron, and aluminum cations. Conductivity normally coinciding with the level of dissolved salt in water and has a strong correlation with salinity. An increasing in level of dissolved salt would result an increased in conductivity. There is slightly difference of salinity based on times and location of the sampling in this study. Based on the result, there is no significant different between conductivity and sampling times but there is significant different between conductivity and different station which means conductivity is influenced by sampling location. Conductivity in streams and rivers is affected primarily by the geology of the area through which the water flows. Streams that run through areas with granite bedrock tend to have lower conductivity because granite is composed of more inert materials that do not ionize when washed into the water. On the other hand, streams that run through areas with clay soils tend to have higher conductivity because of the presence of materials that ionize when washed into the water. Soil and rocks release dissolved solids into the waters that flow through or over them. Therefore, the geology of a certain area will determine the conductivity. Most streams have a fairly constant range of conductivity under normal circumstances. Therefore, significant changes in conductivity can be an indicator that a discharge or some other source of pollution has entered the water. This might explain the reason why conductivity for different sampling times is constant and the difference can be seen mostly based on the sampling location. The conductivity was ranged from 0.013 mS/cm to 0.017 mS/cm. This is considered low and does not affect the water quality (Suki et al., 1988; Rashid et al., 2013).

Dissolved oxygen (DO) measures the amount of gaseous oxygen dissolved in an aqueous solution. Oxygen gets into water by diffusion from the surrounding air, by aeration and as a waste product of photosynthesis from phytoplankton, algae, seaweed and other aquatic plants (Othman et al., 2012). The amount of dissolved oxygen varies depending on temperature, pressure and salinity. Solubility of oxygen decreases as temperature increases. This is due to the fact that warmer surface water requires less dissolved oxygen to reach 100% air saturation compared to cooler water. There is negative correlation of DO and temperature which means higher temperature will reduce DO level. Temperature slightly increased from station 1 to station 6 and the DO level recorded is slightly reduced from station 1 to station 6. Dissolved oxygen decreases exponentially as salt levels increase. There is significance difference between DO and station. In freshwater systems such as lakes, rivers and streams, dissolved oxygen concentrations will vary by season, location and water depth and it will decrease with higher temperature, salinity and elevation (Beale et al., 2000). Oxygen concentrations vary with the volume and velocity of water flowing in a stream. Faster flowing water areas, such as in a mountain stream or large river tend to be more oxygen rich because more oxygen enters the water from the atmosphere in those areas than in slower and stagnant areas. Oxygen is more easily dissolved into water at low altitudes that at high altitudes. Oxygen is also more easily dissolved into water with low levels of dissolved or suspended solids. Removal of riparian vegetation may lower oxygen concentrations due to increased water temperature resulting from a lack of canopy shade and increased suspended solids resulting from erosion of bare soil. Upstream area has
faster flowing water compared to lower area thus this might explain why level of DO is higher in upstream area.

Salinity is the total concentration of all dissolved salts in water. Freshwater contains few salts and thus has low salinity. Most of natural freshwater do not show much different on level of salinity unless there are additional discharged from surrounding environment into the river. The salinity value recorded from this study is low thus it would not have huge impact in the environment.

Total dissolved solid (TDS) is a measure of the amount of material dissolved in a water sample. This material includes dissolved minerals and organic matter but can also include contaminants. There is significant difference found between TDS with station. TDS is influenced by sampling location. TDS can vary spatially and temporally due to natural and anthropogenic factors such as climate, soil type, relief, land use and human activities (Augustijn et al., 2011). TDS in water supplies originated from natural sources, sewage, urban and agricultural run-off and industrial wastewater. Water containing TDS concentrations below 1,000 mg/l is usually acceptable but water with extremely low concentrations of TDS may also be unacceptable to consumers because of its flat, insipid taste to consumers. No recent data on health effects associated with the ingestion of TDS in drinking-water appear to exist (WHO, 2004).

Water quality index (WQI) provides relative indication of the quality of water. A river with WQI value in the ranges 0-59 is considered polluted, 60-80 is considered slightly polluted and 81-100 is considered the water body is clean. A higher value indicates a better quality of water (Suki et al., 1988; Rashid et al., 2013). From this study, the average WQI calculated at each sampling station for both sampling times are described as follows: Station S1 (94.29: Class I), S2 (94.86: Class I), S3 (90.23: Class II), S4 (85.45: Class II), S5 (87.32: Class II) and S6 (87.17: Class II) (Fig. 4). The results show that WQI obtained from this study fall under Class I (clean category). Overall, the WQI value of Tekala River was classified under Class II which indicated as clean. Generally, WQI tends to decrease from upstream to downstream of river due to increasing of pollutant. WQI recorded slightly reduced from station 1 to station 6.

**Biological parameter**

*E. coli* is mainly originated from the feces of warm blooded organism. This type of bacteria usually entered into water surface by run-off from agricultural and residential areas (Pearson et al., 1987; Inatsuka et al., 2010). From the first sampling, there is a decrease of *E. coli* from station 1 to station 6. Second sampling recorded less number of *E. coli*. There is correlation relationship found between *E. coli* and NH₃-N, temperature and DO. It is reported that fecal coliform levels were lower at higher temperature, high dissolved oxygen and high pH implying that one of the fecal coliform sources could be related to human recreational activities (Al-Badaii et al., 2013). This might explain the downtrend of bacteria from first station to the last station. Since this river is a recreational area, it probably potentially exposed to fecal contamination sources from human activities (swimming, camping) that cannot be detected by measuring physico-chemical parameters alone.

From the first and second samplings, there is a decrease of coliform from station 1 to station 6. Second sampling recorded less number of coliform. There is correlation relationship found between coliform with pH and BOD. Water with lower pH value helps in maintaining the survival of indicator bacteria (Parhad et al., 1974; Mara, 2013). A study done by Šolić and Krstulović (1992) also suggested that the optimal pH value
for coliform survival is between pH 6 and pH 7 and other pH value above or below these range can cause rapid decline in fecal coliform population. The bacteria population is influenced by favourable condition and food sources provided by the sediments for the bacteria (Davies et al., 1995). Lower temperature of surface water can cause rapid decline in coliform and E. coli population in the water (McDaniels et al., 1985; Pope et al., 2003).

**Macroinvertebrate**

Macroinvertebrates and water quality are interrelated to each other, as macroinvertebrates are a potential biological indicator of water quality. They are most frequently used in biomonitoring studies (Harikumar et al., 2014). Often many species within a community with varying sensitivities to stresses and relatively quick reaction times, resulting in a spectrum of graded, recognisable responses to environmental perturbation. The responses to different types of pollution have been established for many common species (Ollis et al., 2006). Stream communities are shaped by environmental influences at multiple spatial scales. Their distribution, interaction and adaptation can be influenced by abiotic factors which vary in space and time. Climate is one of the factors that influence population dynamics of aquatic insects (Nor Zaiha et al., 2015). In this study, the number of macroinvertebrate found is low. Most of the macroinvertebrate found belong to the same family. A study on surber net was conducted at the forested stream site by Wan Mohd Hafezul et al. (2016). They found very low species abundance was recorded by the surber net. Its small size makes it difficult to set on rough substrates in deep water and often results in the loss of large organisms that are fast enough to crawl out of the front of the sampler. Usually area disturbed within the surber net is exposed, providing an opportunity for aquatic fauna to escape (Wan Mohd Hafezul et al., 2016).

The most dominant insect found in all stations is water strider from family Gerridae. Based on the correlation test, there is relationship between abundance of macroinvertebrate and temperature. Water striders most prefer waters around 25 °C. Large numbers can indicate moderate, or slightly polluted water quality. In the upstream area, a few mayfly from order Ephemeroptera was caught in station 4. Ephemeroptera is a group of aquatic insects which is sensitive to environmental interference and can only survive in clean and oxygen-rich waters. Therefore, they are often considered as good biological indicators of water quality (Nor Zaiha et al., 2015). The most significant environmental factor affecting life-history patterns, especially growth rates and the seasonal timing of aquatic insects, is water temperature. Mayfly is widely distributed in peninsular Malaysia. It inhabits upstream rivers of peninsular Malaysia. Low water temperature seemed to favor mayfly abundance. Predators, particularly fish, play a role in the mayfly community (Suhaila et al., 2016). In station 4, there is no fish found in fast flowing water thus mayfly might prefer the area. From station 3 to 6, there is no more mayfly found. Station 3 has fish community, thus this area might not be preferred by mayfly. There is less human contact from station 1 to station 3, where visitor usually stay in area station 4 to station 6.

There are several other organisms that were found in the river such as freshwater shrimp, freshwater crabs and water penny. Freshwater crabs are one of the most ecologically important macro-invertebrate groups in tropical inland waters worldwide. These strictly freshwater decapods are found in almost all clean freshwater bodies in the tropics from moist lowland forests to rugged mountains. Crabs live in rivers, streams,
waterfalls, wetlands, karsts, caves and many are semi-terrestrial. Almost all require pristine water conditions to survive and are excellent indicators of good water quality (Cumberlidge et al., 2009). Generally, it can be concluded that more abundant macroinvertebrate communities exist in the littoral zone than in the deeper river bed. The individual numbers are usually very low in the deep water zone which is dominated by continually shifting sand (Csányi et al., 2012). Type of food sources and physical properties of river determine the distribution of aquatic organisms (Cummins and Merritt, 1996).

Conclusions

From this study, physico-chemical parameters and biomonitoring analysis have been conducted to determine the status of water quality at Tekala River, Selangor. Water quality index for Tekala River were ranged from 85.45 to 94.86, thus indicated that the water quality status for both sampling times and six sampling stations are classified under Class I and Class II. It was found that there is significance different between ammoniacal nitrogen, pH and temperature. Correlation was also found between several physico-chemical parameters with biological parameter. From the Pearson’s correlation test, there is relationship between E. coli with ammoniacal nitrogen, temperature and DO. For coliform, there is relationship between coliform with pH and BOD. The data from this study was also compared with the National water quality standard (NWQS) and most parameters were classified under Class I to Class IIb, implying that this river is safe to be used for recreational purposes. It is recommended that further research on other parameters on this river to be conducted in order to get better assessment on the water quality.

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Conflict of interest. The authors confirm that this article content has no conflict of interest.

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