Modelling of Water Turbidity Parameters in a Water Treatment Plant

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Abstract

The high cost of chemical analysis of water has necessitated various researches into finding alternative method of determining portable water quality. This paper is aimed at modelling the turbidity value as a water quality parameter. Mathematical models for turbidity removal were developed based on the relationships between water turbidity and other water criteria. Results showed that the turbidity of water is the cumulative effect of the individual parameters/factors affecting the system. A model equation for the evaluation and prediction of a clarifier’s performance was developed:

Model: \[ T = T_0(-1.36729 + 0.037101 \cdot 10^{\text{pH}} + 0.048928t + 0.00741387 \cdot \text{alk}) \]

The developed model will aid the predictive assessment of water treatment plant performance. The limitations of the models are as a result of insufficient variable considered during the conceptualization.

Keywords

Performance evaluation, pH, Alkalinity, Mathematic model, Simulation

1. Introduction

Tebbutt (1983) describes Water as the World’s most important natural resource since without it life cannot exist and industry operate. Unlike many other resources, it has no substitute in many of its uses. Water plays an essential role in community development since a reliable supply is a prerequisite for establishing a permanent settlement.
There is a vast amount of water present in the earth and its’ surrounding atmosphere. About 7% of the earth’s mass is made up of water. 97% of this is found as saline water in oceans, about 2.3% is in the polar caps and only 0.7% exists in fresh water lakes, rivers, aquifers and in the atmosphere.

Water that is pure is not found in nature; even water vapour condensing in air contains solids and dissolved gases (Tebbutt, 1983). As it condenses and falls, it sweeps up other materials from the air, it becomes still more contaminated on reaching the ground, as it runs running over soil surface and percolate the soil strata.

Waste from human activities, either industrial or domestic, according to Tchobanoglous and Burton (1991), introduce even more pollutants, than any natural sources, into water bodies. Since the available water is seldom found in conditions that meet potable specifications, a treatment plant is clearly necessary to improve the water quality. Water supply to a community goes through the following stages; the community water demand is carefully estimated with allowances for population growth; the most suitable raw water source is identified and analysed; then a water treatment plant to effect the required changes is designed, constructed and operated along with it’s distribution network.

When online, a periodic review of plant performance is undertaken to ascertain if, or otherwise, the plant works according to prediction. Hammer (1973) and Cairncross (1980) both agree that record–keeping and periodic reviews of plant performance are necessary decision tools when the plant requires expansion or when operational problems arise.

The aim of this paper is to model water turbidity as function of several other parameters that will aid in assessing the performance of a water treatment plant over a period of time and to suggest ways of improving plant performance

2. Experiments

**Determination of Temperature**

The temperature of the sample of water was determined with the aid of thermometer, beakers, and electrodes. The water samples were collected in beakers and labelled appropriately to avoid mixing up. The thermometer is switched on and the electrode is dipped
into the beaker in turn, the display is allowed to attain a steady value in each and the readings are recorded.

**Determination of Turbidity**

This involves the use of turbid meter. The sample cells were washing with sample water, and was discarded and refilled with the same sample. The sides of the cells were wiped with a dry cloth. The power source of the turbid meter was switch on and adjusted to read the turbidity in NTU. The cell holder was opened and the sample cell place in it and the lid replaced. The reading shows up on a digital display and the units are in NTU.

**Determination of pH**

The pH of the sample was determined with aid of colometric comparator. Two cells were rinsed with sample water and shaken, one cell was filled to mark “B” and 10 drops of indicator were added. The second cell was filled to mark “A” and placed in the comparator on the side marked “blank”. Slide the colorimeter through the slot in the comparator until the colour in both cells appears identical. The number in the matching colour slot is the pH.

**Determination of Total Alkalinity**

The total alkalinity was determined using titration. The burette was filled alkalimetric reagent up to the zero mark. 100ml of water sample was transferred to a breaker and add 4 to 6 drops of helianthin indicator was added. The Colour changes from pink to yellow. Titrate with the alkalimetric reagent until orange colour appears. Titrate with the alkalimetric reagent until orange colour appears. Read the alkalinity directly from the burette

3. **Modelling for Turbidity Removal**

For a constant flow rate and a fixed set of conditions the suspended solids (and hence turbidity) removal is a fixed fraction of inlet suspended solid. Assuming also, in accordance with Belan (1988), that for mildly contaminated rivers the suspended solid is linear with turbidity i.e. that the particle size and type distribution is constant over a time period.

Then Turbidity removal

\[
\frac{T_0 - T}{T_0} = k_0
\]
where \( T = \text{turbidity of clarified water}, \ T_0 = \text{Turbidity of aerated water}, \ k_0 = \text{constant} \)

\[
1 - \frac{T}{T_0} = k_0, \ T/T_0 = 1 - k_0 = k', \ T = T_0 k' \tag{2}
\]

In reality operating conditions do vary. Therefore \( k' \) is only one of the function by which water quality may be measured. Other essential parameters according to Nikoladze et al (1989) are temperature \((t)\), pH and alkalinity \((\text{alk})\). It is well established that these parameters exhibit independent and cumulative effect on \( T \).

**Effect of Water Temperature**

Water molecules and impurity particles are in thermal Brownian motion whose intensity is directly proportional to temperature (Nikoladze et al, 1989). It is clear that the probability of collision of individual particles with one another and their consequent aggregation depend on their relative velocities i.e. on thermal Brownian motion (and therefore on water temperature).

Again, it shall be assumed that the temperature profile within the clarifier is constant and is equal to that of the clarified water samples.

\[
\frac{(T_0 - T)}{T_0} \sim t \tag{3}
\]

where \( t = \text{temperature in °C} \).

**Effect of Alkalinity**

Turbidity removal is preceded by and dependent on the formation of chemical flock. The equation of the overall process as simplified by Hammer (1975) is

\[
\text{Al}_2(\text{SO}_4)_3 + 3\text{Ca(HCO}_3)_2 \rightarrow 2\text{Al(OH})_3 + 3\text{CaSO}_4 + 6\text{CO}_2
\]

Obviously, good coagulation is dependent on the presence of sufficient alkalinity \((\text{HCO}_3^-)\) and therefore

\[
\frac{(T_0-T)}{T_0} \sim \text{alk} \tag{4}
\]

where \( \text{alk} = \text{alkalinity of water in mg/l as CaCO}_3 \).

**Effect of pH**

Suspended matters in water are surface–charged particles and it is the function of the coagulant to neutralize the charges. Different particles types have been seen to have a particular pH at which the net charge on them is zero and coagulation optimum (Nikoladze et al 1989). This pH is the isoelectric point \((\text{pH}_{ie})\). A large difference between pH of the water
medium and pH$_{is}$ confers greater anticoagulation properties. For clay and humus pH$_{is}$ = 7.1 and 7.0 respectively. This effect is confirmed by graphs obtained by Tebbutt (1983) where turbidity (color) removal is reduced as the water pH deviates from an optimum value of about 7.0.

Therefore it can be concluded that:

\[(T_0 - T)/T_0 \sim 1/(\text{pH} - \text{pH}_{is})\] (5)

Assuming pH$_{is}$ as 7 then pH - 7 = $\Delta$PH has a positive value.

\[(T_0 - T)/T_0 \sim 1/( - \Delta\text{PH})\] (6)

pH = - log[H$^+$], therefore a more appropriate form of the equation would be:

\[(T_0 - T)/T_0 \sim 1/10^{-\Delta\text{PH}}\] (7)

An expression similar to this was used by Belan (1988) to express the lime dosage required to bring about a desired change in the pH of water to pH$_{is}$ of it’s contaminants.

Introducing constants into equations 3, 4 and 7:

\[(T_0 - T)/T_0 = k_1/10^{-\Delta\text{PH}}\] (8)

\[(T_0 - T)/T_0 = k_2t\] (9)

\[(T_0 - T)/T_0 = k_3\text{alk}\] (10)

Other factors, such as organic content do exert influence on turbidity removal; their contribution can be accounted for by the introduction of corrective coefficient.

Assuming that hydrodynamic conditions are approximately constant, the influences of temperature, pH, and alkalinity on the turbidity of water in a clarifier basin may be additive or multiplicative (Lucey, 2000). However a change in one will cause dis-equilibrium in the overall turbidity. Such changes are accounted for by the various constants. Using the additive method:

\[(T_0 - T)/T_0 = k_0 + k_1/10^{-\Delta\text{PH}} + k_2t + k_3\text{alk}\] (11)

from which:

\[T = T_0(1 - k_0 + k_1/10^{-\Delta\text{PH}} + k_2t + k_3\text{alk})\] (12)

Because 1 - $k_0$ is a constant, let us say $k_0$ in place of:

\[T = T_0(k_0 + k_1/10^{-\Delta\text{PH}} + k_2t + k_3\text{alk})\] (13)

The values of these constants maybe determined using the least – square method for multiple regression as outlined below adapted from Stroud (1995).

Let $T_i$ be an observed value of outlet turbidity; the value predicted by the model equation is then:
Then the error of prediction $e_i$ will be given by:

$$E_i = T_i - T_{0i}(k_0 - (k_i/10^{APHI}) - k_2T_i - k_3alk_i)$$

(15)

The square of the error is:

$$E_i^2 = [T_i - T_{0i}(k_0 - (k_i/10^{APHI}) - k_2T_i - k_3alk_i)]^2$$

(16)

For all data sets, by summing:

$$\Sigma E_i^2 = \Sigma [T_i - T_{0i}(k_0 - (k_i/10^{APHI}) - k_2T_i - k_3alk_i)]^2$$

(17)

Expanding the brackets gives:

$$\Sigma E_i^2 = \Sigma T_i^2 + k_0^2\Sigma T_{0i}^2 + k_1^2\Sigma T_{0i}^2/(10^{APHI})^2 + k_2^2\Sigma T_{0i}^2T_i^2 + k_3^2\Sigma T_{0i}^2alk_i^2 - 2k_0k_1\Sigma T_{0i}^2/(10^{APHI}) - 2k_0k_2\Sigma T_{0i}^2T_i - 2k_0k_3\Sigma T_{0i}^2alk_i + 2k_1k_2\Sigma T_{0i}^2/(10^{APHI}) + 2k_1k_3\Sigma T_{0i}^2alk_i(10^{APHI}) + 2k_2k_3\Sigma T_{0i}^2T_ialk_i - 2k_0\Sigma T_iT_{0i} + 2K_1\Sigma T_iT_{0i}/(10^{APHI}) + 2k_2\Sigma T_{0i}T_iT_{0i} + 2k_3\Sigma T_{0i}T_{0i}alk_i$$

Partial differentiation of this equation with respect to the constants $k_0$, $k_1$, $k_2$, and $k_3$ gives:

$$\delta(\Sigma E_i^2)/\delta k_0 = 2\Sigma T_i(\delta T_i/\delta k_0) + 2k_0\Sigma T_{0i}^2 - 2k_1\Sigma T_{0i}^2/(10^{APHI}) - 2k_2\Sigma T_{0i}^2T_i - 2k_3\Sigma T_{0i}^2alk_i - 2\Sigma T_iT_{0i}$$

(18)

$$\delta(\Sigma E_i^2)/\delta k_1 = 2\Sigma T_i(\delta T_i/\delta k_1) + 2k_1\Sigma T_{0i}^2/(10^{APHI}) - 2k_0\Sigma T_{0i}^2/(10^{APHI}) + 2k_2\Sigma T_{0i}^2T_i/(10^{APHI}) + 2k_3\Sigma T_{0i}^2alk_i/(10^{APHI}) + 2\Sigma T_iT_{0i}$$

(19)

$$\delta(\Sigma E_i^2)/\delta k_2 = 2\Sigma T_i(\delta T_i/\delta k_2) + 2k_2\Sigma T_{0i}^2T_i - 2k_0\Sigma T_{0i}^2T_i + 2k_3\Sigma T_{0i}^2alk_i + 2\Sigma T_iT_{0i}$$

(20)

$$\delta(\Sigma E_i^2)/\delta k_3 = 2\Sigma T_i(\delta T_i/\delta k_3) + 2k_3\Sigma T_{0i}^2alk_i - 2k_0\Sigma T_{0i}^2alk_i + 2k_2\Sigma T_{0i}^2T_ialk_i + 2\Sigma T_iT_{0i}$$

(21)

Equating all the derivatives to zero yields the following system of equations

$$k_0\Sigma T_{0i}^2 - k_1\Sigma T_{0i}^2/(10^{APHI}) - k_2\Sigma T_{0i}^2T_i - k_3\Sigma T_{0i}^2alk_i = \Sigma T_iT_{0i}$$

(22)

$$k_0\Sigma T_{0i}^2/(10^{APHI}) - k_1\Sigma T_{0i}^2/(10^{APHI}) - k_2\Sigma T_{0i}^2T_i/(10^{APHI}) - k_3\Sigma T_{0i}^2alk_i/(10^{APHI}) = \Sigma T_iT_{0i}/(10^{APHI})$$

(23)

$$k_0\Sigma T_{0i}^2T_i - k_1\Sigma T_{0i}^2T_i/(10^{APHI}) - k_2\Sigma T_{0i}^2T_i^2 - k_3\Sigma T_{0i}^2T_ialk_i = \Sigma T_iT_{0i}T_i$$

(24)

$$k_0\Sigma T_{0i}^2alk_i - k_1\Sigma T_{0i}^2alk_i/(10^{APHI}) - k_2\Sigma T_{0i}^2T_ialk_i - k_3\Sigma T_{0i}^2alk_i^2 = \Sigma T_iT_{0i}alk_i.$$ 

(25)

The various effects of pH, temperature and alkalinity on turbidity removal may also be combined in a multiplicative or exponential manner since no evidence exists to suggest the
relationship used. Following from the same premise as before, a possible second model equation such as $61$ could be developed.

$$T = k_0(T_0 \cdot 10^{k_1 \Delta pH \cdot k_2 \cdot \text{alk}^{k_3}})$$   \hspace{1cm} (26)

Comparison of the models results indicates their extent of validity. The constants $k_0$, $k_1$, $k_2$, $k_3$ could be obtained from the solution of the generated $4 \times 4$ matrix.

The alternatively the developed models can be solved software packages such as Polymath3 (Himmelblau, 1996). The package was used to find the coefficients in both models. The developed models are presented below:

$$T = T_0(-1.36729 + 0.037101 \cdot 10^{\text{pH}} + 0.048928 \cdot t + 0.00741387 \cdot \text{alk}) \quad \text{(Model 1)}$$

$$T = 2.20673 \cdot 10^{-11} \cdot T_0 \cdot 10^{0.045606 \Delta pH \cdot 5.35007 \cdot \text{alk}^{1.55059}} \quad \text{(Model 2)}$$

The graphical error analysis technique was then used as the criteria for selecting the best-fit model.

4. Results

The results of the various experimental methods are as presented in Table 1.

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</table>
The simulated results from the models are presented in figure 1.

![Figure 1. Comparative analyses of the experimental and simulated values from models 1 and 2](image_url)

5. Discussions

Results from table 1 showed that turbidity in the treated water fell short of standards (5NTU max.) on one occasion (5.75NTU) but the mean value of 1.56NTU is well within the guide line value (WHO, 1985). The cause of this is most likely the high $T_0$ and the consequent clarifier’s overload. This may also suggest that the clarifier was overdue for desludging.

Mathematical modeling of the clarifier’s performance based on additive and multiplicative models using the pre-determined parameters gave models 1 and 2 respectively.

Model 1: $T = T_0(-1.36729 + 0.037101x_{10^{pH}} + 0.048928t + 0.00741387.alk)$

Model 2: $T = 2.20673 \cdot 10^{-11}T_0 \cdot 10^{0.045606_{pH}} \cdot t^{5.35007} \cdot \text{alk}^{1.55059}$

Simulation results of the models showed that model 1 to a large extent will give a better turbidity prediction. It showed also that factors affecting turbidity values are mainly independent in operation. The turbidity of water is the cumulative effect of the individual parameters/factors affecting the system.
From model 1 the change in turbidity caused by one of the independent variables is the cumulative effects of individual contributions of $T_0$, $t$, and alk. In model 2 the relationship of the turbidity parameters is as the synergy or interdependence of the variables. The nature of the model 1 equation gives turbidity change a dimensionless significance. Therefore $k_2$ the coefficient of temperature $t$ will have a unit of $1/^\circ C$. It could be defined as the change in ratio of inlet and outlet turbidity caused by changing the temperature of the clarifier by $1^\circ C$.

The discrepancies between the experimental and simulated values could be attributed to the assumptions made during the formulating of the model. The numbers of variables considered were quite small. The influence of human activities, impact of environmental pollution/contamination etc. on the coagulation and settling processes and much other need to be considered to enhance the reliability of the model.

**Conclusions**

Developed a model equation for the evaluation and prediction of a clarifier’s performance as is presented below:

(Model 1) $T = T_0(-1.36729 + 0.037101 \cdot 10^{\rho H} + 0.048928 \cdot t + 0.00741387 \cdot alk)$

The limitations of the models are as a result of insufficient variables considered during the conceptualization process.

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References


