A standard test for filter media cleanliness

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Abstract

Rapid sand filters are expected to produce clean, safe water, without interruption, for many years. Unfortunately, filters very often develop some problems during this time, most of which only become apparent when the damage is already done. Routine measurement of filter media cleanliness could provide the necessary early warning, which is the reason why the American Water Works Association (AWWA) suggested a media cleanliness test for inclusion in a structured filter assessment programme at drinking water treatment plants. After performing such assessments at 3 South African water treatment plants, the authors found that the results were not consistent, the turbidity could not be measured easily and the guideline values seemed to be excessively conservative. This led to an investigation to find a method for *stripping* the filter deposits from the media grains with an easy, reproducible method, and for *characterising* the stripped deposits.

After a series of tests on various filter media, using 6 stripping methods, 2 methods were identified that met the criteria in the first objective. The first, a magnetic stirrer method is a mechanical agitation method and the second, referred to as the cylinder inversion method, is a manual agitation method. Each of these methods was chosen on the basis of their operator and speed-of-agitation independence.

The criteria in the second objective were met by an in-depth suspended solids (SS) analysis performed on the filter media residue, with the total mass of solids removed from the media quantitatively separated on the grounds of acid solubility and volatility at 550°C. *Standard Methods* (1985) 209C and 209D were applied to the sample, with and without acid addition, in order to characterise the total filter media residue using 4 groupings: Soluble, non-volatile; soluble, volatile; non-soluble, non-volatile; and non-soluble, volatile.

By meeting the criteria of these 2 objectives, the previous non-specific suggestions in the literature have been improved to suggest the performing of tests in a well-specified, uniform way, the results of which can be internally compared.

Keywords: filter deposits, filter residue, filter media, cleanliness, floc retention

Introduction

In almost all South African water treatment plants, rapid sand filtration is the backbone of the process, providing the primary barrier against turbidity and protozoan cysts and oocysts in the final water.

Rapid sand filters almost inevitably develop some technical problems during the many years they are expected to continuously produce clean, safe water. With time, it is observed that media is lost, mudballs are formed, cracks appear, media grains grow by chemical deposition and backwash rates gradually decline due to, e.g. worn pump impellers and leaking valves. Unfortunately, most of these problems become apparent when the damage is already done. However, in all these cases an early warning is given by filter media that are not properly cleaned after every backwash cycle. If the cleanliness of the media grains is routinely measured, the potential for some of the more serious problems could be detected at an early stage and the problem possibly arrested before serious damage is done.

It is such thinking that led the American Water Works Association (AWWA) to include a media cleanliness test (called a floc retention test in their publications) in a battery of tests suggested for a structured filter assessment programme at drinking water treat-

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ment plants (AWWA, 2000). The test is simply described as taking approximately 50 g of filter sand, adding 100 m ℓ of water, shaking it vigorously and decanting the resultant suspension into a beaker. After 5 repetitions, the turbidity of the suspension is measured and reported as NTU (nephelometric turbidity units) / 100 g of sand. A value of 30 to 60 NTU indicates a clean bed, 60 to 120 NTU indicates a slightly clogged bed, 120 to 300 NTU indicates a clogged filter with mudball formation potential, and values above 300 NTU indicate a serious problem with highly probable mudball formation.

The Water Research Group at the Rand Afrikaans University was involved in such filter assessment programmes at 3 South African water treatment plants during 2000 and 2001 and performed this media cleanliness test with all the other prescribed tests (Ceronio et al., 2002a;b). Some difficulties quickly became apparent. Results were not consistent, the turbidity could not be measured easily and the guideline values seemed to be excessively conservative. The media cleanliness test thus had to be improved and standardised before it could be widely adopted as a comparison standard. The project comprised 2 different parts, which will be reported on in this paper:

- To find a method for stripping the filter deposits from the media grains with an easy, reproducible method, and
- To find a method for characterising the stripped deposits in a meaningful way to provide some clues for corrective action should the media be clogged.

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Experimental work

Sample and media preparation

The filter media are sampled according to the needs of the test. If a combined sample through the entire bed is required, a core sampler can be used. If media at a specific depth are required, undisturbed samples have to be taken upon careful excavation of the bed from top to bottom. Because the filter deposits can be surprisingly "patchy", it is always better to sample at 3 or more locations where the same results are anticipated and to make sample composites. The plant from which the media for this investigation was obtained was selected on the basis of the type of raw water treated, i.e. high organic load, high algal mass content and sewage contribution. It is generally known that water of this type would cause media to be "sticky" and difficult to clean. Hence, this choice of media ensures that the methodology would be relevant for all types of treatment plant media.

After sampling, the entire sample has to be homogenised without stripping the filter deposits prematurely from the media grains. For this study, a riffle splitter (such as used in standard engineering soils laboratories) was used to split the sample 5 consecutive times. Thereafter, the sample was sealed in a plastic bag and refrigerated until analysis, which was always done within 5 d of sampling.

For all the tests described further on, 50 g of sample was required. To obtain exactly 50 g of media, small quantities had to be added or taken away a few times, leading to possible disturbance of the filter deposit; a 60 ml measuring cup was used to facilitate this. In this way, a volume of sample could be taken and weighed in one easy step. It was also necessary to take into account the moisture content of the sample. This is done by simply weighing a separate sample of the same media both before and after drying to obtain a mass difference and, thereby, the moisture content (expressed as a percentage) of the sample. After analysis, the final values were mathematically adjusted (taking moisture content into account) to exactly 50 g of media.

Initial screening of media strip tests

Three methods were devised with mechanical agitation, which would be operator-independent, namely the shaker table method, the jar test method and the magnetic stirrer method. In view of the fact that not all treatment plants had access to this equipment, 3 further manual agitation tests were devised which could be performed with the absolute minimum of laboratory equipment, namely the hand-swirl method, the cylinder inversion method and the violent shake method. Three different operators repeated these 3 methods, to check for degree of operator dependency. The same media sample was subjected to all 6 tests.

Shaker table method

Two media samples were placed in 250 m ℓ Erlenmeyer flasks, each with 100 m ℓ tap water. One flask was shaken at 55 U/min and the other at 200 U/min, using a ROTABIT orbital-swinging shaker, for 30 s. The resultant suspensions were drained off into 2 × 500 m ℓ Erlenmeyer flasks and the procedure was repeated 4 more times, resulting in 2 × 500 m ℓ suspensions.

Jar test method

Two media samples were placed in 1 000 ml beakers with 100 ml tap water each. One sample was stirred at 30 r/min and the other at 200 r/min, both using a LEETECH LTS-630III paddle stirrer, for

30 s. The resultant suspensions were drained off into 2×500 m ℓ Erlenmeyer flasks. The procedure was repeated 4 more times to obtain 2×500 m ℓ suspensions.

Magnetic stirrer method

Two media samples were placed in 1 000 m ℓ beakers with 100 m ℓ tap water each. The samples were then stirred using a HANNA HI 290N magnetic stirrer. One sample was stirred at slow speed and one at high speed (these speeds were difficult to define numerically, even though they are measured in rpm units) for 30s. The resultant suspensions were drained off into 2 × 500 m ℓ Erlenmeyer flasks. The test was repeated 4 more times to obtain 2 × 500 m ℓ suspensions.

Hand-swirl method

A media sample was placed in a 500 m ℓ Erlenmeyer flask with 100 m ℓ tap water. The flask was swirled with a gentle wrist turning action for 30s and the resultant suspension drained off into a clean 500 m ℓ Erlenmeyer flask. This procedure was performed 4 times to obtain a single 500 m ℓ suspension for each sample.

Cylinder inversion method

A media sample was placed in a 250 m ℓ measuring cylinder with $100\,\text{m}\ell$ clean water. The cylinder was sealed at the top and inverted 20 times, as illustrated in Fig. 1 below. After every inversion, the test was momentarily paused to allow the media to settle at the bottom. The resultant suspension was drained off into a 500 m ℓ Erlenmeyer flask. The procedure was repeated 4 more times to obtain a 500 m ℓ suspension.

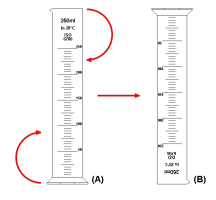


Figure 1

Illustration to show a single inversion of the measuring cylinder, i.e. from position (A) to position (B)

Violent shake method

A media sample was placed in a 500 m ℓ Erlenmeyer flask with 100 m ℓ clean water. The flask was vigorously shaken for 30 s and the resultant suspension drained off into a clean 500 m ℓ Erlenmeyer flask. The procedure was repeated 4 more times to obtain a single 500 m ℓ suspension.

Turbidity measurement

Turbidity, measured using a HACH ratio turbidimeter (Model 18900: range 0-200 NTU) and expressed as NTU/100 g media, was the response parameter used to compare the 6 methods. However, the suspension of filter residue in the samples presented 2 problems for turbidity measurement, namely that the relatively large particles caused some "bounce" in the turbidimeter reading, and that the

TABLE 1 Single-factor ANOVA results for 6 methods, testing for the dependence on operator/ speed, at α = 0.01

		mean	std. deviation	c.o.v.	F value	F _{crit} @ α=0.01
Hand swirl	Operator A Operator B	66.4 32.7	3.2 2.4	4.8 7.2	429.4	5.49
	Operator C	30.9	3.5	11.3		
Violent shake	Operator A Operator B Operator C	135.5 640.1 460.8	18.6 153.9 167.0	13.7 24.0 36.2	37.8	
Cylinder inversion	Operator A Operator B Operator C	98.2 105.1 119.6	7.6 12.9 11.5	7.7 12.2 9.7	10.1	
Orbital shaker	Slow Fast	43.8 66.4	3.7 5.4	8.5 8.1	118.5	8.29
Jar tester	Slow Fast	70.2 83.1	7.3 13.7	10.5 16.5	6.9	
Magnetic stirrer	Slow Fast	132.4 131.1	24.4 13.3	18.4 10.1	0.022	

suspension settled fairly quickly in the sample cell. Stirring the sample vigorously, then transferring it rapidly into the sample cell and immediately into the turbidimeter overcame these problems.

In addition, the first turbidity reading was taken after 10 s with consecutive readings every 5 s up to 30 s, since significant settling a flocculation is not expected to occur over 30 s. The average of these 5 readings is taken as the representative turbidity.

The samples also required dilution before they were within the measuring range of the turbidimeter, therefore requiring the use of a dilution factor, as follows:

Dilution factor =
$$\frac{m\ell \text{ sample} + m\ell \text{ clean water}}{m\ell \text{ sample}}$$
 (1)

Total NTU = measured NTU
$$\times$$
 dilution factor (2)

NTU per
$$100g = \frac{100g}{\text{mass of media sample (g)}} \times \text{total NTU (3)}$$

Comparison of results

For all tests with mechanical agitation, 2 different speeds were selected, and 10 identical samples tested at each speed. For each method, there were therefore $2\times10=20$ tests performed. For the tests with manual agitation, 3 different operators each tested 10 identical samples, yielding $3\times10=30$ tests for each method. For these initial screening tests, only one response parameter was measured, namely the turbidity of the supernatant, expressed as NTU/100 g of media, which was determined using Eqs. (1) to (3) above (Spaleck, 2001).

Single factor analysis-of-variance (ANOVA) tests were performed for each method and the results are summarised in Table 1. When comparing the statistical results, the null hypothesis, i.e. the method is operator- or speed-independent, is rejected at the α =0.01 level of significance when the F_{crit} value is larger than the calculated F value. The value for F_{crit} is read from a table of critical values of the F distribution ($F_{0.01}$), using the degrees of freedom, $v_1(k-1)$ and $v_2(k(n-1))$, where k represents the number of variables tested and n represents the number of tests performed for each variable (Johnson, 1994). Therefore, in terms of mechanical agitation, the jar tester and magnetic stirrer methods showed speed independence, and the magnetic stirrer method was chosen as the favoured mechanical agitation method due to the very low F value for the results. However, none of the manual agitation methods were found to statistically show complete operator independence and, therefore, other generated results need to be compared.

The hand-swirl method was rejected due to the very high F value for the results. The violent shake method was also rejected as it was shown, by plotting the turbidity for consecutive repeats (Fig. 2), that there was a trend for the operator to tire when performing consecutive tests and, therefore, inconsistent results were obtained. Therefore, the cylinder inversion method was chosen as the favoured manual agitation method. In addition, the mean value obtained using the cylinder inversion method is closer to that obtained using the magnetic stirrer method.

The two favoured methods were then subjected to further testing and refinement in the next section.

Refinement of the cylinder inversion method

Three experimental variables in this method were changed, resulting in 7 tests being devised. The variables tested were the number of inversions, the size of the cylinder and the ratio of the number of inversions to the number of washes. The same media sample was subjected to all 7 tests.

TABLE 2 Single-factor ANOVA results for 7 cylinder inversion methodologies

		Mean	Std. deviation	C.O.V.	F value	F _{crit} @ a=0.01
No. of inversions	10 20 40	597.0 774.0 790.7	161.1 32.0 74.9	27.0 4.1 9.5	3.2	10.92
Cylinder size	250 ml 500 ml	755.0 876.3	16.0 104.7	2.1 11.9	3.9	21.18
Inversion:wash ratio	20 :5 50 :2	787.3 786.7	56.6 52.5	7.2 6.7	0.00022	

Violent shake method turbidities

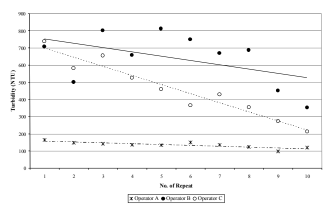


Figure 2

Violent shake method turbidity results for 3 operators, plotted with trendlines. This illustrates that the turbidity decreases over consecutive tests and, therefore, that the operators tire.

Number of inversions

Three media samples were placed in 250 m ℓ measuring cylinders, each with 100 m ℓ tap water. One flask was inverted 10 times, the second 20 times, and the third 40 times. The resultant suspensions were drained off into 3 \times 500 m ℓ Erlenmeyer flasks and the procedure was repeated 4 more times, resulting in 3 \times 500 m ℓ suspensions.

Size of the measuring cylinder

One media sample was placed in a 250 m ℓ measuring cylinder, and a second media sample was placed in a 500 m ℓ measuring cylinder, each with 100 m ℓ tap water. Each sample was inverted 20 times and the resultant suspensions drained off into 2 × 500 m ℓ Erlenmeyer flasks. The procedure was repeated 4 more times to obtain 2 × 500 m ℓ suspensions.

Ratio of number of inversions to number of washes

Two media samples were placed in 500 m ℓ measuring cylinders with 100 m ℓ tap water each. One sample was inverted 20 times and the other 50 times. The resultant suspensions were drained off into 2 × 500 m ℓ Erlenmeyer flasks. The test was repeated 4 times for the sample that was inverted 20 times and once more for the sample that was inverted 50 times, resulting in 1 × 500 m ℓ suspension and 1 × 200 m ℓ suspension respectively.

Comparison of cylinder inversion method results

For the first variable tested, i.e. the number of inversions, 3 identical samples were tested for each number of inversions. Therefore, there were $3 \times 3 = 9$ tests for this variable. For the second and third variables tested, i.e. the size of the measuring cylinder and the ratio of the number of inversions to number of washes, 2 different sizes and ratios were used respectively and 3 identical samples were tested for each. Therefore, there were $2 \times 3 = 6$ tests for each of these 2 variables.

Single-factor ANOVA tests were performed for each method, with the null hypothesis stating that there is no difference in the results due to a change in the variables. The results of these analyses are summarised in Table 2.

The results for all 3 tests were found to comply with the null hypothesis and, therefore, comparisons of the coefficient of variation (c.o.v.) results were performed to choose which method was favoured.

The c.o.v. value for the 20 inversions was found to be considerably lower than the other 2 values (just under half of that of the 40 inversions, with the 10 inversions value nearly 7 times greater). It was, therefore, concluded that 20 inversions of the measuring cylinder should be used for the final method.

The c.o.v. value for the 500 ml measuring cylinder was found to be almost 6 times that of the 250 ml measuring cylinder and, therefore, the 250 ml measuring cylinder was selected for use in the final method.

Both the c.o.v. values for the inversion:wash ratio methods were approximately 7 and, therefore, further analysis was necessary. After further consideration, it was concluded that a 500 ml supernatant would prove to be better, since a dilution factor for a 200 ml supernatant would be greater than that for a 500 ml supernatant and, therefore, may result in less reliable results.

Therefore, the optimal cylinder inversion method would make use of a 250 m ℓ measuring cylinder, with the sample inverted 20 times and a total of 500 m ℓ supernatant being produced, i.e. the sample would be washed 5 times.

Refinement of the magnetic stirrer method

Four experimental variables in this method were changed, resulting in 8 tests being devised. The variables tested were the size of the beaker, the stirring speed, the stirring duration and the size of the magnetic stirrer. The same media sample was subjected to all 8 tests.

Size of the beaker

One media sample was placed in a 250 m ℓ beaker (internal diam. 65 mm) and a second in a 500 m ℓ beaker (internal diam. 75 mm), each with 100 m ℓ tap water. Each sample was stirred at high speed for 60 s using a large magnetic stirrer (see Fig. 3 below). The resultant suspensions were drained off into 2 × 500 m ℓ Erlenmeyer flasks and the procedure was repeated 4 more times, resulting in 2 × 500 m ℓ suspensions.

Stirring speed

Two media samples were placed in 250 m ℓ beakers, each with 100 m ℓ tap water. One sample was stirred at slow speed and the other at high speed, both for 60s, using a large magnetic stirrer (Fig. 3 below). The resultant suspensions were drained off into 2×500 m ℓ Erlenmeyer flasks. The procedure was repeated 4 more times, resulting in 2×500 m ℓ suspensions.

Stirring duration

Two media samples were placed in 250 ml beakers, with 100 ml tap water each. One sample was stirred for 30s and the other for 60s, both on high speed and using a large magnetic stirrer (Fig. 3 below). The resultant suspensions were drained off

into 2×500 m ℓ Erlenmeyer flasks and the procedure was repeated 4 more times, resulting in 2×500 m ℓ resultant suspensions.

Magnetic stirrer size

Two media samples were placed in 250 m ℓ beakers, each with 100 m ℓ tap water. Each sample was stirred for 60s at high speed, one using a small magnetic stirrer and one using a large magnetic stirrer (see Fig. 3 below for size classifications). The resultant suspensions were drained off into 2 × 500 m ℓ Erlenmeyer flasks. The procedure was repeated 4 more times, resulting in 2 × 500 m ℓ suspensions.

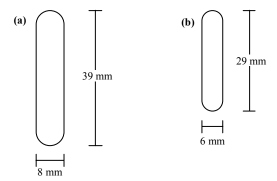


Figure 3

Illustration of size differences of magnetic stirrers used for the refinement of the magnetic stirrer methodology. a = large magnetic stirrer; b = small magnetic stirrer.

Comparison of magnetic stirrer method results

For each variable tested, i.e. the size of the beaker, the stirring speed, the stirring duration and the magnetic stirrer size, 2 different sizes, speeds, durations and stirrer sizes were used respectively and 3 identical samples were tested for each. Therefore, there were $2 \times 3 = 6$ tests for each of the 4 variables.

Single-factor ANOVA tests were performed for each method, with the null hypothesis stating that there is no difference in the results due to a change in the variables. The results of these analyses are summarised in Table 3.

As in the case of the cylinder inversion method, the results from all 4 tests were found to comply with the null hypothesis and, therefore, comparisons of the c.o.v. results were necessary to select which method was favoured.

TABLE 3 Single-factor ANOVA results for 8 magnetic stirrer methodologies

		Mean	Std. deviation	c.o.v.	F value	F _{crit} @ α=0.01
Beaker size	250 ml 500 ml	1476.5 1431.0	139.3 154.1	9.4 10.8	0.1	98.50
Stirring speed	Slow Fast	1110.5 1183.5	103.9 156.3	9.4 13.2	0.3	
Stirring duration	30 s 60 s	799.5 1230.5	101.1 71.4	12.6 5.8	24.2	
Stirrer size	Small Large	1297.0 1434.0	14.1 45.3	1.1 3.2	16.7	

Comparing the c.o.v. values for the beaker size tests, there is very little difference between the 2 values. However, the 250 ml beaker was selected for the final method as it had the lower c.o.v. value.

When comparing the results from the stirring speed tests, the c.o.v. values for the 2 were found to be very similar and, once again, the test with the lower c.o.v. value, namely the slow stirring speed, was chosen for the final method.

At the a=0.01 level of significance there was no difference between the results of the stirring duration tests. However, at the a=0.05 level of significance ($F_{0.05}$ = 18.5), there is a difference and the null hypothesis is rejected. The c.o.v. values also supported the statistics, with the c.o.v. value of the 30s tests being more than double that of the 60s tests. Therefore, a 60s stirring duration was selected for the final method.

The c.o.v. values for the stirrer size methods were both very low and, therefore, any stirrer size could be selected. However, the mean value for the large stirrer tests was closer to the mean values for the beaker size methods, which in turn showed little variation. In terms of practical considerations, the large magnetic stirrer is also easier to handle and more difficult to lose or misplace. Therefore, the large stirrer size was chosen for the final method.

Therefore, the optimal magnetic stirrer method would make use of a 250 m ℓ beaker, with the sample stirred at a slow speed for 60 s, using a large magnetic stirrer.

Comparison of the cylinder inversion method with the magnetic stirrer method

In taking the mean values obtained from each set of refinement tests, it can be seen (Table 4 below) that the magnetic stirrer method is more effective in removing the solids deposited on the filter media than the cylinder inversion method. However, not all laboratories may have the equipment required for this method readily available and, therefore, the cylinder inversion method could be used in such cases.

Characterising the filter grain residue

In order to determine the nature of the solids removed during the preceding tests, a number of simple tests were attempted. The basic premise was that the total mass of solids removed from the grains (measured as suspended solids (SS) in the standard way) could be

TABLE 4 Comparison of turbidity for 2 methodologies used to measure flocculation retention

Method used	Mean turbidity (NTU)	Difference in turbidity (%)	
Cylinder inversion Magnetic stirrer	767 1245	38	

TABLE 5 Suspended solids composition matrix					
	soluble	non-soluble	Total		
Volatile	Unknown A - B - C + D	Unknown B – D	Unknown A – C		
Non-volatile	Unknown C - D	Known (D) Eq. (7)	Known (C) Eq.(6)		
Total	Unknown A - B	Known (B) Eq.(5)	Known (A) Eq.(4)		

quantitatively separated on the grounds of acid solubility as well as volatility at 550°C. In this way, the total solids mass could be split into 4 fractions: Non-volatile and non-soluble, non-volatile and soluble, volatile and non-soluble, and volatile and soluble. A number of brief descriptions of these tests follow.

Total suspended solids (TSS)

A volume of 20 m ℓ to 30 m ℓ of the water sample was drawn off whilst vigorously stirred to prevent settling of the solids, and then filtered through a clean, weighed Millipore glass-fibre prefilter, using Millipore filtration apparatus. Total suspended solids (in mg/ ℓ) is then determined using *Standard Methods* (1985) procedure 209C and Eq. (4) below:

mg Total solids/
$$\ell = \frac{\left(A - B\right) \times 1000}{\text{Sample volume } \left(m\ell\right)}$$

where $A = mass\ of\ watchglass\ and\ filter\ with\ dried\ residue\ \big(mg\big)$

B = mass of clean filter (mg) + mass of watchglass (mg)

Non-soluble SS

"Non-soluble solids" is the term applied to the residue of total, suspended, or dissolved solids that remains after treatment with an acid solution.

A 32% dilution of hydrochloric acid (HCl) was further diluted by 20%, i.e. $25\,\text{ml}$ of 32% HCl is mixed for a minimum of 1 minute with $100\,\text{ml}$ water, and $10\,\text{ml}$ of this acid was mixed with $100\,\text{ml}$ of the resultant suspension. The same procedure as used for the TSS determination is then used to determine the total non-soluble solids (in mg/l) for the sample, together with Eq. (5) below:

mg Non-soluble solids/
$$\ell = \frac{(A-B)\times 1000}{\text{Sample volume } (m\ell)}$$

where A = mass of watchglass and filter with dried acid residue (mg)

$$B = mass of clean filter (mg) + mass of watchglass (mg)$$
 (5)

Non-volatile SS

"Non-volatile or fixed solids" is the term applied to the residue of total, suspended, or dissolved solids remaining after ignition for a specified time at a specified temperature. The mass loss on ignition is called "volatile solids". However, the determinations of non-volatile and volatile solids do not distinguish between organic and inorganic matter, as the loss on ignition cannot be confined to organic matter, but also includes losses due to decomposition or volatilisation of some mineral salts (*Standard Methods*, 1985).

The determination of the total non-volatile solids (in mg/ ℓ) was done using *Standard Methods* (1985) procedure 209D and Eq. (6) below:

mg Non-volatile solids/
$$\ell = \frac{(A - B) \times 1000}{\text{Sample volume } (m\ell)}$$
 (6)

where A = mass of watchglass and filter with dried residue before ignition (mg)

B = mass of watchglass and filter with dried residue after ignition (mg)

This same procedure was also used to determine non-volatile, non-soluble solids using the acid-sample residue filters and Eq. (7) below:

mg Non - volatile non - soluble solids/
$$\ell = \frac{(A-B)\times 1000}{Sample volume (m\ell)}$$

where A = mass of watchglass and filter with dried acid residue before ignition (mg)

B = mass of watchglass and filter with dried acid residue after ignition (mg)

The values obtained using Eqs. (4) to (7) can then be placed in a matrix as follows (Table 5) to determine various unknown values:

The components in Table 5 can roughly be interpreted as follows Haarhoff (2002):

Soluble, non-volatile originates from the carry-over of

chemical precipitates.

Non-soluble, non-volatile corresponds to inorganic par-

ticles present in the raw water.

Non-soluble, volatile bacterial and algal biomass,

organic detritus

Comparison of suspended solids results

Tests performed on filter sand and granular activated carbon (GAC)

Suspended solids analyses were performed on filter sand and GAC media, using the cylinder inversion method, for both before and after backwash samples. The samples analysed were taken from various filters at various visits to the plant so that some of the samples represented clogged filters (not recently regenerated or washed), some represented regenerated filters (as in the case of the GAC filters) and some represented recently washed filters.

In each case, samples were taken at 4 depths in the media, at 2 locations in the filter, and were evenly spaced through the profile of the filter. These samples were then combined to form a sample that represented the full profile of the filter.

Each analysis was repeated 3 times for each sample and the results were statistically analysed using a Q test at the 90% confidence level. This was used to determine if any questionable (or outlying) results should be rejected. In each case, the questionable result is rejected with 90% statistical confidence if the Q value obtained is larger than the $Q_{0.90}$ value (Fritz and Schenk, 1979).

Turbidity versus Total Solids

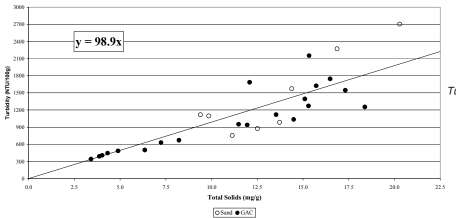


Figure 4
Turbidity vs. SS (mg/g) for various media types

After the questionable results had been discarded, the means of the remaining values were determined and used in Eqs. (4) to (7) and to draw up a matrix of SS composition (as in Table 5 above).

The TSS were measured in mg/ ℓ and then converted to mg/g using Eq. (8) below and plotted against turbidity (NTU/100g) for comparison of results (see Fig. 4):

$$\text{mg suspended solids } / g = \frac{\text{mg suspended solids } / \ell}{2} \times \frac{1}{\text{weight of media(g)}}$$
(8)

As can be seen from the graph above, the relationship between turbidity and SS (in mg/g) is very close to turbidity = $100 \times SS$ (mg/g).

Discussion and conclusions

The testing of 3 mechanical agitation and 3 manual agitation methods for the removal of solid deposits from filter media resulted in the choice of the magnetic stirrer method (mechanical agitation) and the cylinder inversion method (manual agitation) and the subsequent refinement of these 2 methods.

The tests performed to refine these 2 methods led to the following 2 procedures to be used for further testing of filter media cleanliness:

Magnetic stirrer method

- Obtain a homogenised 60 ml sample (approximately) of filtration media.
- 2. Weigh the sample and record this value in grams.
- 3. Perform a moisture content test by weighing a second sample of the sand, both before and after drying. The difference in weight divided by the weight before drying will give the fraction of moisture content of the sample. This value should be used to correct the weight in (2) above.
- 4. Place the sample in a 250 ml beaker.
- 5. Add 100 ml clean water to the sample.
- Using a large magnetic stirrer and magnetic stirring apparatus, continuously stir the media at slow speed for 60 s.
- Drain off the resultant suspension into a clean, separate 500 ml. Erlenmeyer flask, as quickly and accurately as possible.
- 8. Repeat Steps 5 to 7 an additional 4 times until the Erlenmeyer flask contains a 500 ml suspension.
- Continuously stir this suspension using additional magnetic stirring apparatus until further analyses can be performed.

Cylinder inversion method

- Obtain a homogenised 60 ml sample (approximately) of filtration media.
- 2. Weigh the sample and record this value in grams.
- 3. Perform a moisture content test by weighing a second sample of the sand, both before and after drying. The difference in weight divided by the weight before drying will give the fraction of moisture content of the sample. This value should be used to correct the weight in (2) above.
- 4. Place the sample in a 250 ml measuring cylinder.
- 5. Add 100 ml clean water to the sample.
- Seal the end of the measuring cylinder and invert the media 20 times, pausing between inversions to allow the media to settle at the bottom.
- Drain off the resultant suspension into a clean, separate 500 ml Erlenmeyer flask.
- 8. Repeat Steps 5 to 7 an additional 4 times until the Erlenmeyer flask contains a 500 ml suspension.
- Continuously stir this suspension using magnetic stirring apparatus until further analyses can be performed.

The turbidity and composition of the resultant suspensions can then be determined by means of a turbidimeter and SS analyses.

Characterisation of the filter media residue

The total SS removed from the media sample can be quantitatively separated on the grounds of acid solubility as well as volatility at 550°C, as follows:

- Whilst vigorously stirring the resultant suspension, draw off 20 ml to 30 ml and filter through a weighed glass-fibre prefilter, using filtration apparatus.
- 2. Using *Standard Methods* (1985) procedure 209C determine the total SS of the sample from this filtrate (*A*).
- Add 10 ml 6.4% hydrochloric acid to 100 ml resultant suspension and stir vigorously.
- 4. Draw off 20 ml to 30 ml of this acid-water solution, whilst stirring, and filter through a weighed glass-fibre prefilter, using filtration apparatus.
- 5. Using *Standard Methods* (1985) procedure 209C determine the total non-soluble solids of the sample from this filtrate (*B*).
- 6. Using the Standard Methods (1985) procedure 209D and the total SS and total non-soluble SS filtrates, obtained from the above 2 procedures, determine the total non-volatile solids (C) and the total non-volatile non-soluble solids (D) respectively.

Using the 4 values determined from the above methods (A, B, C) and (A, B, C), determine the unknown values in Table 5, thereby, characterising the filter media residue in terms of 4 values: soluble, non-volatile; soluble, volatile; non-soluble, non-volatile; and non-soluble, volatile.

Interpretation of the results

It is important to realise throughout that the numerical values obtained from the above tests are only relative. Aggressively stirring or inverting a small media sample in the laboratory can strip much more residue from the media than any backwash system possibly can. The total solids stripped from the media by the backwash system will therefore be much less. However, by now being able to perform the tests in a specified, reproducible way, the results obtained from a filter assessment programme can at least be internally compared, a significant improvement on the previous non-specific suggestions in the literature.

Recommendations

The focus of this paper is to suggest methods for sampling the media, stripping the residue from the media and characterising the filter residue, which was comprehensively reported on. Further work is required to provide guidelines for the interpretation of these results, and to provide typical benchmarks against which such results can be compared.

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