**Performance of a water defluoridation plant in a rural area in South Africa**

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**Abstract**

The fluoride concentration of a borehole water supply in a rural area (Madibeng Local Municipality, North West Province, South Africa) varies between 5 and 6 mg/ℓ. This water is therefore not suitable for potable purposes because its high fluoride concentration will cause mottling of the tooth enamel in children and fluorosis in adults (Holden, 1970; Bishop and Sancoucy, 1978). Therefore, the fluoride concentration should be reduced to less than 1.5 mg/ℓ to make the water suitable for potable purposes. The activated alumina and reverse osmosis processes are both processes that can be very effectively applied for water defluoridation. The activated alumina process, however, is considered to be a more simple and robust process for water defluoridation, especially in a rural area. No reduction in plant output was experienced over 6 service cycles. Therefore, it appears that fouling of the activated alumina should not be a problem. Plant output varied between 940 and 1 296 m³ to a fluoride breakthrough of approximately 2.0 mg/ℓ.

The high selectivity of activated alumina for fluoride compared relatively simple to use. The process can remove fluoride in the feed-water concentration range from 4 to 20 mg/ℓ to <1 mg/ℓ. The activated alumina process was therefore selected for water defluoridation in the study area.

The objectives of this paper are to report on the performance of the defluoridation plant during commissioning and on the economics of the process.

**Fluoride quality requirements**

The US Environmental Protection Agency has made the maximum allowable fluoride concentration for drinking purposes dependent on climatic conditions (Drinking Water Standards, 1962) because the amount of water consumed and, consequently, the amount of fluoride ingested, is temperature-related.

The maximum allowable concentrations are shown in Table 1 (National Interim Primary Drinking Water Regulations, 1975) as are the recommended fluoride concentrations established by the US Public Health Service (Drinking Water Standards, 1962).

The SA Bureau of Standards (1971) has recommended a limit of 1 mg/ℓ for fluoride in drinking water supplies and has set the maximum allowable concentration at 1.5 mg/ℓ.

**The activated alumina process**

Activated alumina (Al₂O₃) functions as an anion exchanger and the common anions are selected in the following order by acid concentration:

\[
\text{OH}^{-} > \text{PO}_{4}^{3-} > \text{F}^{-} > \text{SO}_{4}^{2-} > \text{SO}_{3}^{2-} > \text{NO}_{3}^{-} > \text{Br}^{-} > \text{Cl}^{-} > \text{NO}_{2}^{-}.
\]

The high selectivity of activated alumina for fluoride compared...
Alumina, in contact with fluoride ions, displaces the sulphate anions. Ion exchange (loading). Once this acidic form is brought into synthetic anion exchangers makes it extremely suitable for fluoride removal from water containing ions such as sulphates, chlorides and bicarbonates, which may compete for the limited adsorption sites.

A typical adsorption-regeneration cycle for the removal of fluoride may be presented as follows (Clifford et al., 1978).

Acidification. When neutral (water washed) alumina, which is represented as alumina, H$_2$O, is treated with sulphuric acid, acidic alumina is formed following the reaction:

$$\text{alumina.H}_2\text{O} + \text{H}_2\text{SO}_4 \rightarrow \text{alumina.H}_2\text{SO}_4 + \text{H}_2\text{O}$$

Ion exchange (loading). Once this acidic form is brought into contact with fluoride ions, they displace the sulphate anions.

$$\text{alumina.H}_2\text{SO}_4 + 2\text{NaF} \rightarrow \text{alumina.H}_2\text{F}_3 + \text{Na}_2\text{SO}_4$$

Backwashing. This is carried out after loading to remove suspended solids, to redistribute the alumina particles and break up any tendency towards wall effects and channeling.

Regeneration. This is usually carried out with a dilute solution of hydroxide as the most preferred anion.

$$\text{alumina.H}_2\text{F}_3 + 3\text{NaOH} \rightarrow \text{alumina.NaOH} + 2\text{NaF} + 2\text{H}_2\text{O}$$

Rinsing. This is necessary to remove excess regenerant from the alumina bed before neutralisation.

Neutralisation. To restore the fluoride removal capacity, the basic alumina is contacted with dilute sulphuric acid.

$$2 \text{alumina.NaOH} + 3\text{H}_2\text{SO}_4 \rightarrow 2 \text{alumina.H}_2\text{SO}_4 + \text{Na}_2\text{SO}_4 + 2\text{H}_2\text{O}$$

The acidic alumina (alumina.H$_2$SO$_4$) is now ready for another adsorption cycle.

Design criteria

Design criteria for the full-scale plant are shown in Table 2 (Schoeman et al., 1985).

Experimental

A simplified flow diagram of the defluoridation plant is shown in Fig. 1. Borehole water is pumped through the defluoridation plant and the treated water is stored in a reservoir for distribution to the community. Spent regenerant and backwash water is stored in a lined evaporation pond.

A number of runs were conducted and breakthrough curves established. The experimental conditions for the runs are summarised in Table 3. The breakpoint for fluoride was taken as 2.0 mg/l and the number of bed volumes (BV, 1 BV = volume of alumina in vessel) produced at breakthrough were determined from these curves. All fluoride analyses were carried out with a fluoride-specific electrode and the other ions were analysed with ICP-OES.

### Results and discussion

#### Plant performance

Six loading-regeneration runs were conducted during the commissioning phase of the plant. Plant performance data for the six runs are summarised in Table 4.
The fluoride concentration in the feed water (5.3 to 10.8 mg/l) could be easily reduced to ~2 mg/l in the treated water. The average concentration of fluoride in the treated water was approximately 1 mg/l. Therefore, an excellent quality water could be produced with the activated alumina defluoridation process.

The loading time to reach a breakthrough of approximately 2 mg/l in the treated water varied from 67.5 to 79.5 h for the 6 runs (Table 4). The throughput varied between 940 and 1 296 m³ and the bed volumes produced between 566 and 781. The differences in throughput and bed volumes of defluoridated water produced could be ascribed to intermittent runs and slight changes in regeneration conditions. It was found that a run achieving a 2 mg/l fluoride breakthrough could be significantly increased by an intermittent run which gives the fluoride more time to adsorb onto the alumina.

Breakthrough curves

A typical breakthrough curves for Run 5 is shown in Fig. 2. The fluoride concentration in the feed water could be reduced from 5 to 6 mg/l to less than 1.5 mg/l in the treated water. The running time to approximately 1.5 mg/l fluoride breakthrough could be significantly increased by an intermittent run which gives the fluoride more time to adsorb onto the alumina.

Composition of spent regenerant

The fluoride and silicon concentration and pH of the spent regenerant (regeneration after Run 4) is shown in Table 5. The fluoride concentration in the spent regenerant showed a peak after 70 min regeneration time (Table 5). The fluoride concentration then dropped and was 370 mg/l when the caustic soda regeneration was completed after 100 min. The regeneration time was long as a result of the low viscosity of the caustic lye during winter. Very little fluoride was removed during the subsequent rinsing and neutralisation steps.

The silicon started to desorb from the alumina at high pH when most of the fluoride had been removed from the alumina. Silicon was also desorbed from the alumina during the neutralisation step when the pH became lower.

pH of the feed, acidified feed and caustic soda adjusted product water

The pH of the feed and treated water (product) is shown in Table 6 (next page). The pH of the feed was lowered from 7.6 to 7.8 to 6.1 to 6.6 in the acidified feed. The pH of the product water varied between 6.3 and 6.5. The pH is on the low side and should be raised to approximately 7. It is also interesting to note that the initial pH of the product water was low (pH 3.8). This is due to acid leaching from the alumina bed after neutralisation of the alumina with acid. Care should be taken not to put too much acid through the bed during neutralisation.

Electrical conductivity of the feed and product water

After 19 h and 282 m³ throughput the electrical conductivity was 1 292 µS/cm for the feed, 1 318 µS/cm for the acid feed and 1 358 µS/cm for the (caustic) product. The electrical conduc-
ity of feed and product water increased somewhat after the addition of chemicals (acid and caustic soda) for pH control.

**Chemical composition of the feed and product water**

The chemical composition of the feed and product water is shown in Table 7. Alkalinity is reduced through the addition of acid and by adsorption on the activated alumina. However, a significant amount of alkalinity is still entering the alumina bed in this case and this alkalinity reduces the capacity of the alumina for fluoride removal because it competes with fluoride for adsorption sites. It is further interesting to note that aluminium is released into the product water (0.1 mg/l) (Table 7). However, the aluminium concentration in the treated water is less than 0.3 mg/l which shows that the water is of acceptable quality according to the South African specifications for drinking water (Schutte, 2006). Care should be taken, however, that the pH of the treated water should not drop too low (< 6) during neutralisation because the aluminium ion could dissolve in the water at a too low pH.

The sodium concentration of the product water increases somewhat as a result of caustic soda addition while the sulphate concentration increases as a result of acid addition and displacement of sulphate ions from the alumina by fluoride adsorption.

**Economics**

The capital costs of a 200 m³/d defluoridation plant are estimated at R1.3 m. Operational costs are estimated as follows:

- Caustic soda regeneration: R0.35 / m³
- Sulphuric acid neutralisation: R0.07 / m³
- Sulphuric acid for pH control: R0.16 / m³
- Caustic soda for pH control: R0.14 / m³
- Electrical energy cost for pumping: R0.06 / m³
- R0.68 / m³

**Note:**
- Acid cost R0.82/kg
- Caustic soda cost R2.98/kg
- Electrical energy cost R0.25/kWh (2003)

**Conclusions**

- The water defluoridation plant was commissioned successfully and the plant is performing satisfactorily. Fluoride in the feed water present in the 6 to 10 mg/l range is reduced to <1.5 mg/l, the recommended maximum concentration.
- No reduction in plant output was experienced over 6 loading-regeneration cycles. Therefore, it appears that fouling of the activated alumina should not be a problem. This matter, however, should be monitored over a more extended period.
- Plant output varied between 940 and 1 296 m³ to a fluoride breakthrough of approximately 2.0 mg/l. It is recommended that the plant be regenerated after at least 940 m³ of defluoridated water has been produced.
- Care should be taken that the pH of the treated water should not drop too low after neutralisation of the alumina bed with acid to ensure that the aluminium ion does not go into solution.
- The capital cost of a 200 m³/d activated alumina defluoridation plant is estimated at R1.3m. Operational costs are estimated at R0.68/m³.
- It was demonstrated that a 1st world technology could be effectively operated and maintained in a rural area with proper training and constant supervision of the operators.

**References**


BISHOP PL and SANCOUCY G (1978) Fluoride removal from drink-
ing water by fluidized activated alumina adsorption. *J. Am. Water Works Assoc.* **70** 554.


DRINKING WATER STANDARDS (1962) US Public Service, Washington, DC.


NATIONAL INTERIM PRIMARY DRINKING WATER REGULATIONS (1975) EPA Federal Register. **40** 51.


