Properties of catalyst for iron and manganese oxidation in filter materials

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Abstract. The aim of this work is to determine the properties of contact layers of grain of different type of filter media in term of their suitability for catalytical manganese oxidation and auto-activation for manganese removal from groundwater. The following oxidative filtration materials were investigated: auto-activated silica sand taken from full-scale filters (oxide coated filter media), Pyrolox – manganese ore – natural material and Hydrolit – artificially activated material. As a reference two natural not chemically active materials were tested: clean silica sand and clean chalcedonite. The investigation resulted in chemical composition and microstructure parameters of catalyst contact layers of grains. The modern analytical methods were used: EDAX, Raman spectroscopy, mercury and helium porosimetry. The data indicated that catalyst from autoactivated filtration material has better chemical composition and porosity parameters in comparison to tested catalytic filtration materials, making it more suitable for adsorption and catalytic manganese oxidation.

1 Introduction

Traditional technology of groundwater treatment consists of aeration followed by filtration. The iron and manganese removal occurs due to heterogenic and autocatalytic oxidation in catalytic-oxidative filtration beds. The catalyst is manganese dioxide – the product of Mn(II) oxidation.

Most common filtration material is silica sand. Clean fresh silica sand has no catalytic properties becoming an auto-activated material after covering grains with manganese oxides. This permanent oxides contact layer may be created with no chemical dosage [1,2]. The rippening of filtration material is often supported with biological oxidation process [3-6].

Another type of filtration materials are natural catalytic filter media (e.g. Pyrolox, Gabon manganese ore). This type of media is often use in small WTPs. Often these media loose in time the high efficiency in manganese removal and new steady state of dynamic balance has to be reached [7].

There are on the market artificially activated oxidative media – industrial products, manufactured with patented technology. It may be expected that the manganese oxidation catalysers of different filtration materials have different characteristics. The difference appear in chemical composition, microstructure parameters and polymorphic forms present compounds [8,9].

Manganese oxidation from the bivalent to quadrivalent form undergo on different pathways. One of the possible pathways of manganese oxidation is given in Fig. 1.

\[
\text{[Mn(H}_2\text{O)}_6]^{2+} \downarrow \quad \text{Mn(OH)}_2 \\
\text{(Pyrochroite)} \quad \downarrow \quad \text{Buserite} \quad \downarrow \quad \gamma\text{-MnO}_2 \\
\text{(Nustite)} \quad \downarrow \quad \beta\text{-MnO}_2 \\
\text{(Pyrolusite)}
\]

Fig. 1. Manganese oxidation pathways [10].

Pyrolusite is the most stable form of manganese dioxide. It occurs naturally or is formed with use of chemical oxidants [6]. Pyrolusite has adsorption capacity to Mn(II) but does not have auto-catalytical properties [11]. Birnessite is the manganese oxide with the valency equal to 3.5 – 3.9, easily undergoes red-ox reactions, takes part in ion exchange, has high adsorption capacity for manganese, ensuring conditions for auto-catalytical oxidation [11]. Birnessite is a product of auto-catalytical
reactions of manganese oxidation taking place in systems of aeration and rapid filtration, without chemical dosage [2,11].

The manganese oxidation, taking place in filter media, is a heterogenic process on the surface of grains. An important factor affecting the adsorption of manganese on contact layer is the surface charge. The values of isoelectric point for quartz and manganese dioxide are shown in Table 1.

Natural groundwater usually has pH within the range of 5.5–7.5. At this pH surfaces of silica sand grains have negative charge. Higher negative charge in lower pH of media, is a heterogenic process on the surface of grains. The manganese oxidation, taking place in filter media, is a heterogenic process on the surface of grains. The oxidation of manganese on contact layer is the surface charge. The values of isoelectric point for quartz and manganese dioxide are shown in Table 1. The manganese oxidation, taking place in filter media, is a heterogenic process on the surface of grains. The values of isoelectric point for quartz and manganese dioxide are shown in Table 1.

Table 1. Point of zero charge for manganese dioxide [10,12].

| Polymorphic variant of MnO2 | Formula | pH of isoelectric point |
|-----------------------------|---------|-------------------------|
| Hollandite                  | α-MnO2  | 5.0                     |
| Pyrolusite                  | β-MnO2  | 7.2                     |
| Nsutite                     | γ-MnO2  | 5.6                     |
| Birnessite                  | δ-MnO2  | 2.8                     |
| Quartz                      | SiO2    | 2.0                     |

The time and effects of auto-activation of media for manganese removal depend on the microstructure parameters of grains filter media. Investigation of specific surface and pore volume allows determination of the potential ability of the catalyst for manganese adsorption and further auto-catalytic Mn oxidation.

2 Goal and scope of the research

The aim of this work is to determine the properties of contact layers of grains in different type of filter media in term of their suitability for catalytical manganese oxidation and auto-activation for manganese removal from groundwater.

Following oxidative filtration materials were investigated (Table 2):
- auto-activated silica sand taken from full-scale filters (oxide coated filter media),
- Pyrolox – manganese ore – natural material,
- Hydrolit – artificially activated material.

As a reference two natural, not chemically active, materials were tested:
- clean silica sand,
- clean chalcedonite.

3 Methods

For catalytic contact layer of filter media grains investigation the following methods were used:
- energy dispersive spectroscopy – EDAX, coupled with SEM,
- Raman spectroscopy,
- mercury and helium porosimetry method.

Raman spectrometry was done with Renishaw apparatus. The SEM-EDAX analysis was done with Quanta FEG 250 (FEI) Microscope coupled with EDS Octane II Silicon Drift Detector (EDAX).

To determine the chemical composition of catalysts the energy dispersive spectroscopy EDAX, coupled with SEM was used. Weight shares of predominant elements were analysed using Microscope Quanta FEG 250 (FEI) with EDS Octane II Silicon Drift Detector (EDAX).

For the research on the identification of different polymorphic forms of oxides of the investigated catalysts, the method of μ-Raman spectroscopy was used. The non-polarised Raman spectra of filtration materials were recorded in the backscattering geometry using a Via Renishaw micro-Raman system. All spectra were recorded in the range of 100 to 2000 cm⁻¹. As an excitation light, the green line of argon laser operating at 514.5 nm (three accumulations of each spectra) was used. A laser beam was focused on the sample through the long working distance of the 50x/0.5 objective. The spatial resolution of the Raman spectra was about 2 μm. The power of the laser beam was less than 5 mW. All measurements were performed at room temperature. The crystalline structure of the catalytic contact layer in the manganese removal zone as well as silica sand beds were examined using the method of X-ray powder diffractometry (XRD). Examinations consisted of a comparison of diffraction diagrams obtained in Brucker D8Advance diffractometer for the analysed samples and the reference diagrams available in specialised databases [13]. The internal structure parameters pore volume and specific surface area was determined with the nitrogen adsorption method with a Tristar II 3020 apparatus.

Mercury porosimetry analysis is the progressive

Table 2. Element composition of tested materials.

| Investigated material | Weight shares of elements, in % |
|-----------------------|---------------------------------|
|                       | Fe     | Mn     | O      | Si     | Ca      | Al      | K       | Mg      | S       | P       |
| Chalcedonite          | 0.6    | –      | 61.5   | 33.3   | 0.36    | 3.54    | 0.32    | –       | 0.32    | –       |
| Silica sand           | 0.17   | –      | 60.7   | 37.4   | –       | 1.42    | 0.24    | –       | –       | –       |
| Catalyst – iron removal zone | 30.1 | 17.5   | 45.1   | 1.57   | 3.75    | 1.15    | –       | –       | 0.08    | 0.56    |
| Catalyst – manganese removal zone | 17.0 | 18.5   | 56.6   | 1.06   | 3.27    | 3.32    | –       | –       | –       | 0.28    |
| Pyrolox               | 3.66   | 48.95  | 40.6   | 1.84   | –       | 2.73    | 1.52    | –       | –       | 0.07    |
| Hydrolit              | 1.4    | 36.57  | 36.88  | 3.93   | 15.22   | 3.33    | 0.71    | 1.46    | 0.47    | 0.02    |
intrusion of mercury into a porous structure under stringently controlled pressures. From the pressure versus intrusion data, the instrument generates volume and size distributions using the Washburn equation. Clearly, the more accurate the pressure measurements, the more accurate the resulting pore size data.

Mercury porosity analysis of tested materials was done using AutoPore9220 apparatus by Micrometrics. Applied instrument allows to determine the pore distribution in the range of pore diameter from 0.003 to 500 μm, with applied maximum pressure of 60000 psi. The helium porosimeter due to penetrating properties of helium allows to determine the submicropore structure of tested materials. It enables to precise measure of specific density of samples.

The sample of filtration material (of given mass-M and given volume V_{ext}) is placed into calibrated chamber of helium porosimeter and then helium is introduced. From the amount of introduced helium the volume of structure of filtration material - V_{structure} is calculated. The total open porosity coefficient m_{open} is calculated according to the formula (1):

\[ m_{open} = \frac{V_{ext} - V_{structure}}{V_{ext}} \] (1)

Next the same sample is introduced into mercury porosimeter and the bulk density \( \rho_b \) is determined taking into account the volume of mercury - \( V_{Hg} \) (non-moisten liquid) driven out by the porous sample.

\[ \rho_b = \frac{M}{V_{Hg}} \] (2)

The dynamic porosity is calculated taking into account the volume of mercury introduced into the sample under pressure up to 60 000 psi. The calculation of pores volume includes only the volume of pores available for mercury: 0.01-360μm. The value of dynamic porosity is smaller than the value of open porosity.

4 Results and discussion

The photographs of investigated materials’ surfaces using Raman spectrometer and scanning electron microscope show that the auto-activated contact layer is not homogenous and consists of centres of iron and manganese oxides.

The comparison of not-chemically active materials shows the more developed surface structure of chalcedonite than silica sand. This creates better suitability of chalcedonite for autoactivation for manganese removal from groundwater [2]. In Table 2 the element composition of investigated media are presented. The data were obtained using apparatus EDAX [14]. The EDAX results (Table 2) indicate, that:

- silica sand and chalcedonite are chemically homogenous, mostly consisting of SiO₂.
- catalyst of iron removal zone consists of iron (30.1%), manganese (17.5%), oxygen (46.0%), calcium (3.75%), indicating presence of iron and manganese oxides in the mass of catalyst.
- catalyst of manganese removal zone consist of manganese (18.5%), iron (17%), calcium (3.25%). The presence of iron and calcium in this catalyst lowers catalytical activity of contact layers.
- in Pyrolox manganese dioxide is the main constituent,
- Hydrolit is a mixture of manganese iron, calcium and magnesium compounds.

For identification of compound structure of investigated media the method of µ-Raman spectroscopy was used. The method is based on comparison of tested materials spectra and reference spectra. In Table 3 the values of characteristic wavenumber for iron, manganese and quartz oxides expected in contact layers are presented. For the investigation the Renishaw spectroscope was used. Raman spectra of tested materials and catalysts are presented in Fig. 2 – 7.

The Raman spectrum of the chalcedonite and silica sand samples, presented in Figures 2 and 3 respectively have the following Raman bands: 127, 206, 464 cm⁻¹. The same values of characteristic wavenumbers can be found in the reference spectrum of quartz (Table 3). Silica sand and chalcedonite are filtration materials with the structure of pure quartz.

Comparison of Raman spectrum for the iron removal zone with the reference plot of characteristic wavenumber (Table 3, Fig. 4) of iron oxides, indicated the presence of hematite \( \alpha Fe_2O_3 \) and maghenite \( \gamma Fe_2O_3 \) [15].

In manganese removal zone the presence of birnessite, was proved (Table 3, Fig. 5).

| Compound          | Wavenumber, [cm⁻¹] |
|-------------------|---------------------|
| Quartz, SiO₂      | 127 206 464         |
| Maghemite \( \gamma Fe_2O_3 \)| 350 512 665 730 |
| Hematite \( \alpha Fe_2O_3 \)| 265 300 345 – 395 515 645 – 670 715 |
| Birnessite \( \delta MnO_2 \)| 495 – 505 570 – 575 635 – 655 |
| Pyrolusite \( \beta MnO_2 \)| 538 665 |
| Nsutite \( \gamma MnO_2 \)| 379 491 520 572 631 738 |
| Ramsdelite        | 294 518 580 630 740 |

Table 3. Raman shifts for silica and iron and manganese oxides [11,15,16].
In Pyrolox the presence of manganese oxides in the form of pyrolusite, ramsdelite and nsutite (Table 3, Fig. 6) were found. Raman spectrum of Hydrolit indicated manganese dioxide in the form of pyrolusite (Table 3, Fig. 7).

In Table 4 porozymetry parameters obtained with mercury porozymetry method and helium porosity method are presented. AutoPore IV 9500 used in investigation allows to determine the pore structure within the range of pore diameter 0.003 – 500 μm.

Manganese removal zone catalyster has the high value of specific surface area 124.75m²/g. The low value of bulk density 0.652 g/cm³ correlates with high value of total porosity 80.552%. Established porosity parameters allow to consider manganese removal zone catalyster as a porous material with good adsorption properties. The specific surface area of Pyrolox and Hydrolit are few times lower than manganese removal zone catalyster.

The lowest value of specific surface area have silica sand. On the other hand the specific surface area of chalcedonite is almost two times higher than Pyrolox and Hydrolit and 200 times higher than silica sand. It gives the evidence of better properties of chalcedonite to auto-activation for manganese removal from groundwater [2].

![Fig. 2. Raman spectrum for chalcedonite.](image)

![Fig. 3. Raman spectrum for silica sand.](image)

![Fig. 4. Raman spectra for iron removal zone.](image)

![Fig. 5. Raman spectrum for manganese removal zone.](image)

| Material                                      | Mass density [g/cm³] | Total open porosity [%] | Bulk density [g/cm³] | Dynamic porosity [%] | Specific surface area [m²/g] |
|-----------------------------------------------|----------------------|-------------------------|----------------------|----------------------|-----------------------------|
| Manganese removal zone of rippened silica sand bed | 3.335                | 80.552                  | 0.652                | 72.056               | 124.75                      |
| Pyrolox                                       | 4.03                 | 23.81                   | 3.08                 | 20.35                | 2.58                        |
| Hydrolit                                      | 2.92                 | 13.01                   | 2.49                 | 11.94                | 2.06                        |
| Silica sand                                   | 2.64                 | 5.15                    | 2.48                 | 5.00                 | 0.02                        |
| Chalcedonite                                  | 2.63                 | 35.68                   | 1.71                 | 29.73                | 4.74                        |
5 Conclusions

In the described investigation on catalysts for catalytic manganese oxidation, the methods of instrumental analysis were used. The obtained data allowed:

- determination of elements’ shares and chemical composition of the investigated catalysts,
- determination of microstructure parameters e.g. specific surface area and porosity of the contact catalytic layer of tested media.

The analysis of catalysts for manganese removal zone pointed that this material is the most capable for effective manganese removal from groundwater in the process of heterogenic autocatalytic oxidation in filter bed. This capability comes from chemical composition of catalysts from auto-activated beds and because of high content of manganese dioxide in the form of birnessite, MnO$_{1.3-1.9}$ – not fully oxidized manganese dioxide with deficiency of oxygen. Such oxide becomes a semi-conductor able to transport electrons, guaranteeing good efficiency of heterogeneous reaction.

Catalyst from iron removal zone contains the iron oxides in the form of $\alpha$Fe$_2$O$_3$ – hematite and $\gamma$Fe$_2$O$_3$ – maghemite.

The analysis of microstructure also proof the capability of auto-activated catalysts for effective adsorption of manganese from treated water due to high values of the specific surface area and pore volume, measured by Autopore apparatus. Catalyst of the manganese removal zone has a specific surface area 124.75 m$^2$/g and total porosity 80.55%.

In Pyrolox the manganese dioxides were detected in the form of ramsdelite, nsutite and pyrolusite. This bed is characterised by much lower specific surface area 2.58 m$^2$/g in comparison to auto-activated catalyst.

Artificially activated Hydrolit is a mixture of manganese, iron, calcium and magnesium compounds. This bed is characterised by the low specific surface area of 2.06 m$^2$/g determined with mercury porosimetry method.

Investigation of two not-chemically active materials demonstrate the higher chalcedonite potential for manganese removal auto-activation due to higher, in comparison to silica sand, values of microstructure parameters.

Further research on relationships between the characteristic of catalysts and technological effects of groundwater treatment should be conducted. It will help the evaluation of mechanisms of heterogenous, catalytic manganese oxidation in filtration bed.

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References

1. J. Jež-Walkowiak, Z. Dymaczewski, J. Wat. Sup. Res. Tech. AQUA 61, 6 (2012)
2. J. Jež-Walkowiak, Wpływ właściwości złoć filtrów pospiesznych na efekty technologii odzelaowania i
3. P. Mouchet, AWWA J. **84**, 4 (1992)
4. K. Olańczuk-Neyman, R. Bray, Pol. J. Env. Stud. **9**, 2 (2000)
5. R. Bray, K. Olańczuk-Neyman, Wat. Sci. Tech. Wat. Sup. **1**, 2 (2001)
6. R. Bray, *II Congress of Environmental Engineering, Lublin, Poland, Monografie PAN, vol.32*, (2005)
7. J. Jeż-Walkowiak, Z. Dymaczewski, A. Szuster-Janiaczyk, A.B. Nowicka, M. Szybowicz, Water **9**, 498 (2017)
8. J. Jeż-Walkowiak, *Intensyfikacja procesu odżelaziania odmanganiania wód podziemnych w złożach oksydacyjnych filtrów pospiesznych (PhD Th., 2000)*
9. J. Jeż-Walkowiak, Z. Dymaczewski, Ł. Weber, J. Wat. Sup. Res. Tech. AQUA **64**, 1 (2015)
10. W. Stumm, J.J. Morgan, *Aquatic Chemistry, Chemical Equilibria and Rates in Natural Waters, Third edition* (Wiley Interscience, 1996)
11. J.H. Bruins, B. Petrusevski, Y.M. Slokar, J.C. Kruithof, M.D. Kennedy, Des. Wat. Tr. **55**, 7 (2015)
12. M. Kosmulski, *Surface charging and points of zero charge*, (CRC Press, Tylor&Francis Group, 2009)
13. S. Jędrychowska, *Możliwości wykorzystania spektroskopii ramanowskiej w branży naftowej, cz. I*, (Nafta-Gaz, 2012).
14. A. Barbací, *Mikroskopia elektronowa*, (Wyd. PP, Poznań, 2007)
15. M. Szybowicz, M. Koralewski, J. Karoń, L. Melnikova, Acta Phys. Pol. O. Acc. **127**, 2 (2015)
16. C. Julien, M. Massot, R. Baddour-Hadjjeans, S. Franger, S. Bach, J.P. Pereira-Ramos, Sol. State Ion. **159** (2003).