Electrodialytic removal of heavy metals and chloride from municipal solid waste incineration fly ash and air pollution control residue in suspension - test of a new two compartment experimental cell

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Highlights

- Acidification of residue suspension is faster in the 2 compartment cell
- pH in residue suspension is the determining factor for remediation success
- Leaching limiting values can be met for fly ash after electrodialytic remediation
Electrodialytic removal of heavy metals and chloride from municipal solid waste incineration fly ash and air pollution control residue in suspension – test of a new two compartment experimental cell

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Abstract

Municipal solid waste incineration (MSWI) residues such as fly ash and air pollution control (APC) residues are classified as hazardous waste and disposed of, although they contain potential resources. The most problematic elements in MSWI residues are leachable heavy metals and salts. For reuse of MSWI residues in for instance concrete, the aim of remediation should be reduction of the heavy metal leaching, while at the same time keeping the alkaline pH, so the residue can replace cement. In this study a MSWI residues were subjected to electrodialytic remediation under various experimental conditions. Also a newly developed 2 compartment experimental cell was tested. The results show that the pH development in the MSWI residue suspension depended on the type of MSWI residue and the experimental cell type. The acidification of the suspension occurred earlier when using the 2 compartment setup and the acidification of the fly ash occurred earlier than for the APC residue but the highest removal was seen with the 3 compartment cell. The lowest final pH for the fly ash and APC residue was 6.4 and 10.9, respectively. The results showed that the leaching of Cd, Cu, Pb and Zn was reduced compared to the initial heavy metal leaching except when the pH was reduced to a level below 8 for the fly ash. On the other hand, Cr leaching increased by the electrodialytic treatment. Cl leaching from the MSWI residues was less dependent on experimental conditions and was reduced in all experiments compared to the initial levels.
Keywords: electrokinetic remediation, fly ash, leaching, heavy metal, chloride

Nomenclature:

APC - Air pollution control
EDR - Electrodialytic remediation
MSW - Municipal solid waste
MSWI - Municipal solid waste incineration
I/S REFA - Waste-to-energy incineration plant, semi-dry APC system
SD – APC residue from REFA
I/S Vestforbrænding - Waste-to-energy incineration plant, wet APC system
W – Fly ash from Vestforbrænding

1. Introduction

Incineration of municipal solid waste (MSW) is commonly used to gain energy and hygenize and minimize the amount of solid waste. The residues from MSWI are flue gas, bottom ash and fly ash or air pollution control (APC) residues depending on the air pollution control system of the incineration plant. In Denmark, all MSWI plants have air pollution control system to minimize the emissions of acidic gases and dioxins. Fly ash is typically removed in electrofilters or cyclones, while the air pollution control process can be either a dry/semidry (injection of lime in dry form or in a slurry to the flue gas) or wet (flue gas is subjected to a scrubber after removal of fly ash). Either treatment method, the resulting residues are classified as hazardous waste due to the high content of contaminants and salts. The main management system of MSWI residues is disposal in hazardous waste facilities [1].

The electrodialytic remediation (EDR) method has been used to remove mainly heavy metals from particulate waste materials such as soil, harbor sediment and different ashes (biomass, sewage sludge, MSW) [2-6]. Ions electromigrate in the applied electric field during EDR and to maximize their desorption and availability, ionic forms are essential for the process. To reuse the MSWI
residues as secondary resources, removal of contaminants and salts is necessary to reduce their toxicity. EDR treated APC residue has indicated potential for substitution of cement in mortar [7], but improvements in reduction of heavy metal leaching and chloride content is still needed.

The removal of heavy metals from MSWI fly ashes using EDR was first applied in a conventional stationary EDR cell with the fly ash saturated with water or desorbing agent [8]. These initial experiments resulted in severe precipitations in the electrolytes, on the ion exchange membranes and in the fly ash. Furthermore, poor control of the pH both in the electrolytes and in the fly ash itself, revealed the need to suspend the fly ash matrix in the cell. Pedersen et al. [9] introduced a stirrer in the suspension cell compartment II, Fig. 1a. After the introduction of the stirrer, several studies about EDR of fly ash were made, especially with the use of assisting agents to further enhance the remediation process. Ammonium citrate and gluconate were mostly used as assisting agents, with the purpose of enhancing solubilisation of the heavy metals and form stable charged complexes that could be removed in the electric field and at the same time keeping the alkaline pH in the fly ash. Allowing acidification of the ash during EDR means dissolution of a high quantity of the ash and is thus not an energy efficient separation of the ash and heavy metals. Ammonium citrate as assisting agent was used in EDR of fly ash and these electro dialytic experiments showed higher removals than just by batch extractions and up to 20 % Cr, 60 % Cd, 6 % Pb, 40 % Zn and 60 % Cu were removed after 2 weeks of remediation [9]. When increasing the remediation time, even higher removals could be obtained, with the lowest still for Cr and Pb [10]. Gluconate was tested as assisting agent for APC residue and high desorption of the metals was seen in batch extraction but much lower removal was seen in EDR experiments, probably due to the size of the metal-gluconate complexes that could not pass the ion exchange membranes [11]. EDR with gluconate was later improved with pre-treatment of the APC residue before EDR, such that 82 % Cd, 10 % Pb, 63 % Zn and 22 % Cu were removed after 2 weeks of remediation [6]. One major drawback of using assisting agents is however the significant matrix changes observed both in the mineralogy and the morphology of the fly ash [12] which could hinder reuse. The addition of assisting agents also adds an extra cost to the process.

Suspending the fly ash in distilled water during EDR also results in removal of heavy metals [9]. The removal process in this case is dependent on the solubility and release of the heavy metals from the fly ash particles at high pH. In the 3 compartment cell (Fig. 1 a), acidification of the suspension is caused by water splitting at the anion exchange membrane and proton leakage through the anion
exchange membrane \[13, 14,\], which leads to enhanced removal of heavy metals during EDR. Some of the few experiment of EDR with fly ash suspended in distilled water in the stirred set-up, showed varying removal results of Cd, Cr, Cu, Pb and Zn and indicate that the removal occurs both towards the anolyte and the catholyte \[9, 15\]. The removal towards the anode can occur due to the complexation between divalent metal ions and chloride to form stable negatively charged complexes (MeCl\(_3^−\), MeCl\(_4^{2−}\)) and for Cr also the presence of negatively charged Cr (VI) ions.

Recently, a new cell design was developed and patented for EDR of suspensions, where the anode is placed directly in the material suspension and thus combines the anolyte and the suspension compartment into compartment I, Fig. 1b \[16, 17\]. This new cell design promotes a more direct acidification of the suspension, where all the produced H\(^+\) ions from the electrolysis at the anode \((2\text{H}_2\text{O} \rightarrow \text{O}_2 \text{(g)} + 4\text{H}^+ + 4\text{e}^-)\) will be supplied directly in the suspension. Anions are in the 3 compartment cell removed to the anolyte and in the case of Cl\(^−\) an oxidation to Cl\(_2\) (g) will occur at the anode. In the 2 compartment cell, it is expected that the oxidation and subsequent removal could be faster, since the electrode is in direct contact with the suspension. Contrarily, if the heavy metals are present as anions in the 2 compartment cell, they will not be removed from the suspension.

The purpose of this study was to investigate 1) the remediation of two types of MSWI residue (fly ash and APC residue) for heavy metals and chloride according to several different experimental variables and 2) to which extent the new 2-compartment cell could improve the remediation of heavy metals and chloride from MSWI residues. The aim was to obtain a stable residue in the sense that it is not leaching heavy metals and chlorides above the Danish limiting values for reuse of waste in constructions after the treatment.

2. Materials and methods

2.1. MSWI residues

Two types of MSWI residues were used in this study from Danish municipal waste incineration plants:

- **SD**: APC residue collected after a semi-dry process after the injection of slaked lime and activated carbon from I/S REFA, where 3,000 tons APC residue is produced annually.
• **W:** A fly ash sample collected prior to the neutralization of acidic components by a wet scrubber process from I/S Vestforbrænding, where 15,000 tons of APC residues, including fly ash is produced annually.

### 2.2 Analytical methods

The MSWI residues characterization and extraction experiments were carried out using dried residue at 105 °C for 24 h and triplicates were made for the analysis. Total concentration of Cd, Cr, Cu, Pb and Zn in the MSWI residues were measured by Inductively Coupled Plasma - Optical Emission Spectrometry (ICP-OES), after pre-treatment according to DS 259 [18], in which 1 g of MSWI residue and 20 mL (1:1) HNO₃ were heated at 200 kPa (120 °C) for 30 min. After digestion, the samples were vacuum filtered through a 0.45 µm filter and diluted till 100 mL. The units used in this paper are mg/kg in dry matter for the concentration.

The pH in the MSWI residues was measured with a Radiometer pH-electrode in 1 M KCl at L/S ratio of 2.5 and in deionised water at liquid to solid (L/S) ratio of 5, after 1 h of agitation at 190 rpm. The electrical conductivity was measured at the same time as the pH with deionised water, with a Radiometer Analytic electrode. Water content was measured as weight loss after 24 h at 105 °C (calculated as weight loss over the weight of the wet sample). Loss on ignition (LOI) was calculated as weight loss after heating for 30 min at 550 °C. Water solubility was made by suspending 100 g MSWI residue in 500 mL deionised water and agitate for 1 min. After settling, the water was decanted and another 500 mL deionised water were added. This was repeated three times to ensure that the MSWI residues were properly washed. The suspension was then filtered thought a 0.165 µm filter and the MSWI dried residue (105 °C) weighed. Carbonate content was determined using a Scheibler apparatus, where a standard curve was first performed using different CaCO₃ contents combined with HCl. The MSWI residues were mixed with HCl and then the CaCO₃ concentration was determined.

Leaching tests on the MSWI residues were made according to DS/EN 12457-3 [19], with slight modifications. The L/S ratio was 2, mixing 10 g of MSWI residue and 20 mL deionised water. The suspension was shaken for 23 h on an end-over-shaker. The pH was measured in the suspension before vacuum filtration through a 0.45 µm nucleo filter and the heavy metal concentrations in the filtrate were determined by ICP-OES. Water soluble chloride was determined by Ion...
Chromatography (IC) after extracting 10 g of MSWI residue with 50 mL of deionised water overnight.

Residue mineralogy was studied by X-ray powder diffraction (XRD), for identification of major crystalline phases. The instrument was a PANalytical X’Pert Pro operating at 45 kV and 40 mA applying Cu Kα radiation with a 2Θ X’Celerator detector. The samples were scanned in the range of 4-100 2Θ within 2.5 hours. The diffractograms were interpreted using the ICDD PDF-4 database for minerals.

2.3 Electrodialytic experiments

Twenty-four EDR experiments were performed with the MSWI residues, according to the experimental conditions presented in Table 1. Twelve of them were performed in an ED cell with 3 compartments (Fig. 1a) and the other 12 were carried out in an ED cell with 2 compartments (Fig. 1b). All experiments were made with a suspension consisting of 350 mL of deionised water and 100 g MSWI residue, corresponding to an L/S ratio of 3.5.

The EDR cells were made of plastic with an internal diameter of 8 cm and the suspension compartment length was 10 cm. Electrode compartments were separated from the suspension compartment by an anion-exchange membrane and/or a cation-exchange membrane from Ionics. The electrodes were platinum coated titanium. A power supply (Hewlett Packard E3612A) was used, maintaining a constant current of 5 mA or 50 mA, corresponding to current densities of 0.1 mA/cm² or 1 mA/cm², respectively. The MSWI residue suspension was stirred in the suspension compartment by a flexible plastic flab, attached to a glass-stick and connected to an overhead stirrer (RW11 basic from IKA) with a rotation velocity rate of up to 2000 rpm.

The electrolytes were 0.01 M NaNO₃ with pH adjusted to 2 with HNO₃ (1:1). In the experiments with the 3 compartment cell, 500 mL of anolyte and 1000 mL of catholyte were used, while in the experiments with the 2 compartment cell, 500 mL of catholyte was used. The recirculation was made using “Pan World” magnetic pumps with a performance rate of 8 L/min. Voltage and electric current were read daily, whereas pH in the electrolytes and the pH and the electrical conductivity of the MSWI residue suspension were measured twice a day. pH in the electrolyte was adjusted with (1:1) HNO₃, if pH > 2.
At the end of the experiments the residue suspension was filtered before measurements for heavy metals and chloride. The dried MSWI residue was digested according to DS 259 and measured for heavy metals. The membranes and stirrer were soaked in 1M HNO₃ and the electrodes in 5M HNO₃, to release precipitated metals. The obtained aqueous phase of the membranes, the stirrer, the electrodes and the electrolytes were all analysed for heavy metals (Cd, Cr, Cu, Pb and Zn) by ICP-OES. The pH, heavy metal and chloride leaching were also determined on the dry MSWI residues after the EDR experiments. The chloride concentrations in the liquid from the filtered MSWI residue suspension were measured according to DS 239 [20] by potentiometric titration with AgNO₃.

3. Results and discussion

The chemical characteristics of the experimental MSWI residues are presented in Table 2, along with leaching limiting values. These values are Danish limiting values that represent the maximum allowed leaching values for reuse of waste materials in construction without restrictions (C1) to restricted use (C3) [21]. Statistical analysis of the residue characteristics was carried out using GraPhad Prim 6 software. T-test was chosen because it is one-sided, since the objective was to compare each individual parameter. This test showed that the two types of MSWI residues are significantly different, with 99% and 95% confidence levels with some exceptions, such as water solubility and water content, due to the type of APC system.

The MSWI residues were rich in heavy metals and with significant leaching of both Pb and Zn above the limiting values. The MSWI residues were also characterized by having a large water solubility which is related to the high amount of soluble salts and minerals in MSWI residues. This finding is supported by the high electrical conductivity. Both MSWI residues are highly alkaline with pH above 12 and with high carbonate content, which was even higher for the APC residue than for the fly ash.

The low LOI obtained for the fly ash indicates a complete and efficient combustion during the incineration process, which means that a higher incineration temperature was achieved. Regarding the carbonate content, the APC residue presents a high value, which was expected as lime (CaO) is used in the semidry gas cleaning process. Once CaO is exposed to moderate amounts of water, hydration occurs forming Ca(OH)₂ that may later react with carbon dioxide from the ambient air to form calcium carbonate.
3.1 Electrodialytic remediation experiments

The pH and conductivity in the MSWI residue suspensions during the EDR experiments is shown in Fig. 2, all measurements for each experimental series are displayed by the same notation. The applied currents of 5 mA and 50 mA were maintained constant during all experiments. The development of the pH and electrical conductivity was consistent within the same experiment, although the developments from the single experiments are not shown in Fig. 2, as the whole experimental series are displayed as one trend line. Thus, there are differences between each experiment series with the same conditions except the duration and these are mainly due to the heterogeneity of the MSWI residues. When 5 mA were applied, the voltage drop was 1.3 - 2.6 V and 1.3 - 5 V for the fly ash and APC residue, respectively. Similarly, the voltage drop when applying 50 mA was 2.5 - 7 V and 2.8 - 7.6 V for the fly ash and APC residue. The low voltage drop is a result of the high conductivity in the MSWI residue suspensions and the voltage levels were slightly lower for the 2 than the 3 compartment cell, especially for the fly ash experiments. However, the general energy consumption in a 3 compartment cell was found to be suspension > membranes > electrolytes [22]. This in addition to the cost of ion exchange membranes would favour the 2 compartment cell when upscaling of the method. The electrical conductivity in the EDR experiments with APC residue was higher than for the fly ash, mainly due to the higher chloride content in the APC residue. The overall trend seen from Fig. 2 is that the electrical conductivity decreased faster in the experiments with the 3 compartment cell than in the 2 compartment cell, and also to a lower level. In the 2 compartment cell, the lowest conductivity was obtained with the lowest current density, whereas in the 3 compartment cell the highest current density resulted in the lowest conductivity. This finding is due to the different setups, where the direct supply of H⁺ ions into the suspension in the 2 compartment cell increases with current density, whereas there is increased removal of ions from the suspension in the 3 compartment cell with increasing current. As the pH in the residue suspension decreases, the result will be a higher dissolution of the residue. This is seen as a higher electrical conductivity in the experiments lasting 14 days. The electrical conductivity in the W-2-1 experiments did not significantly decrease during the experimental duration, as H⁺ ions are continuously being introduced to the suspension by the electrode reaction and continuous dissolution of residue at the decreasing pH. Contrarily, for the W-3-1 experiments a different trend was seen, with an initial decreasing electrical conductivity until the pH decreases to where the main residue dissolution starts. It is expected that the electrical
conductivity will continue to increase in both cell types, also for the 3 compartment cell, which was seen for soil [23] and sewage sludge ash [17] at longer duration.

Regarding pH, the buffering capacity was much higher in the APC residue than in the fly ash, which is in accordance to what was expected from the carbonate content. It was also expected that the pH would generally decrease faster in the 2 compartment cell than in the 3 compartment cell. However, this was mainly seen for the fly ash at high current density, because these were the only experiments in which the buffering capacity was exceeded. Although after 12 days of remediation, the pH also decreased slightly in the APC residue suspension using the 2 compartment cell at high current density. In ash with a lower buffering capacity, e.g. sewage sludge ash, the pH decrease was significantly faster with the 2 compartment cell regardless of experimental conditions [17]. Furthermore, the higher buffering capacity observed by the slower pH decrease in the experiments with the APC residue could be due to the difference in mineralogy between the two residues, see Fig. 3. Especially by the presence of the main calcium phases; calcite (CaCO₃), portlandite (Ca(OH)₂), anhydrite (CaSO₄) and calcium hydroxychloride in the APC residue as opposed to only carbonate and anhydrite in the fly ash before remediation. In the APC residue, the Ca-containing minerals were also changed during the electrodialytic experiments, to also include bassinite (CaSO₄•0.5(H₂O)), thus keeping the high buffering capacity.

### 3.2 Heavy metal removal

The final pH measured in the residue and the removal of heavy metals, is shown in Table 3, for the experiments lasting 14 days. The removal is defined as the amount of metal measured in the electrolytes and electrodes for the 3 compartment cell and in the catholyte and at the cathode for the 2 compartment cell. The highest total removals of metals were obtained in the experiments with the lowest final pH in the MSWI residue, as expected [15], and the metal removal was dependent on current density and type of MSWI residue, but less on the cell type. The removal was lower for the experiments lasting 3 and 7 days (results not shown), so increasing remediation time increased removal, due to the lower pH. The pH given in Table 3 is measured after drying the residues and this residue pH was higher than the suspension pH shown in Fig. 2. Generally, the removals were below 5 % for the heavy metals from the MSWI residues, with some clear exceptions; up to 61 %
Cd and 53 % Zn were removed from the fly ash and up to 66 % Pb was removed from the APC residue.

In experiment W-3-0.1-14D the predominant removal was seen towards the anode. When the current density increased (W-3-1-14D) the speciation of the metals seemed to change, indicated by the lower removal towards the anode and 13 % Cd and Zn remained in the suspension compartment. This is probably caused by the decreasing pH and simultaneous removal of Cl in the suspension compartment [15] and also indicates uncharged species of Cd and Zn. Increasing the current density in the 2 compartment cell for the fly ash, increased the amount of especially Cd, Cu and Zn in the suspension compartment, but only increased the removal of Cd and Cu. The final pH in the 2 compartment cell experiments was about 2 pH units lower than in the 3 compartment cell, so this suggests a different metal speciation. Compared to the 3 compartment cell results, it suggests that the predominant speciation of Cd, Cu and Zn is cationic and cations cannot be removed in the 2 compartment cell.

In the experiments with the APC residue a high Pb removal (22 – 66 %) was achieved. The Pb removal was generally higher in the 3 compartment cell, than in the 2 compartment cell. The amount of Pb in the liquid suspension after the experiments with the 2 compartment cell was close to zero, meaning that there was no uncharged or negatively charged Pb species that could not be removed in this cell setup. A higher Pb removal was obtained in the SD-2-0.1-14D experiment, in which the resulting pH was higher than in experiment SD-2-1-14D. The higher pH might be the reason for higher removal as Pb is amphoteric and soluble at very high pH and at low pH. Previous pH dependent desorption experiments with the APC residue showed immobilization of Pb in the pH range 6-11 [24].

Even though the removal was highest in the experiments with 14 days duration, the final heavy metal concentrations in the treated MSWI residues (results not shown) were higher or similar to the initial concentrations in Table 2. This is due to a significant dissolution of the residue; 13-27 % for the fly ash and 19 – 65 % for the APC residue during the experiments, highly dependent on the pH of the suspensions. According to Lima et al. [25] the removal of heavy metals from MSWI residues should also be assessed in relation to the leachability of the heavy metals, especially if the treated ash should be reused [7].

3.3 Heavy metal leaching
The heavy metal leaching before and after the electrodialytic experiments in relation to pH measured in the leachate is shown in Fig. 4, along with limiting values C1 and C3.

The results showed significantly reduced leaching of heavy metals from ED treated fly ash, even with higher total heavy metal content. The leaching level was significantly different between the two MSWI residues after the electrodialytic treatment. The only similar patterns were increased leaching of Cr and reduced leaching of Pb in all the experiments. In this study, it was not investigated whether Cr was present as Cr (IV) or Cr (III), but it is important to include its speciation in future studies. However, even if Cr leaching increased, studies of reusing EDR treated APC residue in mortar has shown that Cr leaching from mortar with cement replacement by EDR treated APC residue is at a similar level as reference mortar without APC residue [7]. For Cd, the leaching increased from the fly ash after EDR treatment, although remaining below limiting values if pH is not reduced below 10; while for the APC residue, the leaching value was below the detection limit for most of the experiments. Cu leaching was below C3 before and after all experiments. Increased leaching of Zn was observed after experiments reaching pH below 8 in the experiments with fly ash. The highest heavy metal removal rates were obtained at the lowest pH, but lower pH resulted in higher leaching values.

The leaching limiting value C3 was successfully met for all heavy metals in some of the experiments meaning that, in what regards leaching, the treated residues could be utilized in construction materials. For the fly ash, these C3 values were met for experiments except W-2-1-7D, W-3-1-14D, W-3-0.1-14D and W-3-1-14D. The leachate pH was between 10.2-11.5 when the C3 values were met. For the APC residue the C3 values for heavy metal leaching was only met in one experiment (SD-3-0.1-14D) at leachate pH 10.5. As the pH decrease was faster in the 2 compartment cell, especially for the fly ash (Fig. 2) the remediation times could possibly be shorter to achieve the desired pH range. This would reduce costs even further for upscaling and making the 2 compartment EDR method an eligible remediation method for fly ash.

3.4 Chloride

The concentration of leached chloride after the electrodialytic experiments is shown in Fig. 5, together with the C1 and C3 values for chloride. Contrarily to the heavy metals, chloride leaching is not pH dependent but availability controlled [26]. The chloride leaching was significantly reduced in all the electrodialytic experiments for both the MSWI residue types and for both cell types. For
the fly ash, which had the lowest chloride content, the chloride leaching was reduced to below C3 in all experiments, which means that for four experiments, the measured heavy metal and chloride leaching levels complied with the regulative limits C3. The Cl limiting value for C3 was fulfilled for two of the experiments (SD-3-1-7D and SD-3-1-14 D) with the APC residue, but the heavy metal leaching was not below the limit for these two experiments.

For Cl leaching there seems to be a difference between the two cell types, with generally lower final leaching of Cl from the residues when using the 3 compartment cell. The amount of Cl in the suspension liquid at the end of the ED experiments was higher in the experiments with the APC residue than with the fly ash (results not shown). There was a general tendency for a higher amount of Cl in the suspensions with the 2 compartment cell, which means that the oxidation of chloride directly at the anode is not as efficient as electromigration for removal of Cl from the suspension.

The main speciation for Cl in APC residues is halite (NaCl), sylvite (KCl) and calcium hydroxochloride (CaOHCl), where the two former are easily soluble minerals and the latter is easily soluble condensate on fly ash surface particles [24]. These minerals were also identified in the APC residue of the present investigation (Fig. 3). The main Cl minerals for the fly ash were NaCl and KCl. It was found that these phases were removed after electrodialytic treatment in both the 2 and the 3 compartment cells and from both MSWI residues, which was also expected according to other studies [7, 12]. Since Cl leaching is availability controlled, increased remediation time would increase the Cl removal and thus Cl decrease leaching, which is also indicated by the results of this study (Fig. 5).

4. Conclusion

- The removal percentages for heavy metals were generally low for both MSWI residues, except for up to 61 % Cd and 53 % Zn from the fly ash and 66 % Pb from the APC residue after 14 days of remediation in the 3 compartment cell.
- The new 2 compartment cell did not positively affect the heavy metal removal. Electrodialytic remediation with the 3 compartment cell could be a method to selectively remove Cd and Zn from fly ash and Pb from APC residue, to possibly extract these metals from the respective residues and process the metals for reuse.
The leaching concentrations of Cd, Cu, Pb and Zn generally decreased in all the electrodialytic experiments. Contrarily, Cr leaching increased after all the electrodialytic experiments and the reasons for this need further investigations.

The most important factor for both the heavy metal removal and leaching after electrodialytic remediation is the final pH of the MSWI residue. For the fly ash the optimum pH range is pH 10.2-11.5, which could be achieved within 3 days of remediation. Whereas the APC residue shows to be much more susceptible towards small changes in pH, which result in higher leaching, with the lowest leaching results at pH 10.5 achieved after 14 days of remediation.

It is possible to reach below the Danish leaching limiting values for possible reuse for Cd, Cu, Cr, Pb and Zn when applying electrodialytic remediation to the MSWI residues using the 3 compartment cell. For the fly ash, Cl leaching also reaches below the leaching limit.

The Cl removal is not higher when using the 2 compartment cell compared to the 3 compartment cell. Also higher concentrations of Cl are present in the liquid in the suspension compartment when using the 2 compartment cell.

The fly ash shows higher potential for reuse after electrodialytic remediation than the APC residue. A separation of fly ash before adding the air pollution control products such as lime and activated carbon could be recommended to produce a particulate residue fraction that could be more suitable for reuse after electrodialytic remediation.

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2483.
Table 1

Experimental conditions for electrodialytic experiments. (W/SD-X-Y-ZD, where W - fly ash, SD- APC residue, X – cell type, 2 or 3 compartment cell, Y – current density (mA/cm²) and ZD- duration in days).

<table>
<thead>
<tr>
<th>Experiment</th>
<th>MSWI residue</th>
<th>Cell type</th>
<th>Current density (mA/cm²)</th>
<th>Duration (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>W-2-0.1-ZD</td>
<td>Fly ash</td>
<td>2</td>
<td>0.1</td>
<td>3, 7, 14</td>
</tr>
<tr>
<td>W-3-0.1-ZD</td>
<td>Fly ash</td>
<td>3</td>
<td>0.1</td>
<td>3, 7, 14</td>
</tr>
<tr>
<td>W-2-1-ZD</td>
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</tr>
<tr>
<td>SD-2-0.1-ZD</td>
<td>APC residue</td>
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<tr>
<td>SD-3-0.1-ZD</td>
<td>APC residue</td>
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<tr>
<td>SD-2-1-ZD</td>
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<tr>
<td>SD-3-1-ZD</td>
<td>APC residue</td>
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<td>1.0</td>
<td>3, 7, 14</td>
</tr>
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Table 2

Characteristics of fly ash and APC residue (mean values ± standard error).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Fly ash (W)</th>
<th>APC residue (SD)</th>
<th>Statistical difference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>pH&lt;sub&gt;KCl&lt;/sub&gt; (L/S=2.5)</strong></td>
<td>12.4</td>
<td>12.2</td>
<td>a</td>
</tr>
<tr>
<td>Conductivity (mS cm&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>55.33 ± 1.11</td>
<td>75.24 ± 1.82</td>
<td>a</td>
</tr>
<tr>
<td>Water content (%)</td>
<td>0.36 ± 0.00</td>
<td>0.87 ± 0.25</td>
<td>-</td>
</tr>
<tr>
<td>Water solubility (%)</td>
<td>0.76 ± 0.06</td>
<td>0.66 ± 0.06</td>
<td>-</td>
</tr>
<tr>
<td>Loss on ignition (%)</td>
<td>0.76 ± 0.14</td>
<td>4.07 ± 0.04</td>
<td>a</td>
</tr>
<tr>
<td>Carbonate content (%)</td>
<td>8.37 ± 0.06</td>
<td>14.55 ± 0.12</td>
<td>a</td>
</tr>
<tr>
<td>Cd (mg kg&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>97.6 ± 0.4</td>
<td>88.7 ± 2.0</td>
<td>b</td>
</tr>
<tr>
<td>Cr (mg kg&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>99.0 ± 1.1</td>
<td>73.1 ± 0.9</td>
<td>a</td>
</tr>
<tr>
<td>Cu (mg kg&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>749.1 ± 7.0</td>
<td>551.2 ± 20.7</td>
<td>a</td>
</tr>
<tr>
<td>Pb (g kg&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>2,710 ± 300</td>
<td>3,110 ± 110</td>
<td>-</td>
</tr>
<tr>
<td>Zn (g kg&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>22,400 ± 410</td>
<td>19,440 ± 370</td>
<td>a</td>
</tr>
</tbody>
</table>

**Leaching pH**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Fly ash (W)</th>
<th>APC residue (SD)</th>
<th>C1 [21]</th>
<th>C3 [21]</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>12.4</td>
<td>11.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cd (mg L&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>n.d.</td>
<td>n.d.</td>
<td>0.002</td>
<td>0.04</td>
</tr>
<tr>
<td>Cr (mg L&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>0.02 ± 0.00</td>
<td>0.03 ± 0.00</td>
<td>0.01</td>
<td>0.5</td>
</tr>
<tr>
<td>Cu (mg L&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>0.01 ± 0.00</td>
<td>0.27 ± 0.02</td>
<td>0.045</td>
<td>2</td>
</tr>
<tr>
<td>Pb (mg L&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>41.9 ± 2.3</td>
<td>450.0 ± 9.6</td>
<td>0.01</td>
<td>0.1</td>
</tr>
<tr>
<td>Zn (mg L&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>2.41 ± 0.23</td>
<td>12.70 ± 0.39</td>
<td>0.1</td>
<td>1.5</td>
</tr>
<tr>
<td>Cl&lt;sup&gt;-&lt;/sup&gt; (mg L&lt;sup&gt;-1&lt;/sup&gt;)</td>
<td>12,100 ± 180</td>
<td>25,600 ± 150</td>
<td>150</td>
<td>3,000</td>
</tr>
</tbody>
</table>

n.d. not detected, <sup>a</sup>Significantly different (p < 0.01), <sup>b</sup>Significantly different (p < 0.05)
Table 3
Removal and distribution of heavy metals in the 14 days experiments.

<table>
<thead>
<tr>
<th></th>
<th>Final pH *</th>
<th>Removal (%)</th>
<th>To anode side of total removal (%)</th>
<th>In liquid suspension (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Cd</td>
<td>Cr</td>
<td>Cu</td>
</tr>
<tr>
<td>W-3-0.1</td>
<td>10.6</td>
<td>3.0</td>
<td>4.5</td>
<td>1.5</td>
</tr>
<tr>
<td>W-2-0.1</td>
<td>10.6</td>
<td>1.1</td>
<td>3.8</td>
<td>3.8</td>
</tr>
<tr>
<td>W-3-1</td>
<td>8.2</td>
<td>61</td>
<td>1.1</td>
<td>3.3</td>
</tr>
<tr>
<td>W-2-1</td>
<td>6.4</td>
<td>56</td>
<td>3.8</td>
<td>15</td>
</tr>
<tr>
<td>SD-3-0.1</td>
<td>11.7</td>
<td>5.2</td>
<td>4.5</td>
<td>1.7</td>
</tr>
<tr>
<td>SD-2-0.1</td>
<td>12.8</td>
<td>0.3</td>
<td>0.3</td>
<td>0.5</td>
</tr>
<tr>
<td>SD-3-1</td>
<td>10.1</td>
<td>0.4</td>
<td>2.4</td>
<td>1.3</td>
</tr>
<tr>
<td>SD-2-1</td>
<td>10.9</td>
<td>0.8</td>
<td>1.4</td>
<td>0.4</td>
</tr>
</tbody>
</table>

* pH measured on dry residue
Fig.1. The experimental set-up of the 3 and 2 compartment electrodialytic cell. AN-anion exchange membrane, CAT-cation exchange membrane.
Fig. 2. pH and electrical conductivity in the ash suspensions during the electrodalytic experiments, W-wet fly ash, SD-semidry APC residue.
Fig. 3. XRD diffractograms of initial and EDR treated MSWI residues, 7 days experiments. Q-Quartz, A – anhydrite, S-sylvite, C- carbonate, H- halite, B – bassanite, Cl- Calcium hydroxychloride.
Fig. 4. Leaching of heavy metals before and after electrodialytic remediation
Fig. 5. Chloride leaching in the MSWI residues after the ED experiments.