Optimization and assessment of an electrochemical advanced oxidation system for synthetic stormwater treatment

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Abstract
Electrochemical advanced oxidation processes (eAOPs) such as the current advanced oxidation system (AOS) are a type of electrochemical wastewater treatment that creates oxidative species, such as iodide species, chloride species, and hydroxyl radicals, that can treat even the most recalcitrant contaminants. It is important to determine the concentrations and locations of oxidative species in eAOPs for optimization of the wastewater treatment process. In this study, a spectrophotometric methodology was used to determine concentrations of iodide and chloride oxidative species (starting at 10, 25, and 50 ppm) within an AOS under various input voltages (6, 12, and 24 V). Overall, it was found that iodate and chlorite were the dominant species created in their respective treatments. Additionally, the concentration of iodide oxidative species increased with increasing voltage, whereas the chloride species decreased with increasing voltage. The optimal conditions for the efficient creation of AOS oxidative species were 12 V and 10 ppm potassium iodide and 6 V and 10 ppm sodium chloride, respectively. In addition, the use of iodide is recommended for wastewater treatment using the AOS to effectively create oxidative species. Following optimization, the AOS performance was tested for synthetic stormwater. Results indicated that the AOS performed well for reduction of Escherichia coli; however, reduction of other contaminants was inconsistent as would be expected given the AOS was optimized for disinfection, not decontamination. Further AOS optimization for decontamination would be expected to result in improved decontamination performance.

Keywords Advanced oxidation processes · Electrochemical treatment · Iodide · Chloride · Process optimization

Introduction
A wide variety of anthropogenically contaminated wastewaters enter receiving water bodies that are of ongoing concern globally due to their negative impacts on human and environmental health. Wastewaters may contain contaminants including dissolved and suspended solids, organic and inorganic matter, pathogens (including Escherichia coli), metals, pharmaceuticals, and polycyclic aromatic hydrocarbons (PAHs). Typical anthropogenic wastewaters include municipal, industrial, and agricultural wastewaters (Moreira et al. 2017; Sirés et al. 2014; Teh et al. 2016); landfill leachates (Moreira et al. 2015a; Oturan et al. 2015); and municipal stormwaters (Feng et al. 2018a; Fraser et al. 2018; Harper and Herr 1987; Mohanty et al. 2014; Taylor et al. 2015). Many technologies have been developed for the treatment of these wastewaters; however, there remain challenges in increasing efficiency and cost-effectiveness for the treatment of these wastewaters having widely varying matrices.

Conventional wastewater treatment processes can be categorized into physical, chemical, and biological processes. Examples of physical treatment methods include bar racks/screens, sedimentation, and filtration (Clark and Pitt 2012; Mickova 2015). Common chemical treatment
processes include coagulation/flocculation (Clark and Pitt 2012; Harper and Herr 1987; Sansalone and Kim 2008; Teh et al. 2016), air stripping towers, activated carbon adsorption, and ion exchange processes (Ding et al. 2017; Ganiyu et al. 2018; Moreira et al. 2015b). In addition, chemical treatments are often used for disinfection with chlorine, ozone, and UV typically used based on their ability to economically kill a variety of pathogens (Clark and Pitt 2012; Hussain et al. 2014; Mickova 2015). Examples of biological treatments include fixed growth, suspended solids growth, trickling filters, activated sludge, rotating biological contactor, facultative ponds, and aerated lagoons (Clark and Pitt 2012; Fraser et al. 2018; Mickova 2015; Mohanty et al. 2014; Moreira et al. 2015a). In general, these conventional processes are reasonably effective when used individually, or in treatment process trains; however, these may not be as effective in treating more complex and recalcitrant wastewater contaminants (Chauhan et al. 2021). For example, biological treatment processes transform complex pollutants into intermediates with the potential of accumulating in the environment (Bilińska et al., 2016). There remains a need for advanced wastewater treatment processes to complete the pollutant removal processes (Ahmadi et al. 2015).

Of more recent interest are advanced wastewater treatment processes, such as electrochemical advanced oxidation process (eAOP) technologies which rely upon oxidative electrochemical species for wastewater treatment (Bergmann et al. 2014; Feng et al. 2016; Ganiyu et al. 2018; Moreira et al. 2015a). Generally, eAOPs have been used as a pre-treatment to increase efficiencies of downstream processes or as a post-treatment polishing step to assist in meeting effluent standards (Chan et al. 2012; Moreira et al. 2015b; Oturan et al. 2015). Advantages of eAOPs include versatility, high efficiency, low resource consumption, and potential cost-effectiveness (Ding et al. 2017; Mickova 2015; Radjenovic and Sedlak 2015; Sirés et al. 2014). However, high electrode costs, low conductance of some wastewaters (e.g., stormwater), and the potential for release of toxic by-products can be challenges for eAOP technologies (Bergmann et al. 2014; Mickova 2015; Radjenovic and Sedlak 2015). Thus, there is a need for further optimization of both electric supply and operational conditions to improve treatment efficiencies, reduce energy consumption, and, in turn, be more cost-effective (Seibert et al. 2020).

A potentially low-cost method to increase treatment effectiveness is the addition of salts, such as iodide and chloride, that reduce a treatment reactor’s potential and increase the oxidation process efficiency (Cañizares et al. 2006; Llanos et al. 2014). This is especially the case for stormwater which can have relatively low electrical conductivity (herein fluctuating between 331 and 2,080 µs/cm). Similarly, previous studies have reported that the addition of salts can lead to electrogenerated strong oxidant formation during electrochemical oxidation resulting in higher pollutant (such as organic carbons, phenolic-based compounds, and pharmaceuticals) removal rates from hospital/industrial wastewater (Fajardo et al. 2017; Lan et al. 2017). Although salt addition may be beneficial for improved electrochemical treatment, it may still not be sufficiently effective for all pollutants. For example, sulphate addition has been reported to have limited effectiveness in the removal of salbutamol (Lan et al. 2017). Therefore, there is a need to investigate the interaction of salt and pollutants to determine optimal salt types and concentrations. Additionally, the reactor conditions need to be optimized for various salt concentrations and salt types to create the most effective oxidants at low voltages for cost efficiency.

Currently, our research group has been assessing a novel iodide-based eAOP known as the AOS (advanced oxidation system: Fig. 1; BioLargo Water Inc., Canada; Moustafa et al. 2021) for the treatment of municipal and synthetic stormwaters given its exceptional performance in
other water treatment applications (Moustafa et al. 2021). To our knowledge, this is the first application of iodide-based eAOP used for stormwater treatment. The determination of oxidative species created while treating real wastewaters is not realistic given their complex matrices and the short-lived nature of the created oxidative species within eAOP reactors. Thus, the first objective of the current study was to assess the AOS potential for the treatment of wastewaters through the determination of concentrations of produced oxidative iodide and chloride species within the reactor to better understand iodide-based eAOP versus a more typical chloride-based eAOP. Therefore, three concentrations (10, 25, and 50 ppm) each of potassium iodide (KI) or sodium chloride (NaCl) prepared in distilled water (DI) were added to the influent AOS water and three voltage levels (6, 12, and 24 V) were applied to the AOS. The KI has been recognized to create strong oxidative species such as iodate (IO$_3^-$) and periodate (IO$_5^-$) (Moustafa et al. 2021), while NaCl was chosen given chloride is frequently found in typical wastewaters and creates oxidative species including hypochlorite (ClO$^-$), chlorite (ClO$_2^-$), and chlorate (ClO$_3^-$).

In addition to DI water, tap water was considered in some experiments given it contains chloride in a simpler matrix than wastewaters; thus, oxidative species formed could be readily determined. Given the short-lived nature of the oxidative species, sampling and analysis via typical instrumentation would not be feasible. Thus, a novel methodology of determination of both iodide and chloride species was used such that the species could be determined quickly via UV–vis spectrometry given its simplicity, accuracy, quick response, and accessibility (Afkhami et al. 2001). The second objective was the assessment of a synthetic stormwater matrix including E. coli and a variety of other known stormwater contaminants to determine the potential for AOS to be used for the treatment of real stormwaters. Results determined from the overall AOS assessment and optimization, in addition to the synthetic stormwater treatment, may be useful in informing the treatment of real stormwaters, and other wastewater matrices, using this eAOP technology.

Methods

The study methodology for the optimization and assessment of the AOS application in the stormwater treatment includes an overview of the AOS configuration and its adjustment (“The AOS”); the determination of iodide and chloride oxidative species during the operation of AOS at 6, 12, and 24 V (“Determination of iodide and chloride oxidative species”); preparation of synthetic stormwater treatment (“Preparation of synthetic stormwater”); and its treatment via the AOS (“Synthetic stormwater treatment using the AOS”). This study excluded (1) the use of real stormwater containing various compounds for the determination of optimized AOS configuration given the difficulty of oxidative species monitoring and (2) the ion chromatography for the determination of both iodide and chloride species given its complexity and the oxidative species short lifespans making sampling and analysis impractical.

The AOS

The laboratory-scale AOS was a cylindrical reactor with a schematic shown in Fig. 1 as provided by BioLargo Water Inc., Canada. A detailed description of the AOS can be found in Moustafa et al. (2021) with a brief overview included herein. The reactor has a 1-inch diameter with a series of alternating proprietary materials used as anodes and cathodes made up of expanded graphite given its high stability, developed specific surface, electrical conductivity, and temperature resistance (Li et al. 2006; Li et al. 2007). The anodes and cathodes were subjected to voltages applied via a DC power supply while separated by glass wool to allow for the increased production of oxidative species. The reactor was operated in an upflow configuration using a peristaltic pump set at a constant 5 mL/min flow rate resulting in a residence time of about 30 min for all experiments. Sampling ports were located throughout the reactor with five anode ports (A1–A5) and three cathode ports (C1–C2, C3 = Outlet). The expectation was that oxidative species would be generated at the anodes and depleted at the cathodes, and the system cathodic cells would, thus, result in the minimization of the potential for release of active chemical discharges into the environment (Moustafa et al. 2021). Thus, the sampling ports were staggered to sample from each of these respective reactor volumes.

Before each experiment, the reactor was “recharged” to remove residual contaminants that could be in the reactor and to ensure the ability of the reactor to create oxidative species before samples were collected. Firstly, the reactor was flushed with distilled (DI) water for 30 min. Then, the voltage was turned on and the AOS run with either the iodide or chloride influent water for 30 min (treatment dependent). Samples were then taken from each port prior to analysis via UV–vis as discussed in the following section. To facilitate the reaction(s) used to measure the oxidative species, 1 mL each of the sample and 0.64 M KI, 2 mL of the buffer, and 6 mL of water were mixed in a 10-mL flask. The mixture was then transferred to a glass cell to be read by the spectrophotometer. It is important to note that the anode pH values typically decrease, while the cathode pH values increase making the reactor pH a dynamic environment (Jadhav and Ghangrekar 2009).
Therefore, several different buffers were used to adjust the pH of the samples in order to determine the individual oxidative species given the need for the associated reactions to each oxidative species at a certain pH to create peaks as shown in the Supporting Information (SI; Table S1). All experimental conditions included for the assessment of the AOS are summarized in Table 1.

### Determination of iodide and chloride oxidative species

The potassium iodide (KI) and sodium chloride (NaCl) were purchased from Fisher Scientific (Thermo Fisher Scientific, MA, USA) and were of ACS grade purity. The distilled water was produced via a Millipore Direct-Q 8 UV system with a conductivity of 18.2 MΩ/cm (Millipore Canada, ON, CA). The 10, 25, and 50 ppm solutions for each salt were produced fresh for each experimental run. A HACH DR 4000 UV–vis spectrophotometer (Hach Canada, ON, CA) with a 1-cm glass cell was used to determine the concentrations of the oxidative species. The periodate and iodate detection limits and ranges were 0.05–8.00 µg/mL and 0.05–5.00 µg/mL, respectively.

Calibration curves for each oxidative species at their corresponding absorbances were developed prior to experimental runs as discussed in the following sections. After recharging, the AOS was operated at the various concentrations and at 6, 12, and 24 V. Samples were taken from each sampling port, while the reactor was in operation, pH adjusted in a 10-mL flask, and ~1 mL was placed into the glass cell which was then immediately tested for absorbance. These experiments were conducted in the Environmental Engineering laboratories at the University of Saskatchewan.

**Iodide species in AOS**

The iodide species considered were IO$_3^-$ and IO$_4^-$ as they are predicted to be created by electrochemical oxidation. Prior to the current experiments, these have been shown to be the dominant species in the AOS based on parallel research done by our group at the HXMA beamline of the Canadian Light Source (CLS, SK, CA) as shown in Supporting Figure S1. Currently, the iodide species were determined using a method described by Afkhami et al. (2001) based on their reaction with excess iodide to create triiodide (I$_3^-$) as follows (Eqs. 1 and 2):

\[
\text{IO}_4^- + 11\text{I}^- + 8\text{H}^+ \leftrightarrow 4 \text{I}_3^- + 4\text{H}_2\text{O} \quad (1)
\]

\[
\text{IO}_3^- + 8\text{I}^- + 6\text{H}^+ \leftrightarrow 3 \text{I}_3^- + 3\text{H}_2\text{O} \quad (2)
\]

The concentration of the species was determined spectrophotometrically by measuring the absorbance of the I$_3^-$ at 352 nm (Afkhami et al. 2001; Feng et al. 2017; Wang et al. 2018). The reactions are pH dependent with IO$_3^-$ reacting at pH 6 and both species reacting at pH 3.3. These reactions will result in the creation of two equations for the absorbance values (Eqs. 3 and 4); therefore, the concentration of each individual species can then be determined when both species are in a mixture by solving these equations simultaneously.

\[
A_1 = a_1 + b_1 C_{\text{periodate}}
\]

\[
A_2 = a_2 + b_2 C_{\text{periodate}} + b'_2 C_{\text{iodate}}
\]

where $A_1$ and $A_2$ are the measured absorbances; $a_1$, $b_1$, $a_2$, $b_2$, and $b'_2$ are constants found when developing the calibration curves; and $C_{\text{periodate}}$ and $C_{\text{iodate}}$ are the concentrations (mg/L) of each species. The absorbance of known concentrations of IO$_3^-$ and IO$_4^-$ were measured to develop the calibration curves (Eqs. 3 and 4) with results presented in the Tables S2 and S3.
Chloride species in AOS

Species tested for chloride speciation included ClO\(^-\), ClO\(_2\)\(^-\), and ClO\(_3\)\(^-\). As for the iodide species, these were chosen because they are predicted to be electrochemically generated in the reactor. Unfortunately, a similar experiment as the iodide species in the CLS was not completed for chloride species. The chloride species were determined using a similar method as for I\(_3\)\(^-\) using Eqs. 5–8 which indicate that the chloride species will react with excess iodide to form I\(_3\)\(^-\) (Epstein and Kustin 1985; Mohammad et al. 2010; Narayana et al. 2005; Nowack and Von Gunten 1999).

\[
\begin{align*}
\text{ClO}_3\(^-\) + 6I^- + 6H^+ &\leftrightarrow 3I_2 + Cl^- + 3H_2O \\
\text{ClO}_2\(^-\) + 4I^- + 4H^+ &\leftrightarrow 2I_2 + Cl^- + 2H_2O \\
\text{ClO}^- + 2I^- + 2H^+ &\leftrightarrow I_2 + Cl^- + H_2O \\
I_2 + I^- &\leftrightarrow I_3^- 
\end{align*}
\]

The reactions are pH-dependent on the absorbance of the I\(_3\)\(^-\) was measured at different wavelengths to ensure that there was no interference between the oxidative chloride species. The concentration of hypochlorite was determined at a pH of 9.7 and a wavelength of 291 nm, ClO\(^-\) and ClO\(_2\)\(^-\) at pH 4 and 381 nm, and all three species at pH 4 and 300 nm (Table S4, S5, and S6). Note that the pH and wavelength for ClO\(^-\) and ClO\(_2\)\(^-\) were chosen by determining their maximum absorbances, while for all species, it was chosen where contributing absorbances for all three species can be measured as a maximum. The three equations used to determine the concentrations of each species in a mixture included:

\[
\begin{align*}
A_1 &= a_1 + b_1C_{\text{hypochlorite}} \\
A_2 &= a_2 + b_2C_{\text{hypochlorite}} + b_2'C_{\text{chlorite}} \\
A_3 &= a_3 + b_3C_{\text{hypochlorite}} + b_3'C_{\text{chlorite}} + b_3'C_{\text{chlorate}}
\end{align*}
\]

where A\(_1\), A\(_2\), and A\(_3\) are measured absorbances; a\(_1\), b\(_1\), a\(_2\), b\(_2\), b\(_2'\), b\(_3\), b\(_3'\), and b\(_3''\) are constants found when developing the calibration curves; and C\(_{\text{hypochlorite}}\), C\(_{\text{chlorite}}\), and C\(_{\text{chlorate}}\) are the concentrations (mg/L) of the each species.

Preparation of synthetic stormwater

Table 2 shows the general recipe for the synthetic stormwater (SSW) matrix that was used for the AOS experimental testing. All chemicals used in the SSW experiments were purchased from Fisher Scientific (Thermo Fisher Scientific, MA, USA) and were of ACS grade purity where applicable. The SSW was used as a representative alternative to real stormwater given it can be easily and reliably replicated in any laboratory, thus allowing for the potential for direct comparisons between the results of various studies. In comparison, real

| Component     | Material          | Concentration       | SSW experiment |
|---------------|-------------------|---------------------|----------------|
| NO\(_3\)\(^-\) | Sodium nitrate    | 2 mg/L              | 1              |
|               |                   | 2 to 20 mg/L        | 2              |
| Organic N     | Glycine           | 4 mg/L              | 1              |
|               |                   | 6 to 40 mg/L        | 2              |
| Phosphorus    | Dibasic sodium phosphate | 0.6 mg/L            | 1              |
|               |                   | 0.6 to 1.8 mg/L     | 2              |
| Dissolved solids | Calcium chloride  | 120 mg/L            | 1              |
|               |                   | 120 to 600 mg/L     | 2              |
| Suspended solids | Bentonite         | 25 mg/L             | 1              |
| E. coli       | E. coli MC4100     | 6×10\(^8\) CFU/100 ml | 1              |
| E. coli       | Epower E. coli ATCC 51,813 | 50 to 300 CFU/100 mL | 2              |
| Copper        | Cupric sulfate    | 0.08 mg/L           | 1              |
| Lead          | Lead chloride     | 0.08 mg/L           | 1              |
| Zinc          | Zinc chloride     | 0.6 mg/L            | 1              |
| Influent salt | Potassium iodide  | 10 ppm              | 1              |
|               |                   | 5 ppm               | 2              |
Stormwater samples taken as part of a parallel study by our research group showed wide-ranging stormwater compositions which would be expected to need differing AOS treatment optimization for each individual stormwater sample. Generally, there are many potential recipes for SSW in the literature that take this variability into consideration while also including differences due to location, time of year, and intensity of storm events. Ultimately, a SSW matrix was chosen based on literature that used a SSW to test urban stormwater treatment which has been used by a variety of researchers over the past 20 years (Davis et al. 1993; Hong et al. 2006; Okochi and McMartin 2012). There were two sets of SSW experiments used to test the AOS treatment process train. Stormwater Experiment 1 focused on disinfection using the AOS given its potential usage for wastewaters, including stormwater, having bacterial contamination including *E. coli*. Stormwater Experiment 2 included *E. coli* while additionally investigating the decontamination capability of the AOS in a treatment train. Note that the AOS was designed and optimized for disinfection and this research only determined the preliminary decontamination potential.

For Stormwater Experiment 1, sodium nitrate, glycine, dibasic sodium phosphate, bentonite, cupric sulfate, lead chloride, zinc chloride, and potassium iodide at the corresponding concentrations in Table 2 were mixed into a total volume of 120 L of reverse osmosis (RO) water (Millipore Canada, ON, CA). In addition, the *E. coli* standard strain was inoculated overnight in a 400-mL Luria Bertani broth (LB) using a pre-streaked plate of the MC4100 strain. The culture was then spun in a centrifuge, resuspended in RO water, and added to the SSW prior to running the experiments. The Stormwater Experiment 1 experiments were completed as part of an internship by N.B. at the laboratory facilities of BioLargo Water Inc. in Edmonton, AB.

For Stormwater Experiment 2, sodium nitrate, glycine, dibasic sodium phosphate, calcium chloride, and potassium iodide at the corresponding concentrations ranges shown in Table 2 (actual concentrations are included with experimental results in Table 5) were mixed into a total volume of 25 L of DI water produced via a Millipore Direct-Q 8 UV system (Millipore Canada, ON, CA). The *E. coli* was prepared using the standard method given by the supplier (Microbiologics Inc., MN, USA) and added to the SSW prior to running the experiments. These experiments were performed in the Environmental Engineering laboratories at the University of Saskatchewan in Saskatoon, SK.

**Synthetic stormwater treatment using the AOS**

Stormwater Experiment 1 focused on testing the disinfection of *E. coli* in a laboratory setup that included a larger version of the AOS reactor with a 2" diameter and 12 chamber configurations (BioLargo Water Inc., Canada), a peristaltic pump, and a DC power supply. The prepared SSW was pumped through the AOS in an upflow direction at a constant 1 L/min flow rate, and the AOS was connected to a DC power supply set at a constant 250 mA current. Samples to test the residual *E. coli* concentration were collected from the treated effluent every 30 min (including at time “0”). The samples were plated on LB agar (LBA) plates using a spiral plater, incubated at 35 °C for 24 h, and the colonies were counted. In addition, a sample from the “stock” SSW prior to treatment was collected and plated to determine the initial *E. coli* concentration. Note that experiments did not include any pre-treatment prior to the AOS for these runs.

The Stormwater Experiment 2 included a pre-treatment coagulation/flocculation step and the AOS treatment step. The coagulation/flocculation step would be expected to be used as an initial treatment process for the treatment of real stormwater to, at a minimum, decrease solids loading onto the AOS which could potentially lead to clogging of the reactor. Alum was chosen as the coagulant based on previous research (Harper and Herr 1987) and its historic use for water and wastewater treatment coagulation/flocculation. Jar tests were conducted to determine the optimum concentration of alum for the SSW pre-treatment process. In addition, this apparatus was also used for the coagulation pre-treatment step for these experiments. The ASTM standard of rapid mixing at 120 RPM for 1 min, slow mixing at 30 RPM for 20 min, and settling for 30 min (ASTM International 2013) was used for all experiments. The AOS laboratory setup consisted of influent pre-treated SSW, a variable speed peristaltic pump (VWR International, Edmonton, Canada) set at a constant pump rate (100 mL/min), a DC power supply (Newark, Canada) set at a constant 40 mA current, and the 1" 6 chamber laboratory scale AOS reactor (Fig. 1) which was used in the optimization iodide and chloride experiments. Samples were taken prior to treatment, after coagulation, and after the AOS.

Note that the current applied was different for each reactor from Stormwater Experiments 1 and 2 due to the differing AOS sizes (12 vs. 6 chambers; 2" vs. 1") which impacted the current density within each AOS. The total current density within the AOS was approximately 12.7 mA/cm² for Stormwater Experiment 1 (1.15 mA/cm² per electrode) as compared to 8.2 mA/cm² for Stormwater Experiment 2 (1.64 mA/cm² per electrode). However, the resultant applied voltages were in the ~7 to 8 V range for both experimental conditions.

**Synthetic stormwater quality parameter monitoring**

The total dissolved solids (TDS), electrical conductivity (EC), and pH were tested using typical laboratory meters. Chemical oxygen demand (COD) was measured using a
spectrophotometer, vials, and digester (HACH USA, CO, USA). The total suspended solids (TSS) were measured using Standard Methods for quantifying solids in wastewaters (“2540 SOLIDS (2017),” 2018). A total organic carbon (TOC) analyzer and related standard method were used to measure the TOC concentrations. Lastly, *E. coli* was measured using m-ColiBlue24 Broth PourRite Ampules (Hach USA, Colorado USA).

Fig. 2 Concentrations of oxidative iodide (A, B, C) and chloride (D, E, F) species present in the AOS at various KI and NaCl doses.
Results and discussion

Iodide results

The concentrations of iodate and periodate formed using 10, 25, and 50 ppm KI concentrations at various voltages are shown in Fig. 2A, B, and C as total oxidative iodide concentrations. The raw data and complementary information for the concentrations of each individual species are presented in Table S7 (10 ppm), S8 (25 ppm), and S9 (50 ppm). The resultant current that was applied to the reactor for each experimental condition is shown in Table 3.

Overall, iodate was the dominant oxidative species at all KI concentrations and all applied voltages, typically having 5 to 10 times higher concentrations than periodate for all sampled anode and cathode ports. Interestingly, periodate started to form at the higher applied voltage of 24 V and 10 ppm KI, and at the higher KI concentrations of 25 ppm and 50 ppm and voltage of 6 V. It is indicative that higher concentrations/voltages could lead to the formation of highly and even very highly oxidized species and demanding more input energy (Figure S1; very highly oxidized species herein were not reported given their very short life-span and associated difficulties in measurement). This higher energy input resulted in much higher currents applied in the AOS (Table 3), which also increased with increasing salt concentrations. These high currents generated excessive heat (~40–45 °C; experiments were performed at room temperature ~22 °C) that could potentially impact the integrity of the AOS. However, higher treatment temperatures may increase the effectiveness of treatment if the AOS was designed to handle these higher temperatures given faster reaction rates would likely result from the temperature increase. The creation of oxidative species increased with increasing voltages for all KI concentrations with 24 V showing the highest generation in all experiments (Fig. 2). However, the excessive heat generation, coupled with the increased costs that would be expected for the higher voltage input, makes the 24 V application unlikely for use in real wastewater treatment applications.

Iodate was found in the anodes and cathodes, while periodate was not found in either of the anodes or cathodes at 10 ppm KI and voltages less than 24 V. The concentration of the oxidative species mostly increased after the anodes and decreased after the cathodes indicating that iodate was being oxidized at the anodes and reduced at the cathodes. This difference between samples between the anodes and cathodes was more prominent at higher voltages and KI concentrations given the higher reduction/oxidation potential at higher voltages and concentrations of salt in the reactor. In contrast, there were some conflicting results of experiments. For example, the oxidative species concentrations reduced after A3 at 25 ppm KI and all voltages (Fig. 2B) and increased after C1 at 50 ppm KI (Fig. 2C). It should be noted that the sampling ports on the reactor were not perfectly aligned with anodes/cathodes while being closely spaced together, thus, complete reduction/oxidation between them would not be expected. These reactor issues can be attributed to the hand-packing process of the AOS which would be remedied by using a commercial reactor design and fabrication process.

In general, iodide oxidative species that can be formed include hypoiodous acid (HIO), iodite, iodate, and periodate with iodate being the most commonly formed and stable species (Wang et al. 2018; Ye et al. 2012). The use of iodide is more desirable than other salts, such as chloride and bromide, because the hazardous oxidative iodide species formed are readily transformed back to nonhazardous iodide (Feng et al. 2017). However, hazardous iodinated disinfection by-products such as iodoforms can also be formed (Wang et al. 2018; Ye et al. 2012).

The percentages of total oxidative species generated based on the input KI concentration are shown in Fig. 3A, B, and C. As would be expected, as the voltage increased there was also an increase in the oxidative iodate and periodate species, particularly at the anodes, with 10 ppm totals of 40–70%, 80–90%, and approximately 100% for 6, 12, and 24 V inputs, respectively. Following the same trend, 25 ppm totals were 20–60%, 50–100%, and 85–100% and 50 ppm totals were 15–30%, 70–90%, and 80–100%, respectively. Clearly, the lower the applied voltage, the lower the formation of oxidative species regardless of the initial KI concentration. Interestingly, there was a possibility that other oxidative species such as HIO and IO2− were being formed at lower voltages (Wang et al. 2018; Ye et al. 2012) but these were not determined in the current study. However, it is unlikely that HIO is formed since acidic conditions are needed for its formation and it is an unstable species (Wang et al. 2018). Based on this analysis, the optimal combination of maximum oxidative species and voltage selection for use

Table 3  Measured current (A) applied to the AOS at constant voltage inputs for various experimental treatments of iodide (KI), chloride (NaCl), and no salt added

| Treatment          | Current (A) |
|--------------------|-------------|
|                    | 6 V   | 12 V  | 24 V  |
| 10 ppm KI          | 0.01  | 0.02  | 0.07  |
| 25 ppm KI          | 0.02  | 0.05  | 0.10  |
| 50 ppm KI          | 0.01  | 0.05  | 0.10  |
| Tap + 50 ppm KI    | 0.08  | 0.22  | 0.65  |
| 10 ppm NaCl        | 0.01  | 0.04  | 0.15  |
| 25 ppm NaCl        | 0.01  | 0.05  | 0.20  |
| 50 ppm NaCl        | 0.02  | 0.07  | 0.25  |
| No salt added      | 0.02  | 0.07  | 0.31  |
in the AOS for treatment purposes would be 10 ppm and 12 V. The 10 ppm concentration had the highest percentage of iodate and periodate created; thus, an increased dosage may be unnecessary for treatment purposes. The 12 V treatments were markedly higher than the 6 V treatments, while the 24 V treatment was only marginally better than the 12 V
In addition, the added cost of applying 24 V versus 12 V and the potential negative impacts of the heat generation at 24 V make the 12 V option recommended.

The next experimental treatment considered tap water with 50 ppm KI treated in the AOS at 6, 12, and 24 V. The concentrations of the oxidative periodate and iodate species for the tap water treatment are shown in Fig. 4 in comparison to the 50 ppm KI DI water results shown previously, while the individual species raw data is included in Table S10. Interestingly, the tap water showed higher oxidative species concentrations for the 6 V and 12 V experiments versus the DI water results. In addition, the difference in concentration between the anodes and cathodes when tap water was used was greater than DI water. The increase in oxidative species may be due to the increased current (about 4 to 8 times higher) that was applied through the reactor at the same KI concentration and voltage applied for the tap water versus DI water treatments (Table 3). The increased current can be attributed to the higher number of ions present in tap water versus DI water. The increased current may have allowed the reactor to approach the threshold potential necessary for maximum production of oxidative species.

Clearly, tap water results indicate that iodide species may be naturally found in the tap water which can be oxidized within the AOS. Alternatively, the tap water may have other oxidative species present, or created in the AOS, that could have enhanced the production of the measured species iodate and periodate. For example, chloride is known to be present in tap water created in the City of Saskatoon (City of Saskatoon 2017) and is a common chemical added for the protection of treated water before distribution. Given this result, chloride was assessed for its potential to create oxidative chloride species within the AOS in the next stage of experiments.

**Chloride results**

The concentrations of chlorite and chlorate formed using 10, 25, and 50 ppm NaCl concentrations at various voltages are shown in Fig. 2D, E, and F as total oxidative chloride concentrations. The raw data and complementary information for each of the individual species are presented in Table S11 (10 ppm), S12 (25 ppm), and S13 (50 ppm). As for the iodide results, the resultant current applied to the reactor for each experimental condition is shown in Table 3.

Overall, chlorite was the dominant oxidative species at all NaCl concentrations and applied voltages to the AOS, whereas chlorate was only found in some experiments when 6 V was applied to the reactor. As for iodide, the current applied to the AOS increased with increasing NaCl concentration and applied voltage (Table 3). Creation of the two measured oxidative species was at a maximum at 6 V for the 10 and 25 ppm NaCl treatments and decreased with...
increasing voltages (Fig. 2). The 50 ppm NaCl treatment showed negligible concentrations of both chlorite and chlorate regardless of the input voltage. As compared to KI experiments, the trends for NaCl were unexpected and some results were conflicting. For example, the concentration of the oxidative species increased at the cathodes and decreased at the anodes with 6 V and 10 ppm NaCl treatment. However, they increased at the anodes and decreased at the cathodes at all voltages for the 25 ppm NaCl treatment. Thus, more research would be needed to better assess the trends for NaCl.

The percentages of total oxidative species generated based on the total input NaCl concentration are shown in Fig. 3D, E, and F. The measured oxidative species were only significantly formed for the 10 ppm NaCl treatment at 6 V with totals of 20–100%. Oxidative species were formed for all voltages for the 25 ppm NaCl treatment, decreasing with input voltages with totals of 25–100%, 15–30%, and 5–20% for 6, 12, and 24 V, respectively. There was a negligible concentration of chlorite and chlorate formed when 50 ppm NaCl was applied to the AOS as indicated previously. Generally, the lack of chlorite and chlorate may be due to the creation of other unmeasured oxidative species, such as perchlorate, being formed within the AOS. These species can be the result of higher currents generated in the reactor when increasing the applied voltage and/or increasing the salt concentrations (Table 3).

Chloride is a potentially viable salt to produce oxidative species as it is commonly found in many wastewaters making addition unnecessary for treatment purposes. Chloride species that can be formed include chlorine gas (Cl₂), hypochlorous acid (HClO), hypochlorite (ClO⁻), chlorite (ClO₂⁻), and chlorate (ClO₃⁻), with each species shown previously to be electrochemically generated (Bergmann et al. 2014; Oturan et al. 2015; Sirés et al. 2014). The HClO is the most powerful oxidant but is only formed in highly acidic conditions; it is also difficult to measure so was not included in the current species being assessed. The voltage applied to any electrochemical reactor system is important because if the anodic potential is too high, chlorate and perchlorate species can be formed. This scenario should be avoided as these species do not have any oxidation capacity and are known to be hazardous (Bergmann et al. 2014; Chaplin 2014; Moreira et al. 2017; Radjenovic and Sedlak 2015; Sirés et al. 2014). Thus, the determination of these species currently would be of little value as they are not valuable for wastewater treatment purposes. In addition, no hypochlorite was found in any of the samples, thus, it is not included in further discussion.

Clearly, the higher the applied voltage and NaCl treatment concentration in the AOS, the lower the formation of chlorite and chlorate, and potentially other species such as perchlorate can be created. Interestingly, Bergmann et al. (2014) used boron-doped diamond (BDD) electrodes and found that chlorate is steadily formed, and if the electrolysis time is long enough, all the chloride was reacted to perchlorate. Potentially using lower input voltages would lead to a decrease in applied current, or lowering the pH of the influent water below 4, which would help to achieve lower oxidation states of chloride, such as hypochlorite, to be formed for safe and effective treatment (Cabeza et al. 2007; Juang et al. 2013). Currently, the comparison between iodide and chloride was done at three input voltages based on preliminary studies; thus, lower voltage experiments were not considered.

Despite somewhat marginal results for NaCl, the optimal combination of NaCl concentration and applied voltage to create the most chloride within the AOS for treatment purposes would be 10 ppm and 6 V, respectively. Increasing the NaCl concentration and/or the applied voltage to the AOS led to decreases in the total concentrations and percentages of chlorite and chlorate formed while also increasing the risk of harmful higher oxidative species, such as perchlorate, being formed within the reactor.

**Comparing iodide and chloride**

Overall, the concentrations of oxidative species formed for each of the various iodide and chloride treatments can be compared in Figs. 2 and 3. The biggest difference between the salts is that the concentration of iodide oxidative species increases with increasing voltage, whereas the chloride species decrease with increasing voltage. In addition, the percentage of measured oxidative species is generally higher for iodide treatments in most experiments. For both salts, when the concentration of salt and/or applied voltage was increased, the resultant current also increases (Table 3). In addition, the higher concentration of ions in the tap water vs. DI water also resulted in an increased in current. Typical wastewaters can contain high concentrations of ions; therefore, the applied current within the AOS will be expected to be high even at low input voltages. This will make it difficult to keep the current low for real wastewater treatment; thus, it is expected that only higher chloride oxidative species will be formed for this treatment. However, the concentration of iodide oxidative species increased with increasing current making the use of iodide with the AOS being the recommended combination for consideration for the assessment of treatment for real wastewaters.

**Synthetic stormwater preliminary results**

Following the assessment of the AOS for KI and NaCl concentrations, an SSW matrix was considered for treatment using the AOS. The optimum conditions for KI and NaCl were 10 ppm and 10 ppm, and 12 V and 6 V, respectively.
Based on these results, the initial Stormwater Experiment 1 used a 10-ppm KI solution for the AOS treatment process. Given the disinfection results (see below) were successful, a 5-ppm KI solution was used for the Stormwater Experiment 2 under the assumption that the added SSW chloride (Table 2) would work synergistically in the treatment of the SSW. Similarly, the optimum voltage would be expected to be in the range of 6 to 12 V based on the AOS optimization experiments. The SSW experiments controlled the current at 250 mA (Experiment 1) and 40 mA (Experiment 2) to maintain similar current densities in each reactor design. The measured voltage was in the 7- to 8-V range for all experiments. Results for *E. coli* reduction by the AOS are shown in Table 4, while preliminary results for the determination of a suite of physicochemical parameters and use of a floculation treatment prior to the AOS are presented in Table 5.

Stormwater Experiment 1 included 13 sets of treatment runs (Table 4). The untreated stormwater had an average *E. coli* concentration of log 6.43 (CFU/100 mL) with a standard deviation of log 0.44 CFU/100 mL. The *T* = 0 min sample had log 3.68 (CFU/100 mL) with a high standard deviation of log 4.08 CFU/100 mL due to about half of these samples having values below the detection limits. This variability could be attributed to operational variables including bacteria present in the tubing or the reactor prior to start-up and therefore *T* = 30 min was considered the first timepoint to assess performance. At *T* = 30 min and later time points, no *E. coli* was detected in up to a total 120 L of SSW treated with the AOS. Thus, the AOS showed a consistent 6 log disinfection of *E. coli* for the SSW using 10 ppm KI and 250 mA applied current indicating that this technology is well suited to disinfection and has the promise to be successful for use in real stormwater treatment processes. This high efficiency for disinfection by the AOS is not surprising, as previous studies have shown consistent disinfection of wastewaters using other eAOPs. For example, Rajab et al. (2015) used a boron-doped diamond eAOP and generated chloride oxidative species to achieve 4–8 log inactivation of *Pseudomonas aeruginosa*. Similarly, Feng et al. (2018b) used a dimensional stable anode (DSA) eAOP and chloride species to achieve a 3 log (or total in this case) disinfection of *E. coli* present in both SSW and real stormwater samples. Additionally, Cano et al. (2012) used a conductive diamond eAOP for treating a wastewater and found a greater than 4 log disinfection of *E. coli*.

Following the disinfection success of Stormwater Experiment 1, Stormwater Experiment 2 was used to test the ability of the AOS to provide for both disinfection

### Table 4 Synthetic stormwater (SSW) results of Stormwater Experiment 1 for *E. coli* reduction using 10 ppm KI and 250 mA (~7 to 8 V)

| Samples       | n  | Average (log CFU/100 mL) | SD (log CFU/100 mL) |
|---------------|----|-------------------------|---------------------|
| SSW stock     | 13 | 6.43                    | 0.44                |
| AOS treated   | 52 | 0.00                    | 0.00                |

### Table 5 Preliminary synthetic stormwater (SSW) results for Stormwater Experiment 2 including all physicochemical parameters using 5 ppm KI and 40 mA (~7 to 8 V). The modified SSW recipes were based on actual stormwater sample data collected as part of a parallel study (data not shown)

| Experiments       | EC (µS/cm) | TDS (mg/L) | pH  | COD (mg/L) | TOC (mg/L) | TSS (mg/L) | *E. coli* (CFU/100 mL) |
|-------------------|------------|------------|-----|------------|------------|------------|------------------------|
| 30 June Initial   | 1409       | 700        | 7.12| 83.1       | 18.8       | 160        | 200                    |
| 60 ppm Alum       | 1575       | 784        | 6.10| 70.2       | 19.2       | 2.5        | 40                     |
| AOS               | 1373       | 680        | 6.15| 68.0       | 16.8       | 6.5        | 6.5                    |
| 3 July Initial    | 2010       | 1010       | 6.88| 329        | 32.0       | 110        | 310                    |
| 60 ppm Alum       | 2090       | 1058       | 6.18| 294        | 30.9       | 3.5        | 15                     |
| AOS               | 1825       | 913        | 5.76| 228        | 9.60       | <1.0       | 7.5                    |
| 7 September Initial | 511       | 248        | 6.90| 39.5       | 7.15       | 110        | 42                     |
| 60 ppm Alum       | 577        | 280        | 6.94| 32.5       | 2.73       | <1.0       | 17                     |
| AOS               | 318        | 153        | 5.86| 17.5       | 0.48       | <1.0       | <1                     |

*Modified SSW recipe* | Sodium nitrate (mg/L) | Glycine (mg/L) | Dibasic sodium phosphate (mg/L) | Calcium chloride (mg/L) | *E. coli* (CFU/100 mL) |
|----------------------|----------------------|----------------|-----------------------------|------------------------|------------------------|
| 30 June              | 6                    | 12             | 1.8                         | 480                    | 200                    |
| 3 July               | 20                   | 40             | 6                           | 600                    | 300                    |
| 7 September          | 2                    | 6              | 0.6                         | 120                    | 50                     |
and decontamination including three preliminary treatment runs. For these experiments, the KI concentration was reduced to 5 ppm to reduce the potential costs associated with the addition of KI in the treatment process. In addition, results prior to these experiments suggested that higher 10 ppm KI concentration may not be necessary for the treatment of more complex matrices such as the current SSW which have other salts already available for creation of oxidative species. The E. coli initial concentrations were also reduced in these experiments (Table 5) to better represent concentrations found in real stormwater samples collected by our research team as part of a parallel study (Table S15). In addition, the sodium nitrate, glycine, dibasic sodium phosphate, and calcium chloride concentrations added to the SSW were varied to better represent different contaminant variations in these real stormwaters (Table 5). Overall, the EC and TDS concentrations remained unchanged through the treatment process train and followed the added concentrations of calcium chloride and sodium nitrate which would be expected to impact these parameters. Generally, the COD, TOC, and TSS all decreased through the coagulation/flocculation and AOS treatment process with the TSS showing the largest reduction due to the coagulation/flocculation process. The initial E. coli concentrations were also reduced during the coagulation/flocculation process to a range of log 17 to 40 CFU/100 mL, while the AOS reduced these further to the range of log <1 to 7.5 CFU/100 mL.

Overall, the Stormwater Experiment 1 indicated that the AOS has a high capacity for disinfection of stormwater, or other wastewaters, having high initial E. coli concentrations. It would be expected that this capability would also be beneficial for the reduction of other wastewater pathogens. The results for Stormwater Experiment 2 showed that the addition of the coagulation/flocculation step prior to the AOS provided for a marked decreases in TSS and E. coli concentrations with the AOS process further “polishing” the SSW for these two parameters. Additionally, it is important to note that the electrode surface structure might contribute to pollutant removal processes; however, the previous study has shown that there is minimum adsorption to the electrode surface without iodine and/or electrical potential (Moustafa et al. 2021). Assessment of the AOS for treatment of real stormwater samples is currently underway in our research labs and will be presented in a future research article as they are beyond the scope of the current study. However, the results here for the SSW indicate that the treatment process train including coagulation/flocculation followed by the AOS shows promise for treatment of real stormwater samples.

Conclusions

In this study, the concentrations of iodate and periodate formed from the potassium iodide addition and chlorite and chlorate formed from the sodium chloride addition were determined in the AOS reactor. Operating parameters analyzed the impacts of different salt concentrations and input voltages on the production of these oxidative species. Overall, the results showed that the addition of salts to SSW can be useful to decrease the applied voltage needed for effective treatment, and the iodine-based AOS produces oxidative species with disinfection and decontamination capabilities indicating that it has the potential for effective treatment of real stormwaters, as well as other wastewaters. The dominant oxidative species were found to be iodate with iodide addition and chlorite with chloride addition. For iodide, the optimal conditions for the creation of oxidative species were found to be 10 ppm KI and 12 V. Similarly, the 10 ppm NaCl and 6 V conditions were optimal for chloride. The AOS showed excellent disinfection of SSW E. coli using 10 ppm KI and a 2" 12 chamber reactor. The AOS also exhibited E. coli disinfection of SSW E. coli using 5 ppm KI and a 1" 6 chamber reactor. Additional studies testing real stormwater treatment and energy cost calculations will have to be done to further test the disinfection and decontamination capability of the AOS and evaluate its cost-effectiveness.

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