

The direct anodic oxidation or electrolysis occurs directly on the anode (M) and involves direct charge transfer reactions between the anode surface and the organic pollutants involved. The mechanism only involves the mediation of the Adsorption electrons, which are capable in oxidizing some organic pollutants at defined potentials more negative the oxygen evolution reaction (OER) potential. The direct electrolysis usually requires prior adsorption of pollutants onto the anode surface, which is the rate-Direct limiting process and does not lead to the overall Charge Transfer combustion of organic pollutants. In direct electrolysis, the pollutants are oxidized after adsorption on the anode surface without the involvement of any substances other than the electron, which is a "clean reagent":

Desorptio

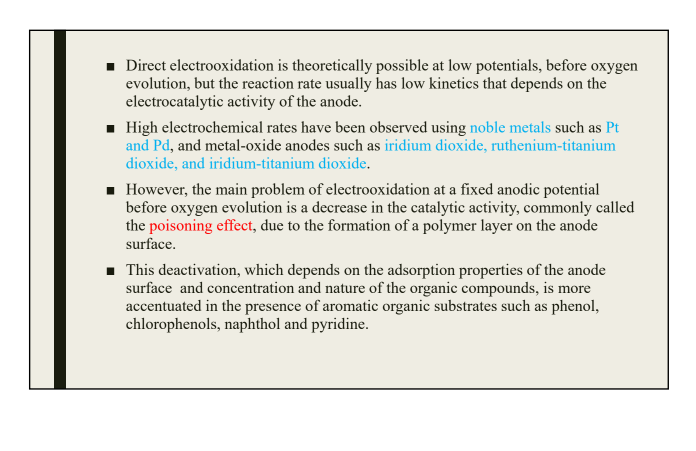
Cathode

(Reduction)

Anode

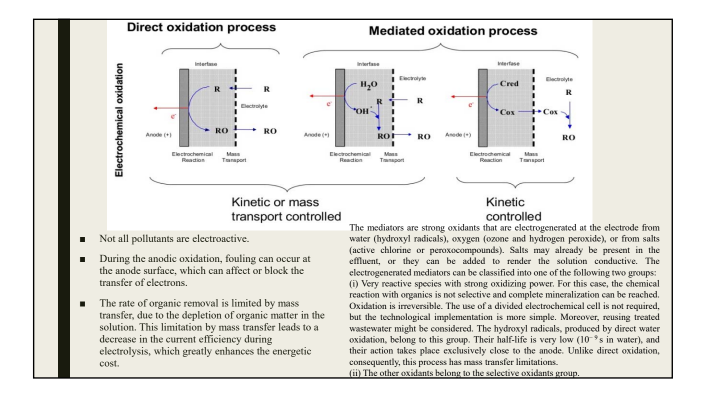
Oxidation)

■ R_{ads} -ze⁻ $\rightarrow P_{ads}$



INDIRECT ELECTROOXIDATION

The indirect EO processes are mediated by the in situ electro-generation of highly oxidant species at the electrode surface. Different kinds of oxidant species can be generated by the EO process (i) reactive oxygen species and (ii) chlorine active species.



Electro-generation of reactive oxygen species

- Reactive oxygen species (ROS) are reactive chemical species containing oxygen such as hydrogen peroxide (H₂O₂), ozone (O₃), or hydroxyl radical (•OH). Their chemical reactivity is due to the oxygen molecule's unpaired electron.
- The generation of such oxidants strongly depends on several key reaction parameters. electrode material,

electrolyte composition,

applied current (or voltage),

pH, and

temperature.

The anode material is the key parameter. Because all oxidants are formed at high potentials, the competitive reaction is the formation of oxygen. An anode material with a high oxygen overpotential is required.

■ The indirect EO by reactive oxygen species is based on the electrogeneration of adsorbed hydroxyl radical (•OH) (E° = 2.8 V/SHE) onto the anode surface as an intermediate of the OER (1)

Anode:
$$M + H_2O \rightarrow M(\bullet OH) + H^+ + e^-$$
 (1)

where, M is referred the anode and M(•OH) is the adsorbed hydroxyl radical.Reaction between an organic compound R and hydroxyl radicals (loosely adsorbed on the anode) takes places close to the anode's surface.(n:the number of electrons involved in the oxidation reaction of R).

$$R_{(aq)} + M (\bullet OH)_{n/2} \rightarrow M + Oxidation \text{ products} + n/2 H^+ + n/2 e^-$$
 (2)

However, the inevitable competitive reactions (3) and (4) that consume the radical species leading to oxygen evolution are also feasible.

$$M(\bullet OH) + H_2O \rightarrow M + O_2 + 3 H^+ + 3 e^-$$
 (3)

 $2M(\bullet OH) \rightarrow 2 M + O_2 + 2 H^+ + 2 e^-$ (4)

- In order to produce greater amounts of M(•OH), anodes with high overpotential for OER should be used to promote reaction (1) and to avoid the parasitic reactions (3) and (4). Electrodes for wastewater treatment can be classified under two groups regarding to high overpotential oxygen evolution : "active" and "non-active" anodes.
- The different performance of these anodes is related to the enthalpy of adsorption of the OH radicals onto the anode surface. Physisorbed species are more oxidant than the strongly chemisorbed ones that are represented by reaction (1) and reaction (5), respectively.

 $M(\bullet OH) \rightarrow MO + H^+ + e^-$

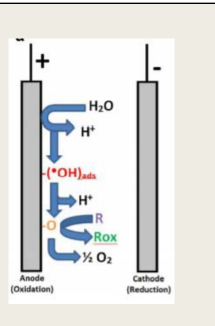
(5)

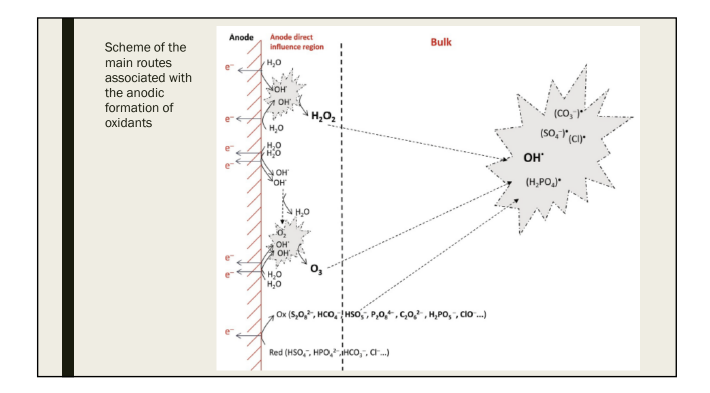
where, \underline{MO} represents the oxidant species of the so-called higher oxide that is generated onto the anode surface by the chemisorption of OH radicals

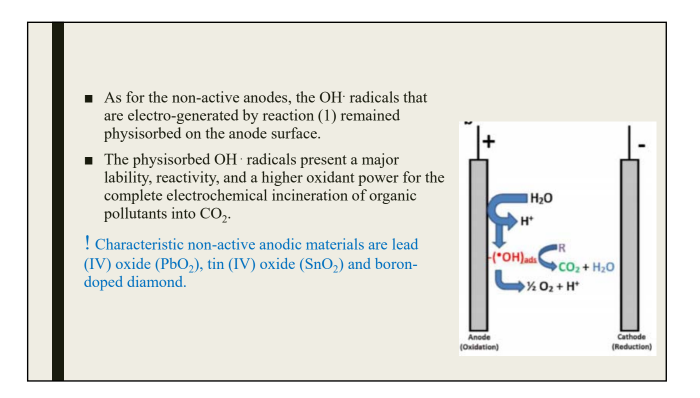
!The active anodes are only capable in inducing the electrochemical conversion of organics into more biodegradable molecules such as short-chain carboxylic acids, but they cannot achieve complete mineralization or organics combustion into carbon dioxide (CO_2)

■ This occurs because higher oxidation states are available for these metal or metal oxide anodes above the standard potential for OER (E° = 1.23 V/SHE), leading to the preferential formation of chemisorbed active oxygen species MO by the concatenated reactions (1) and (5). Following this, the oxidation is mediated by the reaction of pollutants with the chemisorbed MO.

! Characteristic <u>active anodic materials</u> are platinum (Pt), dimensionally stable anodes (DSA) of ruthenium (IV) oxide (RuO₂), iridium (IV) oxide (IrO₂), and other mixtures of metal oxides.







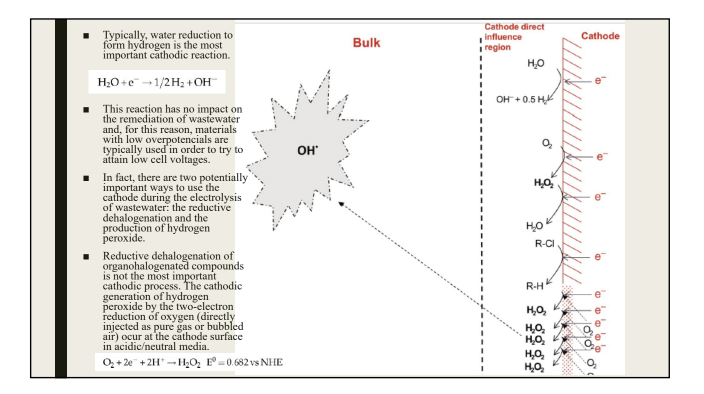
Other reactive oxygen species

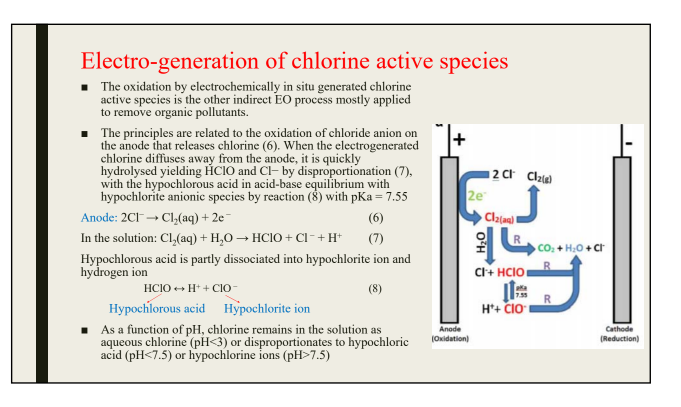
- One of the main disadvantages of EO of organics with electrogenerated ClO- ions is the formation of toxic chlorinated intermediates especially while working in acidic medium. Therefore, EO by other reactive species is of interest.
- Indirect EO of organic compounds can be enhanced by electro-Fenton reactions, electrogeneration of hydrogen peroxide, peroxodisulfate, peroxidiphosphate, and ozone.

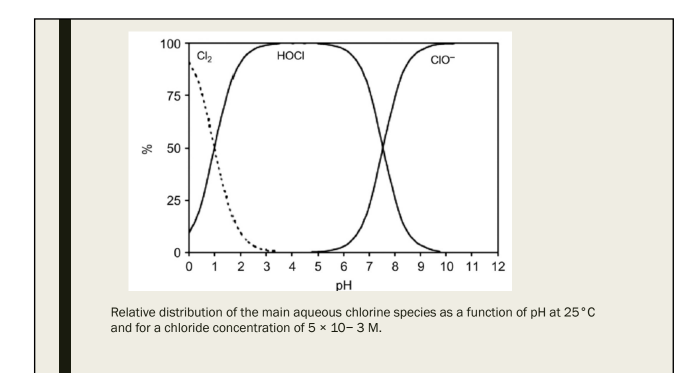
 $2SO_{4}^{2-} \rightarrow S_{2}O_{8}^{2-} + 2e^{-}$ $H_{2}PO_{4}^{-} + (H_{2}PO_{4})^{\bullet} + e^{-}$ $H_{2}PO_{4}^{-} + OH \rightarrow (H_{2}PO_{4})^{\bullet} + OH^{-}$ $HPO_{4}^{2-} \rightarrow (HPO_{4}^{-})^{\bullet} + e^{-}$ $HPO_{4}^{2-} + HO^{\bullet} \rightarrow (HPO_{4}^{-})^{\bullet} + OH^{-}$ $O_{2} + 2H^{+} + 2e^{-} \rightarrow H_{2}O_{2}$ $H_{2}O_{2} + 2H^{+} + 2e^{-} \rightarrow 2H_{2}O$

Cathode

Typically, water reduction to form hydrogen is the most important cathodic reaction. This reaction has no impact on the remediation of wastewater and, for this reason, materials with low overpotencials are typically used in order to try to attain low cell voltages and, at least, to not increase the operation cost of the wastewater treatment unnecessarily.







Standard Potential (V)	Reduction Half-Reaction			
2.87	$F_2(g) + 2e^- \longrightarrow 2F^-(ag)$		The species near the prefer to be reduced	
1.51	$MnO_{a}^{-}(aq) + 8H^{+}(aq) + 5e^{-} \longrightarrow Mn^{2+}(aq) +$	4H,O(1)	agents)	(Bood ovidizing
1.36	$Cl_2(g) + 2e^- \longrightarrow 2Cl^-(ag)$	2	while the species ne	ar the bottom
1.33	$Cr_2O_2^{2-}(aq) + 14H^+(aq) + 6e^- \longrightarrow 2Cr^{3+}(aq)$	+ 7H,O(l)	prefer to be oxidized	
1.23	$O_2(g) + 4H^+(aq) + 4e^- \longrightarrow 2H_2O(l)$	2	agents.	
1.06	$Br_2(l) + 2e^- \longrightarrow 2Br^-(aq)$			
0.96	$NO_3^{-}(aq) + 4H^+(aq) + 3e^- \longrightarrow NO(g) + H_2O_3^{-}(aq) + H_2O_3^{-}(aq)$	O(l)		
0.80	$Ag^{+}(aq) + e^{-} \longrightarrow Ag(s)$		 Note that the standard 	l reduction
0.77	$ \begin{array}{l} \operatorname{Fe}^{3+}(aq) + e^{-} \longrightarrow \operatorname{Fe}^{2+}(aq) \\ \operatorname{O}_{2}(g) + 2\mathrm{H}^{+}(aq) + 2e^{-} \longrightarrow \mathrm{H}_{2}\mathrm{O}_{2}(aq) \\ \operatorname{MnO}_{2^{-}}(aq) + 2\mathrm{H}_{2}\mathrm{O}(l) + 3e^{-} \longrightarrow \mathrm{MnO}_{2}(s) + 4\mathrm{OH}^{-}(aq) \end{array} $		potential of $Cl_2(aq)$ (E° = 1.36 V/SHE)	
0.68			and HClO ($E^{\circ} = 1.49$ V/SHE) are considerably higher than ClO ⁻ ($E^{\circ} =$	
0.59				
0.54	$I_2(s) + 2e^- \longrightarrow 2I^-(aq)$		0.89 V/SHE), indicati	
0.40	$\tilde{O}_2(g) + 2H_2O(l) + 4e^- \longrightarrow 4OH^-(aq)$		oxidation of organics	
0.34	$Cu^{2^+}(aq) + 2e^- \longrightarrow Cu(s)$		chlorine active specie acidic pH conditions	s is obtained und
0	$2H^+(aq) + 2e^- \longrightarrow H_2(g)$		-	
- 0.28	$Ni^{2+}(ag) + 2e^{-} \longrightarrow Ni(s)$	$Cl_{2}(g) + 2$	$2 e \rightleftharpoons 2 Cl^{-}$	1.35827
- 0.44	$Fe^{2+}(aq) + 2e^{-} \longrightarrow Fe(s)$	HClO+	$H^+ + e \rightleftharpoons 1/2 Cl_2 + H_0$	1.611
- 0.76	$Zn^{2+}(aq) + 2e^{-} \longrightarrow Zn(s)$		$H' + 2e \neq Cl' + H,O$	1.482
- 0.83	$2H_2O(l) + 2e^- \longrightarrow H_2(g) + 2OH^-(aq)$			
- 1.66	$Al^{3+}(aq) \div 3e^{-} \longrightarrow Al(s)$	CIO" + 1	$L_0 + 2e \rightleftharpoons Cl^- + 2OH^-$	0.81
-2.71	$Na^+(ag) \div e^- \longrightarrow Na(s)$			

The formation of hypochlorite ions

• The formation of hypochlorite ions is dependent on many parameters such as

*concentration of chloride ions in the electrolyte solution,

*the temperature of the electrolyte solution,

*applied current density,

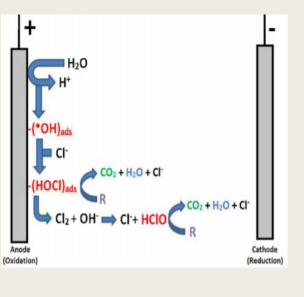
*material of anode, and cathode.

- For example, higher concentration of chloride ions (usually higher than 3 g/L) in the electrolyte solution leads to the higher formation of hypochlorite ions; higher temperature of electrolyte solution leads to the higher evolution of oxygen, which is the waste reaction, and results to the lower formation of ClO⁻.
- In this regard, the temperature control should be carried out during indirect EO and addition of sodium chloride salt can be required in the case of low chloride ions concentration in the treated wastewater. Anodic materials should have low overpotential toward chlorine evolution, which in turn leads to higher ClO⁻ production. Cathodic materials should have inert properties toward ClO⁻ ions reduction.

The possible simultaneous electrogeneration of chlorine active species mediated by the reactive oxygen species is such that the oxygen transfer reactions will be carried out by OH)ads oxychlorinated adsorbed species generated by reaction (9)as intermediates of the chlorine evolution (10).HOCI)

 $\blacksquare M(OH) + Cl \rightarrow M(HOCl)$ (9)

$$\blacksquare M(HOCl) \rightarrow \frac{1}{2}Cl_2 + OH - (10)$$



- The active anode materials present better performances than the non-active anode materials in the electrochemical anodic oxidation mediated by chlorine active species are
- The better electrocatalytic properties for chlorine evolution is described for active anodes (i.e., Pt, RuO₂, TiO₂ and IrO₂) where chloride is oxidized to chlorine active species
- Conversely, the non-active anodic materials result to the further oxidation of Cl₂ and HClO/ClO⁻ to undesired <u>non-oxidising chlorine species</u>.
- The degradation of organic pollutants mediated by electrogenerated chlorine active species is a process of great industrial interest due to the ubiquitous presence of chlorine in mainstream water bodies and industrial effluents.
- Even though the oxidation of organic pollutants is considerably faster with homogeneous chlorine active species than the mediated ones by adsorbed hydroxyl radicals, they also could yield undesirable organo-chlorinated by-products (e.g. haloacetic acids, halomethanes, etc) and noxious ionic species, such as chlorate and perchlorate

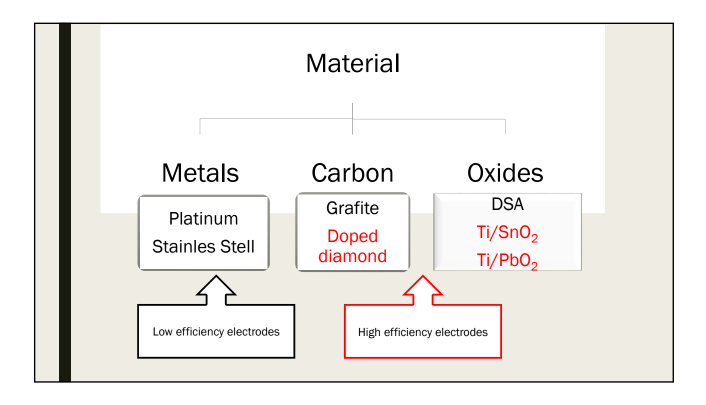
Side reactions:

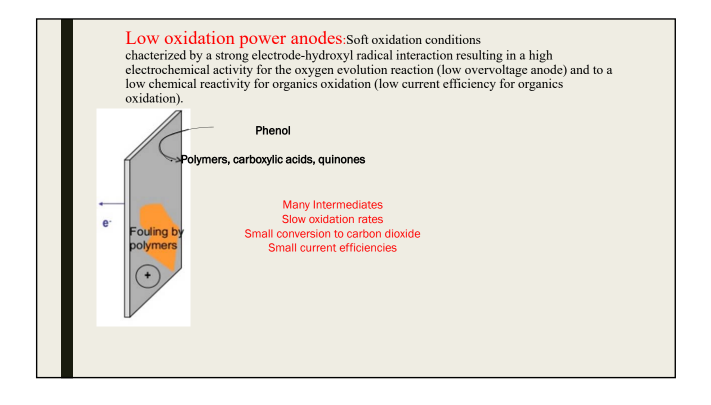
- The process of hypochlorite ions formation can be interrupted by side reactions. For example, ClO⁻ ions can be lost in anodic and cathodic transformation reactions with the formation of chlorate and chloride ions, respectively (R. 11-12).
- At anode: $12\text{ClO}^- + 6\text{H}_2\text{O} 12\text{e}^- \rightarrow 4\text{ClO}_3^- + 8\text{Cl}^- + 3\text{O}_2 + 12\text{H}^+$ (11)
- At cathode: $ClO^+ H_2O + 2e^- \rightarrow Cl^- + 2OH^-$ (12)

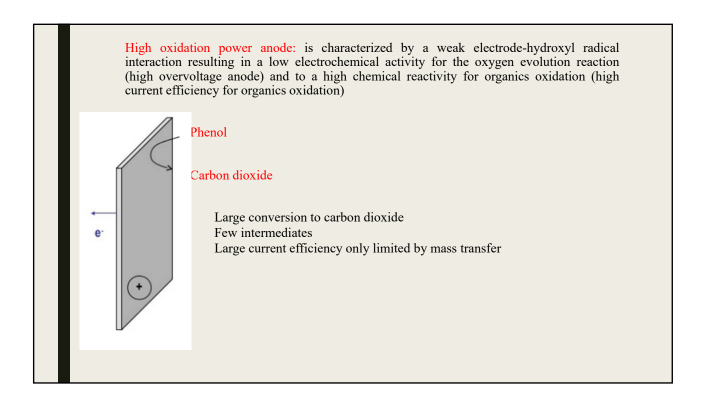
Electrode Materials

Several electrodes have been used for water treatment by electrochemical oxidation.

- Mechanical stability
- Chemical stability
- Morphology
- Electrical Conductivity
- Catalytic properties
- Ratio price/lifetime
- For generating reactive oxygen species, a high oxygen overpotential is required; otherwise, a large portion of the applied current will be wasted to produce oxygen as a side reaction, thereby reducing the efficiency of the electrochemical process. Several electrode materials such as PbO₂, SnO₂-based DSA, and BDD have been reported to exhibit high oxygen overpotential. These electrode materials are able to hinder the thermodynamically favored oxygen evolution reaction, thus improving the ROS generation efficiency.







• Active electrodes (have low oxygen evolution overpotential and aconsequently are good electrocatalysts for the oxgen evolution reaction: only permit the partial oxidation of organics)

Carbon and Graphite

Pt based anodes

IrO₂ Iridium-based oxides

RuO₂ Ruthenium-based oxides

■ **Non-active electrodes** (have high oxygen evolution overpotential and consequently are poor electrocatalysts for the oxygen evolution reaction and favor the complete oxidation of organics to CO₂ and so)

Ti/SnO₂ (Antimony-doped tin oxide)

Ti/PbO₂ (lead dioxide)

Boron-doped diamond (BDD)

Drawbacks of non-active electrodes:

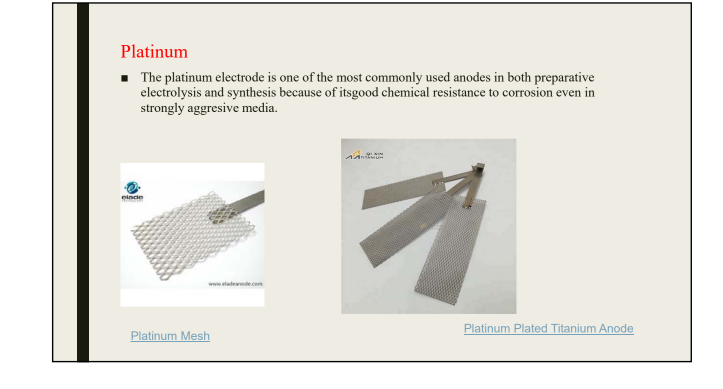
Conductive diamond: large price

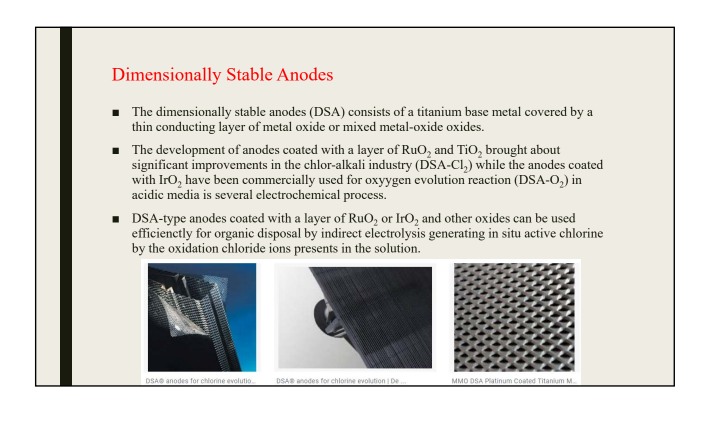
PbO₂/SnO₂: Dissolution of toxic species

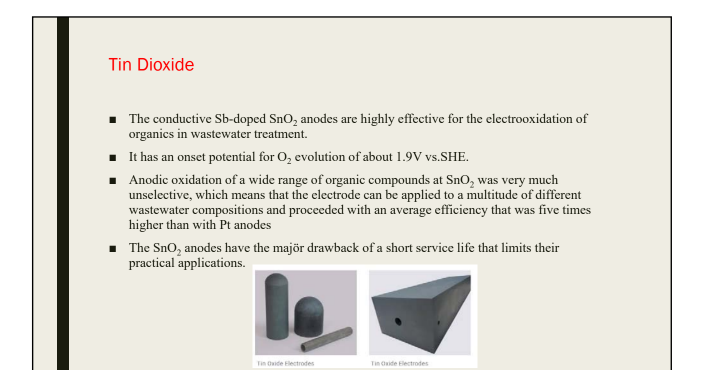
Carbon and Graphite

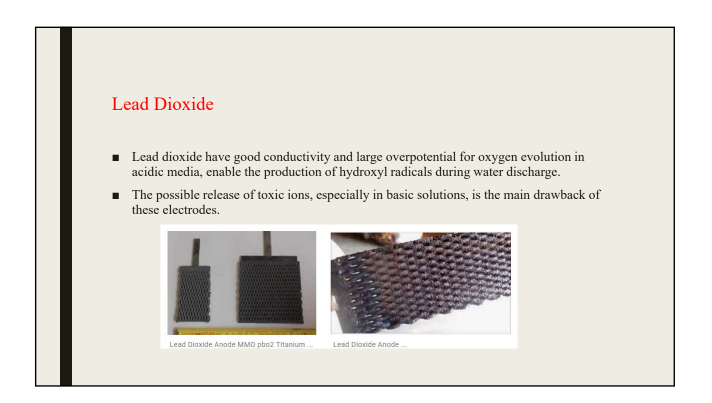
- Carbon and graphite electrodes are very cheap and have a large surface area.
- They have been used widely for the removal of organics with three-dimensional electrodes (e.g. Packed bed, fluidized bed, carbon particles, porous electrode, etc.)
- During the electrolysis there was a rapid decrease in the reaction rate due to the blocking of the anode surface with insoluble polymeric products that were slow to oxidize or desorb.





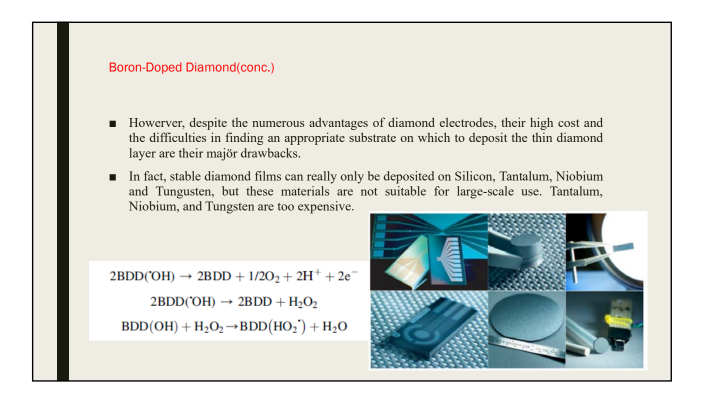






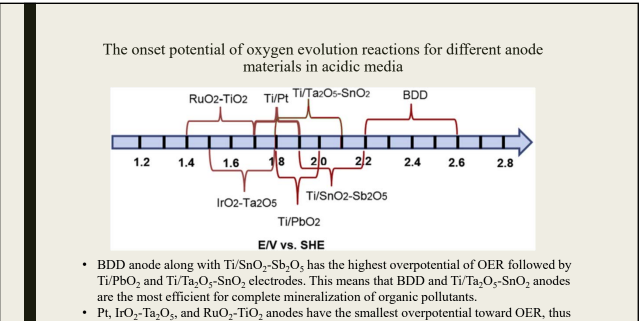
Boron-Doped Diamond

- High quality BDD electrodes possess several technologically important characteristics including an inert surface with low adsorption properties, remarkable corrosion stability even in strong acidic media, and extremely high oxygen evolution overpotential.
- During the electrolysis in the region of water discharge, a BDD anode produces a large quality of the OH that is weakly adsorbed on its surface, and consequently it has high reactivity for organic oxidation, providing the possibility of efficient application to water treatment.
- BDD anodes allow complete minerilization of several types of organic compounds.
- It has been shown that the oxidation is controlled by the diffusion of the pollutants toward the electrode surface, where the hydroxyl radicals are produced, and the current efficiency is favored by high mass-transport coefficient, high organic concentration, and low current density.
- Performing electrolysis under optimum conditions, without diffusion limitation, the current efficiency approaches 100%.



Summary

- Anodes with low oxygen evolution overpotential, such as graphite, IrO₂, RuO₂, or Pt only permit the primary oxidation of organics, but not to complete mineralization, due to the accumulation of oxidation intermediates, mainly aliphatic acids, which are quite stable against further attack at these electrodes.
- The complete mineralization of the organics to CO₂ and good Faradic efficiency can be obtained using high oxygen overpotential anodes, such as SnO₂, PbO₂, and BDD, because these electrodes involve the production of oxygen evolution intermediates, mainly hydroxyl radicals, that nonselectively oxidize the organic pollutants and their intermediates.
- Despite their notable ability to remove organics, doped-SnO₂ anodes have the major drawback of a short service life that limits their practical applications.
- Even the application of Ti/PbO₂ anodes to wastewater treatment may be limited by the possible release of toxic lead ions, due to their dissolutions under specific anodic polarization and solution composition.
- On the contrary, conducting diamonds offer significant advantages over other electrodes in terms of current efficiency and stability. However, further improvements, such as finding an appropriate substrate on which to deposit the thin diamond layer and reduction of production costs, are required before their wide indutrial application.



Electrode	Oxidation potential (V)	Overpotential of OER (V)	Adsorption enthalpy of M-OH	Oxidation power of anode	
tuO ₂ - TiO ₂ DSA® - Cl ₂)	1.4-1.7	0.18	Chemisorption of OH radical	Lower	
$rO_2 - Ta_2O_5$ DSA [®] - O ₂)	1.5-1.8	0.25			
Γi/Pt	1.7-1.9	0.3			
Гі/РbO ₂	1.8-2.0	0.5			
Fi/SnO ₂ - Sb ₂ O ₅	1.9-2.2	0.7		マフ	
o-Si/BDD	2.2-2.6	1.3	Physisorption of OH radical	Higher	

Table 1. Oxidation power anodes and potential for oxygen evolution of various anode materials in acidic media. Standard potential for oxygen evolution is 1.23 V vs normal hydrogen electrode. Adapted from refs. 6 and 7

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	ElectrodeType Noble metal electrodes (Pt, Au)	Advantages Stable in a wide range or potentials and pH Excellent repeatability properties Intensive use in laboratory scale for new process investigations	Disadvantages Expensive Low mineralization efficiency Low overpotential toward OER Poor use in industrial wastewater treatment application	
Advantages and Disadvantages of Different	PbO ₂	Cheap Relatively high overpotential toward OER Relatively high ability to mineralize organics	Potential leaching of toxic Pb Poor performance in industrial wastewater treatment application	
Group Anodes Used in Electrochemical Oxidation Applications of	Carbon and graphite electrodes	Cheap Intensive use in laboratory scale for new process investigations	High electrode corrosion rates Low mineralization efficiency Low overpotential toward OER	
Organic Compounds	$\begin{array}{l} \mbox{MMO} \\ (Ti/TiO_2-RuO_2, \ Ti/ \\ Ta_2O_5-IrO_2, \ Ti/ \\ TiO_2-RuO_2-IrO_2, \ Ti/ \\ IrO_2-RuO_2, \ Ti/ \\ SnO_2-Sb_2O_5, \ etc.) \end{array}$	High stability Good conductivity properties Acceptable price Possibility to regenerate catalytic oxide coating	Sometimes it is difficult to reproduce quality of the catalyst layer Potential leaching of toxic compounds such as Sb	
	BDD	High overpotential toward OER High ability to mineralize organics Excellent conducting properties even at low temperatures High electrochemical stability and corrosion resistance	Expensive Reduced efficiency in diluted solutions and at increasing current density higher than a limiting current	
	BBD, boron-doped diamond; MMO, mixed-metal oxide; OER, oxygen evolution reactions.			

References

- Sergi Garcia-Segura, Joey D. Ocon, Meng Nan Chong, <u>Electrochemical oxidation remediation of real</u> <u>wastewater effluents</u> — <u>A review</u>, Process Safety and Environmental Protection, 1 1 3 (2 0 1 8) 48–67
- Heikki Särkkä, Amit Bhatnagar. Mika Sillanpää, <u>Recent developments of electro-oxidation in water</u> <u>treatment — A review</u>, Journal of Electroanalytical Chemistry, Volume 754, 1 October 2015, Pages 46-56
- Comninellis, Christos, Chen, Guohua (Eds.), Electrochemistry for the Environment, Springer, 2010.
- Carlos Alberto Martínez-Huitle, Manuel A Rodrigo, Onofrio Scialdone (Eds), <u>Electrochemical Water</u> and Wastewater Treatment, 1st Edition, Elsevier, 2018
- Mika Sillanpää, Marina Shestakova, <u>Electrochemical Water Treatment Methods</u>, 1st Ed., Elsevier, 2017
- O'Brien, Thomas F., Bommaraju, Tilak V., Hine, Fumio, <u>Handbook of Chlor-Alkali Technology</u>, Springer, 2005

Videos about Electrocoagulation

https://youtu.be/duj6qDrHIM