# United States Patent [19]

### Feige, Jr.

### [54] METHOD OF PRODUCING A COATED ANODE

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- [52] U.S. Cl. ..... 204/38 R, 204/96, 204/290 F, 204/290 R, 204/280
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- [58] Field of Search...... 204/38 R, 290 R, 290 F, 204/56 R, 96, 83

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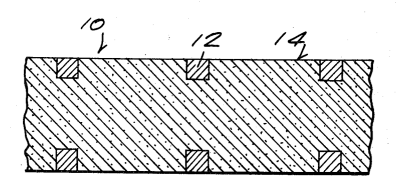
## [11] **3,855,084** [45] **Dec. 17, 1974**

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### [57] ABSTRACT

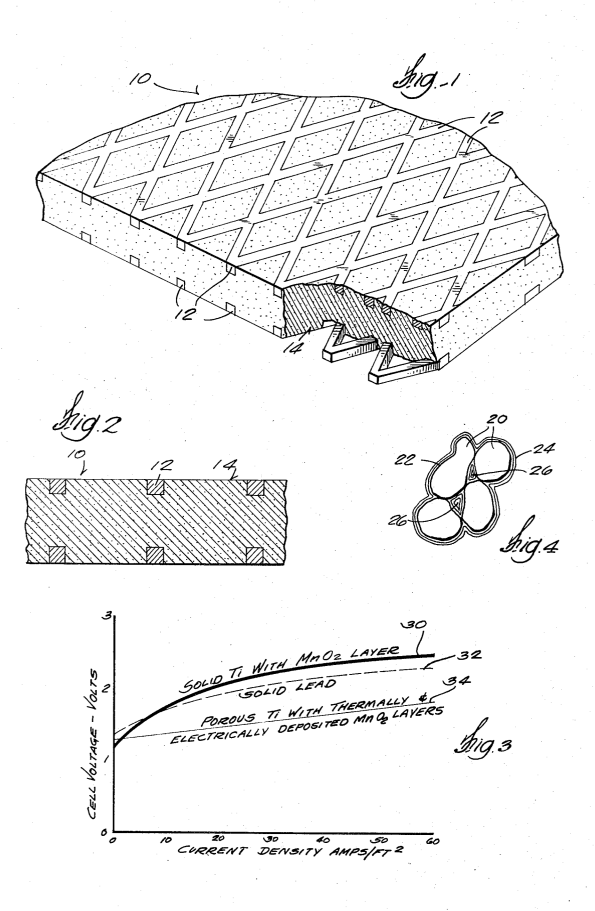
An electrode useful for electrowinning and other processes including evolution of oxygen or chlorine or for plating is formed of titanium particles, compacted cold, and coated and cemented with a first layer of manganese dioxide thermally deposited from Mn (NO<sub>3</sub>)<sub>2</sub> on the grains to form a coating of a combination of manganese dioxide and titanium oxide having a rutile crystal structure, and a further outer layer of manganese dioxide which is electrodeposited, the two coatings not being limited to the surface of the electrode but extending to all exposed surfaces of the grains including those which are walls of channels between the grains. The compression of the titanium powder is to a density between 30 and 70 percent of the density of solid metal. Solid metal such as metal mesh may be included in the titanium powder prior to compacting, such as expanded titanium metal, to assist in strengthening the form. In an alternate embodiment the metal substrate is lead.

### 14 Claims, 4 Drawing Figures



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### METHOD OF PRODUCING A COATED ANODE

### BACKGROUND OF THE INVENTION

It has long been known that titanium metal has superior properties for use as an electrode in cells, baths or 5 solutions which would corrode and be contaminated by metal from electrodes of other compositions. Nevertheless there are difficulties in the use of a titanium electrode as well. Solid titanium leads to a requirement U.S. Pat. Nos. 2,631,115 and 2,608,531 discuss some of the difficulties in connection with specific systems, for instance the passive surface coating on the titanium, and indicate that at least in the production of oxides such as manganese dioxide for use in batteries as a depolarizer, the use of a porous titanium anode made of chips is of assistance. Under certain special conditions described by Fox, the plating of MnO<sub>2</sub> or in a battery, the use of a specific form of titanium chips of about 35 tionships during electrolytic formation of the surface of the electrode prior to use in the other patent, improved results are observed using porous titanium anodes. Fox discloses coating the surfaces of the porous titanium mass with graphite, gold or iron, or with any good con- 25 ductor which is inert to the electrolyte in which the electrode is to be used. Among other things Fox discloses that his electrodes are useful for the preparation of electrolytic manganese dioxide. His described electrode does not function as an oxygen evolution anode, 30 but passivates under those conditions.

In industry it is most important to select a suitable anode that does not contaminate the electrolyte or contaminate the cathode deposit, that has a long life, and has a low oxygen overvoltage during electrolysis. Plati- 35 num is an excellent known anode material which satisfies the above mentioned characteristics.

Recently platinum and other precious metals have been applied to a titanium substrate to retain their attractive electrical characteristics and further reduce manufacturing cost. However such anodes are expensive and are not suitable for some industrial uses. Thus carbon and lead alloy electrodes have generally been used. The carbon anode, however, has the disadvantage that it greatly contaminates the electrolyte, wears fast, and has high electrical resistance which results in an increase in cell voltage. It may also be degraded to CO<sub>2</sub> during oxygen evolution. The disadvantage of the lead alloy anode is that PbO<sub>2</sub> changes to Pb<sub>3</sub>O<sub>4</sub> which is poorly conductive. O<sub>2</sub> gets below this layer and flakes off the film. These particles become trapped in the deposited copper at the cathode, degrading it.

In order to overcome these disadvantages it has recently been proposed to plate the surface of a titanium 55 substrate with platinum and to electrodeposit either lead dioxide or manganese dioxide on the plated surface. Such anodes have the disadvantage of comparatively high oxygen overvoltage. In addition both coatings have high internal stress when electrolytically deposited and are liable to be detached from the surface during commercial usage, both contaminating the electrolyte and, in the case of lead, being deposited on the cathode to reduce its value. Thus current density with such anodes is very limited.

To improve the high oxygen overvoltage it has also been proposed to compact and sinter titanium chips to increase the apparent surface area. Such an anode has somewhat improved characteristics but does not properly receive and retain the electrodeposited manganese dioxide coating.

This invention is based on recognition that deposited manganese dioxide is both insoluble and electrically conductive, and cannot readily be deposited as a reduced product on the cathode. Experiments have shown that my electrode has a low oxygen overvoltage during electrolysis, economizes on electric power necfor high voltage, and gives poor current efficiency. Fox 10 essary for electrolysis, reduces the loss of manganese from the electrode to the bath to a very low figure, and is believed to be the optimum electrode for electrolytic winning of copper, zinc and nickel in sulfate electrolytes. It is also useful for evolving oxygen in sulfate sys-15 tems and chlorine in chloride systems.

### SUMMARY OF THE INVENTION

The preferred electrode of my invention is made of titanium powder cold compacted to the shape of an mesh in one case and the use of specific voltage rela- 20 electrode, the compaction being sufficient to produce a density in the powder between 30 and 70 percent of the density of the solid metal of which the powder is composed. A layer of manganese dioxide is produced on the surface of the grains throughout the mass of the electrode by thermal decomposition of Mn  $(NO_3)_2$ . That layer is modified by ion exchange with the titanium dioxide surface layer and probably with the titanium metal of the powder grains to contain titanium atoms as well as manganese atoms. These do not significantly modify the crystal structure which is essentially that of rutile. This mixed structure is formed while the material of the coating is in the intermediate manganeous-manganite form during the decomposition from manganeous nitrate to manganese dioxide.

> A final coating of manganese dioxide is electrodeposited over the hybrid coating.

> Preferably the structure is strengthened by including solid metal in the electrode. A preferred form is an expanded metal lattice of titanium although other shapes may be used. The preferred size range of the titanium powder is +100 mesh to -325 mesh. The preferred thickness of the outer coating of mananese dioxide is 100 microns. The density referred to is calculated on the weight of the powder alone. The upper limit is the loss of connected pores between the particles.

> The preparation of an electrode in this manner produces a stronger electrode without the necessity of sintering the powder because the initial thermally deposited layer is extremely effective in cementing the grains of the powder. It has been found that even loose titanium powder may be cemented by this method to produce a coherent shape. Powder which has been cold compacted to between 30 and 70 percent of metallic density and then coated as described produces an extremely durable electrode suitable for use as the anode in an electrowinning process. The anode evolves  $O_2$  in sulfates and Cl<sub>2</sub> in chlorides, and accordingly may also be used for the evoluation of chlorine as well.

It has been observed that manganese dioxide coating of prior art electrodes used in electrowinning is lost during periods of shutdown, at which time it goes into solution and becomes a part of the bath, degrading the electrode. The loss is largely in the form of conversion to manganese cations and permanganate. With the 65 anode of my invention, the ions and permanganate are largely limited to the pores or channels between the grains of titanium powder and a very high percentage

is redeposited as manganese dioxide upon application of current for the electrowinning process, in the same manner that the outer layer was originally deposited. Thus degradation of the anode and pollution of the bath are both avoided.

Finally the electrical and operating characteristics of my electrode compare favorably with prior art electrodes.

The chief presently known use of my electrode is as an anode for electrolysis. So used, an advantage of the <sup>10</sup> anode is low oxygen overvoltage during electrolysis, thus economizing the electrical powder necessary.

A similar electrode may be made with lead particles, at lower cost. As in the case of the titanium electrode a first coating of  $MnO_2$  is thermally deposited and forms a hybrid with the lead oxide surface layer normally present. A further layer of  $MnO_2$  is electrodeposited. A reinforcement of solid metal, which may be titanium expanded metal mesh, may be used.

#### DRAWINGS

FIG. 1 is a perspective view of a rectangular electrode made according to my invention with portions broken away to show interior structure.

FIG. 2 is an enlarged cross-sectional view on line 2-2 of FIG. 1.

FIG. 3 is a graph of cell voltage against current density for an anode according to my invention compared with a lead anode and a solid titanium anode coated  $_{30}$  with MnO<sub>2</sub>.

FIG. 4 is an idealized cross-section of several grains showing the two layers and the pores highly magnified.

### DESCRIPTION OF THE PREFERRED EMBODIMENTS

Although the disclosure herein is detailed in order to enable those skilled in the art to practice the invention, the embodiments disclosed merely exemplify the invention which may take other forms. The scope of the invention is defined in the claims.

FIG. 1 is a broken away perspective view showing a portion of an electrode 10 formed according to my invention. An expanded metal mesh 12 of titanium metal 45 has been placed with -100 mesh titanium powder in a mold and compacted in a press until the powder has a density of 30 to 70 percent of theoretical density of the metal. The precise shape is not a part of my invention 50 and may conform to specifications for a particular use, so the outline is not shown. FIG. 2 is a cross-sectional view. As shown in FIG. 4, which is a highly magnified and idealized cross-sectional view through several particles of titanium, each grain 20 is coated with a first 55 layer 22 and a second layer 24, leaving interconnected pores 26 between the grains.

The first layer 22 is basically  $MnO_2$  which has been deposited by thermal breakdown of  $Mn (NO_3)_2$  and which has exchanged ions with the metal and surface layer of titanium oxide normally present on the metal and not separately shown. Thus layer 22 has a modified rutile character. It is strongly adherent and cements the grains 20 and the mesh 12 together.

The second layer 24 is electrodeposited  $MnO_2$ . It is not known to interchange ions with the first layer and is believed therefore to be identical with commercially electrodeposited manganese dioxide.

The pores 26 are the spaces between the coated grains. As more fully described elsewhere, they are highly interconnected. This not only creates high available surface area (as opposed to surface in a closed cavity not connected through pores 26 with the exterior of the electrode 10) thus improving the apparent current density, it also means that a substantial part of the first and second layers 22-24 are in the pores. During inactivity of the immersed electrode  $MnO_2$  can break down to manganese cations and permanganate. In my electrode these are largely trapped in pores 26. Upon reapplication of current to the anode,  $MnO_2$  is re-formed on the second layer with very little loss.

The method of my invention thus comprises the basic 15 steps of compacting metal particles to a density sufficient to make a coherent final product but not so high as to destroy the high interconnection between the pores, thermally breaking down Mn (NO<sub>3</sub>)<sub>2</sub> at a temperature between about 120° C. and 475° C. to form a 20 first layer on the particles or grains, and depositing a second layer of MnO<sub>2</sub> over the first layer, to form an electrode. The metal may preferably be titanium but may also be lead.

The density of the compacted titanium powder used <sup>25</sup> in my electrode has both a lower and upper limits for satisfactory performance. The freely poured titanium powder of a typical sieve analysis as follows:

TABLE I

30	Ti Powder Analysis					
	Typical Sieve An	alysis				
	Size	All powder 1 -100,	ess than -100 mesh. +150 mesh 25 percent			
35		-150, 200,	+200 mesh +270 mesh	25 percent 21 percent 26 percent		
		270, 325 mesh	+325 mesh	17 percent 11 percent		
	Chemistry (%)	Oxygen Nitrogen	0.30 max 0.04 max			
40		Carbon Chlorine	0.04 max 0.20 max			
		Iron Magnesium	0.50 max 0.30 max			
		Total others Titanium	0.12 max 98.50 min			

has a theoretical density of 25 percent, that is, its weight is 25 percent of the weight of an equal volume of solid titanium metal having the same anlaysis. Using great care it is possible to compact such powder to only 30 percent density in the shape of an anode and cement the grains of the powder together by thermal decomposition of Mn  $(NO_3)_2$  as described elsewhere in this specification. Thus 30 percent density is believed to be a practical lower limit of density. With greater compacting force the density of the powder may be increased to about 70 percent while retaining highly interconnected passageways between the grains of metal of sufficient size for the application of the coatings of my electrode. Above 70 percent theoretical density the pores lose interconnection and surface area is lost to the extent that following the process described in this specification does not result in deposition of sufficient MnO<sub>2</sub> either in the layer produced by thermal decomposition or in the electrodeposited layer. Above 70 percent density the oxygen overvoltage rises unduly as shown in table 2 below:

TA	BLE	NO.	2	

% Den.	Anode Thickness	% MnO₂	Back EMF	Cell Voltage	
				at 24 amps/ft <sup>2</sup>	at 120 amps/ft <sup>2</sup>
72%	1.0cm	2%	1.15 volts	1.55	1.95
63%	1.1cm	3%	1.20 volts	1.54	1.90
49%	1.0cm	15%	1.06 volts	1.43	1.87
46%	1.6cm	6%	1.20 volts	1.48	1.84
38%	2.1cm	14%	1.11 volts	1.35	1.69
37%	1.2cm	17%	1.08 volts	1.30	1.71

Another method of defining the upper limit is by blowing air through the compacted mass. Substantial resistance to the passage of air indicates that few pas- 15 sages are interconnected, showing that compaction is too great. The upper limit may vary with particle size and shape but includes only a degree of compaction which leaves the pores highly interconnected.

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In my electrode, the grains of titanium powder are 20 cemented together by the first coating of MnO<sub>2</sub> which is produced by thermal decomposition of Mn (NO<sub>3</sub>)<sub>2</sub> between 120° C. and 475° C. During the thermal decomposition there is an extensive reaction with the surface titanium oxide layer which is normally present on 25 titanium metal by ion exchange, during the phase when the coating is black liquid manganeous-manganite. The result is an adherent coating bonding the grains of titanium powder with a highly conductive oxide film. Subsequently a further layer of MnO<sub>2</sub> is electrolytically de- 30 posited, resulting in an electrode having extensively interconnected passages between the grains, the surface of each grain both on the surface of the electrode and in the passages being coated with a first layer of modi-35 fied rutile containing titanium, and then with MnO<sub>2</sub>.

Electrodeposited manganese dioxide is brittle, and has large internal stresses. It is readily detached from a substrate when deposited to an appropriate thickness for electrode use, making it difficult to form an effective and long lived electrode from such manganese dioxide alone. My invention permits the application of heavy deposits in excess of 100 microns thick of manganese dioxide within the pores between the grains of titanium powder, thus taking advantage of the large internal stresses of the coating to improve its adherence 45 rather than to cause failure as in the case of a flat sheet electrode.

Finally, by depositing much of the coating internally it is protected both from mechanical dislodgement and from loss into the solution when the electrode is inac- 50 tive.

Although under anodic potential as applied during electrolysis, manganese dioxide is insoluble, under an open circuit permanganate is observed in solution. As the potential is again applied to the anode the perman-55 ganate is redeposited on the anode as manganese dioxide. By coating the porous substrate internally the dissolution of manganese dioxide is reduced and the concentration of permanganate is sufficiently high in the pores for redeposition to reduce significantly the loss of manganese from the anode and the pollution of the bath.

### EXAMPLE 1

An example showing the comparison between my electrode and a similar electrode made with solid titanium plate is as follows:

Two sheets of expanded titanium mesh, each 1.5 millimeters thick, were placed in a mold 500 centimeters by 7.50 centimeters by 3 millimeters thick with 82 grams of titanium powder having the sieve analysis and chemical analysis shown in Table 1. These were subjected to a pressure of 5,400 kilograms, resulting in a composite structure in which the powder component had a theoretical density of 52 percent of the density of solid titanium. The electrode was impregnated with an aqueous solution of Mn  $(NO_3)_2$  and was baked at 176° Centigrade. The electrode developed a gray adherent coating. The electrode was then placed in a bath of MnSO<sub>4</sub> and H<sub>2</sub>SO<sub>4</sub> at 90° Centigrade and current was applied to electrolytically deposit a coating of manganese dioxide following accepted techniques. It was found that during this step current density could be varied from 0.1 ampere to 6.7 amperes per square foot, which was the observed oxygen evolution value for the anode. A black layer of MnO<sub>2</sub> developed over the first laver.

In the same manner a titanium plate 500 centimeters by 750 centimeters by 3 millimeters thick having a sandblasted surface was coated with two layers as described above.

The two electrodes thus produced are compared in FIG. 3 along with an electrode of solid lead. The bath was a copper plating bath containing  $CuSO_4$  and  $H_2SO_4$  at 60° Centigrade and a gap of  $\frac{1}{2}$  inch. The anode made from titanium plate was less durable and required higher cell voltage, despite the use of both thermal and electrolytic deposition of  $MnO_2$  in accordance with a portion of my invention. Line 34 is the curve for the porous anode with two layers made according to my invention. Line 32 is the curve for lead. Line 30 is the curve for the solid titanium anode with two  $MnO_2$  layers.

A porous anode as described in this example was used for electrolytic winning of copper, zinc and nickel in the respective commercial sulfate-sulfuric acid electrolytes. The anode performed satisfactorily and exhibited a substantial improvement in cell voltage in the system as compared to lead anodes in each electrolyte.

The porous electrode described has also been tested as an anode for evolution of oxygen, and as an anode for evolution of chlorine, both with good efficiencies and service life. The anode is useful for chlorination of water, for instance.

#### EXAMPLE 2

The effect of changes in the density of the powder component of my electrode may be seen in table 2 below. Anodes prepared according to the first paragraph of Example 1, with the exception that the compaction and amount of  $MnO_2$  varied, are compared.

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TABLE NO. 2

The Relationship Between Theoretical Density And Anode Performance					
% Den.	Anode Thickness	% MnO2	Back EMF	Cell Voltage	
		-		at 24 amps/ft <sup>2</sup>	at 120 amps/ft2
72%	1.0cm	2%	1.15 volts	1.55	1.95
63%	1.1cm	3%	1.20 volts	1.54	1.90
49%	1.0cm	15%	1.06 volts	1.43	1.87
46%	1.6cm	6%	1.20 volts	1.48	1.84
38%	2.1cm	14%	1.11 volts	1.35	1.69
37%	1.2cm	17%	1.08 volts	1.30	1.71

### **EXAMPLE 3**

In a manner like that of Example 1, lead electrodes were prepared using lead shot particles and titanium expanded metal mesh but using lower pressure. In respective trials lead paraticles ranging from no. 6 shot (0.110 in.) to no. 11 shot (0.065 in.) were used. After the two layers of MnO<sub>2</sub> are deposited as in Example 1, the electrode was tested as an anode in a copper sulphate and sulfuric acid conventional electrolyte. The graph of cell voltage vs. amperes/sq. ft. for this electrode was intermediate between such a graph for a solid lead anode and that for the anode of Example 1 for the particle sizes tested. 15 a modifi-7. The ited man thick. 20 8. The compact the densi 9. A m 25 steps of:

It will be understood that the invention is not limited to the examples described and that many modifications may be introduced therein. The scope of the invention is intended to be limited only by the scope of the ap- $_{30}$  pended claims.

I claim:

1. A method of manufacturing an electrode comprising the steps of:

- first, cold compacting metal particles chosen from 35 the group consisting of titanium and lead to form an electrode having an extensively interconnected pore system between the particles;
- second, depositing a first layer of manganese dioxide on the exterior surface of the compact and within the pores between particles by thermal decomposition of a solution of Mn  $(NO_3)_2$ ;

and third, electrodepositing a layer of manganese dioxide on said first layer.

**2.** The method of claim 1 in which the particles are  $_{45}$  titanium.

3. The method of claim 1 in which the particles are lead.

4. The method of claim 2 in which the particles comprise powder predominantly in the size range of +100 50 mesh to -325 mesh.

5. The method of claim 1 in which the particles are compacted along with at least one solid titanium metal reinforcement to increase the mechanical strength of the electrode.

6. The method of claim 2 in which during the second step the forming manganese dioxide coating inter-

changes atoms with the titanium and titanium oxide normally present on the surface of the particles to form a modified rutile crystal structure which is electrically conductive.

7. The method of claim 1 in which the electrodeposited manganese dioxide is approximately 100 microns thick.

8. The method of claim 1 wherein the density of the compacted particles is between 30 and 70 percent of the density of the solid metal of which the particles are composed.

**9.** A method of forming an electrode comprising the steps of:

- first, forming the body of the electrode from particles of titanium;
- second, thermally decomposing manganeous nitrate in contact with the electrode to form a layer of the intermediate product manganeous-manganite;
- third, causing ion exchange between the body of the electrode and the coating to replace manganese ions with titanium ions;

fourth, causing the product to coat the surfaces of the particles to form an adherent coating of a highly conductive film of manganese-titanium oxide;

and fifth, thereafter electrodepositing a layer of manganese dioxide on the surfaces of the coated particles.

10. The method of claim 9 in which the second step cements the grains or particles together in the body of the electrode thereby forming a mechanically sound electrode for use as an anode.

11. The method of claim 9 in which the powder is in the size range of +100 mesh to -325 mesh.

12. The method of claim 9 in which the powder is compacted along with at least one solid titanium metal reinforcement to increase the mechanical strength of the electrode.

13. The method of claim 9 in which during step 2 the manganese dioxide coating is reacted with the titanium oxide normally present on the surface of the particles to form a modified rutile crystal structure which is electrically conductive.

55 14. The method of claim 9 in which the electrodeposited manganese dioxide is approximately 100 microns thick.

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