(12) UK Patent Application (19) GB (11) 2 110 242

- (21) Application No 8134777
- (22) Date of filing 18 Nov 1981
- Application published 15 Jun 1983
- (51) INT CL3
- C25D 3/10 3/06 (52) Domestic classification .C7B 120 431 444 447 450
- 452 458 464 718 DM (56) Documents cited
- - GB A 2071151 GB A 2038361
 - GB A 2033427

 - GB A 2018292
 - US 4184929
 - GB 1596995
 - GB 1562188
 - GB 1551609
 - US 4142948
 - GB 1488381
 - GB 1383524
 - GB 1301478 =
 - US 1922853
 - GB 0965686 =
 - US 3006823
 - Chemical abstract vol 91(1979) abstract No 0184048e
- (58) Field of search
 - **C7B**
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(54) Electroplating chromium

(57) A chromium electroplating electrolyte contains a source of trivalent chromium ions, a complexant, a buffer and a sulphur species having S-O or S-S bonds for promoting chromium deposition, the complexant being selected so that the stability constant K₁ of the chromium complex is in the range $10^6 < K_1 < 10^{12} \, M^{-1}$ and the sulphur species being selected from thiosulphates, thionates, polythionates and sulfoxylates. Complexants specified are aspartic acid, iminodiacetic acid, nitrilotriacetic acid, 5-sulphosalicylic acid and citric acid.

SPECIFICATION

Electrodeposition of chromium and its alloys

5 The invention relates to the electrodeposition of chromium and its alloys from electrolytes containing trivalent chromium ions.

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Background art

Commercially chromium is electroplated from electrolytes containing hexavalent chromium, but many attempts over the last fifty years have been made to develop a commercially acceptable process for electroplating chromium using electrolytes containing trivalent chromium salts. The incentive to use electrolytes containing trivalent chromium salts arises because hexavalent chromium presents serious health and environmental hazards - it is known to cause ulcers and is believed to cause cancer, and, in addition, has technical limitations including the cost of disposing of plating baths and rinse water.

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The problems associated with electroplating chromium from solutions containing trivalent chromium ions are primarily concerned with reactions at both the anode and cathode. Other factors which are important for commercial processes are the material, equipment and operational costs.

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In order to achieve a commercial process, the precipitation of chromium hydroxy species at the cathode surface must be minimised to the extent that there is sufficient supply of dissolved ie. solution - free,

chromium (III) complexes at the plating surface; and the reduction of chromium ions promoted. United Kingdom patent specification 1,431,639 describes a trivalent chromium electroplating process in which the electrolyte comprises aquo chromium (III) thiocyanato complexes. The thiocyanate ligand stabilises the chromium ions inhibiting the formation of precipitated chromium (III) salts at the cathode surface during plating and also promotes the reduction of chromium (III) ions. United Kingdom patent specification

1,591,051 described an electrolyte comprising chromium thiocyanato complexes in which the source of chromium was a cheap and readily available chromium (III) salt such as chromium sulphate.

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Improvements in performance i.e., efficiency or plating rate, plating range and temperature range were achieved by the addition of a complexant which provided one of the ligands for the chromium thiocyanato complex. These complexants, described in United Kingdom patent specification 1,596,995, comprised amino acids such as glycine and aspartic acid, formates, acetates or hypophosphites. The improvement in performance depended on the complexant ligand used. The complexant ligand was effective at the cathode surface to further inhibit the formation of precipitated chromium (III) species. In specification 1,596,995 it was noticed that the improvement in performance permitted a substantial reduction in the concentration of chromium ions in the electrolyte without ceasing to be a commercially viable process. In United Kingdom patent specifications 2,033,427 and 2,038,361 practical electrolytes comprising chromium thiocyanato complexes were described which contained less than 30 mM chromium - the thiocyanate and complexant

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treatment of rinse waters was greatly simplified and, secondly, the colour of the chromium deposit was much lighter.

Oxidation of chromium and other constituents of the electrolyte at the anode are known to progressively and rapidly inhibit plating. Additionally some electrolytes result in anodic evolution of toxic gases. An electroplating bath having an anolyte separated from a catholyte by a perfluorinated cation exchange membrane, described in United Kingdom patent specification 1,602,404, successfully overcomes these problems. Alternatively an additive, which undergoes oxidation at the anode in preference to chromium or

being reduced in proportion. The reduction in chromium concentration had two desirable effects, firstly the

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other constituents, can be made to the electrolyte. A suitable additive is described in United Kingdom patent specification 2,034,354. The disadvantage of using an additive is the ongoing expense.

Japan published patent application 54-87643 describes an electrolyte for electroplating chromium in which oxalic acid, a hypophospite or a formate is suggested as a complexant for stabilising trivalent chromium ions. To improve stability and deposition rate a compound characterised as having a S—O bond in the molecule is added to the electrolyte. The compound is selected from the group consisting of thiosulphates, thionates, sulfoxylates and dithionites. However the concentration of chromium ions and

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Disclosure of the invention

complexant was very high, that is greater than 0.4M.

Three related factors are responsible for many of the problems associated with attempts to plate chromium from trivalent electrolytes. These are, a negative plating potential which results in hydrogen evolution accompanying the plating reaction, slow electrode kinetics and the propensity of chromium (III) to precipitate as hydroxy species in the high pH environment which exists at the electrode surface. The formulation of the plating electrolytes of the present invention described here in are based on an understanding of how these factors could be contained.

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Cr (III) ions can form a number of complexes with ligands, L, characterised by a series of reactions which may be summarised as:

	$Cr + L = CrL$ $CrL + L = CrL_2$	K ₁ K ₂
5	etc.	
	where charges are omitted for conve K ₂ , K ₂ , etc. are the stability const	ience and
10	$\begin{aligned} K_1 &= [CrL]/[Cr][L] \\ K_2 &= [CrL_2]/[CrL][L] \end{aligned}$	10
15	etc.	
20	"Stability Constants of Metal-lon Con 1964 - L.G. Sillén and A.E. Martell; (2 Special Publication No. 25, The Cher Stability Constants" Vol. 1 and 2, Ple	t concentrations. Numerical values may be obtained from (1) plexes", Special Publication No. 17, The Chemical Society, London 'Stability Constants of Metal-Ion Complexes", Supplement No. 1, cal Society, London 1971 - L.G. Sillen and A.E. Martell; (3) "Critical um Press, New York 1975 - R.M. Smith and A.E. Martell;
25	acidity constant, pKa, and concentra higher than the pH in the bulk of the precipitate. The value of K_1, K_2, \dots et ligand determine the extent to which	ce pH can rise to a value determined by the current density and the con of the buffer agent (e.g. boric acid). This pH will be significantly ectrolyte and under these conditions chromium-hydroxy species may and the total concentrations of chromium (III) and the complexant precipitation occurs; the higher the values of K ₁ , K ₂ etc. the less are pH. As plating will occur from solution-free (i.e. non-precipitated)
30	chromium species higher plating eff However, a second consideration the K values are too high plating will complexes. Thus selection of the op- chromium and the ligand, is a comp	iencies may be expected from ligands with high K values. related to the electrode potential adopted during the plating process. If action is inhibited because of the thermodynamic stability of the chromium mum range for the stability constants, and of the concentrations of mise between these two opposing effects: a weak complexant results
35	whereas too strong a complexant in A third consideration is concerned (H.E.R.) and of chromium reduction. kinetics for the H.E.R. Thus additives	low efficiency (or even blocking of plating by hydroxy species), bits plating for reasons of excessive stability. with the electrochemical kinetics of the hydrogen evolution reaction lating will be favoured by fast kinetics for the latter reaction and slow which enhance the chromium reduction process or retard the H.E.R. will blating rates. It has been found that many sulphur containing species
40	having S—S or S—O bonds accelerated. The present invention provides a conformium ions, a complexant, a buf promoting chromium deposition, the	the reduction of chromium (III) to chromium metal. From ium electroplating electrolyte containing a source of trivalent agent and a sulphur species having $S-O$ or $S-S$ bonds for complexant being selected so that the stability constant K_1 of the is in the range $10^6 < K_1 \ 10^{12} \ M^{-1}$ and the sulphur species being
45	selected from thiosulphates, thionat By way of example complexant lig aspartic acid, iminodiacetic acid, nit The sulphur species are provided	s, polythionates and sulfoxylates. At a polythionates and sulfoxylates. At a having K_1 values within the range $10^6 < K_1 < 10^{12} \text{M}^{-1}$ include otriacetic acid, 5-sulphosalicylic acid and citric acid. At a following in the electrolyte: sodium
50	potassium polythionate, sodium	hur species are needed to promote reduction of the trivalent chromium of the electrolyte is relatively high a commercial trivalent chromium nromium. This removes the need for expensive rinse water treatment ag-out' from the plating electrolyte is extremely low.
55	In general the concentration of the Chromium (III) ions	constituents in the electrolyte are as follows: 5
		10 ⁻⁵ to 10 ⁻² M
60	Sulphur species A practical chromium/complexan	6

Above a minimum concentration necessary for acceptable plating ranges, it is unnecessary to increase the amount of the sulphur species in proportion to the concentration of chromium in the electrolyte. Excess of the sulphur species may not be harmful to the plating process but can result in an increased amount of sulphur being co-deposited with the chromium metal. This has two effects, firstly to produce a progressively 65

A practical chromium/complexant ligand ratio is approximately 1:1.

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darker deposit and, secondly, to produce a more ductile deposit.

The preferred source of trivalent chromium is chromium sulphate which can be in the form of a commercially available mixture of chromium and sodium sulphates known as tanning liquor or chrometan. Other trivalent chromium salts, which are more expensive than the sulphate, can be used, and include chromium chloride, carbonate and perchlorate.

The preferred buffer agent used to maintain the pH of the bulk electrolyte comprises boric acid in high concentrations i.e., near saturation. Typical pH range for the electrolyte is in the range 2.5 to 4.5.

The conductivity of the electrolyte should be as high as possible to minimise both voltage and power consumption. Voltage is often critical in practical plating environments since rectifiers are often limited to a low voltage, e.g. 8 volts. In an electrolyte in which chromium sulphate is the source of the trivalent chromium ions a mixture of sodium and potassium sulphate is the optimum. Such a mixture is described in United Kingdom patent specification 2,071,151.

A wetting agent is desirable and a suitable wetting agent is FC98, a product of the 3M Corporation. However other wetting agents such as sulphosuccinates or alcohol sulphates may be used.

It is preferred to use a perfluorinated cation exchange membrane to separate the anode from the plating 15 electrolyte as described in United Kingdom patent specification 1,602,404. A suitable perfluorinated cation exchange membrane is Nafion (Trade mark) a product of the Du Pont Corporation. It is particularly advantageous to employ an anolyte which has sulphate ions when the catholyte uses chromium sulphate as the source of chromium since inexpensive lead or lead alloy anodes can be used. In a sulphate anolyte a thin 20 conducting layer of lead oxide is formed on the anode. Chloride salts in the catholyte should be avoided 20 since the chloride anions are small enough to pass through the membrane in sufficient amount to cause both the evolution of chlorine at the anode and the formation of a highly resistive film of lead chloride on lead or lead alloy anodes. Cation exchange membranes have the additional advantage in sulphate electrolytes that the pH of the catholyte can be stabilised by adjusting the pH of the anolyte to allow hydrogen ion transport 25 through the membrane to compensate for the increase in pH of the catholyte by hydrogen evolution at the 25 cathode. Using the combination of a membrane, and sulphate based anolyte and catholyte a plating bath has been operated for over 40 Amphours/litre without pH adjustment.

Detailed description

The invention will now be described with reference to detailed Examples. In each Example a bath consisting of anolyte separated from a catholyte by a Nafion cation exchange membrane is used. The anolyte comprises an aqueous solution of sulphuric acid in 2% by volume concentration (pH 1.6). The anode is a flat bar of a lead alloy of the type conventionally used in hexavalent chromium plating processes.

The catholyte for each Example was prepared by making up a base electrolyte and adding appropriate amounts of chromium (III), complexant and the sulphur species.

The base electrolyte consisted of the following constituents dissolved in 1 litre of water:

Potassium sulphate 1 M
Sodium sulphate 0.5 M
40 Boric acid 1 M
Wetting agent FC98 0.1 gram

Example :

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The following constituents were dissolved in the base electrolyte:

Chromium (III) 10 mM (from chrometan)
DL aspartic acid 10 mM
Sodium thiosulphate 1 mM

at pH 3.5

Although equilibrium will occur quickly in normal use, initially the electrolyte is preferably equilibrated ntil there are no spectroscopic changes which can be detected. The bath was found to operate over a

until there are no spectroscopic changes which can be detected. The bath was found to operate over a temperature range of 25 to 60°C. Good bright deposits of chromium were obtained over a current density range of 10 to 800 mA/cm².

Example 2

The following constituents were dissolved in the base electrolyte:

60	Chromium (III) Iminodiacetic acid Sodium thionate	10 mM (from chrometan) 10 mM 1 mM 3.5	60
	at pH	3.3	

The electrolyte is preferably equilibrated until there are no spectroscopic changes. The bath was found to operate over a temperature range of 25 to 60°C. Good bright deposits of chromium were obtained.

Example 3

The following constituents were dissolved in the base electrolyte:

Chromium (III) 100 mM (from chrometan)

DL Aspartic acid 100 mM

Sodium thiosulphate 1 mM

at pH 3.5

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The electrolyte is preferably equilibrated until there are no spectroscopic changes. The bath was found to operate over a temperature range of 25 to 60°C. Good bright deposits were obtained.

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Example 4

The following constituents were dissolved in the base electrolyte:

15 Chromium (III)
DL Aspartic acid

100 mM (from chrometan)

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DL Aspartic acid Sodium thionate at pH 100 mM 1 mM 3.5

The electrolyte is preferably equilibrated until there are no spectroscopic changes. The bath was found to operate over a temperature range of 25 to 60°C. Good bright deposits were obtained over a current density range of 10 to 800 mA/cm².

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CLAIMS

A chromium electroplating electrolyte containing a source of trivalent chromium ions, a complexant, a buffer agent and a sulphur species having S-O or S-S bonds for promoting chromium deposition, the complexant being selected so that the stability constant K₁ of the chromium complex as defined herein is an the range 10⁶ < K₁ < 10₁₂ M⁻¹ and the sulphur species being selected from thiosulphates, thionates, polythionates and sulfoxylates.

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- 2. An electrolyte as claimed in claim 1, in which the complexant is selected from aspartic acid, iminodiacetic acid, nitrolotriacetic acid 5-sulphosalicylic acid or citric acid.
 - 3. An electrolyte as claimed in claims 1 or 2, in which the buffer agent is boric acid.
- 4. An electrolyte as claimed in any one of the preceding claims, in which the source of chromium is chromium sulphate and including conductivity ions selected from sulphate salts.

35 Id potassium

- 5. An electrolyte as claimed in claim 4, in which the sulphate salts are a mixture of sodium and potassium sulphate.
- 6. A bath for electroplating chromium comprising an anolyte separated from a catholyte by a
 perfluorinated cation exchange membrane, the catholyte consisting of the electrolyte claimed in any one of
 the preceding claims.

7. A bath as claimed in claim 6, in which the analyte comprises sulphate ions.

- 8. A bath as claimed in claim 6 or 7 including a lead or lead alloy anode immersed therein.
- 9. A process for electroplating chromium or a chromium alloy comprising passing an electric current between an anode and a cathode in the electrolyte claimed in any one of claims 1 to 5 or in a bath as claimed in claims 6, 7 or 8.

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