

ELECTROREDUCTION PROCESSES INVOLVING TITANIUM AND BORON SPECIES IN HALIDE MELTS

Victor MALYSHEV,^{a, b} Angelina GAB,^{a, b} Diana-Maria BRUSKOVA,^a Igor ASTRELIN,^a
Ana-Maria POPESCU^{c*} and Virgil CONSTANTIN^c

^aNational Technical University of Ukraine “Kyiv Polytechnical Institute”, Faculty of Chemistry & Technology,
37 Peremoga Ave., Kyiv 05056, Ukraine

^bOpen International University of Human Development “Ukraine”

1g Khoriva St., Kyiv 04071, Ukraine, Email: victor_malyshev@mail.ru, lina_gab@mail.ru

^cRoumanian Academy, “Ilie Murgulescu” Institute of Physical Chemistry, Laboratory of Molten Salts,
Splaiul Independentei 202, Bucharest-006021, Roumania

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The current state of research on the titanium and boron electrochemistry as well as high-temperature electrochemical synthesis of titanium diboride and titanium intermetallic compounds with the iron triad metals is analyzed. The prospects for the development of these studies are discussed. Primary attention is paid to the electrochemical properties and stability of halide systems containing titanium in various oxidation states together with boron species.

INTRODUCTION

Studies on the electrochemical properties of titanium and boron and the processes of joint electroreduction of two or more components from ionic melts present interest for both theoretical and practical purposes. On the one hand, the results of these studies help one to understand characteristic features of multi-electron processes and to follow the development of high-temperature electrochemical synthesis; on the other hand, they underlie the development of new technologies for the production of nontraditional materials with unique properties.¹⁻³ Despite the numerous publications devoted to the electrochemistry of titanium and boron as well as titanium diboride, the data on species existence and mechanisms of joint and separate electroreduction in ionic melts (especially chloride and fluoride melts) are quite scarce and contradictory. These processes serve as the basis for the technology of the electrochemical production of titanium, boron, titanium diboride and titanium intermetallic compounds; therefore, studies in this field are topical for the development of modern high-temperature electrochemistry.

ELECTROCHEMICAL BEHAVIOUR OF TITANIUM SPECIES

Electrolysis of titanium melts is a promising method for the production of metallic titanium and synthesis of high-melting compounds (carbides, borides, silicides, intermetallic compounds). However, the complex pattern of the behaviour of titanium-containing melts accounts for the stringent requirements in the process engineering and selection of the salt systems. Thus, it is necessary to study systematically and comprehensively the electrochemical and chemical properties of titanium in various oxidation states in molten electrolytes.

1. Electrolysis of titanium-containing melts

Titanium metal is mainly produced either by electrolysis of the melts of titanium oxygen compounds, fluorides or chlorides or by electrolytic refinement of crude titanium and its compounds.

Electrolysis of oxygen compounds of titanium has several advantages over electrolysis of other compounds such as ready availability and low cost of

* Corresponding author: popescuamj@yahoo.com

the raw materials, namely, titanium dioxide and alkali metal titanates. These advantages also include simplicity of the equipment and environmental safety of the process. Numerous publications have been devoted to TiO_2 electrolysis in melts of alkali metal chlorides or fluorides or potassium (sodium) hexafluorotitanate. The low solubility of titanium dioxide in various melts (0.02%-0.07%) precludes its use for the production of titanium. In addition, during electrolysis of the $\text{NaCl} - \text{K}_2\text{TiF}_6$ melt containing titanium dioxide, the latter is reduced at the cathode to Ti_2O_3 . The process is accompanied by the formation of a non-decomposing precipitate near the cathode. Electrolytic preparation of pure titanium in this system is practically impossible because of the formation of titanium monoxide at the cathode.

Study of a similar system based on calcium chloride showed that in this case, titanium metal is formed in a secondary process, namely, reduction of TiO_2 at the cathode by calcium metal produced in electrochemical processes. Normal operation of the electrolytic cell requires a low concentration of TiO_2 in the electrolyte, its continuous feeding to the cell and replenishment of the electrolyte after its saturation with CaO . Electrolysis of TiO_2 dissolved in molten alkali metal borates or phosphates has been also considered.^{4,5} However, researchers point out to the necessity of additional purification from oxygen of the resulting metal.

Thus, the use of oxygen compounds for the electrolytic preparation of titanium and synthesis of its derivatives is hardly promising, because lower titanium oxides, which contaminate the product, are formed at the cathode.

Both calculations and experimental results show that electrolysis in molten salts of oxygen-containing acids, alkalis or oxides does not ensure the preparation of pure titanium. The range of molten media is mainly limited to metal halides. However, these compounds are not all suitable for electrolysis. Indeed, metals of Group IV and of subsequent Groups of the Periodic Table are usually much more electropositive than titanium and hence, halides of these metals are, mostly often, thermally unstable or possess low electrical conductivity. Alkali and alkaline earth metal chlorides and fluorides are best suited for this purpose.

Numerous studies have been devoted to the development of a method for the production of titanium by electrolysis of fluoride melts. Potassium and sodium fluorotitanates are recognized as the most promising titanium raw materials.⁵⁻⁹ However

it is not expedient to subject pure hexafluorotitanate to electrolysis because during electrolysis it is reduced to K_2TiF_5 and, after that, the electrodes become passivated.

Usually, the system used for electrolysis is K_2TiF_6 dissolved in molten salts. An electrolyte consisting of potassium, lithium and sodium fluorides has been proposed as the medium for the electrolytic production of titanium.⁶ However, a process carried out in a purely fluoride melt can result in passivation of the anode by a gas film (anodic effect). In addition, since this melt is highly corrosive, the equipment soon goes out of service. Therefore, a mixture of alkali metal chlorides and potassium fluoro-titanate has been proposed as electrolyte.⁵⁻⁹ Since titanium metal reacts with a chloride-fluoride melt, a black crust insoluble is formed gradually on melt surface. The existence of a fluoride complex of divalent titanium, $\text{M}_2[\text{TiF}_4]$, has been reported.⁸ Insoluble compounds formed at the cathode contaminate the cathodic product, this process being a disadvantage of using chloride-fluoride-fluorotitanate melts for producing pure titanium.

Preparation of titanium from titanium chlorides presents substantial interest. Titanium, having a variable valence, forms several chloride species, namely, TiCl_4 , TiCl_3 and TiCl_2 . The most readily available compound is TiCl_4 , the production and purification of this compound on an industrial scale being now a routine practice. In the electrolysis of alkali or alkaline earth metal chlorides, TiCl_4 is either supplied onto the electrolyte surface or bubbled within electrolyte by flowing of an inert gas (see, for example, Refs. 10, 11). Titanium tetrachloride is poorly soluble in molten alkali metal chlorides (the solubility is up to 2% at 1073 K); therefore, this technology does not provide high output. If the current density at the cathode is high, the alkali metal will be produced together with titanium. It has been proposed to make use of this phenomenon, *i.e.* to carry out electrolysis at current densities where titanium tetrachloride is partially reduced by the reducing metal.

The solubility of titanium chlorides in which the metal oxidation number is less than four is substantially higher than the solubility of titanium tetrachloride. This fact has been taken into account in the development of a method for preparation of titanium by electrolysis in chloride melts.¹⁰ Salts used in this process should be relatively pure. Lower valence titanium chlorides can be easily prepared from titanium tetrachloride; they are

readily soluble in molten salts and fairly stable at temperatures of electrolysis. Also, these compounds possess high electrical conductivities and relatively low volatilities. However, the lower valence titanium chlorides react with almost any standard refractory lining material with a reaction rate that depends substantially on the presence of oxygen in the cell atmosphere.

A method has been proposed¹⁰ for continuous production of titanium by electrolysis of TiCl_3 and TiCl_2 , these compounds being formed in a separate reactor upon the reaction of titanium tetrachloride with hydrogen. To simplify this method, the synthesis of lower valence titanium chlorides and electrolysis can be carried out in a single vessel. In this case, the electrolysis occurs in two stages, namely, reduction of TiCl_4 to TiCl_2 and TiCl_3 at an auxiliary cathode and reduction of lower titanium chlorides to the metal on the main cathode.¹¹

It has been considered until recently that the cathodic process is a secondary reaction, consisting in the reduction of titanium by the alkali metal produced at cathode. However, recent studies demonstrated that, under certain conditions, direct electrochemical discharge of the titanium ions occurs at the cathode. For this discharge to be possible, it is required, first of all, that the concentration of titanium ions in low oxidation states in the near-cathode layer is sufficiently high.

The cathode materials used in these processes are mainly tungsten, molybdenum, silver, steel or nickel. The best adhesion of the titanium deposits has been noticed for silver cathodes; however, this metal is expensive. Liquid cathodes made of zinc or lead can hardly find wide application because they form intermetallic compounds with titanium, which are difficult to be decomposed.

Electrolysis of titanium chlorides is carried out under the atmosphere of an inert gas (argon, helium).

For rough purification of the electrolyte from moisture and occluded gases, the electrolytic cell is usually evacuated at 400-650°C to a residual pressure of 10-20 mm Hg. Preliminary electrolysis at an auxiliary cathode with a voltage on the cell terminals equal to 1.8-2.0 V is also employed for this purpose. There are also other methods for purification of the electrolyte. To remove moisture, dry hydrochloride is passed through the melt for a short period of time. Oxygen can be removed by passing tetrachloromethane through the melt.

During electrolysis of titanium chlorides, chlorine formed at the anode oxidises lower titanium chlorides present in the anode compartment

to give TiCl_4 , which is distilled off together with chlorine. In addition, electrolytic oxidation of TiCl_2 and TiCl_3 at the anode, leading to the same consequences, is possible. Moreover, titanium tetrachloride entering the cathode compartment would react with titanium metal, which would result in the loss of the pure metal. Therefore, the cathode and anode compartments should be separated by a diaphragm. The problem of selection of the diaphragm materials has not been solved yet due to the high chemical reactivity of lower titanium chlorides; research along this line is now in progress. For instance, it has been proposed to manufacture the diaphragm from alumina (these diaphragms are not virtually metallised under certain conditions) or graphite (in this case, half-sintered metal is formed at the cathode at 1305°C).^{4,5} It should also be noted that working with titanium chloride requires a high voltage at the electrolytic cell terminals (usually 9-12 V) leading to a high power consumption per unit product.

Whereas the selection of one or another method for electrolytic production of titanium is still debated and has not been adopted on an industrial scale, the efficiency of electrolytic refining of titanium is beyond doubt. The soluble anode can be manufactured from any material possessing electronic conduction, capable of reacting with the electrolyte and insoluble in it.

Titanium powders with various chemical and grain-size compositions are currently produced by electrolysis with a soluble anode in melts of alkali and/or alkaline earth metal chlorides. Electrochemical refinement of titanium is carried out using various electrolytes, namely: NaCl , NaCl-KCl , NaCl-SrCl_2 , NaCl-BaCl_2 , KCl-LiCl and KCl-NaCl-MgCl_2 . Electrolytes based on sodium chloride, an equimolar mixture of sodium and potassium chlorides and the NaCl-KCl-MgCl_2 ternary system are the mostly widespread. Electrochemical investigations show that in the temperature range 750-850°C, which is the best range for the refinement, titanium passes from the anode to the melt as Ti^{2+} ions. An increase in the temperature or in anodic current density increases the amount of the Ti^{3+} ions. Upon anodic dissolution, titanium passes into the melt mostly in an average oxidation state of 2.02-2.13.

The total content of impurities which are to be eliminated during titanium refinement (oxygen, nitrogen, hydrogen, carbon, iron) is normally relatively low, of 1-2 wt %. In the studies dealing with electrochemical refinement of titanium, a

special role belongs to the purification from oxygen. When the process is carried out under adequate conditions, the only way by which oxygen can pass from the anode to the cathode is the formation of the titanyl ion TiO^{2+} . Thermodynamic calculations indicate that the probability of formation of this ion is fairly low even at high concentrations of oxygen in the anode material. A significant role in the production of a high-quality powder at the cathode is played by the ratio of the titanium species concentration in the electrolyte to the current density at the cathode. The best powders are obtained if this ratio ranges from 2 to 3. The anodic current density during electrochemical refinement should be 4-8 times lower than the cathodic current density. In this case, divalent titanium disproportionates in the electrolyte (especially in the near-anode layer), resulting in the formation of highly dispersed metallic titanium in the bulk of the melt. The use of the refining tank for the synthesis of titanium borides is not expedient due to the formation of an uncontrollable amount of dispersed titanium, which disrupts the stoichiometry of the product.

Various opinions concerning the role of fluoride ions have been expressed in studies dealing with the electrolytic refinement of titanium in mixed chloride-fluoride electrolytes (see, for example, Ref. 12).

Some researchers believe that the addition of fluorides has a negative effect on the electrolysis characteristics (the anode material disintegrates, whereas the current yield decreases). Conversely, some other investigators believe that fluoride additives are favourable for the performance characteristics of the process. It has to be noted that anodic dissolution of titanium in a mixed chloride and fluoride electrolyte affords poorly soluble compounds of divalent titanium which contaminate the cathodic product.

The foregoing demonstrates that halide melts are the most suitable systems for high-temperature electrochemical synthesis of titanium compounds and production of the pure metal. However, when using chloride melts, one should create conditions for increasing the solubility of TiCl_4 and suppressing disproportionation. When mixed chloride-fluoride electrolytes are used, the formation of poorly soluble divalent titanium fluoride should be prevented and electroreduction of titanium complex ions be performed in a single stage.

2. State of titanium species in halide melts

a. Thermodynamics of titanium-containing halide systems

Titanium halides react with titanium metal in molten alkali metal halides, the following heterogeneous equilibrium being established:



As a rule, one oxidation state of titanium predominates in the melt and the ratio of various oxidation states depends on the temperature and nature of the salt acting as the solvent. The heterogeneous equilibrium (1) can be characterized by several parameters such as equilibrium constant, apparent standard potentials $E^*(\text{Ti}^{2+}/\text{Ti})$, $E^*(\text{Ti}^{3+}/\text{Ti})$ and formal redox potential $E^*(\text{Ti}^{3+}/\text{Ti}^{2+})$.^{13,14} The equilibrium potential of the titanium electrode (E) would be simultaneously the potential of the Ti/TiCl_2 , Ti/TiCl_3 and $\text{TiCl}_2/\text{TiCl}_3$ systems. The apparent standard potentials and formal redox potential are related (in accordance with the Luter rule) by the equation.¹³

$$3E^*(\text{Ti}^{3+}/\text{Ti}) = E^*(\text{Ti}^{3+}/\text{Ti}^{2+}) + 2E^*(\text{Ti}^{2+}/\text{Ti}) \quad (2)$$

Thus, the task of finding these potential values reduces to a determination of the number of ionic forms of titanium in the potential measurement.

The equilibrium potentials of titanium in an equimolar mixture of potassium and sodium chlorides in the temperature range 973-1148 K have been reported.¹⁴ The temperature variations of $E^*(\text{Ti}^{2+}/\text{Ti})$ and $E^*(\text{Ti}^{3+}/\text{Ti})$ are described by the following empirical equations:

$$E^*(\text{Ti}^{2+}/\text{Ti}) = -2.38 + 4.8 \times 10^{-4} T, \quad (3)$$

$$E^*(\text{Ti}^{3+}/\text{Ti}) = -2.16 + 3.2 \times 10^{-4} T. \quad (4)$$

The values of equilibrium potentials of titanium measured at 843 K vs. chlorine reference electrode are $E^*(\text{Ti}^{2+}/\text{Ti}) = -1.97$ V and $E^*(\text{Ti}^{3+}/\text{Ti}) = -1.90$ V.

In order to identify the full picture of the equilibrium between metallic titanium and a chloride melt containing minor amounts of Ti^{4+} in addition to Ti^{2+} and Ti^{3+} , the $E^*(\text{Ti}^{4+}/\text{Ti}^{3+})$ redox potential, and hence, standard $E^*(\text{Ti}^{4+}/\text{Ti})$ potential, should be determined. The temperature variation of this potential in the KCl-NaCl melt is described by the equation

$$E^*(\text{Ti}^{4+}/\text{Ti}) = -1.91 + 3.84 \times 10^{-4} T \quad (5)$$

The following values for the arbitrary formal redox potential $E^*(\text{Ti}^{3+}/\text{Ti}^{2+})$ in a molten equimolar

mixture of sodium and potassium chlorides, determined experimentally by potentiometric titration, have been reported in the literature: -1.807 (T=973K), -1.762 (T=943K), -1.726 V (T=984-1130 K). The $E^*(\text{Ti}^{3+}/\text{Ti}^{2+})$ value calculated in accordance with the Luter rule from Eqn. (2) using known experimental $E^*(\text{Ti}^{3+}/\text{Ti})$ and $E^*(\text{Ti}^{2+}/\text{Ti})$ values is described by the equation:

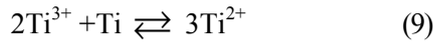
$$E^*(\text{Ti}^{3+}/\text{Ti}^{2+}) = -1.710 - 0.17 \times 10^{-4} T \quad (6)$$

$$E^*(\text{Ti}^{4+}/\text{Ti}^{3+}) = (-1.473 + 5.86 \times 10^{-4} T) \pm 0.007 \quad (7)$$

The variation of the equilibrium potential of titanium as a function of the concentration of titanium ions in the melt has shown that in alkali metal chloride melts titanium content is

$$\log K = \log \frac{[\text{Ti}^{2+}]^3}{[\text{Ti}^{3+}]^2} = \frac{6 \times 10^4}{1.984 T} [E^*(\text{Ti}^{3+}/\text{Ti}) - E^*(\text{Ti}^{2+}/\text{Ti})] \quad (8)$$

Thus, the equilibrium constants for the reaction



can be calculated in various melts. For lithium chloride,

$$\log K = -4.234 + \frac{6351}{T} \quad (10)$$

for potassium chloride,

$$\log K = -6.956 + \frac{8165}{T} \quad (11)$$

for cesium chloride,

$$\log K = -6.956 + \frac{7440}{T} \quad (12)$$

and for the KCl-NaCl melt,

$$\log K = -5.038 + \frac{6774}{T} \quad (13)$$

As the temperature increases, the equilibrium constant decreases, a fact which points out the disproportionation of Ti^{2+} ions. If the temperature remains constant, the equilibrium constant decreases with an increase in the radius (r) of the alkali metal cation in the molten salt, *i.e.* the equilibrium shifts towards disproportionation of Ti^{2+} species. This dependence has the form

$$\log K = \frac{8468}{T} - \frac{1512}{Tr} - 9.073 - \frac{3.175}{r} \quad (14)$$

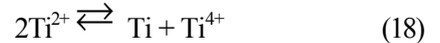
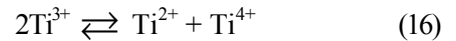
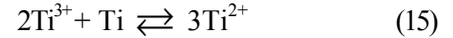
A substantial number of studies have been devoted to the process of disproportionation (see,

It can be seen that the theoretical value of the potential $E^*(\text{Ti}^{3+}/\text{Ti}^{2+})$ depends slightly on the temperature: at 1000 K it is equal to -0.727 V vs. chlorine reference electrode.

The known standard potentials for the Ti^{3+}/Ti and Ti^{4+}/Ti systems were used to calculate the $\text{Ti}^{4+}/\text{Ti}^{3+}$ redox potential leading to its temperature dependence, as:

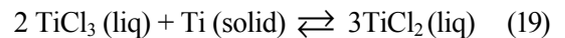
equilibrated with the Ti^{2+} and Ti^{3+} ions. Knowing the standard electrode potentials of the Ti^{2+}/Ti and Ti^{3+}/Ti couples, it can be written.¹⁵

for example, Refs 16-18). The oxidation states of titanium can be described by four disproportionation reactions



These reactions occur simultaneously and are heterogeneous. The outcome depends on the experimental conditions, namely, temperature, composition of the atmosphere, surface area and so on. It has also been noted that Ti^{3+} ions rapidly disproportionate in the lithium chloride melt. When lithium chloride is replaced by cesium chloride, the stability of Ti^{3+} increases. Usually, in halide melts, titanium metal reacts with the components of the melt. When the radius of the alkali metal cation in the melt is greater than the radius of potassium, Ti^{2+} ions appear and they disproportionate to give highly dispersed titanium metal and therefore, a high concentration of Ti^{2+} species cannot be attained. Correspondingly, in the case where the LiCl melt is used, the solution becomes rich in Ti^{2+} ions.

The main reaction which proceeds at a substantial rate in molten potassium chloride is



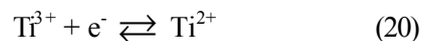
Thus, vast experimental data have been accumulated and a number of important practical

conclusions have been drawn concerning the thermodynamics of titanium-containing chloride melts. The equilibrium and standard potentials of titanium ions in various oxidation states were determined, equilibrium constants for chemical reactions in the melts calculated and disproportionation of titanium ions in low oxidation states was detected. However, in some cases, the thermodynamic functions can be roughly estimated. For example, the measurement of the equilibrium potentials of the Ti^{4+}/Ti , Ti^{3+}/Ti and Ti^{2+}/Ti systems did not appear quite correct because actually, the near-electrode layer would contain titanium ions in various oxidation states formed in chemical reactions. In addition, thermodynamic characteristics do not provide exhaustive information on the mechanism of electro-reduction of titanium, which needs to be known for performing successful electrolysis of titanium-containing melts. Therefore, in addition to the thermodynamic characteristics, the mechanism of the electroreduction of titanium has also been studied.

b. Mechanism of the electroreduction of titanium species

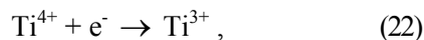
It is not always possible to interpret unambiguously experimental data characterizing the mechanism and kinetics of electroreduction and electrooxidation of titanium complexes. The complexity of interpretation of these data is due to the fact that side processes, not related directly to electrode reactions, proceed in the melt. In some studies, it is suggested that titanium metal is obtained in the reaction of titanium-containing complex ions with an alkali metal resulting from reduction at the cathode. In other studies, two simultaneous processes of titanium deposition were suggested: titanium is formed in the primary electrochemical process at the cathode, $Ti^{2+} + 2e^- \rightarrow Ti^0$, and in a secondary chemical process, its thermal reduction by sodium occurs in the bulk of the electrolyte. Thus, some researchers believe that, in the case of chloride melts, an alkali metal reduces Ti^{2+} ions (see, for example, Refs 19, 20) whereas in the chloride-fluoride melts, the species to be reduced is Ti^{3+} . Titanium in higher oxidation states is preliminarily reduced to lower oxidation states by either electrochemical or chemical reactions. However, voltammograms characterizing the cathodic formation of titanium display a wave corresponding to the discharge of titanium complexes.¹⁸ In several studies (see, for example, Ref. 21), the mechanism of direct reduction of

$TiCl_3$ to the metal without the intermediate formation of Ti^{2+} has been assumed. A study by linear sweep voltammetry of the electroreduction of $TiCl_3$ in the LiCl, CsCl and KCl-LiCl melts at 973K demonstrated that the reduction of Ti^{3+} ions in all three electrolytes involves a couple of reactions:



The stepwise character of the process was pointed out in several other publications.^{18, 22-26}

Electroreduction of titanium tetrachloride in the $BaCl_2$ - $CaCl_2$ - $NaCl$,²² $LiCl$ - KCl , KCl - $LiCl$ - $NaCl$ ²³ and $BaCl_2$ - KCl - $LiCl$ ²⁴ melts occurs as three consecutive stages:



The voltammograms for the electroreduction of Ti^{2+} in the KCl - $NaCl$ - $TiCl$ - NaF - Ti (solid) melt exhibit one clear-cut wave at potentials of $-(2.53 \div 2.6)$ V vs. chlorine reference electrode, which corresponds to the electroreduction of titanium.²⁰

The diffusion coefficients for titanium in chloride melts were calculated^{21,26,27} in KCl - $NaCl$ at 973K and values of 3.72×10^{-5} and $1.94 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$ for Ti^{3+} and Ti^{4+} , respectively, were reported. The diffusion coefficient for divalent titanium in the $LiCl$ - KCl melt at 773K has a value of $(1.54 \pm 0.17) \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$ (see Ref. 28).

Thus, certain progress in the study of chloride and chloride-fluoride titanium-containing melts has been achieved by now; however, the overall picture of the electrochemical and chemical behaviour of titanium in various oxidation states does not appear clear and unambiguous. There is no consensus of opinions on the mechanism of the cathodic process. Several aspects of the stability of systems containing titanium in various oxidation states are poorly documented, namely, which of the lower titanium chlorides is more stable and what is the limiting concentration up to which this stability is retained what are the concentration limits for the stability of other titanium-containing species and also what are the ratios of Ti^{2+} to Ti^{3+} in the melt at which the system remains stable, *i.e.* no disproportionation reactions occur, and what are the ways of suppressing these reactions. Answers to these questions presented considerable interest

from the practical viewpoint, because they would make it possible to create conditions for long continuous operation of an electrolysis cell and estimate the influence of unstable titanium compounds on the technological parameters during production of titanium metal and electrochemical synthesis of titanium compounds.

ELECTROCHEMICAL BEHAVIOUR OF BORON SPECIES

The electrochemical method permits preparation of high-purity boron in large amounts. In the first studies carried out 70 years ago, elementary boron was produced by the electrolysis of borides. The product contained not more than 60%-70% of the target material, whereas 20%-30% was oxygen and 5%-6% was the metal that served as cathode material. When magnesium boride was used, elementary boron was formed at the cathode together with low-stable magnesium borides; the latter can be partially removed by treatment of the cathodic product with an acid.

Currently, oxygen-containing compounds of boron and boron halides are mainly used for the electrolytic production of boron. Numerous studies have been devoted to the electrolysis of oxygen compounds of boron. In all systems, boron oxide serves as the source of boron.²⁹ When oxygen-containing melts are used, the process does not give high-purity boron because the product is contaminated by substantial amounts of oxygen. This is the major drawback of this method.

The use of purely fluoride electrolytes is limited because fluoride melts are corrosive and toxic.³⁰ The use of mixed chloride-fluoride electrolytes presents considerable interest; in this case, high-purity boron free from oxygen impurities can be obtained at relatively low temperatures, the corrosiveness of the melts being relatively low.³⁰⁻³⁶

Several studies have been devoted to the mechanism of electroreduction of boron.³⁷⁻⁴⁶ The cathodic processes occurring during the electrodeposition of boron from purely fluoride (LiF-KF-KBF₄) and fluoride-oxide (LiF-KF-B₂O₃) electrolytes have been studied.^{36,37} It was shown that in a purely fluoride melt, boron is reduced directly from the tetrafluoroborate complex anion in one step involving three electrons. In the LiF-KF-B₂O₃ system, boron oxide reacts with fluorine present in the electrolyte to give the [BF₄]⁻ anion and oxyfluoride complexes, mainly those with a molar ratio of oxygen to boron equal to 1.66. The

ratio of the concentrations of [BF₄]⁻ anion and oxyfluoride complexes is 0.1. The potentials of boron deposition for these two electrochemically active species differ by 0.47 V, and the value of diffusion coefficient for [BF₄]⁻ calculated from experimental data is $4.4 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$ at 973 K.

In a study of the electrochemical properties of boron in chloride and chloride-fluoride melts, the solubility of BCl₃ in the CsCl melt was found to be $1.74 \times 10^{-1} \text{ mol } \%$ at 973 K. Thus, the thermodynamic functions in BCl₃ containing CsCl melts were calculated and the expression of equilibrium potentials of boron reported.³² It was shown that equilibrium potentials of boron shift to the negative region in chloride-fluoride electrolytes, indicating that the reactivity of boron ions in fluoride complexes is lower than in chloride complexes.^{31,32,34} The calculation of the average stoichiometric number of fluoride ions (three or four) in the complex ions by the procedure proposed by Ivanovsky *et al.*³¹ attests indirectly that, in the temperature range studied, most of the boron ions in the chloride-fluoride melts occur as [BF₄]⁻ and [BF₃Cl]⁻ species. Based on the results of voltammetric studies, Chemezov³² concluded that electroreduction of boron in a chloride-fluoride electrolyte proceeds by a one-stage mechanism; conversely, Tsiklauri *et al.*⁴¹ have supposed a redox stage, B³⁺ → B⁺, followed by the discharge of B⁺ at the cathode to give elementary boron.

At low current densities, the charge transfer is the rate-determining step during electrodeposition of boron, whereas for high current densities, the rate of the process is limited by the diffusion of boron from the melt bulk towards the surface. This is typical for both chloride and chloride-fluoride melts. A somewhat different opinion on the kinetics of boron electroreduction is held by other authors³⁹ who claim that the rate of this process is limited by mass transfer over a broad range of polarization rates and that the proper electrochemical stage is irreversible.

Examination of the switching-on curves (potential-time) for various polarization current densities demonstrated³² that neither a homogeneous chemical reaction (*e.g.*, dissociation of the [BF₄]⁻ or [BCl₄]⁻ complexes), nor chemisorption of electrically active species on the electrode surface retards the reduction of boron in

the studied melts. The $\left[\log D - \frac{1}{T} \right]$ plots (D is the diffusion coefficient of boron calculated from the Sand equation using chronopotentiometric

measurements, T is the absolute temperature) in the temperature range 932-1123 K are straight lines; the calculation of the activation energy gives a value of $35.5 \pm 2.3 \text{ kJ mol}^{-1}$ (see Refs. 31, 32). However, in another study,⁴¹ it has been stated that electrochemically active species (mainly molecular boron fluoride) are produced in a chloride-fluoride melt upon dissociation of boron complexes coupled with homogeneous exchange reactions between the initial complexes and the electrolyte.

Study of the solubility and kinetics of electroreduction of BF_3 in the KCl-NaCl melt at 973 K in the range of BF_3 pressures of $(1.01-5.05) \times 10^5 \text{ Pa}$ showed the presence of two sorts of electrochemically active species.⁴⁴ This accounts for the two waves displayed on the voltammograms whose heights increase proportionally to the pressure. In the opinion of the researcher cited, the first wave corresponds to a two-electron transfer involving BF_3 ; at the potential of the second wave, the product formed in the first stage is reduced, the mixed chloride-fluoride complex $[\text{BCl}_x\text{F}_3]^{x-}$ being involved in the cathodic process.

Thus, despite the large number of publications dealing with the electrochemical properties of boron, no clear views on the mechanism and kinetics of its electroreduction from halide molten electrolytes have been developed so far.

SYNTHESIS OF TITANIUM DIBORIDE AND TITANIUM INTERMETALLIC COMPOUNDS

1. Consideration about high-temperature electrochemical synthesis of chemical compounds

The principal methods used to synthesize high-melting compounds have been classified²³ into the following groups: (1) direct synthesis from elements, (2) synthesis from solutions in melts, (3) thermal reduction of metal oxides by metals, (4) reduction of oxides and other compounds by nonmetals and their derivatives, (5) gas-phase synthesis, and (6) electrolysis of molten salt. This classification is also applicable to the methods used to prepare titanium diboride.

Direct synthesis from elements is based on the reaction of a metal (titanium) with a nonmetal (boron) which can be carried out (at various physical states of the elements) by fusing, sintering or hot pressing. In accordance with this, some researchers suggest conducting this synthesis by keeping samples of pure titanium for 5 h in

amorphous boron at a temperature of 1273 K and a residual pressure of 0.133 Pa; according to other researchers, mixtures of titanium and boron powders should be heated by an electric discharge.⁴⁷ The advantages of this method include the possibility of preparing large amounts of the product, possibility of hot moulding the product in order to manufacture the required articles, simplicity of the equipment and relatively short time needed for the synthesis. The facts that it is difficult to prepare borides with a precise composition and that the temperature of the process is too high are drawbacks of the direct synthesis.

The synthesis from solutions in melts involves a chemical reaction between transition metal atoms and nonmetal atoms (or molecules of a nonmetallic compound) occurring in molten salts or metals. An example is provided by the synthesis of titanium borides in an iron melt at temperatures of 1673-2273 K. This route can afford both pure, doped and complex (mixed) compounds. In addition, it ensures the recovery of metal serving as the solvent and is relatively simple. The drawbacks of this method include low yields and high costs of the product, caused by the large amounts of salts and metallic solvents consumed, possible contamination of the product by the solvent, necessity of vigorous stirring of solutions in melts in order to obtain homogeneous compounds and high temperatures of the process.

The thermal reduction of oxides consists in the reduction of titanium and boron oxides by a reducing metal (in the presence of oxygen-evolving compounds) to give borides at temperatures of 2273-3273 K over time periods of 2-3 min. This is the simplest method among those listed above. However, it is not in general use because of the low quality and inhomogeneous composition of the products obtained and also because it is difficult to isolate the product from the slag.

The fourth method is based on the reduction of oxides and other compounds by metals and their compounds. For instance TiB_2 can be prepared by the reaction of BCl_3 with highly dispersed titanium at 873-1373 K. The advantages of this method include the possibility of preparing large quantities of the product and the fact that oxides and other readily available compounds of titanium and boron are suitable for the synthesis. The main drawback of this method is that the products of the synthesis are contaminated by metal and nonmetal oxides.

The gas-phase synthesis includes a decomposition of chemical compounds followed by reactions of the fragments (both ions or radicals and the atoms or molecules produced upon their reduction) occurring in the gas (vapour) phase to give highly-melting compounds. A method for synthesis of titanium diboride by the reduction using hydrogen of a vapour-gas mixture consisting of TiCl_3 and BCl_3 at temperatures of 1273-3773 K and pressures of 1-3 atm has been described.⁴⁷ The major advantage of this method is the possibility to prepare single crystals and coatings on various materials. The inhomogeneity of the phase composition and relatively low yields of the products as well as complexity of the equipment and processes involved are drawbacks of this technique.

High-temperature electrochemical synthesis from molten salts can be regarded as being one of the most promising methods for preparation of highly-melting compounds. The advantages of this method are relative simplicity of the equipment, ready accessibility to the initial compounds, possibility of preparing simple or complex composite coatings of highly-melting compounds, relatively low temperature of the process, and possibility of controlling the morphology and composition of the cathodic deposit by varying electrolysis parameters. The theoretical grounds of this method have been described in detail,⁴⁸ and examples of its practical implementation can be found in reviews.⁴⁹

Electrochemical behaviour of titanium and boron species as well as electrodeposition of titanium intermetallic compounds have been studied in detail.⁵⁰

Depending on the nature of the electrolyte used, two variants of electrochemical synthesis of titanium diboride can be distinguished, namely, from oxygen-containing systems and from halide systems.⁵¹ The former variant commonly employs systems based on alkali and alkaline earth metal borates, which differ in the manner of saturation of the melt by the synthesis components. Boron can be introduced in melt as B_2O_3 , and titanium can be added as TiO_2 ; the concentration of titanium in the electrolyte can be maintained by anodic dissolution of titanium metal.⁵¹ However, though seemingly advantageous, the synthesis based on oxygen-containing systems suffers from serious drawbacks. Specifically, the reaction temperatures are too high (> 1023 K), and a pure product (free from oxygen) cannot be prepared in this way.

The second variant can be implemented using either purely fluoride electrolytes, for example LiF , NaF , KF , LiF-KF , RbF , CsF and cryolite and mixed chloride-fluoride electrolytes. The sources of titanium and boron in this case are either titanium and boron fluorides or soluble anodes made of boron, titanium and boron carbides or titanium diboride. The use of mixed chloride-fluoride electrolytes with the lowest possible content of fluorine is of the most interest because it decreases the corrosion activity and melt cost. In the case where a soluble anode is used, the formation of fluorocarbons can be avoided and the stage of hydrotreating of the cathodic deposits can be simplified.

The mechanism of the cathodic process in electrochemical synthesis of titanium diboride in the LiF-KF melt has been studied.^{52,53} Potassium hexafluorotitanate and tetrafluoroborate were used as sources of titanium and boron. It was found that reduction of Ti^{4+} in the $\text{LiF-KF-K}_2\text{TiF}_6\text{-KBF}_4$ system occurs in two steps: $\text{Ti}^{4+} \rightarrow \text{Ti}^{3+} \rightarrow \text{Ti}$.

It follows a reduction of boron complexes to elementary boron, which then react with the reduced titanium to yield titanium diboride. This process is accompanied by an electrode depolarisation which is due, in the opinion of the researchers, to the great negative value of Gibbs energy for the formation of TiB_2 .

Thus, the analysis of the studies devoted to the methods of electrochemical synthesis of titanium diboride shows that they are mainly applied studies. The information available from the literature is inadequate to interpret the synthesis mechanism which needs to be studied more comprehensively.

2. Electrochemical properties and stability of halide systems containing titanium in various oxidation states and boron species

2.a. Electrochemical properties and stability of titanium-containing melts

Experiments both on the stability of halide systems containing titanium in various oxidation states and boron and on the electrochemical synthesis of TiB_2 and titanium intermetallic compounds have been performed under an argon atmosphere.

The voltammograms characterising the electroreduction of Ti^{3+} at polarisation rates of $> 0.5 \text{ Vs}^{-1}$ (Fig. 1) exhibit two persistent waves with clear-cut peak currents.

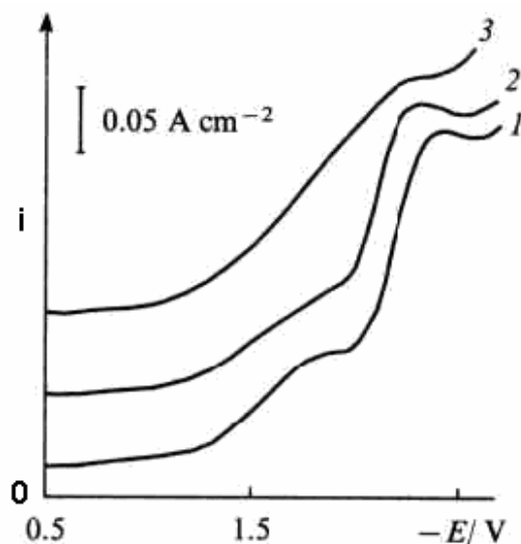
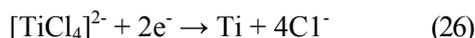


Fig. 1 – Voltammograms for the electroreduction Ti^{3+} ions in the KCl-NaCl-TiCl_3 melt⁵⁰
 $T=1000\text{ K}$, $[\text{TiCl}_3] = 8.3 \times 10^{-5}\text{ mol cm}^{-3}$, chlorine reference electrode; polarization rate/ V s^{-1} : (1) 1.0; (2) 0.1; (3) 0.05.

The waves are due to the following steps of reduction of titanium ions:



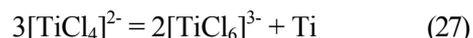
To confirm the assumption that the potentials of the second wave correspond to the reduction to the metal, potentiostatic electrolysis was carried out at these potentials and titanium was identified by X-Ray diffraction on the cathode.

When the potential sweep rate is $< 0.1\text{ V s}^{-1}$ the peak current of process (25) becomes feebly defined and at polarization rates of $< 0.05\text{ V s}^{-1}$ the waves of processes (25) and (26) are superposed, the overall process appearing as occurring in one stage (see Fig. 1, curve 3). This is the reason for the disputes concerning the number of stages in the reduction of Ti^{3+} (see Refs 47, 51), because most of the studies have been carried out at polarisation rates close to the steady-state values (not higher than 10 mV s^{-1}).

Analysis of voltammograms of the type presented in Fig. 2 shows that at high polarisation rates, processes (25) and (26) are not complicated by any additional reactions and, in the range of TiCl_3 concentrations studied (up to 10 mass %), they completely obey the Faraday law. This is confirmed by the fact that the peak currents follow a linear dependence on the concentration of TiCl_3 .

A value of the diffusion coefficient equal to $(3.5 \pm 0.3) \times 10^{-5}\text{ cm}^2\text{ s}^{-1}$ was estimated. It was shown that at $\nu < 0.5\text{ V s}^{-1}$ (see Fig. 2) the process (25) is complicated by a catalytic reaction, *i.e.* electroreduction is accompanied by regeneration of

electrochemically active species and this might be a disproportionation of the divalent titanium formed. It is known that for a redox process not accompanied by catalytic reactions, the ratio of the peak currents (i_p) in the forward (cathodic) and in the reverse (anodic) recording of voltammograms should be unity. If this ratio is greater than unity, the product of electroreduction enters into a subsequent chemical reaction; when there is a disproportionation, this increases the current and changes correspondingly the $i_p/\nu^{1/2}$ value. Study of cyclic voltammograms for process (25) has confirmed⁵⁴ that disproportionation of divalent titanium occurs in the near-electrode layer



The minimum lifetime of divalent titanium species is 0.08-0.8 s for the TiCl_3 concentration ranging from 5×10^{-4} to $5 \times 10^{-5}\text{ mol cm}^{-3}$, the rate constant for disproportionation being $(2.2 \pm 0.2) \times 10^4\text{ cm}^2\text{ mol s}^{-1}$.

Methods for controlling the electroreduction by varying the acid-base properties of the medium have been considered⁵³⁻⁵⁷ in relation to oxygen-containing complex ions. According to the idea of these studies, the electrochemical activity of oxygen-containing anions is due to the acid-base interactions in the melt. The formation and interactions of electrochemically active species in the near-electrode layer occur at a limited rate because these processes require rearrangement of the structure of reacting species. Thus, by changing the acid-base properties of the medium, one can influence the mechanism and rate of the process in a specified manner.

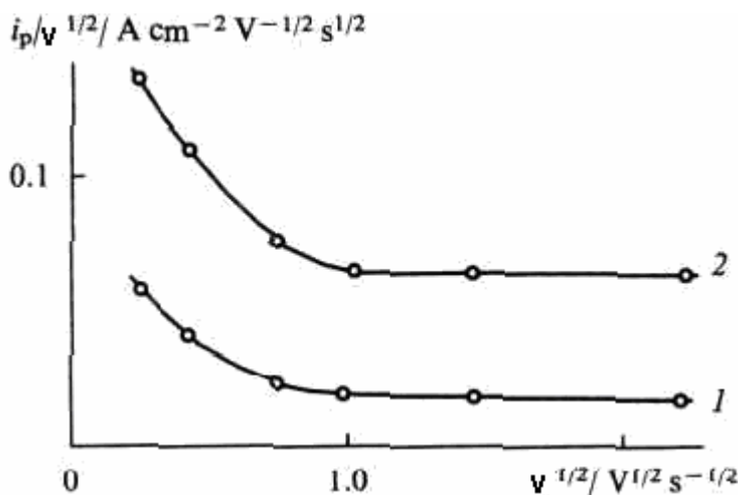


Fig. 2 – Variation of $i_p/v^{1/2}$ vs. $v^{1/2}$ (i_p is the peak current, v is the polarization rate) for process (25) (1) and (26) (2) occurring in the KCl- NaCl- TiCl_3 melt⁵⁰
 $T=1000\text{ K}$, $[\text{TiCl}_3] = 5.5 \times 10^{-5}\text{ mol cm}^{-3}$

Elimination of disproportionation reactions by changing the mechanism of the process can be attained in two ways, namely, by addition of either cations possessing greater polarising ability than potassium and sodium or anions having stronger basic properties than the chloride anion. In the former case, the acidity of the melt diminishes, while in the latter case, the basic properties of the electrolyte are enhanced (for instance, when sodium fluoride is introduced).

The mechanism of the cathodic process in electroreduction of titanium trichloride using chloride melts with different cationic compositions has been studied.⁵⁸ It was found that reduction of titanium in the cesium chloride melt occurs *via* recharge stage (25), whereas in an eutectic LiCl-KCl melt, when the temperature diminishes to 673 K, trivalent titanium is reduced to the metal in one stage: $\text{Ti}^{3+} + 3e^- \rightarrow \text{Ti}$.

However, the LiCl-KCl system cannot be used for the titanium diboride synthesis because the minimum temperature for the onset of the electrochemical synthesis markedly exceeds 673 K.

It has been found⁵⁹ that fluoride ions added to the melt replace the chlorine in the titanium complex species and, as a consequence, the electroreduction process switches to a one-stage mechanism.

In order to elucidate the potential for the synthesis of titanium diboride from chloride and fluoride electrolytes, it is significant to study the influence of fluoride ions, in particular $[\text{Ti}^{3+}]:[\text{F}^-]$ molar ratio, on the mechanism of titanium electrodeposition and the type of titanium species in the melt. It is also necessary to determine the optimum electrolyte composition⁶⁰ which would

preclude both disproportionation reactions and formation of lower titanium fluorides, insoluble in the melt.

As shown above, electroreduction of trivalent titanium in the chloride melt includes the recharge to the divalent state. Upon introduction into the chloride melt of fluoride ions in the ratio $[\text{Ti}^{3+}]:[\text{F}^-] > 1:2$, the peak current of the recharge wave decreases proportionally to the content of fluoride ions in the electrolyte. In addition, as the current of the recharge wave decreases, the peak current of the $\text{Ti}^{2+} \rightarrow \text{Ti}$ discharge wave increases proportionally. When the molar ratio $[\text{Ti}^{3+}]:[\text{F}^-] = 1:2$ is attained, the recharge wave is no longer displayed on the voltammogram and discharge of the complex ions of titanium becomes a one-stage process (Fig. 3).

The switching to a different mechanism is accompanied by a simultaneous shift of electrode potentials towards more negative values. Experimental results imply that the change in the anionic composition of the melt upon the introduction of fluoride ions results in the formation of mixed chloride- and fluoride-containing complexes. The replacement of chlorine in the $[\text{TiCl}_6]^{3-}$ complex (this is the species in which TiCl_3 occurs in the KCl-NaCl melt) by the smaller fluoride ion makes this complex stronger and changes its reduction mechanism. The fact that the mechanism changes at $[\text{Ti}^{3+}]:[\text{F}^-] = 1:2$ suggests that the composition of the resulting titanium complex is $[\text{TiF}_2\text{Cl}_4]^{3-}$.

Titanium chloride-fluoride complexes occur in equilibrium with components of the melt; generally, an equilibrium described by the equation



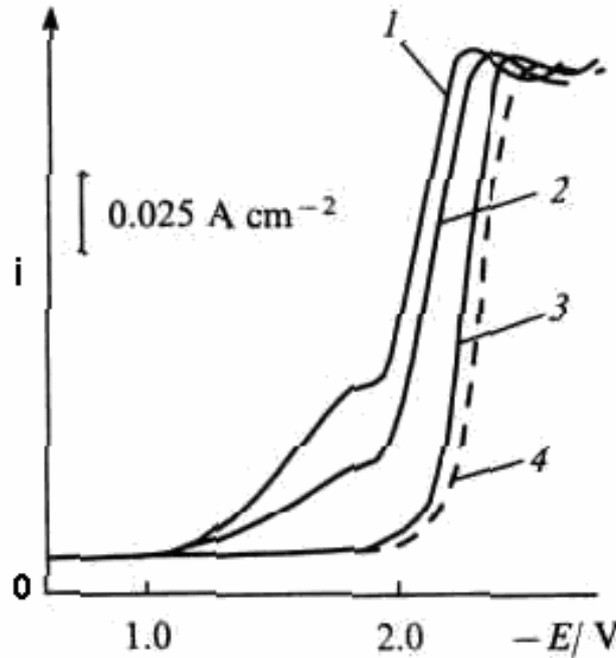


Fig. 3 – Voltammograms¹⁹ for the electroreduction of titanium ions in the KCl-NaCl-NaF-TiCl₃ melt at a [Ti³⁺]:[F⁻] ratio, equal to: 1:0 (1), 1:1 (2), 1:2 (3), <1:2 (4).
T=1000 K, polarization rate 1.0 V s⁻¹, [Ti³⁺] = 1.55 × 10⁻⁴ mol cm⁻³, chlorine electrode as the reference electrode.

Electroreduction involves dissociated species [TiF_x]^{3-y}, the concentration of which is lower than the total concentration of titanium in the electrolyte.

The dependence of the peak potential of the reduction wave on the polarisation rate in the broad range of sweep rates and also the value $\alpha n = 1.0 \pm 0.05$ found using the Matsuda-Ayabe equation, imply that the charge transfer step is slow; α represents the charge transfer coefficient and n is the number of transferred electrons.

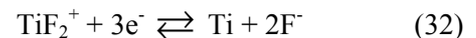
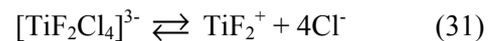
The diffusion coefficient calculated from the Delahay equation using peak currents corresponding to the total concentration of titanium in the melt is equal to $(3.0 \pm 0.4) \times 10^{-5}$ cm² s⁻¹.

The underestimated value of the diffusion coefficient, $(0.7 \pm 0.4) \times 10^{-5}$ cm² s⁻¹, computed by using the peak currents to high polarization rates confirms the assumption that the concentration of electrochemically active species under these conditions does not coincide with the total titanium concentration in the melt.

The number of ligands attached to the initial [TiF₂Cl₄]³⁻ species determined on the basis of the slope of dependence of the straight line plotted in the shift in the wave peak-depolarizer concentration coordinates ($\Delta E - \ln c_F^-$) (Fig. 4) is equal to two.

With allowance for the composition of the initial species, the composition of the titanium complex in

the bulk of a melt containing an excess of fluoride ions should be [TiF₄Cl₂]³⁻. Then the formation and discharge of electrochemically active titanium complexes can be described by the following scheme:



Thus, the electroreduction of trivalent titanium in the near-electrode layer of a chloride melt involves disproportionation of divalent titanium. In a chloride melt, the $\text{Ti}^{3+} \rightleftharpoons \text{Ti}^{2+}$ recharge is a catalysed process and the charge transfer step is retarded over a broad range of polarisation rates. Stabilization of the KCl-NaCl-TiCl₃ molten system can be attained by changing the acid-base properties of the melt by introducing fluoride ions into the chloride electrolyte which results in the formation of mixed chloride and fluoride complexes, such as [TiF₂Cl₄]³⁻. The reduction of these complexes at the molar ratio [Ti³⁺]:[F⁻] < 1:2 occurs in one stage involving the transfer of three electrons.

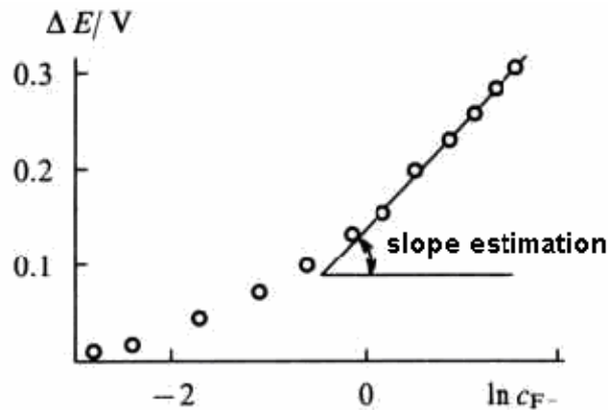


Fig. 4 – Calculation of the coordination number of titanium in titanium complexes occurring in KCl-NaCl-NaF-TiCl₃ melt⁵⁰

2.b. Electrochemical properties and stability of boron-containing melts

In order to perform a high-temperature electrochemical synthesis of titanium diboride, it is necessary that the potentials of the reactive species used in the synthesis to be close. Therefore, investigations of electrochemical properties and stability of boron-containing chloride-fluoride melts are topical. Analysis of the published data indicates that, despite the fact that numerous studies have been devoted to electroreduction of boron complexes, there are still no clear views on the electroreduction mechanism or composition of the boron-containing species existing in chloride-fluoride melts.

The voltammograms of the electroreduction of boron exhibit only one wave in the potential range

from -2.35 to -2.7 V with a clear-cut peak current (Fig. 5). Furthermore cathodic polarisation results in an increase in the current caused by the discharge of alkali metal ions.

The potentiostatic electrolysis at the potential corresponding to the reduction wave produced elementary boron. The dependence of the peak potential of the reduction wave of boron complexes on the polarization rate, substantial difference between the peak potentials found in the forward and reverse recordings of voltammograms, and values $\alpha n = 0.96-1.08$ for the three-electron process calculated using the Matsuda - Ayabe equation demonstrate that in the range of polarization rates from 0.1 to 20.0 Vs⁻¹, the discharge of boron complexes is irreversible, and controlled by the rate of charge transfer.⁶¹

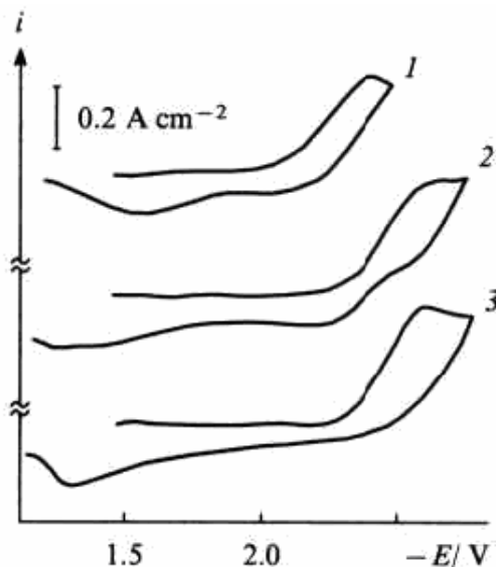


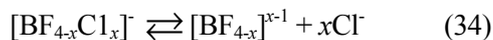
Fig. 5 – Cyclic voltammograms¹⁹ for the B³⁺ → B process occurring in the KCl-NaCl-NaF-NaBF₄ melt (potentials vs. chlorine reference electrode).

[B³⁺]:[F⁻] ratios: [B³⁺] only (1), 1:7 (2), 1:14 (3).

The $[\text{BF}_4]^-$ ions added to a chloride melt enter into an exchange reaction with chloride anions to give mixed chloride-fluoride complexes:



The unstable boron complexes undergo a thermal dissociation which can be generally represented as:



The electroreduction involves species resulting from dissociation of initial complexes, the concentration of these species in the melt bulk being small compared to the total concentration of boron ions. Thus, the melt contains simultaneously two sorts of species, namely, the initial $[\text{BF}_{4-x}\text{Cl}_x]^-$ complexes and electrochemically active species $[\text{BF}_{4-x}]^{x-1}$. Upon the discharge of boron complexes, the near-electrode layer becomes depleted in the $[\text{BF}_{4-x}\text{Cl}_x]^-$ complexes, displacing the equilibrium towards the formation of electrochemically active species. Analysis of the $i_p/v^{1/2} - v^{1/2}$ dependence shows⁶² that a decrease in the potential sweep rate diminishes the influence of dissociation on the electroreduction of boron complexes. The dependence of $i_p/v^{1/2}$ on $v^{1/2}$ can be extrapolated if the peak current corresponds to the total concentration of boron in the melt. The linear dependence of the peak currents on the boron concentration in the melt implies that the preceding chemical reaction exerts no retarding influence under these conditions. The diffusion coefficient calculated from the Delahay equation using the peak current values corresponding to the total concentration of boron is equal to $(3.3 \pm 0.4) \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$.

Thermal dissociation of alkali metal fluoroborates added to a chloride melt yields volatile boron fluorides, poorly soluble in the melt

and evolved as the gas phase. This results in a lower concentration of electrochemically active species and hence, in a lower rate of boron electroreduction.⁶³ The non-controllable change in the concentration of one component hampers the selection of operating conditions suitable for electrochemical synthesis of titanium diboride. Therefore, it is necessary to find a method for stabilization of the chloride melt containing an alkali metal fluoroborate. It has been shown⁶¹⁻⁶² that the introduction of NaF into the KCl-NaCl-NaBF₄ melt induces a shift of the peak potential of the wave corresponding to the discharge of mixed chloride-fluoride boron complexes due to the replacement of chloride ions by fluoride ions. For concentration ratios $[\text{B}^{3+}]:[\text{F}^-] \geq 1:2$, the melt is stabilized upon the formation of mixed chloride-fluoride complexes of boron and the rate of the reduction of boron no longer varies in time. It is of interest to determine the compositions of boron-containing complexes in the initial chloride melt and those of complexes formed after adding fluoride ions.

The coordination number of boron in the chloride-fluoride complex was calculated using a procedure taking into account the dependence of the shift of the peak potential for the reduction of boron complexes on the free ligand concentration in the melt. The concentration of the free ligand was assumed, as a first approximation, to be equal to total concentration of fluorine added to the melt. The coordination number determined from the slope of the straight line plotted in the $\Delta E - \ln c_{\text{F}^-}$ coordinates (Fig. 6) is equal to unity, *i.e.* during titration of the KCl-NaCl-NaBF₄ melt by fluoride ions, one fluoride ion adds to the initial species.

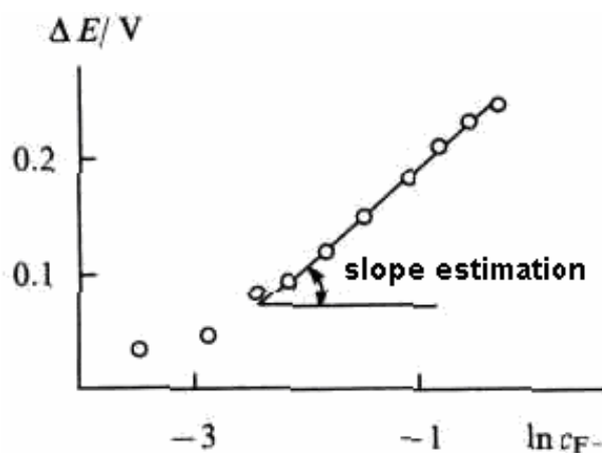
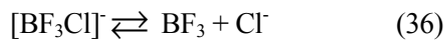


Fig. 6 – Calculation of the coordination number of boron in the complex occurring in the KCl-NaCl-NaF-NaBF₄ melt⁵⁰

Hence, the mixed chloride-fluoride complex of boron in the NaCl-KCl melt has a stoichiometry $[\text{BF}_3\text{Cl}]^-$ and the species occurring in the melt with excess fluoride ions is BF_4^- . The application of a correction for the concentration of the bound ligand, determined by the molar ratio $[\text{B}^{3+}]:[\text{F}^-] = 1:1$, did not alter the results.

Based on the foregoing, it can be concluded that the formation of mixed chloride-fluoride complexes of boron can be described by the following equations:

– in the absence of excess fluoride ions in the melt



– in the presence of excess fluoride ions in the melt



3. Joint electroreduction of titanium and boron species in the chloride-fluoride melt and electrodeposition of titanium diboride powders

It has been shown⁶⁴ that the addition to the KCl-NaCl-NaF-KBF₄ melt of titanium trichloride in a concentration close to half than that of

potassium tetrafluoroborate results in a sharp increase in the current at potentials 0.25 V more positive than the potential of reduction of titanium complexes. This brings about a wave characterized by a steeper slope than the waves for titanium and boron complex ion discharges.

This wave is displayed on voltammograms over broad ranges of polarisation rates and TiCl₃ and MBF₄ concentrations (Fig. 7). If the ratio $[\text{Ti}^{3+}]:[\text{B}^{3+}] = 1:2$ does not hold, then electroreduction of TiB₂ is followed (before the discharge of the alkali metal) by the discharge of the excess (with respect to this ratio) of one component, either titanium or boron.

When this ratio is lower than 1:2, the wave is exhibited at less negative potentials than in the case where the ratio is greater than 1:2. The cyclic voltammograms (with reverse recording) for $[\text{Ti}^{3+}]:[\text{B}^{3+}] = 1:2$ were found to contain one peak, while those for $[\text{Ti}^{3+}]:[\text{B}^{3+}] \neq 1:2$ exhibit at least two peaks. Identification of the products of potentiostatic electrolysis demonstrated that the electrolysis product obtained at the potential of the first cathodic wave was titanium diboride, while that deposited at the second cathodic wave potential were titanium diboride together with the component (either titanium or boron) in excess with respect to the ratio $[\text{Ti}^{3+}]:[\text{B}^{3+}] = 1:2$.

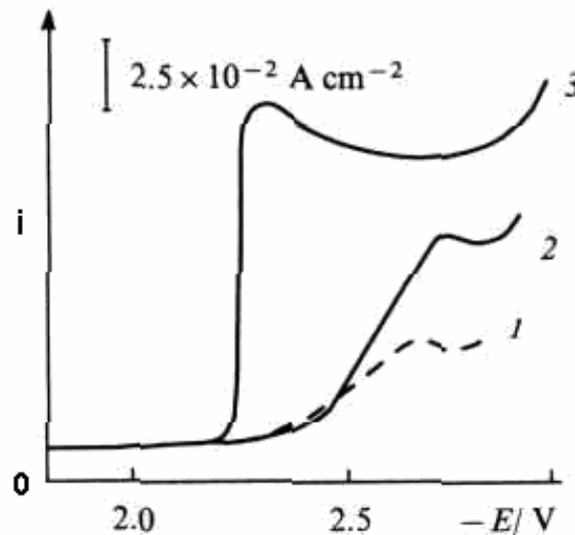


Fig. 7 – Voltammograms¹⁹ for various cathodic processes ($T=1000$ K, polarization rate 1.0 V s^{-1} , chlorine reference electrode); (1) electroreduction of titanium from the NaCl-KCl-NaF melt containing $4.5 \times 10^{-5} \text{ mol cm}^{-3} \text{ TiCl}_3$; (2) electroreduction of boron from the NaCl-KCl-NaF- melt containing $8 \times 10^{-5} \text{ mol cm}^{-3} \text{ NaBF}_4$; (3) electrochemical synthesis of titanium diboride from the NaCl-KCl-NaF-TiCl₃ NaBF₄ melt $[\text{Ti}] + [\text{B}] = 1.2 \times 10^{-4} \text{ mol cm}^{-3}$; $[\text{Ti}^{3+}]:[\text{B}^{3+}] = 1:2$.

The formation of titanium diboride is accompanied by dissipation of a substantial amount of energy and therefore, its synthesis proceeds at more positive potentials than the separate reduction of other component of the synthesis. Depolarization is equal to 0.15-0.3 V (with respect to the initial components), depending on the electrolyte composition. Comparison of the dependences of $i_p/v^{1/2}$ on $v^{1/2}$ for the joint and individual electroreduction of titanium and boron

shows that at high polarization rates the peak current of the synthesis wave is much lower than the sum of the partial currents corresponding to the discharge of the titanium and boron complexes. This fact implies that the process is retarded due to the preceding chemical reaction: the higher the polarization rate, the more pronounced the retardation (Fig. 8). This is due to the fact that the titanium and boron complex ions present in the melt form mixed complexes such as $[B_2TiF_xCl_y]$.⁶⁵

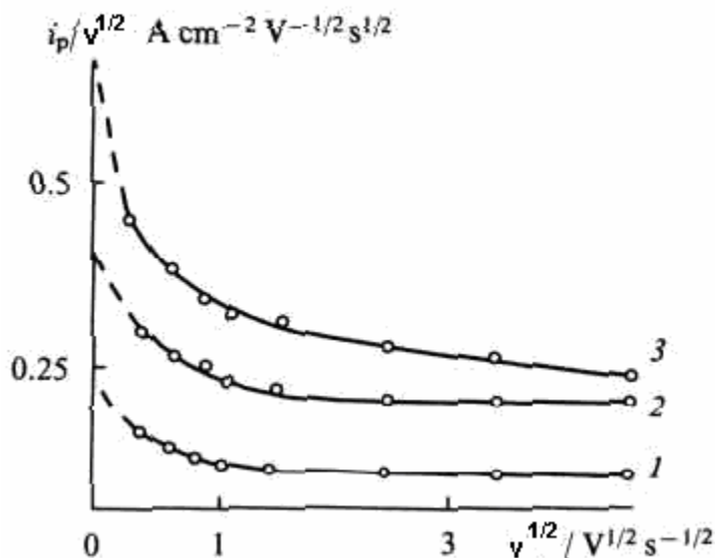


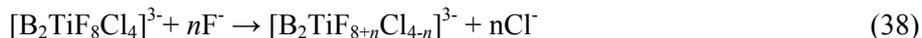
Fig. 8 – Variation of $i_p/v^{1/2}$ vs. $v^{1/2}$ for cathodic processes¹⁹; (1) electroreduction of titanium complexes, $[Ti] = 6 \times 10^{-5} \text{ mol cm}^{-3}$; (2) electroreduction of boron complexes, $[B] = 1.2 \times 10^{-4} \text{ mol cm}^{-3}$; (3) electrochemical synthesis of titanium diboride, $[Ti] + [B] = 1.8 \times 10^{-4} \text{ mol cm}^{-3}$, $[Ti^{3+}]:[B^{3+}] = 1:2$.

The absence of a horizontal section on the $i_p/v^{1/2}$ vs. $v^{1/2}$ plot (as well as on the plots for the partial currents for titanium or boron), which is due to the flow of electrochemically active species moving from the bulk melt, provides grounds for considering, according to the diagnostic electrochemical criteria, that the lifetime of the arising species is short with respect to the lifetime of species formed by titanium and boron in the corresponding systems. A decrease in the polarization rate results in a lower contribution of the formation of joint complexes. Extrapolation of this dependence gives the current corresponding to the total concentrations of titanium and boron in the melt.

The introduction of boron as BF_4^- ions into the chloride melt containing $[TiCl_6]^{3-}$ ions gives rise to exchange reaction (35). The fluoride ion thus liberated is added to the titanium complex, $[TiCl_6]^{3-} + 2F^- \rightarrow [TiF_2Cl_4]^{3-} + 2Cl^-$.

When there is no excess fluoride ions in the melt, the composition of the joint complex is $[B_2TiF_8Cl_4]^{3-}$. The wave of the joint electroreduction (synthesis) is extended along the potential axis and thus the peak width reaches 0.5 V and the potential difference between the peaks of the waves displayed during the forward and reverse recordings of voltammograms is 1.0V. The introduction of fluoride ions into the melt decreases the width of the wave and shifts the potential to the negative region: the difference between the potentials of the peaks of the cathodic and anodic waves decreases simultaneously. For the molar ratio $([Ti^{3+}] + [B^{3+}]):[F^-] \geq 1:15$, the wave width and the difference between the peak potentials barely change.

The shift of the peak potentials of the synthesis wave towards negative values is due to the formation of stronger complexes upon the attachment of the fluoride ions in the melt to the initial complex species.



The fact that the peak and half-peak potentials of the synthesis waves do not depend on the polarization rate as well as the S-shape of voltammograms imply a reversible step of charge transfer at polarization rates $< 2.0 \text{ V s}^{-1}$. The number of electrons involved in the elementary electrochemical step, calculated using the Matsuda-Ayabe equation, is equal to 9. This means that the discharge of the arising joint complexes at a polarisation rate $< 2.0 \text{ V s}^{-1}$ involves simultaneous transfer of nine electrons without kinetic limitations. Study of the electrochemical behaviour of a titanium diboride electrode in a chloride-fluoride melt shows that there is no equilibrium between the electrode and the melt and that the proper electrochemical step is reversible. During dissolution of titanium diboride, limitations appear associated with the chemical interaction at the electrode-melt interface.

As noted above, the charge transfer steps in the separate discharge of titanium and boron complexes are retarded and the electrochemical stage of the synthesis of titanium diboride is reversible. This confirms the assumption that the complexes react chemically in the bulk of the melt to give electrochemically active species, the discharge of which is reversible.

The calculation of the coordination number for a mixed titanium and boron complexes shows that during titration of the melt with fluoride ions, four fluoride ions are attached to the initial species, $[\text{TiB}_2\text{F}_8\text{Cl}_4]^{3-}$, to give $[\text{TiB}_2\text{F}_{12}]^{3-}$ species.

It has been established relying on chronovoltammetric measurements^{72,73} that for the discharge of joint titanium and boron complexes to persist, the molar ratio $([\text{Ti}^{3+}] + [\text{B}^{3+}]):([\text{F}^-]) = 1:4$ (for $[\text{Ti}^{3+}]:[\text{B}^{3+}] = 1:2$) should be maintained in the melt.

The results obtained were taken as the basis for practical implementation of the high-temperature electrochemical synthesis of titanium diboride. The influence of temperature, current density and concentrations of the synthesis components on the composition and properties of the product were studied. A crucial condition for this process is to maintain a constant ratio $[\text{Ti}^{3+}]:[\text{B}^{3+}] = 1:2$ in the melt. In this case, a homogeneous phase of titanium diboride is deposited at cathode over a broad range of current densities. If this condition is not fulfilled, titanium diboride is deposited only at relatively low current densities (up to 0.5 A cm^{-2}),

while at higher current densities it is deposited together with the component present in excess with respect to this ratio. The total concentration of titanium and boron in the molten electrolyte in the 0.6-30 mol % range does not influence significantly the product composition or economical characteristics of the process. A decrease in the total concentration to $< 0.6 \text{ mol } \%$ is not expedient for technological reasons (frequent correction of the cell, low current density and low output), whereas an increase in the concentration to more than 30 mol % brings a greater loss of boron due to the thermal dissociation of potassium tetrafluoroborate, enhanced corrosiveness of the melt and difficulty in washing out the cathodic deposit.

The temperature threshold for the onset of the electrochemical synthesis of titanium diboride in the molten system studied is 873-893K.⁶⁶ The optimum temperature range is 963-1123K. An increase in the temperature above 1123 K is not expedient as it diminishes the thermal stability of the cell and deteriorates the conditions of the formation of the metal salt precipitate. In the temperature range of 963-1123 K, highly dispersed titanium diboride powders with a specific surface area of up to $10 \text{ m}^2 \text{ g}^{-1}$ are produced.

Dispersed powders of titanium diboride were obtained using the KCl-NaCl-NaF (30 mol %)- TiCl_3 (5 mol %) - KBF_4 (10 mol %) system as electrolyte, at current densities of 0.5 to 1.6 A cm^{-2} and a temperature of 1023 K. The degree of dispersion of the powders increased with an increase in the current density if the current density is less than 0.5 A cm^{-2} , dendrites are formed, while at a current density higher than 1.6 A cm^{-2} , titanium diboride is deposited at the cathode together with an alkali metal.

4. Joint electroreduction of titanium with iron, nickel and cobalt species and electrodeposition of titanium intermetallic compounds

In the electrolysis of melts containing simultaneously titanium and iron (nickel, cobalt) chlorides, powdered intermetallic compounds (IMC) of titanium, TiFe , TiFe_2 , Ti_2Ni , TiNi_3 and TiCo , are formed at the cathode.⁶⁷⁻⁶⁹ The ways of controlling the processes of formation of IMC have been studied allowing for the mechanisms proposed for high-temperature electrochemical synthesis.

Electrochemical behaviour of titanium and iron chlorides occurring in the melt has been studied by linear chronovoltammetry with cyclic potential sweep.⁶⁷ The dependence of the limiting current on the concentration was non-linear (unlike this dependence for simple reversible or irreversible processes). The fact that the height of the wave remains constant when the potential sweep rate increases above 0.5 Vs^{-1} implies a retarding influence of the preceding chemical reaction. The illegibility of the peak also confirms that the chemical and electrochemical stages are coupled.⁷⁰ It was noted⁶⁷ that potential of the cathode shifted to the negative region and that the difference between the peak and half-peak potentials increased with sweep rate. In view of the foregoing, with the assumption of first-order chemical reactions, two stages of high-temperature electrochemical synthesis can be distinguished.

A first stage is a chemical reaction to give the equilibrium



(where $z = 3x + 2y$) which shifts towards the formation of the heteronuclear complex $[\text{Ti}_x\text{Fe}_y]^{z+}$; the heat of formation of the corresponding intermetallic compound increases.

The second stage is an electrochemical reduction of the heteronuclear complex cation after the discharge potential has been attained



Analysis of the diffusion coefficients shows that charge numbers of 3 and 4 are preferred for complex IMC (the particular number depends on the particular compound). Therefore, the intermetallic chloride complexes discharged at the cathode are $[\text{TiFe}_2]\text{Cl}_3$ and $[\text{TiFe}]\text{Cl}_3$.

A series of experiments dealing with the influence of the concentration of titanium on the peak potential of nickel discharge has been carried out.⁶⁸ Titanium is formally considered as a "quasi-ligand", although, in accordance with the views of the coordination chemistry of melts,⁷¹ it is incorporated in the nucleus of the heteronuclear complex $[\text{Ti}_x\text{Ni}_y]^{n+}$. The variation of the difference between the potentials of nickel reduction ΔE_{pNi} in the presence and in the absence of TiCl_3 as a function of the logarithm of the ratio of TiCl_3 and NiCl_2 concentrations has three linear sections characterized by different slopes and a horizontal section for a large excess of titanium (Fig. 9).

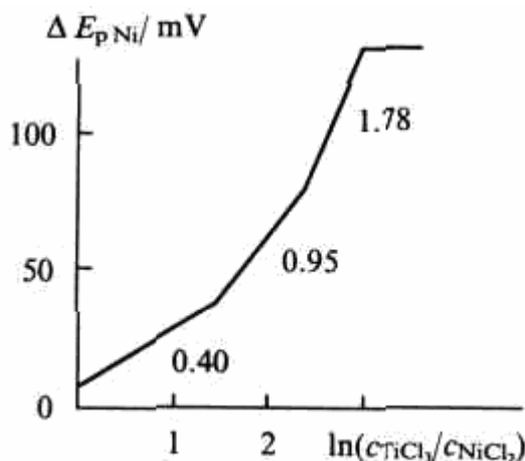


Fig. 9 – ΔE_{pNi} vs. $\ln(c_{\text{TiCl}_3}/c_{\text{NiCl}_2})$ plot⁵⁰, for the KCl-NaCl-NiCl₂-TiCl₃ melt; T = 973 K
The numbers on the curve are explained in the text.

The formal coordination numbers of nickel with respect to titanium (the number of titanium atoms per nickel atom in the complex), determined from the slopes of the linear sections, are 0.4, 0.95 and 1.78. They are close to theoretical values of 0.33, 1.0 and 2.0 respectively, which correspond to the heteronuclear complexes $[\text{TiNi}_3]^{n+}$, $[\text{TiNi}]^{n+}$, $[\text{Ti}_2\text{Ni}]^{n+}$. The assumed formation of these

complexes has been confirmed⁶⁸ by X-ray diffraction analysis of the products of electrolysis at the potential of the intermediate wave of the synthesis. In addition to nickel, the powder contained intermetallic compounds TiNi_3 , TiNi and Ti_2Ni .

The reaction of NiCl_2 and TiCl_3 can be represented in the following way. According to

spectroscopic measurements⁷², Ni²⁺ and Ti³⁺ cations form octahedral complexes [NiCl₆]⁴⁻ and [TiCl₆]³⁻ with chloride anions. Obviously, if the titanium concentrations are low (5×10⁻⁵ molcm⁻³) and the ratio is [Ti]:[Ni] =1:1, only nickel and titanium complexes exist. Upon an increase in the



When titanium is present in a 25-fold (or higher) excess with respect to nickel, the potential of nickel discharge does not shift towards negative values. Apparently, this situation corresponds to that in the systems with excess complex-forming ions. In these systems, the structures of complexes are distorted and they are destroyed or turn into chain structures or into structures of individual molten salts.⁷¹

A study of the electrochemical behaviour of titanium and cobalt species present simultaneously in chloride melts⁶⁹ has proved the possibility of producing intermetallic compounds of titanium with iron group elements by high-temperature electrochemical synthesis. The appearance of the synthesis wave and the character of its variation for Ti-Co and Ti-Ni systems are similar.⁶⁸ The negative slope ratios for the variation of the product of the limiting cathode current by the polarization rate vs. the limiting cathode current of the synthesis wave makes it possible to propose a process of electrochemical reduction of the heteronuclear complex [TiCo]ⁿ⁺. The electrolysis yields a finely dispersed black metallic powder, TiCo. The powder is based on aggregates of dendrites with an average size of 20 μm. The specific surface area of the powder is 30m² g⁻¹.

CONCLUSIONS

The works that are worth mentioning among the most known recent studies devoted to the electrochemistry of titanium and its alloys include investigation of the influence of the cationic and anionic compositions of the melts on the corrosion of titanium in alkali and alkaline earth metal halides,^{73,74} deposition of silicon,⁷⁵ silver^{76,77} and copper^{78,79}, protective coatings onto titanium articles and preparation of titanium and its alloys for deposition.⁸⁰

Studies about electroreduction process in halide melts containing titanium and boron species have allowed for a detailed insight into the equilibria

titanium concentration, a wave for the discharge of the intermetallic compound [Ti_xNi_y]ⁿ⁺ appears between the discharge waves for titanium and nickel which was explained⁶⁸ by a reaction between the two species to give heteronuclear complexes according to the following equations:

existing in these media as well as mechanism of deposition of titanium diboride. The joint electroreduction of titanium species with species of iron group (Fe, Ni, Co) is another way for producing chemical compounds with practical importance.

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