Analysis of the Corrosion Behavior of the TiNi Alloy in the Coarse-Grained State

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Abstract. In this work was investigate the corrosion behavior of the TiNi alloy in a coarse-grained state in inorganic field with different concentration and holding time. An increase in the concentration of the solution leads to a significant acceleration of corrosion processes in the Ti\textsubscript{49.1}Ni\textsubscript{50.9} alloy with a high Ni content, including until the samples are completely dissolved. It was revealed that solutions of 1 M sulfuric and hydrochloric acids after a month's exposure did not change in color and no precipitations were found, while solutions of 5 M hydrochloric and sulfuric acids acquired a violet and then green color, which is due to the predominant release of titanium ions (+4) and nickel (+2).

Introduction
In recent decades, materials with an ultrafine-grained structure have been created, having a grain size of 100 - 300 nm, which have a unique structure and properties, they change fundamental characteristics, such as the Debye and Curie temperatures, saturation magnetization, etc. Shape memory materials (SME) have already found wide application in medicine as long-term functioning materials implanted into the body. A special class of shape memory alloys are nickel and titanium alloys - alloys (NiTi). The range of their application depends on the temperature of martensitic transformation and mechanical properties [1-6]. They exhibit high elastic properties, were able to change their shape when the temperature changes and do not collapse under conditions of alternating load. Phase transitions in such alloys are characterized by a wide hysteresis and a long temperature range, in which the material exhibits shape memory and superelasticity effects. It is known that various metal alloys have the ability to change their shape as a result of temperature changes. This effect is called "austenitic transformation", and another transformation, from austenitic state to martensitic state, is called "martensitic transformation" [7-8]. The biochemical compatibility of physiological fluids and metal implants is largely determined by the electrochemical interaction between them, which usually leads to the transfer of metal ions into tissue fluids. In this case, the implant may also contain heavy elements that are toxic to the body. However, it is impossible to assess biochemical compatibility by the concentration of toxic elements, especially if their introduction into the composition of the implant leads to a significant increase in its corrosion resistance [9-14]. Since the corrosive properties of the implant are the
most important indicators of biochemical compatibility, it is necessary to take into account both
the direct corrosion processes associated with the transition of ions through the interface, and the
reactions leading to the formation of poorly conducting protective films. As a rule, such protective
films delay the release of toxic ions into the tissue, as a result, the implant containing toxic
elements weakly interacts with the surrounding tissues and becomes practically inert with respect
to biological media [15-20].

Corrosion resistance is largely determined by the degree of defectiveness of the material and
the peculiarities of its implantation into the human body as one of the most active carriers of
aggressive media. Tissues are a complex biological system that reacts to the introduction of an
implant by changing its own structure up to physical and mechanical destruction [21-23].
Therefore, it is important to know the features of corrosive behavior, affecting biochemical and
biomechanical compatibility with body tissues.

Material and methods
The study material was a two-component alloy: Ti_{49.1}Ni_{50.9} alloy, which has a B2 austenite
structure at room temperature, with a bcc lattice of the CsCl type. To form a solid solution based
on TiNi and to exclude the prehistory of obtaining the material, the alloy was quenched from the
homogeneity region (from 800 °C) into water. The average grain size of the hardened alloy was
200 μm. To reveal the microstructure of the initial titanium nickelide, an etchant of the following
composition was used: 60% H_2O + 35% HNO_3 + 5% HF. The study of the microstructure was
carried out on an optical metallographic microscope OLYMPUS GX51, as well as using scanning
electron microscopy (SEM) JEOL JSM-6490LV, an inverted microscope AXIO OBSERVER
Z1M in the dark field mode (the studies were carried out in the laboratory of solid state physics of
the Institute of Physics of Molecules and Center of the Russian Academy of Sciences). Before
testing, pre-prepared samples were weighed on an analytical balance, the samples were placed in
a desiccator, where the test sample was in contact with an aggressive medium at a temperature (25
°C). The samples were placed in the solution for a certain time until the appearance of noticeable
traces of corrosion. After testing, the samples were washed with water, treated with alcohol, dried,
and weighed on an analytical balance.

Results
Figure 1 shows optical microscopy of the surface of samples of the Ti_{49.1}Ni_{50.9} alloy in a coarse-
grained state.

![Fig. 1. Photographs of the surface of Ti_{49.1}Ni_{50.9} alloy specimens before corrosion tests in
course-grained state](image)

Figure 2 shows the surface structures of alloy samples in a coarse-grained state after corrosion
tests in a 1 M H_2SO_4 solution with different holding times. At a given concentration of the solution,
no noticeable corrosion damage is observed; only corrosion products are observed on the surface of the samples. At the same time, with an increase in the exposure time in the solution, the amount and volume fraction of corrosion products increase linearly.

Fig. 2. Photographs of the surface of Ti$_{49.1}$Ni$_{50.9}$ alloy samples obtained using an optical microscope (a-d), SEM (e) and an inverted microscope (dark field) (f) in a solution of 1 M H$_2$SO$_4$ in a coarse-grained state: a) 1 day; b) 3 days; c-d) 15 days; e-f) 30 days

Corrosion tests in 1 M HCl also do not lead to damage to the samples, accompanied by an insignificant loss of sample weight, corrosion products are observed on the surface, the volume fraction is maximum at the maximum exposure time, small pitting is observed in some areas (Fig. 3, e).
Fig. 3. Photographs of the surface of Ti$_{49.1}$Ni$_{50.9}$ alloy samples obtained using an optical microscope (a-d), SEM (e) and an inverted microscope (dark field) (f) in a solution of 1 M HCl in a coarse-grained state: a) 1 day; b) 3 days; c-d) 15 days; e-f) 30 days.

Studies in 3% NaCl showed that, compared to tests in acid solutions, there is a smaller amount of corrosion products (Fig. 4). In this case, in the structure of the alloy, regions of different contrast with a size of about 400 ± 30 µm are distinguished at the maximum exposure time in the solution (Fig. 4, e). The maximum amount of corrosion products on the surface of the samples is observed after 15 days of exposure (Fig. 4, c-d).
Investigations of samples after corrosion tests in 5 M HCl on an inverted microscope (Fig. 5, f-g) made it possible to determine the nature of corrosion damage. In the case of a coarse-grained state, deep pits are observed, occupying more than 50% of the sample surface. Studies using scanning electron microscopy also made it possible to estimate the average size and depth of pitting corrosion pits in the coarse-grained state (Fig. 5, e). The average size of the holes is about 100 ± 10 µm. After 1 day of holding the samples in the solution, a large number of corrosion products are observed on the surface, significant damage with the formation of corrosion pits is observed on the 15th day of exposure (Fig. 5, c-d).

Fig. 4. Photographs of the surface of Ti49.1Ni50.9 alloy samples obtained using an optical microscope (a-d), SEM (e) and an inverted microscope (dark field) (f) in a solution of 3% NaCl in a coarse-grained state: a) 1 day; b) 3 days; c-d) 15 days; e-f) 30 days
Fig. 5. Photographs of the surface of Ti_{49.1}Ni_{50.9} alloy samples obtained using an optical microscope (a-d), SEM (e) and an inverted microscope (dark field) (f-g) in a solution of 5 M HCl in a coarse-grained state: a) 1 day; b) 3 days; c-d) 15 days; e-g) 30 days
Studies in solution showed that after 1 day, significant corrosion damage to the sample is observed (Fig. 6, a); after 3 days, the sample weight decreased by 50%, followed by complete dissolution of the alloy sample in a 5 M H₂SO₄ solution.

![Photographs of the surface of Ti₄⁹.₁Ni₅⁰.⁹ alloy samples obtained using an optical microscope in a solution of 5 M H₂SO₄ in a coarse-grained state: a, b) 1 day; c) 3 days](image)

**Fig. 6.** Photographs of the surface of Ti₄⁹.₁Ni₅⁰.⁹ alloy samples obtained using an optical microscope in a solution of 5 M H₂SO₄ in a coarse-grained state: a, b) 1 day; c) 3 days

**Conclusions**

An increase in the concentration of the solution leads to a significant acceleration of corrosion processes in the Ti₄⁹.₁Ni₅⁰.⁹ alloy with a high Ni content, including until the samples are completely dissolved (5 M H₂SO₄ after 3 days of exposure; 5 M HCl is a significant proportion of corrosion damage after 30 days). It was revealed that solutions of 1 M sulfuric and hydrochloric acids after a month's exposure did not change in color and no precipitations were found, while solutions of 5 M hydrochloric and sulfuric acids acquired a violet and then green color, which is due to the predominant release of titanium ions (+4) and nickel (+2). The process of dissolution of the Ti₄⁹.₁Ni₅⁰.⁹ alloy in acid solutions at high concentrations follows the mechanism of pitting, which is confirmed by the photographs of the microstructures presented.

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**References**
