- Corrosion behavior of welded joints of X70 pipeline steel produced by high-frequency welding
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6 Abstract: Corrosion behavior of welded joints of pipe steel category X70 produce by high-7 frequency welding, in different conditions, in which it is possible to initiate stress-corrosion cracking, 8 hydrogen or sulfide cracking, is investigated. According to the results of electrochemical researches, the 9 kinetic parameters of the cathode and anode processes on the surface of the welded joint of pipe in the 10 investigated solutions are determined. Corrosion resistance of the welded joint HFW-pipe is similar to 11 corrosion resistance of the base metal.

Key words: pipe, high-frequency welding, corrosion, stress-corrosion cracking, corrosion
 potential

#### 14 Introduction.

It is well-known that pipes of big diameter from 530 to 1420 mm for pipelines are manufactured 15 using arc welding under flux [1-3]. In this case the effectivity of such type of welding significantly 16 decreasing with decreasing of the thick of pipe wall. Moreover, taking into the account the obligatory 17 implementation of double-sided welding, the equipment for arc welding is considerably complicated, due 18 to the need to transport flux inside the pipe. This contributed to the intensive development of high-19 20 frequency welding (HFW) method, primarily for small and medium diameter pipes. Up to date, in the world, with the HFW technology produces pipes of the categories X65 ... X80 with the diameter from 219 21 to 660 mm and wall thickness up to 25,4 mm. Despite the rapid development of this technology, at the 22 present time in the scientific and technical literature there is virtually no information on the corrosion 23 resistance of the welded joints of HFW-pipes. 24

The purpose of this work was to investigate the corrosion behavior of welded joints of HFW-25 26 pipes, on the example of  $\emptyset$ 508.0 × 8.7 mm pipes from a steel of category X70, in the conditions where initiation of stress-corrosion cracking, hydrogen or sulfide cracking is possible. This researches were 27 conducted in two directions. The first, were evaluation of electrochemical properties in the environments 28 of different aggressiveness. The second, is corrosion resistance in hydrogen sulfide solutions. Also, took 29 into the account the peculiarities of the design of the connections of HFW-pipes, that determined by the 30 31 implementing of the pressure welding, and also the presence of heat treatment in a mode close to normalization, including significantly less, in comparison to the arc processes, chemical and structural 32 heterogeneity. 33

#### 34 Materials and methods.

С

0,08

The chemical composition of the investigated material was determined by the spectral method on the device "Spectrovak-1000" produced by the company "Baird". The chemical composition of rolling steel is shown in Table 1.

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	1 4010			position	01 110		ai 01 111	·· P·P·		
			Chemica	l compo	sition,	wt-% ma	ax			
Si	Mn	P	S	V	Nb	Ti	Ni	Cr	Al	Cu

0,06

0,014

0,19

0,22

0,037

0,02

Table 1 - Chemical composition	of the base metal of HFW-pipes
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39	Metallographic research of the microstructure	e were	carried	out on	the	microscope	"Neofot-21"
40	after the microsection etching in the nital.						

0,040

41

### Electrochemical measurements.

1,34

0,014

0,006

0,22

The evaluation of the corrosion resistance of the welded joint of the HFW-pipe in various media were carried out by the method of polarization curves. As a rule, from the experience of operating oil and gas pipelines, stress-corrosion cracking occurs in the environment with a hydrogen index close to neutral [4-6]. Thereby, the tests were carried out in three solutions with a hydrogen index in the range from 5.4 up to 8.4: 3% NaCl 3 pH 6,8 [4] (solution N1); 0,037 g/l KCl+0,559 g/l NaHCO<sub>3</sub>+0,008 g/l CaCl<sub>2</sub>+0,089 g/l MgSO<sub>4</sub>·7H<sub>2</sub>O, pH 8,2-8,4 [5-6] (solution N2); 1,92 g/l HCOOH + 54,5 g/l HCOONa + 10 g/l KCNS, pH 5,4 [7] (solution N2).

Electrochemical researches were carried out on the microsection of the welded joints of the HFW-49 pipe  $\emptyset$  508,0 × 8,7 mm in the size of 50×20×8,7 mm, with the fusion zone in the middle. Before research, 50 samples were sanded with sandpaper of different grain sizes from 320 to 1000, then washed with running 51 and distilled water and rubbed with alcohol. On the surface of the welded joint area of  $5 \text{ mm}^2$  installed 52 clamping electrochemical cell, the other surface was protected by an insulating coating. Measurement was 53 54 carried out according to the three electrode scheme: working electrode (WE) - basic metal and weld joint; auxiliary electrode (AE) - platinum, comparison electrode (CE) - chlorine-silver electrode. The 55 polarization curves were measured in dynamical mode at the speed of potential sweep of 0.0005 V/s. The 56 57 polarization range from the corrosion potential was 1.5 V to the cathode side and 2.0 V to the anode. The 58 time for measuring the potential of the corrosion potential of the working sample was 60 minutes, after 59 which the potential for scanning was switched on. From the received polarization curves were determined corrosion potential, the current in the region of active anode dissolution for the potential -0.5 V ( $i_{-0.5 \text{ V}}$ ), 60 the Tafel slope of the initial part of the anode curve  $(b_a)$ , the limiting diffusion current  $(i_d)$  and the 61

potential for initiating the release of hydrogen ( $E_{H_{2}}$ ). In analyzing the influence of the protective potential 62 on the corrosion of the surface of the pipe under the protective cover, it is proposed to determine the 63 current density of the cathode protection  $(j_{cp})$  and compare it with the density of the limiting current of 64 oxygen recovery  $(j_{o_1})$  [8]. If the ratio of currents is less than 1, corrosion of the pipe wall in the coating 65 defect is possible; if the ratio of currents is in the range from 1 to 3 - protective effect is achieved. Further 66 increase in the current density of the cathode protection does not lead to a significant reduction in the rate 67 of corrosion, but is accompanied by a sharp increase in the volume of hydrogen which is formed during 68 69 the decomposition of the electrolyte.

# Investigation of the resistance of the welded joint of HFW-pipes against hydrogen cracking and hydrogen sulfide corrosion cracking.

According to the requirements of ANSI / API specification 5L [9], corrosion researches are carried out, by the results of which corrosion rate of the metal in an environment that containing hydrogen sulfide, as well as its resistance to hydrogen cracking (HC) and hydrogen sulfide cracking (HSC) is determined. Tests for resistance to HC were conducted in accordance with NACE TM 0284-2003 [10] for 96 hours. After the tests, each sample was cut into 3 equal parts. Metallographic specimen was made, and the rate of total corrosion, Crack Length Ratio (CLR), Crack Thickness Ratio (CTR), which, accordingly, should not exceed 0.5 mm/year, were determined, CLR-6 % and CTR - 3%.

%

$$CLR = \frac{\sum a}{W} \cdot 100$$

$$CLR = \frac{\sum b}{T} \cdot 100 \% .$$

81 where a - length crack, b - thickness crack, W - section width, T - test specimen thickness.

For tests of resistance to hydrogen sulfide cracking (HSC), the samples were made in such a way that one of the surfaces is located at the minimum possible distance from the inner surface of the pipe. Tests were conducted into the accordance to NACE TM 0177-2005 (method B, ISO 7539-2-89) for 720 hours [11]. The load level was equal. The criterion of resistance to HSC is the absence of cracks.

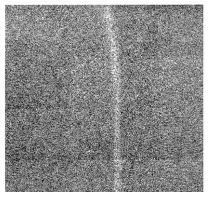
86 **Results and discussion.** 

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The microstructure of the fusion zone of the HFW pipe  $Ø508.0 \times 8.7$  mm from steel of category X70 is shown in Figure 1. The fusion line represents a distinct continuous carbon-free light band. Due to the local thermal treatment of coarse-grained lamellar or needle structure that formed after welding in the

- 90 zone of thermal influence, transforms into fine-grained ferrite-perlite. The amount of perlite component in
- 91 the structure is determined by the carbon content of the steel.





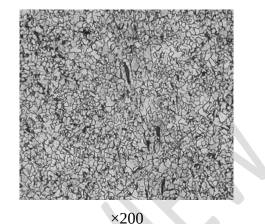
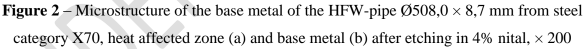


Figure 1 - Microstructure of the fusion zone of the HFW-pipe welded joint  $Ø508.0 \times 8.7$  mmfrom steel category X70 after etching in the nital

92 The microstructure of the base metal and the heat affected zone of the pipe is a fine-grained ferrite-

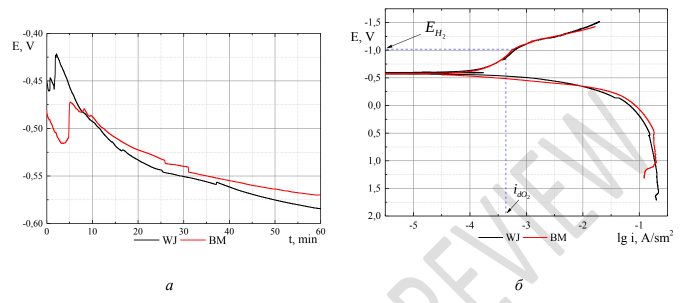
93 perlite mixture with enough highlighted strips (Figure 2).





Figures 3, 4 and 5 shows the dynamics of changes in the corrosion potential (a) and the 94 polarization curves (b) of the base metal (BM) and the welding joint (WJ) of the HFW-pipe in solutions № 95 1,  $N_{2}$  2 and  $N_{2}$  3, respectively. As can be seen from the curves presented Fig. 3a, the corrosion potentials 96 on BM and WJ are close in value and are -0.570 V and -0.584 V respectively. The behavior of the anode 97 and cathode curves of the two welded zones is similar Fig. 3b. The numerical values of the 98 electrochemical parameters of the cathode and anode processes, that determined graphically from Fig. 3b, 99 is shown in Table 2. From the corrosion potential up to 0.5 V for both zones in this solution, is 100 characterized a wide area of active dissolution. There is a slightly smaller angle of the Tafel slope of the 101 102 starting zone of the anode curve of WJ zone in comparison the BM, which indicates a higher rate of anode 103 dissolution under the same conditions. For potentials larger than -0.25 V, the deceleration of the anode

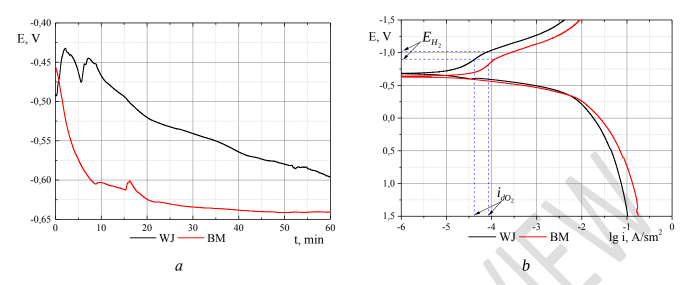
104 dissolution was observed due to passivation of the surface of the sample.



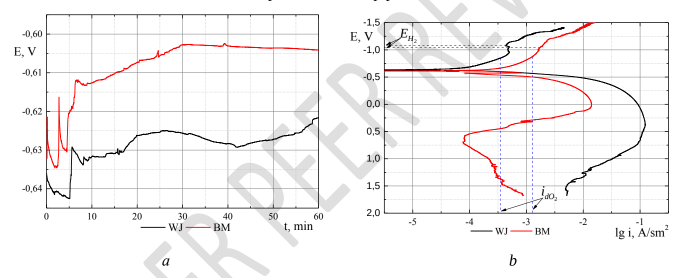
*Figure 3* - Changes in corrosion potential in time (a) and the polarization curves (b) BM and WJ of the sample of the HFW-pipe in solution №1.

In the solution №2, BM is characterized by a sharp shift of the corrosion potential in the region of 105 negative values from -0.455 V that stabilizes after 60 minutes at -0.641 V, and on the WJ were shifted 106 smoothly from -0.490 V, and stabilizes at -0.596 V, Fig. 4a. This difference in potentials is significant, but 107 considering the much smaller area of the WJ as compared to the BM, it does not pose any danger in terms 108 of the development of local corrosion processes. The area of active dissolution is long enough for both 109 zones - from the potential of corrosion to the potential is about -0.25 V, above which there is a slowing 110 down of the anode process due to the surface shielding by products of corrosion. The Tafel's angles of 111 slope of the initial regions of the anode curves indicate a higher rate of dissolution of the BM in 112 comparison with the WJ. 113

In solution N  $\Im$  3, the dynamics of the change in corrosion potentials within 60 minutes is characterized by a slight displacement in the region of positive values for both on the BM and the WJ, varying from -0.632 V up to -0.604 V and -0.636 V up to -0.622 V, respectively. The electrochemical characteristics of the anode and cathode processes on the BM and the WJ are shown in Table 2. The analysis of the course of the anode curves shows that the rate of dissolution of the WJ at the potential -0.5 V is almost 6 times higher than the BM (Table 2). Both zones are characterized by the area of active dissolution: for the BM in the range -0.543 ... -0.03 V; for the WJ - -0.588 ... 0.25 V.



*Figure 4* - Changes in corrosion potential in time (a) and the polarization curves (b) of the BM and the WJ of the sample of the HFW-pipe in solution №2



*Figure 5* - Changes in corrosion potential in time (a) and polarization curves (b) of the BM and the WJ of the sample of the HFW-pipe in solution №3

121 The analysis of the ratio of currents  $j_{cp}/j_{o_2}$  for the region of the WJ and the BM were conducted 122 in order to determine the possible formation on the surface of the pipe under peeling coating of local areas 123 with different electrochemical activity. The results of the analysis are shown in Table 2.

From the analysis of experimental data it can be concluded that in the range of protective potentials (from -0.85 V up to -1.15 V relative to copper-sulfate electrode comparison, which corresponds the value from -0.75 V to -1.05 V relative to the chlorine-silver electrode comparison), the ratio of currents  $j_{cp} / j_{o_2}$  in solutions No 1-3, both on the BM and on metal of the WJ is more than 1 by the polarization potential, which is approaching the maximum protective value. Such conditions contribute to the decomposition of the electrolyte with hydrogen release. But it should be noted that the ratio of currents  $j_{cp}/j_{o_2}$  for the BM and the WJ practically does not differ. This indicates that when working under the conditions of cathodic protection of hydrogen absorption of the weld and base metal is not expected.

- The results of the study of the resistance of the welded joint of the HFW-pipe to the HC and theHSC are given in Table 3.
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Table 2 - Electrochemical characteristics of different zones of the HFW-pipes in solutions №1-3

	Zone	Electro	ochemica	al character	ristics	Currents	of cathode	<b>Ratio</b> $j_{cp} / j_{o_2}$		
		anoc	le	cathode		protection, A/m <sup>2</sup>				
tion				<sup><i>i</i><sub>d</sub></sup> , A/m <sup>2</sup>	<sup>Е</sup> <sub>н2</sub> , В	for	for	for cathode	for	
Solution		i <sub>-0.5 B</sub>	<sup><i>b</i></sup> <sup><i>a</i></sup> , B			cathode	cathode potentials	potentials -0,75 V	cathode	
		A/m <sup>2</sup>				potentials			potentials	
						-0,75 V	-1,05 V		-1,05 V	
<b>№</b> 1	BM	1,1×10 <sup>-3</sup>	0,069	0,0007	-1,03	2,8×10 <sup>-4</sup>	7,6×10 <sup>-4</sup>	0,4	1,08	
J (± 1	WJ	3,1×10 <sup>-4</sup>	0,054	0,0007	-1,03	$2,8 \times 10^{-4}$	7,6×10 <sup>-4</sup>	0,4	1,08	
<u>№</u> 2	BM	1,2×10 <sup>-4</sup>	0,074	8,2×10 <sup>-5</sup>	-0,94	4,5×10 <sup>-5</sup>	2,6×10 <sup>-4</sup>	0,5	3,2	
J 1=2	WJ	8,1×10 <sup>-4</sup>	0,087	3,8×10 <sup>-5</sup>	-1,0	$1,5 \times 10^{-5}$	1,1×10 <sup>-4</sup>	0,4	2,9	
№3	BM	1,2×10 <sup>-3</sup>	0,057	1,5×10 <sup>-3</sup>	-1,2	7,1×10 <sup>-4</sup>	1,8×10 <sup>-3</sup>	0,5	1,2	
	WJ	6,8×10 <sup>-3</sup>	0,069	5,0×10 <sup>-4</sup>	-1,2	2,5×10 <sup>-4</sup>	4,9×10 <sup>-4</sup>	0,5	1,0	

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**Table 3** - Indicators of corrosion resistance of the BM and the WJ metal of HFW-pipes

Indicators of corrosion resistance 136								
Corrosion	Н	C	HSC at <sup>37</sup>					
rate,			$0,7\sigma_{0,2}^{138}$					
mm/year	CLR, %	CTR, %	139					
0.5	32-37	2.0-2.3	Cracks are					
0,5	5,2 5,7	2,0 2,5	not detected					
	0.0	0.0	Cracks are					
	0,0	0,0	not detected					
	Corrosion rate,	Corrosion H rate, mm/year CLR, %	Corrosion         HC           rate,         CLR, %         CTR, %           0,5         3,2-3,7         2,0-2,3					

The results obtained in Table 4 show that the welded joint of the HFW-pipe is resistant to HC and HSC by the determined parameters (corrosion rate,

145 CLR, CTR, HSC resistance at  $0.7\sigma 0.2$  for 720 hours).

## 146 **Conclusions**

The mechanism of anodic dissolution of both the base metal and the welding joint is controlled by
 diffusion of oxygen, which is confirmed by the values of the Constants of Tafel (from 0.054 V to 0.087

V). Differences in the values of the observed Tafel's slopes, indicate a greater or lesser speed of this
process. Such anode behavior indicates that the corrosion resistance of the welded joints of the HFWpipes in general is similar to the corrosion resistance of the base metal.

152 2. The potential for the initiation of hydrogen release in the range of pH from 5.4 to 8.2 is approximately153 similar for the base metal and for the weld joint.

154 3. In the range of protective potentials in solutions of different aggressiveness, the ratio of the current of 155 cathode protection to the boundary diffusion current  $j_{ep} / j_{o_2}$  for the base metal and the weld joint does not 156 differ. Thus, in the conditions of cathodic protection, flooding of the base metal and the weld is not 157 expected.

Summing up the above, it can be argued that when operation a pipeline constructed using high-frequency welding technology under pressure in the conditions of cathodic protection, the resistance of welded joints to stress-corrosion cracking will depend only on the protective properties of the insulating coating.

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