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A Calculation Method for Determining the Concentration of Potentially Acid Substances in Feedwater of Once-Through Boilers

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Abstract—The problem of chemically monitoring the products resulted from destruction of organic substances in the steam–water path of a supercritical-pressure power unit is considered. The level at which the concentrations of acetic and formic acids have to be measured is determined, and the results from measurements of electric conductivity and pH in feedwater and steam are given. A procedure that allows the conditional concentration of acetic acid in steam to be calculated from measured electric conductivity in feedwater and steam is presented.

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The feedwater of once-through boilers for supercritical steam conditions contains almost no mineral admixtures (the electric conductivity $\chi_H < 0.3 \,\mu$ S/cm) and a small quantity of organic substances. When exposed to the conditions of oxidizing water chemistry with the oxygen concentration ranging from 100 to 400 ppb and with the coolant temperature rising from 300°C at the boiler inlet to 550°C at the boiler outlet, these organic substances undergo deep thermal destruction (thermolysis), during which acidic substances are frequently generated.

Organic substances enter into the feedwater of powergenerating boilers together with main condensate [petroleum products, products from destruction of ion-exchange resins used in the condensate polishing unit (CPU), and inleakages of cooling water in turbine condensers and heat network water in delivery-water heaters] and with makeup water from the condensate storage tank. It has been found that acetic acid is the most probable product from thermolysis of such substances [1–3]. When acetic acid and other acidic substances appear in the steam of once-through boilers, the medium circulating in them becomes more corrosive and frequently results in corrosion damage to the tubes of high-pressure heaters and delivery-water heaters that use steam of extractions from the turbine flow path [4, 5].

At present, the Operational Regulations for powergenerating boilers installed at Russian thermal power stations (TPSs) do not prescribe the need to monitor organic impurities such as potentially acid substances in feedwater and steam for the lack of reliable measurement tools [6]. The staff of foreign power stations frequently carry out such monitoring using the total organic carbon and total inorganic carbon as indicating parameters. The standard level for the first parameter is equal to 100 ppb [3, 5, 7]. The operating services of Russian power stations make attempts to estimate the quantity of potentially acid substances from measurements of χ and pH of feedwater and steam in oncethrough boilers [5, 8]. Although easy to implement, such a method provides insufficient information, since the effect on these parameters of impurities contained in water and steam is multifactorial in nature.

The authors of this paper have measured χ and pH of feedwater and steam in once-through boilers at several TPSs. An analysis of the obtained results and data available from the literature (Table 1) shows that the electric conductivity of samples (or the electric conductivity of H-cationated samples if ammonia is metered into feedwater) increases from 0.05 to 0.20 µS/cm and the pH value of samples decreases by 0.1–0.3 as a transition occurs from feedwater to steam. Such changes are stable in nature and occur in all likelihood as a result of thermolysis of organic impurities contained in feedwater [9]. In Table 2 we find some results from analyses of feedwater and steam sampled in powergenerating boilers for the content of products from thermolysis of organic impurities in coolant according to the data of [10-12].

The results from the investigations specialists of the All-Russia Thermal Engineering Institute carried out on CPUs at two district power stations have shown that the average quantity of lactic and acetic acids that enters into condensate downstream of these units is in the range from 20 to $40 \,\mu g/dm^3$ for each of them. However, the major portion of sorbed organic substances escaped from the CPU in the second half of the filtration cycle if mixed-bed filters were installed in the CPU. These substances were observed escaping from the polishing unit during almost the entire filtering cycle if separate H–OH ion-exchange treatment was used in the CPU. The results from analyses of feedwater and steam in supercritical-pressure drum boilers

Power unit	Feedwater		Ste	am	Steam-water difference		
rowerumt	χ, μS/cm	рН	χ, μS/cm	рН	Δ χ, μS/cm	ΔрН	
Reftinsk district power station	0.15	7.9	0.25	7.6	+0.10	-0.3	
Mosenergo's TETs-26 cogeneration station, the 250-MW power unit:							
before the replacement of CPU ion-exchange filters (1997)	_	_	_	_	+0.10	-0.40	
after replacement of CPU ion- exchange filters (2005)	0.10	7.0	0.15	6.8	+0.05	-0.20	
Perm district power station	0.1	8.08	0.16	7.87	+0.06	-0.19	

Table 1. Electric conductivities and pH values of feedwater and steam and their differences for TPS units equipped with oncethrough boilers

Table 2.	Results	form	analyses	of sam	ples for t	the content	of acetic	and	formic	acids
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Characteristic of a sample	Concentration, $\mu g/dm^3$ of			
Characteristic of a sample	acetic acid	formic acid		
Turbine condensate of a 300-MW power unit (after the CPU) [10]	1060	_		
Cogeneration power unit equipped with a drum boiler ($p = 14$ MPa) with distillate from a multistage evaporating plant (MEP*) used as makeup water [3]:				
feedwater	18.7	2.5		
superheated steam	28.1	4.8		
Cogeneration power unit with a drum boiler ($p = 14$ MPa) with chemical demineralization of makeup water [3]:				
feedwater	500750	10100		
superheated steam	7590	24		
Superheated steam at the cogeneration station of Moscow Power Engineering Institute [11]	060	025		
Power unit at the Saransk TETs-2 cogeneration station equipped with a drum boiler ($p = 15$ MPa) during abnormal operation of the MEP [12]:				
feedwater	6090	34		
superheated steam	40100	2		
Kazan TETs-3 cogeneration station (1997) [12]:				
chemically demineralized water	215	2		
distilled water from the MEP	54	2		
superheated steam ($p = 14$ MPa)	82	2		

* MEP – multistage evaporating plant.

[11, 12] have revealed that the concentrations of acetic acid in water and steam are considerably higher in them (50 to 100 μ g/dm³) than they are in once-through boilers. The generalized results of studies carried out by foreign researchers [9] are in agreement with the data obtained in Russia: the concentration of acetic acid in steam is in the range from 1 to 7 μ g/dm³ if a CPU is in operation and from 9 to 120 μ g/dm³ if a CPU is not used.

We proposed a method for calculating the concentration of products from destruction of organic compounds in a once-through boiler recalculated for acetic acid, which is the most probable product of thermolysis. It is assumed that there are no other sources from which acidic substances could enter into boiler steam. The values of χ , $\chi_{\rm H}$, and pH of feedwater and steam measured by means of regular instrumentation are used for calculations [13, 14]. The concentrations of mineral impurities in feedwater are thus determined.

The equations of electric neutrality can be written as follows:

for an H-cationated sample of live steam,

$$[H^{+}]_{H}^{l.s} + [Na^{+}]_{H}^{l.s} = [Cl^{-}]_{H}^{l.s} + [HCO_{3}^{-}]_{H}^{l.s} + [CH_{3}COO^{+}];$$
(1)

and for an H-cationated sample of feedwater,

$$[H^{+}]_{H}^{fw} + [Na^{+}]_{H}^{fw} = [Cl^{-}]_{H} + [HCO_{3}^{-}]_{H}^{fw}.$$
 (2)

In these equations, the concentrations of sodium ions and chloride ions in feedwater and steam are equal to each other, i.e., $[Na^+]_{H}^{1.s} = [Na^+]_{H}^{fw}$ and $[Cl^-]_{H}^{1.s} = [Cl^-]_{H}^{fw}$ provided that these impurities transfer completely from water to steam. Then, the difference of Eqs. (1) and (2) gives the expression

$$[H^{+}]_{H}^{l.s} + [H^{+}]_{H}^{fw}$$

$$= [CH_{3}COO^{-}] - ([HCO_{3}^{-}]_{H}^{fw} - [HCO_{3}^{-}]_{H}^{l.s}).$$
(3)

The concentration of acetate ion in Eq. (3) can be taken approximately equal to the total concentration of acetic acid, i.e., $[CH_3COO^-] = C_{ac}$. The value the ratio of the forms $[CH_3COO^-]/C_{ac}$ has for an H-cationated cooled sample for the usual values of pH of around 6.0 or higher is equal to 20 : 1 or higher; hence, the adopted assumption is valid within the limits of a 5% error.

The equations of electric conductivity in an H-cationated sample can be written in the following form:

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Fig. 1. Circuit arrangement for metering ammonia and acetic acid at the laboratory setup.

for live steam,

$$1000\chi_{\rm H}^{\rm l.s} = \lambda_{\rm H^{+}}[{\rm H^{+}}]_{\rm H}^{\rm l.s} + \lambda_{\rm Na^{+}}[{\rm Na^{+}}]_{\rm H}^{\rm l.s}$$
$$+ \lambda_{\rm CI^{-}}[{\rm CI^{-}}]_{\rm H}^{\rm l.s} + \lambda_{\rm HCO_{3}^{-}}[{\rm HCO_{3}^{-}}]_{\rm H}^{\rm l.s} \qquad (4)$$
$$+ \lambda_{\rm CH_{3}COO^{-}}[{\rm CH_{3}COO^{-}}]_{\rm H}^{\rm l.s};$$

and for feedwater,

$$1000\chi_{\rm H}^{\rm fw} = \lambda_{\rm H^{+}}[{\rm H}^{+}]_{\rm H}^{\rm fw} + \lambda_{\rm Na^{+}}[{\rm Na}^{+}]_{\rm H}^{\rm fw} + \lambda_{\rm CI^{-}}[{\rm CI}^{-}]_{\rm H}^{\rm fw} + \lambda_{\rm HCO_{3}^{-}}[{\rm HCO_{3}^{-}}]_{\rm H}^{\rm fw} .$$
(5)

Subtracting expression (5) from Eq. (4), substituting (3) into the result, and taking the adopted assumptions into account, we obtain:

$$1000(\chi_{\rm H}^{\rm l.s} - \chi_{\rm H}^{\rm rw}) = (\lambda_{\rm H^{+}} + \lambda_{\rm CH_{3}COO^{-}})C_{\rm ac}$$

$$\cdot (\lambda_{\rm H^{+}} + \lambda_{\rm HCO_{3}^{-}})([\rm HCO_{3}^{-}]_{\rm H}^{\rm fw} - [\rm HCO_{3}^{-}]_{\rm H}^{\rm l.s}),$$
(6)

where $\chi_{\rm H}^{\rm l.s}$ and $\chi_{\rm H}^{\rm fw}$ are the measured values of the electric conductivity of H-cationated samples of live steam and feedwater, μ S/cm; and $\lambda_{\rm H^+}$, $\lambda_{\rm HCO_3^-}$, and $\lambda_{\rm CH_3COO^-}$ are the reference values of the limiting electric mobilities of H⁺, HCO_3^-, and CH_3COO^- ions.

If the concentration of acetic acid in a cooled H-cationated sample is considerably higher than the change that takes place in the concentration of hydrocarbonates as a transition occurs from feedwater to steam, i.e.,

$$C_{\rm ac} \ge [\rm HCO_3^-]_{\rm H}^{\rm tw} - [\rm HCO_3^-]_{\rm H}^{\rm l.s},$$
 (7)

the second term on the right-hand side of Eq. (6) can be ignored. The solution of this equation with respect to the concentration of acetic acid $(\mu mol/dm^3)$ is given by

$$C_{\rm ac} = 1000 \frac{\chi_{\rm H}^{\rm l.s} - \chi_{\rm H}^{\rm fw}}{\lambda_{\rm H^+} + \lambda_{\rm CH_3 COO^-}}.$$
 (8)



Fig. 2. Electric conductivities of the direct sample (a) and H-cationated sample (b) vs. the concentration of acetic acid. Value of $C_{\rm NH_*}$, μ g/dm³: (1) 3, (2) 20, and (3) 45.

For a temperature of 25°C, Eq. (8) becomes

$$C_{\rm ac} = 2.56(\chi_{\rm H}^{\rm 1.s} - \chi_{\rm H}^{\rm fw}).$$
 (9)

Equation (9) allows us to determine the concentration of potentially acid substances in feedwater of a once-through boiler expressed in terms of $C_{\rm ac}$ in the steam of this boiler from measured values of the electric conductivity of cooled H-cationated samples of steam and feedwater. If no ammonia NH₃ is metered into feedwater, the calculation using Eq. (9) can be carried out for the electric conductivities of direct samples (unlike H-cationated ones) of live steam $\chi^{\rm l.s}$ and feedwater $\chi^{\rm fw}$.

If inequality (7) is not satisfied, the expression for $C_{\rm ac}$ obtained from (6) has a more complicated form, which we do not consider here. Then, Eq. (9) for the concentration of conditional acetic acid (μ g/dm³) becomes

$$C_{\rm ac} = 153.6a(\chi_{\rm H}^{\rm l.s} - \chi_{\rm H}^{\rm fw}),$$
 (10)

 Table 3. Results from measurements of electric conductivity and pH and calculation of the concentrations of acetic acid and ammonia from laboratory analyses of water samples

Instrument readings						Measured	concentra	Calculate	d concen	
S	ource wate	r	after metering of NH ₃ or CH ₃ COOH		tions of		trations of		$\Delta C_{\rm ac},\%$	
χ	Хн	рН	χ	χн	рН	$C_{\rm NH_3},$ $\mu g/{\rm dm}^3$	$C_{\rm ac},$ µg/dm ³	$C_{\rm NH_3},$ µg/dm ³	$C_{\rm ac},$ $\mu g/{\rm dm}^3$	
0.23	0.30	7.95	0.18	0.46	6.96	-	15	_	24.6	+63.8
0.27	0.25	7.94	0.22	0.48	6.94	_	25	—	34.7	+38.8
0.28	0.27	7.76	0.21	0.45	6.85	_	24	—	27.9	+16.4
0.19	0.15	7.30	0.28	0.57	6.53	_	60	—	63.9	+6.48
0.15	0.24	6.87	0.46	0.85	6.05	_	90	—	93.2	+3.58
0.14	0.25	6.73	0.29	0.31	7.00	20	-	19.2	—	-4.10
0.14	0.23	6.48	0.50	0.30	6.94	40	—	42.6	—	+6.58
0.14	0.22	6.72	0.79	0.35	7.75	80	-	89.6	—	+12.1
0.12	0.22	6.75	0.25	0.36	6.95	19.2	7.7	19.2	8.3	+7.66
0.13	0.23	6.68	0.27	0.41	6.76	19.2	23	9.93	27.3	+18.8
0.15	0.24	6.40	0.32	0.65	6.32	19.2	78	19.0	62.5	-19.9
0.20	0.32	6.84	0.33	0.84	6.56	25	120	22.1	80.1	-33.2
0.15	0.32	6.95	0.53	0.52	7.89	45	27.2	12.6	29.9	+10.1
0.14	0.31	7.24	0.42	0.93	7.55	45	108	35.0	96.1	-11.3

		Calculated values				
pH^{fw} χ_H^{fw} , $\mu S/cm$		χ^{fw}_{H} , pH ^{l.s} μ S/cm		$\chi_{\rm H}^{\rm l.s},$ $\mu S/cm$	C _{NH3} , μg/dm	$C_{\rm ac},$ $\mu g/{\rm dm}^3$
8.179	0.090	0.40	7.79	0.13	21.1	6.14
8.083	0.083	0.28	7.74	0.14	10.3	8.76
8.08	0.100	0.31	7.87	0.16	13.8	9.22
8.30	0.103	0.593	8.01	0.125	38.0	3.38
7.99	0.080	0.28	7.9	0.11	14.9	4.62
6.60	0.188	0.153	6.5	0.230	0	6.40
6.60	0.214	0.176	6.5	0.272	0	8.90

 Table 4. Results from calculation of the concentrations of acetic acid and ammonia in power units equipped with oncethrough boilers obtained using the authors procedure

where *a* is an empirical coefficient, the value of which in a first approximation is equal to unity.

Experiments were carried out on a test facility (Fig. 1) to verify the procedure for calculating the indicators characterizing the quality of feedwater and steam in a once-through boiler. Distilled water is fed into an H-cation-exchange filter and then into an OH-anionexchange filter. The demineralized water downstream of the H and OH ion-exchange filters has high quality ($\chi = 0.2-0.15 \ \mu$ S/cm) and very low concentration of mineral salts. A solution of ammonia or acetic acid is then metered (to simulate a steam sample), after which χ and pH of mixed solution are measured (a direct sample). Further on, water flows to an H-column, downstream of which $\chi_{\rm H}$ (an H-cationated sample) is measured, and to the demineralized water tank.

The test results are shown in Fig. 2 (for different values of $C_{\rm NH_3}$). We see that the electric conductivity of a direct sample (Fig. 2a) varies very considerably depending on the combination of acetic acid and ammonia concentrations. Different values of electric conductivity may correspond to the same value of $C_{\rm ac}$, a circumstance that makes it difficult to determine it by calculation. In contrast, the electric conductivity of an H-cationated sample (Fig. 2b) gives a stable linear dependence on $C_{\rm ac}$ and depends only slightly on $C_{\rm NH_3}$ in the sample.

At a = 1, the calculated values of C_{ac} are in satisfactory agreement (with an average error of 18%) with the measured values (Table 3).

The obtained equation (10) was used for calculating the concentration of the products from thermolysis of potentially acid substances in feedwater of several power units at district power stations equipped with once-through boilers (Table 4).

Thus, a simple calculation formula has been proposed for determining the concentration of potentially acid substances in feedwater of a once-through boiler

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from the measured electric conductivity of H-cationated samples of feedwater and steam. The obtained calculation results differ from measured data by 18% on the average. The proposed method was presented at the International Conference on Instrumentation for Power Plant Chemistry in Zurich (Switzerland) in September 2006 [15] and received positive appraisal from experts.

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