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CARBON NANOWALLS IN FIELD EMISSION CATHODES

The carbon nanowall (CNW) layers were grown from a gas mixture of hydrogen and methane, activated by a DC glow discharge, on Si substrates (Si/CNW layered structure). The second layer of CNW was grown either on the first layer (Si/CNW/CNW structure) or on Ni or NiO films deposited on the first CNW layer (Si/CNW/Ni/CNW and Si/CNW/NiO/CNW structures). The composition and structure of the resulting layered structures were studied using scanning electron microscopy, Raman spectroscopy, and X-ray diffractometry. It was found that annealing of Si/CNW structure in vacuum, growing of the second CNW layer on Si/CNW, as well as deposition of Ni or NiO films prior to the growing of the second CNW layer improve functional properties of field emission cathodes based on the electron-emitting CNW layers.

Keywords: carbon nanowalls, layered structures, electron microscopy, Raman spectroscopy, field emission cathodes.

Carbon materials, including various crystalline (diamond, graphite) and noncrystalline (fullerene, nanotubes, graphene, etc.) ordered substances with unique physicochemical properties are of practical interest. Some carbon materials due to the autoemission property are promising for use as an emitting layer of field emission cathodes (autocathodes). The presence of field emission means a decrease in the electric field strength to 1–10 V/μm, which is required for the onset of field emission of electrons. Autocathodes are used in the development of X-ray tubes, microwave devices, electron guns for exciting lasers, cathodoluminescent lighting devices, flat displays and other devices [1–5]. The most promising for the creation of autocathodes with a low electron emission barrier are the so-called carbon nanowalls (CNW) — layers of a plate-like carbon material with a predominant orientation of the plates perpendicular to the substrate [1–3].

The layers of carbon materials formed by plasma methods, including CNW, are as a rule multiphase layers [6–8]. The structure and concentration of crystalline and X-ray amorphous phases depend on the conditions for carbon materials formation and affect their emission properties. The problems of using CNW in autocathodes are associated with the instability of emission parameters (magnitude and density of the cathode current, as well as the degree of electrical current uniformity over the

cathode area) due to changes in composition and structure during testing and operation [4, 6, 8].

Before being placed into electrovacuum devices and soldered, autocathodes are always preliminarily tested in a vacuum chamber for compatibility with the parameters of the device. In some cases, preliminary tests are carried out to achieve such required parameters as autoemission current and its stability in time. Stability tests can be performed both in the voltage stabilization mode [4] and in the current stabilization mode. In the first case we consider cathode current dependence (decrease) on time at a fixed stabilized voltage, in the second case — voltage dependence (growth) on time at a fixed stabilized current. In both cases, the graphs of the dependencies (hereinafter aging curves) objectively characterize the degradation (aging) of the autocathode, regardless of what causes it.

Storing the tested autocathodes with CNW layers on open air also leads to a deterioration of their emission properties. This is caused by the fact that during vacuum testing on the surface of CNW plates, the layer of adsorbed hydrocarbons is destroyed. This layer normally prevents adsorption of the components of the air mixture (water and nitrogen molecules, etc.) that impair the emission characteristics of the autocathodes [4, 15]. To recover the emission properties of autocathodes that had passed preliminary tests, they are annealing in vacuum or in an inert gas

atmosphere at a temperature of about 720 K [6]. Autocathodes that had not passed preliminary tests (even without vacuum breakdown), as a rule are not further used.

We can assume that autocathodes with a second CNW layer should have better emission properties than autocathodes with only the first CNW layer annealed. Moreover, for autocathodes that had not been preliminarily tested, growing of the second CNW layer on top of the first one should increase the yield ratio for vacuum electronics.

This study researches how vacuum annealing and growing of a second CNW layer affects the emission properties of layered autocathodes based on carbon nanostructures.

Samples used in the research

The CNW samples were grown on Si substrates from a gas mixture of hydrogen (H_2) and methane (CH_4) activated by a DC glow discharge [3, 6]. Before growing CNW, priming carbon centers were created on the substrate. For this purpose, at a temperature of 1020 K, the surface of the substrate was bombarded with H^+ and $C_xH_y^+$ ions (high frequency discharge, 13.56 MHz, 40 W, 20 min), formed in the microwave plasma mixture of hydrogen and methane (8–10 volume % CH_4) at pressure of $6,6 \cdot 10^3$ Pa. Silicon substrates with seed particles were treated in H_2 plasma, after which a CNW layer was grown at a substrate temperature of 800–1300 K and a deposition rate of $6 \mu m/h$. Emission characteristics of the obtained Si/CNW layered structures were tested for 0.5 hour.

Si/CNW structures that were tested and/or stored in the open air for a long time (1 to 3 years) were either annealed in vacuum, or a second CNW layer was grown on their surface under the same conditions as the first layer. In a vacuum (10^{-3} – 10^{-5} Pa), the samples were annealed for 1.5 hours at 720 K (Si/CNW(ann) structure). When the second layer was grown, the crystallites of the first layer acted as seed centers of the second layer. The second layer of CNW was also grown on the surface of the first layer of CNW coated with Ni or NiO.

Ni films were obtained by magnetron sputtering from a Ni target with a direct current in an argon atmosphere (Si/CNW/Ni structure). The conditions for obtaining Ni films are as follows: Ar pressure 1.2–1.5 Pa; discharge power 900 W; substrate temperature 420–570 K; deposition rate $1.5 \mu m/h$. The thickness h of the resulting Ni films was 10, 40, 80, and 160 nm (Si/CNW/Ni^h structures).

NiO films were formed in two stages. At the first stage, a 0.25% solution of $Ni(NO_3)_2$ was applied to the CNW layer in a 50% hydroalcoholic

($H_2O + C_2H_5OH$) mixture at room temperature, followed by heat treatment at 420 K. The heat treatment caused the crystalline hydrate $Ni(NO_3)_2 \cdot 6H_2O$ formed during heat treatment to decompose to NiO form at a temperature of 370–410 K.

$Ni(NO_3)_2$ was deposited at atmospheric pressure either by immersing the substrate with a CNW layer in a solution, followed by heat treatment (the result was the Si/CNW/NiO structure) or by aerosol precipitation (5–10 cycles of 1 minute with heat treatment after each cycle, the result was the Si/CNW/NiO* structure). To generate the aerosol, the Albedo IN-8 (Альбе́до ИИ-8) halogenator was used with an average mass median aerodynamic diameter of the aerosol particles of $3.94 \mu m$.

Research technique

The CNW composition and the layered structures were studied using a Carl Zeiss Supra 40-30-87 scanning electron microscope (SEM), a Rigaku D/MAX-2500/PC X-ray diffractometer (CuK radiation) and a LabRAM HR800 (HORIBA Jobin-Yvon) laser Raman scattering spectrometer (632.8 nm line of He-Ne laser, beam spot diameter $4 \mu m^2$, depth of the analyzed layer $3 \mu m$).

Current-voltage (I-V) characteristics and aging curves that determine the dependence of voltage on time during long-term emission tests at a given current in the regime of constant current stabilization were obtained using a Pw2500_v2_3kV_1a source of stabilized pulsed current produced by SINP MSU and a Spellmann SI30 source of stabilized direct current. The measurements were carried out in diode cells at a pressure of $5 \cdot 10^{-5}$ Pa.

To correctly compare the structures with the second layer of CNW, each sample with the first CNW layer was divided into two parts (A, B) equal in area which were used to co-grow a second layer of CNW in one charge. Part A was used as control, and part B was covered with a Ni or NiO layer, then the A:Si/CNW/CNW and B:Si/CNW/Ni/CNW (or B:Si/CNW/NiO/CNW) structures were compared. In the tests with annealing, the second CNW layer was not grown onto part B, while the first CNW layer was annealed, then the A:Si/CNW/CNW and B:Si/CNW(ann) structures were compared.

The I-V characteristic and the autoemission parameters of the samples were recorded in the pulsed mode of electric current measurement. The aging curves and their parameters were measured in the constant stabilized current mode.

Field emission tests were carried out on samples with a surface having intrinsic conductivity.

During measuring the I-V characteristic in the pulsed mode, a glass plate with a conductive layer of mixed indium-tin oxide (ITO, chemical formula: $(\text{In}_2\text{O}_3)_{0.9}(\text{SnO}_2)_{0.1}$) was used as an anode. The plate was covered with a luminophore layer. This anode completely covered the emitting surface of the sample. When measuring the aging curves in the constant stabilized current mode (10 mA), a water-cooled thick-walled (5 mm) copper anode was used with a polished working surface in the form of a 5×2×2 mm strip located above the 4×2 mm rectangular area of the autocathode.

The gap (Δ) between the surface of the autocathode and the anode was 250 μm when measured in pulsed mode and 125 μm — in direct current mode. The I-V characteristics were plotted in the coordinates (E, J), where $E = U/\Delta$ is the electric field strength in the gap between the anode and the autocathode, $J = I/S$ is the current density, U is the potential difference between the electrodes; I is the current of the autocathode, S is the working area of the autocathode. According to the I-V characteristics, Fowler–Nordheim diagrams were plotted in ($E^{-1}, \ln(J/E^2)$) coordinates.

Composition and structure of CNW

Carbon nanowalls are a porous material formed by curved lamellar (scaly) clusters of X-ray amorphous and crystalline phases of carbon (Fig. 1).

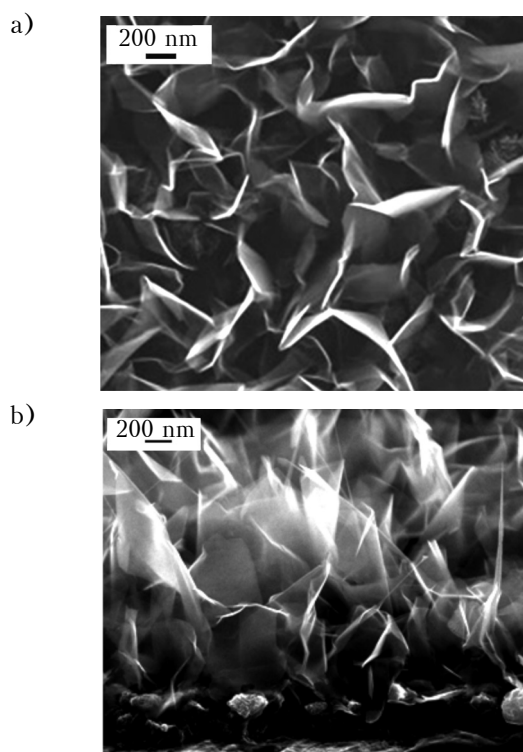


Fig. 1. SEM image of the Si/CNW layer structure: *a* — CNW growth surface; *b* — Si/CNW cleavage

The CNW plates were 3–10 nm thick [3]. Apart from bent carbon plates, the structure of CNW samples also contain rods (plates folded into tubes), nanotubes and equiaxed particles with an average size of 40–50 nm (Fig. 1, *b*).

X-ray diffractometry shows that CNW contains mainly graphite ($P6_3/mmc$ spatial group) and carbyne (hexagonal syngony), as well as phases of diamond ($Fd\bar{3}m$), chaotite ($P6/mmm$ spatial group) and graphite modifications ($R3$ and $P3$ spatial groups) [6, 8]. The thickness of the CNW plates corresponds to the size of the crystallites (X-ray coherent scattering regions, L_{CSR}) equal to 8.5–9.5 nm and calculated from the broadening on X-rays of diffraction peaks of 0002 graphite.

In the Raman spectra of the Si/CNW layer structure, which explicitly reflect the composition and structure of CNW [9, 10], we can observe intense D , G and $2D$ bands, located at the Raman shift $\Delta\nu$, equal to 1330–1343, 1577–1591 and 2660–2673 cm^{-1} , respectively. At the same time, weak bands are fixed at $\Delta\nu$ equal to 233–243, 863–879 and 1081–1167 cm^{-1} (x band); 1612–1627 (D' band); 2449–2482 ($x+D$ band); 2909–2934 ($D+G$ band) and 3221–3248 cm^{-1} ($2D'$ band). (In this study we denote CNW Raman spectral bands as D , G , x , D' , $x+D$, $2D$, $D+G$ и $2D'$ [11–13].)

Fig. 2 (curve 1) shows the Raman spectrum of one of the Si/CNW samples. The values of the intensity ratio of the main CNW Raman spectral bands depending on their formation conditions have a considerable spread: $I_D/I_G = 0.32–2.03$; $I_D/I_{2D} = 0.98–1.23$; $I_D/I_{D+G} = 14.1–17.6$; $I_D/I_{2D} = 13.0–16.1$ [6, 8].

The CNW Raman spectra were compared with similar spectra of highly oriented pyrolytic UPV-1T (УПВ-1Т) graphite (Fig. 2, curve 2). The $2D$ graphite band consists of two components: $2D_1$ and $2D_2$ (Fig. 2, curves 3, 4) with an intensity proportional to the one of the G band. In contrast to the $2D$ graphite band, the $2D$ CNW band is symmetrical, which is characteristic of graphene [9, 13]. The differences in the $2D$ band in the CNW and graphite Raman spectra are caused by a significant curvature of individual regions of the graphite atomic layers $\{0001\}$, which disrupts atomic bonds inside and between the layers.

Depending on the degree of the CNW plates curvature (curvature radius 590–770 nm), the $2D$ band changes shape, which reflects changes in the electronic bands corresponding to the positions of atoms in the lattice. The layers in such a crystallite (CNW plates) form a hexagonal lattice (two-layer stacking of carbon atoms) [1, 6, 14]. If we consider the CNW crystallites as graphite plates, then their size (plate thickness) calculated

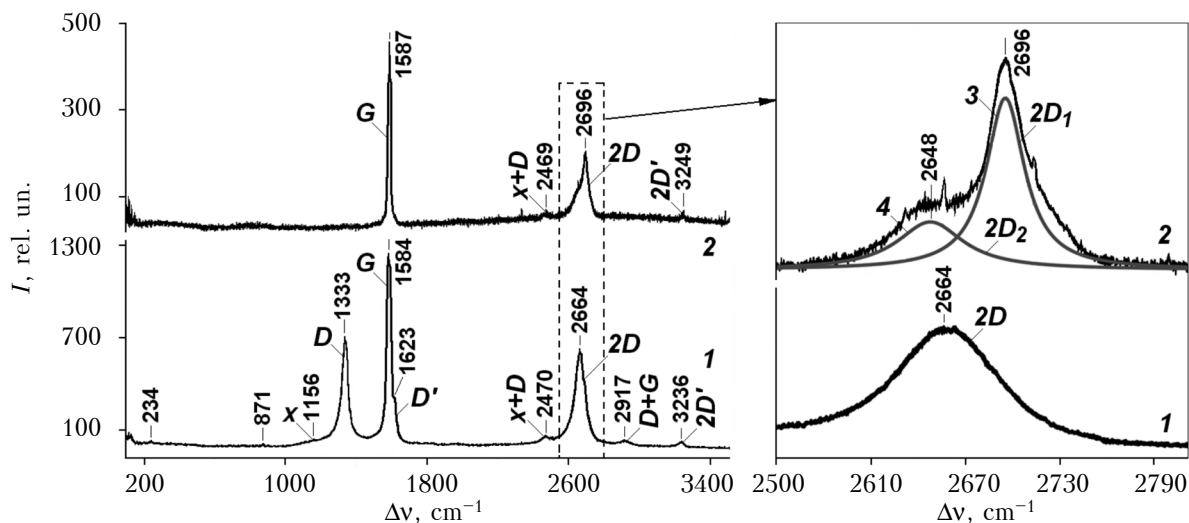


Fig. 2. Raman spectra of Si/CNW (1) and graphite (2) layered structures. Inset graph shows a fragment of the spectra in the $\Delta\nu$ range of 2550–2800 cm^{-1} (3 and 4 are Lorentz distribution functions, which in sum approximate curve 2)

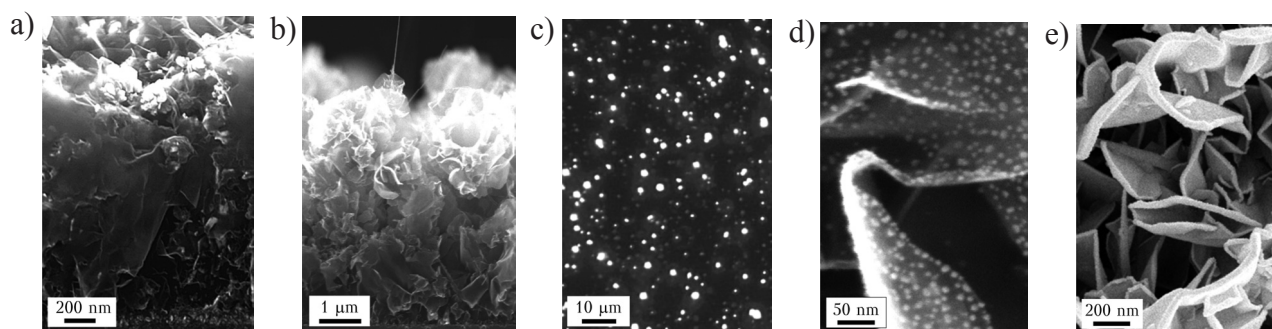


Fig. 3. SEM images of the samples:

a – Si/CNW (cleavage); *b* – Si/CNW/NiO*/CNW (cleavage); *c* – Si/CNW/NiO*; *d* – Si/CNW/Ni¹⁰; *e* – Si/CNW/Ni⁸⁰

from the intensity ratio of the *D* and *G* (I_D/I_G) Raman spectral bands would be $L = 3.3 - 9.9$ nm. The obtained size is close to the values calculated from the X-rays. Taking into account that the interplanar distance of graphite (0001 plane) is 0.335 nm, one can state that there are about 10–30 layers of graphene in a CNW plate.

The maximum height of the first CNW layer starting from the substrate is 2–4 μm , the total height of the first and second layers is approximately 8.5 μm (Fig. 3, *a*, *b*). After the heat treatment of the Si/CNW layered structure covered with $\text{Ni}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ crystal hydrate, NiO crystallites less than 2 μm in size were formed on the surface of the CNW (Fig. 3, *c*). The deposition of a 10 nm thick Ni film on the CNW layer (Si/CNW/Ni¹⁰ structure) by magnetron sputtering resulted in the formation of an islet structure with a cluster size less than 10 nm (Fig. 3, *d*). A continuous Ni^{*h*} film (thickness $h > 40$ nm) was formed mainly on the CNW edges located at a 90°

angle to the Ni particles flux during magnetron sputtering (Fig. 3, *d*). On all other CNW surfaces, the Ni film thickness was by orders of magnitude smaller.

Fig. 4, *a*, *b* shows the globular structure of the CNW layers with a globule diameter of 1.5–2 μm . The second layer of CNW, deposited on a NiO film, has a more dense packing of globules and a larger thickness of the plates (Fig. 4, *d*). Samples A:Si/CNW/CNW and B:Si/CNW/Ni¹⁰/CNW (Fig. 4, *a*, *b*) contain a large number of nanotubes with a diameter of 10–40 nm, while in samples with Ni or NiO films their number does not exceed 1–2 per globule.

The structure of the second CNW layer is characterized by the presence of carbon plates on the crystallites (plates) of the first layer, including multiwall nanotubes (Fig. 4, *c*). In the Si/CNW/Ni¹⁶⁰/CNW and Si/CNW/NiO*/CNW samples, thickened crystallites of carbon plates with rounded edges (not typical) were found (Fig. 4, *d*, *e*). It was discovered that on average

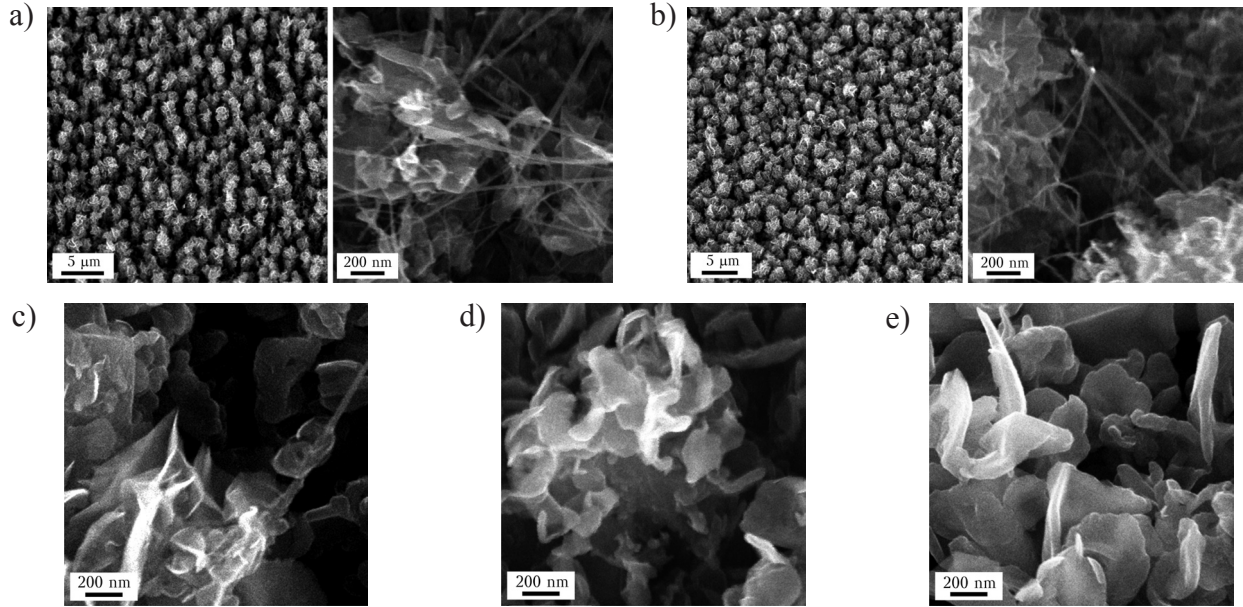


Fig. 4. SEM images of surfaces of layered structures:

$a - A_1\text{:Si/CNW/CNW}$; $b - B_1\text{:Si/CNW/Ni}^{10}\text{/CNW}$;
 $c - B_2\text{:Si/CNW/Ni}^{10}\text{/CNW}$; $d - \text{Si/CNW/Ni}^{160}\text{/CNW}$; $e - \text{Si/CNW/NiO}^*\text{/CNW}$
 (right-hand images in a and b are scaled-up fragments of the surface of the second CNW layer)

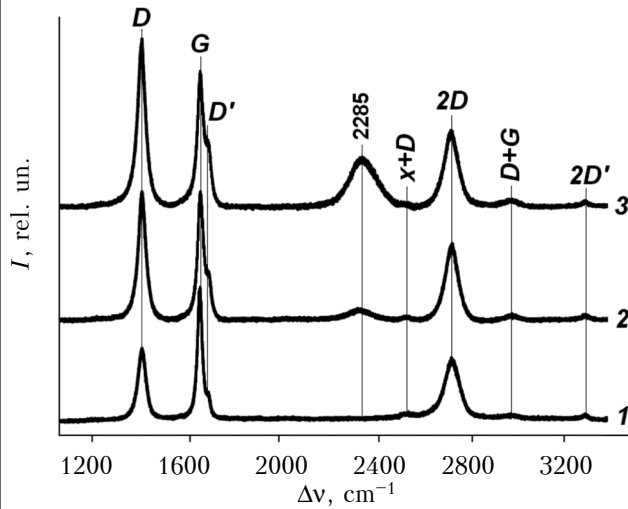


Fig. 5. CNW Raman spectra before and after growing the second CNW layer:

1 – $A\text{:Si/CNW}$; 2 – $A\text{:Si/CNW/CNW}$;
 3 – $B\text{:Si/CNW/NiO}^*\text{/CNW}$

the maximum height of the second CNW layer was 2.4 times greater than that of the first; the globular structure of the second CNW layer became more dense and homogeneous; the number of multiwall nanotubes decreased; on the Raman spectrum of the second CNW layer with a globular structure, a band appeared at $\Delta\nu = 2285 \text{ cm}^{-1}$ (**Fig. 5, Table 1**).

The Raman spectra shown in Fig. 5 are normalized to the intensity of the $2D$ (I_{2D}) band. On the Raman spectra of the $\text{Si/CNW/NiO}^*\text{/CNW}$ structure, the band intensity at $\Delta\nu = 2285 \text{ cm}^{-1}$ increased almost 4-fold. The wide band at $\Delta\nu = 2262 - 2286 \text{ cm}^{-1}$ (broadening of the $\Delta\nu_{1/2} = 120 - 160 \text{ cm}^{-1}$ band) was manifested in the Raman spectra of CNW after annealing at temperatures above 870 K [7]. A similar band was also observed on the Raman spectra of polycrystalline diamond films produced by the microwave discharge method [3].

The crystallite size (L_{CSR}), the number of graphene layers (N), and the I_{2D}/I_D parameter

Table 1

Structural parameters of the upper layer of CNW layered structures

Layered structure	$\Delta\nu_{1/2}, \text{ cm}^{-1}$	I_D	I_G	$I_{D'}$	I_{2D}	I_D/I_G	L_{CSR}	N	$I_{D'}/I_G$	I_{2D}/I_D	I_{2D}/I_G	I_{2285}
Si/CNW	19.4	228.3	388.8	113.4	202	0.6	7.5	22	0.3	0.9	0.5	0
Si/CNW/CNW	17.7	340.6	338.1	130.5	202	1.0	4.4	13	0.4	0.6	0.6	337
Si/CNW/NiO*/CNW	19.9	442.4	356.9	176.5	202	1.2	3.6	10	0.5	0.5	0.6	130.2

of the samples $A:Si/CNW$; $A:Si/CNW/CNW$ and $B:Si/CNW/NiO^*/CNW$ (in this order) decreased, while the I_D/I_G , I_{2D}/I_G parameters and I_{2285} band intensity increased (Table 1).

Emission properties of CNW-based autocathodes

Simultaneously with the I-V measurement, a sequence of images was registered on an luminophore anode screen. Obviously, in this sequence, you should select an image with the worst visual uniformity, which is located in the middle part of the sequence, while at its beginning and at its end there are images that are visually evaluated as homogeneous: almost black ones, obtained at small electric fields, at the beginning, and lighted ones, obtained at large electric fields, at the end (Fig. 6).

To assess the uniformity of images, subjective (visual) criteria are often used. Of the many objective homogeneity criteria based on digital image processing, we chose the simplest criterion based on variation coefficient. We used the intensity (brightness) of a pixel in a gray raster image as the random variable. Uniformity of the image (in percent) was calculated from a sample consisting of all pixels of the image, according to the formula

$$H = 100V - 33,$$

where V is variation coefficient, $V = \sigma/\bar{X}$; σ and $\bar{X}$ are the mean square deviation and average linear deviation in the sample, respectively.

In statistical data processing, the sample is considered homogeneous if $V \leq 0.33$ ($H \leq 0\%$). For the sequence of images obtained from the luminophore anode screen, maximum of H (the worst homogeneity) was taken as homogeneity H^A . As can be seen from Fig. 6, the visual estimation of homogeneity coincides with the maximum at $H^{250} = -16.8\%$. The H^A parameter can be regarded as an estimate of the homogeneity of the cathode current distribution in the working region of the autocathode at the Δ gap.

It is known [4, 15] that the electric field around a pointed conductor is amplified and can be represented as βE_0 , where β is the field gain

near a single emitter and approximately equals to the aspect ratio (height/transverse dimension) of the conductor; E_0 is the ideal electric field strength equal to U/Δ . Assuming that all emission centers have regular geometry (the same sizes and relative position), the dependencies on the Fowler–Nordheim (FN) diagrams are described by the equation of the straight line $y = Bx + C$, where $x = 1/E$, $y = \ln(J/E^2)$. The slope ratio of the straight line B is a value proportional to β , while the density of the emissive centers D_E is proportional to the $\exp(C)$ value (C is the segment cut off by the straight line on the ordinate axis).

For the given films, a linear region can be distinguished on the curves of the FN diagrams [17]. For a relative comparison of the values characterizing emission properties of the autocathodes, it is sufficient to assume that in this linear region only emission centers with regular geometry generate the electrical current, while the contribution of the others to the resulting current is negligible [4, 15]. As emission characteristics of the autocathodes, we considered the following: the autoemission threshold, E_T , is the minimum value of E at which the emission current is registered; the estimation of the aspect number of a single emitter in the regular geometry β ; the estimation of density of emission centers in the regular geometry D_E ; homogeneity H^{250} in a sequence of emission images.

The aging curves (AC) for the structures with the second CNW layer were obtained as the voltage U on time T dependence at a constant current of 10 mA, measured in the current stabilization mode. The 10 mA value of current was chosen due to the capabilities of the equipment available, as well as to current density limitations ($J \approx 0.12$ mA/cm²), at which undesirable vacuum breakdowns are unlikely in the test cell. The aging curves allowed determining the aging rate for 6 hours (V_{6h}), for 3 hours (V_{3h}) and for the last hour (V_{1h}) of tests. A comparison of the aging curves of A and B parts of samples with a second CNW layer was carried out using the Q_{AB} parameter that takes into account the relative position of these curves

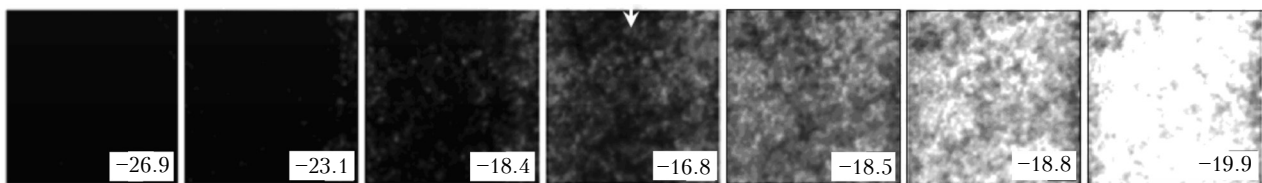


Fig. 6. Sequence of 2x2 mm emission images with different homogeneity H^{250} (the values in % are given in the images)

along the ordinate (U) and the ratio of the areas bounded by these curves and the abscissa axis:

$$Q_{AB} = (S_A / (S_A + S_B)) - 0.5,$$

where S_B is the area under the AC_B for $B:Si/CNW/Ni^h/CNW$, $B:Si/CNW/NiO/CNW$, $B:Si/CNW(ann)$;

S_A is the area under the AC_A of the corresponding B control structures $A:Si/CNW/CNW$ (or $A:Si/CNW$ in the case of annealing).

If the AC_B is located above the AC_A , then $-0.5 < Q_{AB} < 0$ (part B has a higher voltage); if AC_B is located below AC_A , then $0.5 > Q_{AB} > 0$ (part B is less high-voltage). The smaller $|Q_{AB}|$, the closer are the AC_A and AC_B areas. At the same time, the $|Q_{AB}| < 0.5$ inequality is valid.

Fig. 7 presents the I-V characteristic curves and the FN lines described above for the parts of a single sample of layered structure, and their parameters are given in **Table 2**. As can be seen from the presented data, the parts without the second CNW layer ($A:Si/CNW$ и $B:Si/CNW$) are characterized by a high autoemission threshold ($E_T \geq 5.6$ V/ μ m), a large aspect ratio β (which confirms the presence of a large number

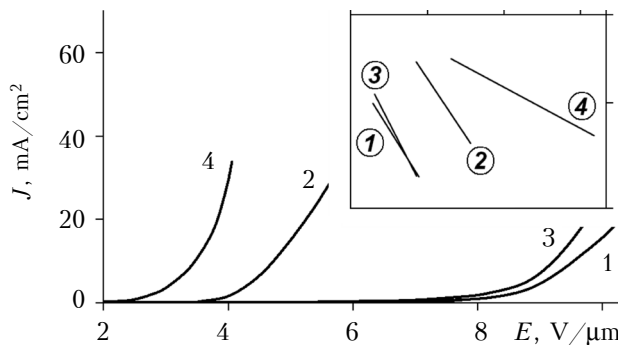


Fig. 7. Example of the I-V characteristics and their linear representations in the FN coordinates (see inset) obtained for the following layered structures:

- 1 – $A:Si/CNW$; 2 – $B:Si/CNW/NiO^*/CNW$;
3 – $B:Si/CNW$; 4 – $A:Si/CNW/CNW$

of multiwall nanotubes on the first CNW layer), a low density of emission centers D_E , and a low homogeneity of field emission images ($H^{250} \geq 0$).

Parts with a second CNW layer (structures $B:Si/CNW/NiO^*/CNW$ and $A:Si/CNW/CNW$) are characterized by a lower autoemission threshold ($E_T \leq 3.6$ V/ μ m), a smaller value of β , a higher D_E density, as well as a better images uniformity ($H^{250} \leq 0$). For a given sample, the part with the $A:Si/CNW/CNW$ structure is characterized by better values of the parameters E_T , β , H^{250} , V_{3h} , V_{1h} than the $B:Si/CNW/NiO^*/CNW$ structure, and the values of the D_E parameter for them are virtually identical.

Examples of the arrangement of the I-V characteristics and the aging curves of the samples with the $Si/CNW/NiO^*/CNW$ (sample 1) and $Si/CNW(ann)$ (sample 2) structures are shown in **Fig. 8, 9**.

As can be seen from Fig. 8, for sample 1, the I-V characteristic of the $B_1:Si/CNW/NiO^*/CNW$

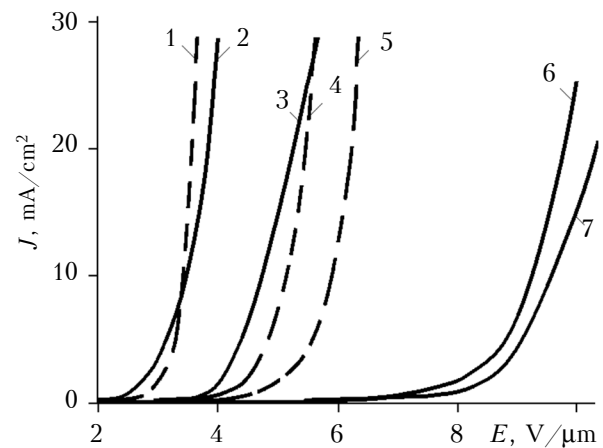


Fig. 8. I-V characteristics of layered structures of two samples:

- Sample 1 (solid lines): 2 – $B_1:Si/CNW/NiO^*/CNW$; 3 – $A_1:Si/CNW/CNW$; 6 – $A_1:Si/CNW$; 7 – $B_1:Si/CNW$;
Sample 2 (dashed lines): 1 – $A_2:Si/CNW/CNW$;
4 – $B_2:Si/CNW(ann)$; 5 – $A_2:Si/CNW$

Table 2

Example of emission characteristics of autocathode parts on layered structures

Structure	Measuring mode							
	Pulse				Continuous			
	E_T , V/ μ m	β	$\ln(D_E)$	H^{250} , %	V_{6h} , V/h	V_{3h} , V/h	V_{1h} , V/h	Q_{AB}
$A:Si/CNW$	6.1	42.7	1.9	39.07	—	—	—	—
$B:Si/CNW/NiO^*/CNW$	3.6	32.9	6.0	−10.52	50	33	10	0.18
$B:Si/CNW$	5.6	47.0	3.0	29.36	—	—	—	—
$A:Si/CNW/CNW$	2.4	21.1	5.9	−7.18	10	0	0	—

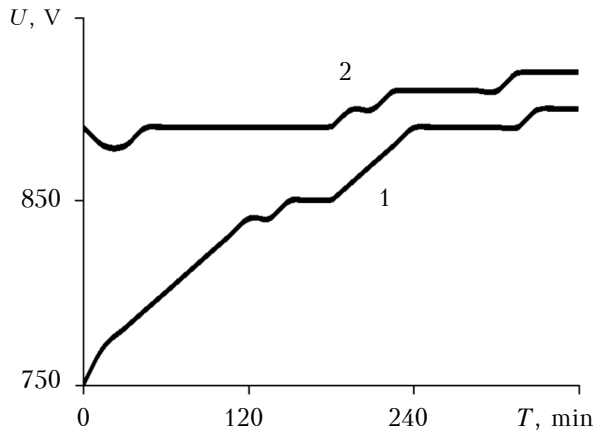


Fig. 9. Aging curves of layered structures of sample 1 for 6 hour tests:

1 – B_1 :Si/CNW/NiO*/CNW; 2 – A_1 :Si/CNW/CNW

structure (curve 2) is shifted to the left by $4 \text{ V}/\mu\text{m}$ relative to the I-V characteristics of the A_1 :Si/CNW and B_1 :Si/CNW structures (curves 6, 7) and by $1.5 \text{ V}/\mu\text{m}$ relative to the I-V characteristic for the A_1 :Si/CNW/CNW (curve 3).

For sample 2, the I-V characteristic of the B_2 :Si/CNW(ann) structure (curve 4) is shifted to the left by $0.8 \text{ V}/\mu\text{m}$ relative to the I-V characteristic of the A_2 :Si/CNW (curve 5) and to the right by $1.8 \text{ V}/\mu\text{m}$ relative to the I-V characteristic of the A_2 :Si/CNW/CNW (curve 1). This indicates that the emission properties of the autocathode

with a second CNW layer deposited on the NiO* oxide layer have improved in comparison with the autocathode with the second CNW layer without the oxide, as well as that the emission properties of the autocathode with the second CNW layer have improved in comparison with an autocathode that had only undergone a restorative annealing of the first CNW layer.

According to the data from Table 2 and Fig. 9, for B_1 :Si/CNW/NiO*/CNW and A_1 :Si/CNW/CNW structures, the Q_{AB} parameter is 0.1 for 6 hours of testing, 0.18 for 3 hours and 0.08 for the last hour. In this case, the AC_B for B_1 :Si/CNW/NiO*/CNW turns out to be less high-voltage than the AC_A for A_1 :Si/CNW/CNW, but it loses to the AC_A in the aging rate. The behavior of the AC_A and AC_B during 6 hour tests ($Q_{AB} = 0.1$) shows their asymptotic convergence at positive values of Q_{AB} , which can be explained by the fact that the second CNW layer on the A_1 and B_1 parts was grown simultaneously during the same charge.

A summary data of the averaged characteristics of the investigated autocathodes is presented in Table 3. The averaging was carried out according to the groups (samples) of the parts of the layered structures indicated in the table. The sample sizes corresponded to the number of parts in each of the groups.

The analysis of the data from Table 3 shows the following. Regardless of which film is used

Table 3

Mean values (M) and standard deviations (S) of the emission characteristics of parts of autocathodes on layered structures

Structure	Sample size	Measurement mode						
		Pulse			Continuous			Q_{AB}
		$E_T, \text{ V}/\mu\text{m}$	β	$\ln(D_E)$	$H^{250}, \%$	$V_{3h}, \text{ V/h}$	$V_{1h}, \text{ V/h}$	
A :Si/CNW; B :Si/CNW	26	M	4.5	60.6	7.2	11.8	—	—
		S	1.3	24.7	2.5	19.9	—	—
A :Si/CNW/CNW	13	M	2.3	37.5	9.2	-2.5	20.0	10.8
		S	1.1	20.5	3.0	17.7	21.3	13.2
B :Si/CNW (ann)	3	M	3.1	53.3	10.2	-14.2	11.1	13.3
		S	0.1	9.1	1.4	10.8	10.2	15.3
B :Si/CNW/NiO/CNW	2	M	3.5	58.2	13.0	-6.0	5.0	15.0
		S	0.3	4.3	0.9	3.6	21.2	21.2
B :Si/CNW/NiO*/CNW	2	M	2.6	33.6	8.5	2.1	16.7	10.0
		S	1.3	1.0	1.7	17.8	23.6	0
B :Si/CNW/Ni ¹⁰ /CNW	3	M	1.9	34.1	10.0	-16.4	14.4	3.3
		S	0.3	11.4	1.7	2.8	5.1	5.8
B :Si/CNW/Ni ⁴⁰ /CNW	1	M	2.2	36.5	6.8	-2.9	26.7	0
B :Si/CNW/Ni ⁸⁰ /CNW	1	M	2.9	35.4	7.4	-7.3	20.0	10.0
B :Si/CNW/Ni ¹⁶⁰ /CNW	1	M	3.2	53.4	13.1	-3.8	13.3	0

(Ni or NiO), the presence of the second CNW layer reduces the average value of the E_T emission threshold to about 2.3 V/ μm as compared to the average value of 4.5 V/ μm for the first CNW layer. The average E_T values for the second CNW layer structures with or without Ni, NiO film differ by approximately 0.1%. On average, the best results for the E_T parameter (1.9 V/ μm) were shown by the Si/CNW/Ni¹⁰/CNW structures, for which the aging rate is also minimal ($V_{1h} \approx 3.3$ V/h) with a positive Q_{AB} value of 0.14. The best V_{1h} results had the Si/CNW/Ni⁴⁰/CNW and Si/CNW/Ni¹⁶⁰/CNW structures ($V_{1h} = 0$) at low E_T values, however for these structures the Q_{AB} parameter turned out to be negative.

Because of the large number of multiwall nanotubes [16], the autocathodes without both the second CNW layer and restorative annealing of the first CNW layer showed the best results for the β parameter (60.5 on average) with a low density of emission centers ($D_E \approx 10^3$, $\ln(D_E) = 7.2$). Then the β parameter was decreasing in structures with a second layer in the following order: Si/CNW/NiO/CNW and Si/CNW/Ni¹⁶⁰/CNW. These same structures with the minimum number of multiwall nanotubes showed the best result for the D_E parameter ($D_E \approx 10^5$, $\ln(D_E) = 13$).

As to the homogeneity of the emission images, the best results on the average were shown by the Si/CNW/Ni¹⁰/CNW structures ($H^{250} = -16.4\%$). Then the H^{250} parameter was getting worse (increasing) in structures in the following order: Si/CNW(ann), Si/CNW/Ni⁸⁰/CNW, Si/CNW/NiO/CNW, Si/CNW/Ni¹⁶⁰/CNW, Si/CNW/Ni⁴⁰/CNW, Si/CNW/NiO*/CNW, and finally the worst were Si/CNW structures.

Conclusion

Thus, the analysis of the I-V characteristics and aging curves of autocathodes based on Si/CNW, Si/CNW/CNW, Si/CNW/Ni/CNW and Si/CNW/NiO/CNW layered structures, as well as the assessment of homogeneity of the images obtained on the luminiferous anode screen allowed establishing the following. On average, the emission properties of autocathodes with a second CNW layer and the presence (optional) of a Ni or NiO film between the CNW layers are better than those of autocathodes that undergo restorative vacuum annealing of the first layer at 720 K. Of all the tested autocathodes with a second layer, the best results on the emission characteristics were shown by the autocathodes with 10 nm thick islet Ni films, which indicates the possibility to use similar cathodes in vacuum electronics. However, for better statistical validity, it is necessary to collect a larger amount of experimental data.

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ВУГЛЕЦЕВІ НАНОСТІНКИ В АВТОЕМІСІЙНИХ КАТОДАХ

Вуглецеві матеріали, що включають різні кристалічні (алмаз, графіт) і некристалічні (фуллерен, нанотрубки, графен та ін.) впорядковані речовини з унікальними фізико-хімічними властивостями, представляють практичний інтерес. Деякі вуглецеві матеріали завдяки властивості автоемісії є перспективними для використання як емітуючого шару автоемісійних катодів (автокатодів). Найбільш перспективними для створення автокатодів з низьким бар'єром емісії електронів вважаються так звані вуглецеві наностінки (C_H) — шари пластинчастого вуглецевого матеріалу з переважним орієнтуванням пластин перпендикулярно підкладці. Роботу присвячено дослідженню впливу відпалу в вакуумі і нарощування другого шару C_H на емісійні властивості шаруватих автокатодів на основі вуглецевих наностінок.

Шари C_H для досліджень вирощували з газової суміші H_2 і CH_4 , активованої тліючим розрядом постійного струму, на підкладках з Si . Перед нарощуванням C_H на підкладках створювалися вуглецеві затравочні центри шляхом обробки поверхні іонами H^+ та C_H^+ . Емісійні характеристики отриманих шаруватих структур Si/C_H контролювали піводинними випробуваннями. Піддані випробуванням та/або тривалому зберіганню на відкритому повітрі шаруваті структури Si/C_H або відпалювали в вакуумі (1,5 години при 720 K), або на їх поверхні нарощували другий шар C_H ($Si/C_H/C_H$) за тих же умов, що і перший. Другий шар C_H нарощували також на поверхні першого шару C_H , вкритого плівкою Ni або NiO (структури $Si/C_H/Ni/CH$ та $Si/C_H/NiO/C_H$). Плівки Ni отримували методом магнетронного розпилення, а плівки NiO — термічною обробкою в розчині $Ni(NO_3)_2$. Максимальна висота першого шару C_H щодо підкладки становила 2–4 мкм, сумарна висота першого і другого шарів — 8,5 мкм. Склад і будову шаруватих структур досліджували з використанням растрової електронної мікроскопії, рентгенівської дифрактометрії і спектрометрії комбінаційного розсіювання світла.

Емісійні властивості представлено у вигляді статистичних оцінок порогу автоемісії, коефіцієнту посилення електричного поля поблизу одиночного автоемітера і щільності емісійних центрів в регулярній геометрії, однорідності емісійних зображень, а також швидкості старіння автоемісійних катодів за тривалих випробувань на постійному стабілізованому струмі. Розроблена методика випробувань дозволила проводити коректне порівняння емісійних характеристик автоемісійних катодів до і після вирощування другого шару C_H , а також після відпалу в вакуумі. Встановлено, що в середньому емісійні властивості автокатодів з другим шаром C_H і плівкою Ni або NiO між шарами C_H є кращими, ніж у автокатодів, які пройшли відновлювальний відпал у вакуумі першого шару за температури 720 K. Відзначено, що з усіх досліджених автокатодів з другим шаром C_H найкращі результати за емісійними характеристиками в середньому показали автокатоди з острівковими плівками Ni товщиною 10 нм. Проведені дослідження підтверджують можливість застосування шаруватих автокатодів на основі вуглецевих наностінок в пристроях вакуумної електроніки.

Ключові слова: вуглецеві наностінки, шаруваті структури, електронна мікроскопія, спектроскопія комбінаційного розсіювання світла, автоемісійні катоди.

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УГЛЕРОДНЫЕ НАНОСТЕНКИ В АВТОЭМИССИОННЫХ КАТОДАХ

Слои углеродных наностенок (C_H) выращивали из газовой смеси водорода и метана, активированной тлеющим разрядом постоянного тока, на подложках из Si (слоистая структура Si/C_H). Второй слой C_H выращивали на первом слое (структура $Si/C_H/C_H$) или на пленках Ni или NiO , осажденных на первом слое C_H (структуры $Si/C_H/Ni/CH$ и $Si/C_H/NiO/C_H$). Методами растровой электронной микроскопии, спектроскопии комбинационного рассеяния света и рентгеновской дифрактометрии исследованы состав и строение полученных слоистых структур. Установлено, что отжиг в вакууме структуры Si/C_H , наращивание на Si/C_H второго слоя C_H , а также нанесение пленок Ni или NiO перед наращиванием второго слоя C_H приводят к улучшению функциональных свойств автоэмиссионных катодов на основе слоев C_H , эмитирующих электроны.

Ключевые слова: углеродные наностенки, слоистые структуры, электронная микроскопия, спектроскопия комбинационного рассеяния света, автоэмиссионные катоды.