

Original Research Article

Clinical and immunological evaluation of application of Ronkoleukin in nonspecific vulvovaginitis at adolescent girls

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Package insert

According to immunological parameters there was found that during the sub-acute there is the secondary immune deficiency and the immunodeficiency is absent during the acute.

During the acute phagocytic function of the local secret is satisfactory, and during the sub-acute against satisfactory absorption function there is a decrease of bacterial growth-inhibitory activity of vaginal secretions, indicating the necessity of correction of phagocytic component of these patients..

The purpose of the research

Clinical and immunological evaluation of Ronkoleukin using depending on the routes of entry in the treatment of non-specific vulvovaginitis among adolescent girls in different variants of the disease state.

Design of there search

Prospective study

Methodology

From 2006 to 2010 years by the assignment of adolescent therapist and upon periodic screening the adolescent girls with a variety of complains for genitalia were examined in the child and adolescent consulting room in Municipal Polyclinic № 11 and «The City Center of Human Reproduction» in Almaty.

124 menstruate adolescent girls from 11 to 18 years, not sexually active, were selected to accomplish the target objective.

I group -20 healthy (control)

II group - 62 with per-acute nonspecific vulvovaginitis

III group - 42 with sub-acute nonspecific vulvovaginitis

In cooperation with the scientific consultant and immunologist, M.D.Professor A. Kurmanova

38 there was developed a dosage schedule of roncoleukine in the complex therapy,
39 based on the results of the analyses of the immune and cytokine status and phagocytic
40 vaginal.system.

41 II group - 62 patients with per-acute nonspecific vulvovaginitis

42 II A– 20 patients with sub-acute nonspecific vulvovaginitis

43 (Standard therapy + roncoleukine 250 000 U/ml vaginal irrigation once a day).

44 II B– 25 adolescent girls with sub-acute

45 (Standard therapy + roncoleukine 250 000 U/ml twice subcutaneously, every other day).

46 II C – 17 adolescent girls with sub-acute (standard therapy)

47 III group - 42 patients with acute nonspecific vulvovaginitis

48 (standard therapy + roncoleukine 250 000 U/ml twice subcutaneously, every other day).

49 Standard therapy of 7-10 days included (1;2):

50 Efficacy of the drug was assessed by patient complaints, inspection, and data of
51 microbiological studies of vaginal discharge, immune state, cytokine status, and local phagocytic
52 system on the 7th and 14th days of treatment.

53 **Keywords:**

54 Nonspecific vulvovaginitis, girl, adolescent, immunity, cytokines, phagocytosis, recombinant
55 human interleukin (roncoleukine).

56 **Introduction**

57 In the structure of gynecological disorders at girls and adolescents vulvovaginitis occupies a
58 leading place (4) .

59 According to scientific studies, the frequency of vulvovaginitis up to 93 per cent for girls,
60 for adolescents it goes up to 53%, while 60% has a recurrent character (5;6;7;8;9;10;11;12).

61 At the present stage there is no doubt that the microbial factor prevails in the pathogenesis of
62 nonspecific vulvovaginitis of the adolescent girls (13;14;15;16;17).

63 Local protection of female genital organs is due to their anatomical and physiological
64 characteristics, the presence of normal microflora, humoral and cellular factors of immunity.

65 Common infectious disease accompanied by decrease in immunity, as well as the hormonal
66 diseasebreak the qualitative and quantitative composition of the vaginalmicroflora that facilitates
67 the invasion of pathogenic microorganisms and can lead to the development of inflammatory
68 processes caused by opportunistic pathogenic bacterium (18).

69 The opportunistic pathogens, involved in the inflammatory process, do not contain highly
70 toxic poisons, but they are dangerous for hypernormal promoting of inflammation mediators of
71 micromycetia microorganism.

72 The practical significance of vulvovaginitis is defined by the fact that they lead to the
73 formation of synechia of the labia, genital infection, disruption of menstrual function, which can
74 cause serious disorders of reproductive function in the future (19;20).

75 The necessity of the clinical relevance of the study of the immunological aspects of
76 vulvovaginitis is that each immune imbalance increases the probability of the progression of the
77 disease, favours the development of complications, the allergization of the organism and the
78 chronization of the process.

79 The evaluation of the general and of the local immunity at vulvovaginitis is one of the
80 challenging issues, taking into account the essential role of the immune-pathological mechanisms at
81 the chronization and at the retrocessions of the process.

82 Opportunistic microorganisms which take part in an inflammatory process do not contain any
83 high-toxic poisons, but they are dangerous due to their excessive activation of the inflammatory
84 mediators of the microorganism.

85 At present time the diagnostic significance of the evaluation of the cytokines concentration
86 level consists in the statement of the very fact of its increasing or reducing during a concrete
87 disease, at that it is appropriate to detect the concentration of pro-inflammatory as well as of anti-
88 inflammatory cytokines in the dynamic of the disease development to be able to evaluate the
89 severity and to make prognosis of the course of disease (21;22;23).

90 In this connection the search of the new immunological drugs in the complex therapy of
91 vulvovaginitis allowed to examine the use of Interleukin-2 human recombinant (Ronkoleukin) in
92 the complex therapy of nonspecific vulvovaginitis at adolescent girls with different kinds of
93 diseases and introduction of the drug.

94 For the diagnosis and treatment of vulvovaginitis it is not possible to rely solely only on a
95 visual assessment of the genitalia and the discharge from the vagina; it is also necessary to provide
96 the microbiological research and study of the immune status, local cytokine, and phagocytic system.

97 **Results and Discussion**

The main complaints and clinical manifestations of nonspecific vulvovaginitis adolescent girls (M ± m)

Indices	Group I	Group II A	Group II B	Group II C	Group III
Complaints					
Discharge	-	3(15,0±3,8)	6(24,0±4,7)	7(41,2±3,9)	19 (45,2±6,0)
Redness	-	11(55,0±7,0)	15(60,0±7,1)	11(64,7±4,8)	30 (71,4±7,1)
Vulvovaginal pruritus	-	7(35,0±5,7)	12(48,0±6,5)	12(70,6±5,0)	24 (57,1±6,6)
Heat	-	3(15,0±3,8)	11(44,0±6,3)	11(64,7±4,8)	22 (52,4±6,4)
Vulvovaginal pain	-	3(15,0±3,8)	8(32,0±5,4)	10(58,8±4,6)	18 (42,9±5,9)
Painfulurination	-	1(5,0±2,2)	3(12,0±3,4)	2(11,8±2,2)	5 (11,9±3,4)
Examination					
Hyperaemia of the labia majoria	-	6(30,0±5,3)	7(28,0±5,1)	5(29,4±3,4)	23(54,8±6,5)
Hyperaemia of the vestibule of vagina	-	11(55,0±7,0)	22(88,0±8,3)	17(100,0±5,8)	39 (92,9±7,5)
Hyperaemiaofthe urethral meatus	-	13(65,0±7,5)	9(36,0±5,7)	7(41,2±3,9)	16 (38,1±5,7)
Oedemataofthelabia majoria	-	4(20,0±4,4)	4(16,0±3,9)	4(23,5±3,0)	13(30,9±5,2)
Oedemataof the vestibular mucous membrane	-	8(40,0±6,1)	11(44,0±6,3)	11(64,7±4,8)	30 (71,4±7,1)
Oedemataoftheurethral meatus	-	4(20,0±4,4)	6(24,0±4,7)	5(29,4±3,4)	11 (26,2±4,8)
Hyperaemiaofthe perineum	-	6(30,0±5,3)	8(32,0±5,4)	6(35,3±3,7)	16 (38,1±5,7)
Pathologicdischargefrom the genital tracts					
The value of discharge					
Low	6(30,0±5,3)	14(70,0±7,8)	18(72,0±7,7)	12(70,6±5,0)	-
Moderate	14(70,0±7,8)	6(30,0±5,3)	5(20,0±4,4)	3(17,6±2,6)	25 (59,5±6,7)
Plethorical	-	-	2(8,0±2,8)	2(11,8±2,2)	17 (40,5±5,8)
Colour of discharge					
Off-white	20(100,0±8,9)	-	-	-	-
Off-white-xanthic	-	13(65,0±7,5)	8(32,0±5,4)	6(35,3±3,7)	8 (19,0±4,2)
Pyromucous (xanthic)	-	7(35,0±5,7)	7(28,0±5,1)	8(47,1±4,2)	17 (40,5±5,8)
Pyogenic (xanthic- greenish)	-	-	10(40,0±6,0)	3(17,6±2,6)	17 (40,5±5,8)

101 The clinical picture of sub-acute and acute nonspecific vulvovaginitis was presented by the
102 symptoms of vulvitis and urethritis.

103 The most common complaints in the acute period were redness in the genital area, asymptomatic,
104 burning and discharge from the genital tract, and on examination there were frequently identified
105 hyperemia and swelling of the prepuce.

106 In the sub-acute the main complaints were redness, asymptomatic, and on examination there was
107 seen hyperemia of the urethral meatus and prepuce.

108 In the clinical picture in acute course of the disease the moderate discharge from the
109 reproductive tracts was more evident, and the sub-acute discharge was low.

110 It drew attention to the fact that the patients in both groups had the discharge of
111 pathological character from the genital tract during several weeks, and only when the asymptomatic,
112 burning, discomfort had appeared in the genitals, they turned to a specialist.

113 On examination of the patients there was noticed the discrepancy in the intensity of
114 inflammatory symptoms in the sub-acute.

115 Objectively sub-acute vulvovaginitis was characterized by isolated hyperemia of internal
116 surfaces of the labia and labia majora, prepuce and injected vessels of the vulva, that indicated
117 the long-term inflammation.

118 The absence of subjective sensations, the limit of inflammation by inflammation of glands
119 and urethral lacunae, hyperemia of stasis and swelling of prepuce confirm that vulvovaginitis is
120 sub-acute in nature.

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Table 2

Immune status and phagocytic function of vaginal discharge in the examined before treatment ($M \pm m$)

Indices	Group I healthy patients (n=20)	Group II with sub-acute form (n=62)	Group III with acute form (n=42)
CD3+(%)	64,5 ± 0,3	59,32±1,9*	69,2 ± 6,4
CD4+(%)	53,9 ± 1,0	49,4 ± 5,1	53,0 ± 8,1
CD8+(%)	23,2 ± 0,7	20,8 ± 2,7	22,0 ± 5,3
CD20+(%)	15,3 ± 0,7	12,4 ± 0,6*	14,9 ± 4,1
ИНФγ (pg/ml)	11,02 ± 0,74	12,49 ± 6,49	17,32 ± 12,2
ΦНОα (pg/ml)	4,48 ± 0,57	3,99 ± 0,70	4,14 ± 1,7
ИЛ-6 (pg/ml)	22,62 ± 1,80	38,24 ± 3,22	36,5 ± 26,7
Phagocytic index, idiopathic, %	22,4±5,4	35,8±5,8	47,2±15,2
Phagocytic number, idiopathic	8,0±0,2	5,9±1,4	6,7±1,5
Phagocytic index, induced by pyrogenal, %	36,3±4,7	48,8±8,4	50,0±10,0
Phagocytic number, induced by pyrogenal	10,0±0,3	6,6±1,2*	8,7±0,7
НСТ idiopathic, %	24,5±13,0	10,1±4,12	14,2±6,2
НСТ stimulated by pyrogenal, %	44,1±5,7	12,4±4,9*	32,3±2,3***
* The disparity is accurate at $P \leq 0.05$ between the Groups I and II . ** The disparity is accurate at $P \leq 0.05$ between the Groups I and III. *** The disparity is accurate at $P \leq 0.05$ between the Groups II and III.			

125 It was found that the patients with sub-acute disease had a significant downstream of the
 126 relative content of Cd3+, indicating the oppression of their differentiation. There was also
 127 recorded a significant reduction of Cd20+. With regard to the value of CD4+ and CD8+
 128 compared to the control group there was a tendency to decrease.

129 Study of immunological parameters with acute disease has shown that there are no reliable
 130 differences in comparison with the control group, although there was watched a trend towards
 131 downstream of CD4 +, Cd8 + and Cd20 + cells and the trend of increase in Cd3 +, due to the
 132 dispersion of individual value.

133 On the basis of the revealed violations by the parameters of the immune status, it can be
 134 assumed that in the sub-acute there is a recurrent immunodeficiency, and there is no deficiency
 135 during acute.

136 In all studied groups the average cytokine profile's value in peripheral blood did not differ
 137 significantly from the average value of the control group in connection with a wide range of
 138 value.

139 Therefore, the further analysis of cytokine output was conducted by the percentage of
 140 occurrence of elevated and high value.

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Table 3

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The frequency of the increased cytokine concentration in the examined patients(M ± m)

IndicesPg/ml	Group I	Group II	Group III
γ-Interferon	0	15,0±8,0*	23,8±6,6**
TNF α	10,0±6,7	20,0±8,9	21,4±6,3
IL-6	0	20,0±8,9*	14,3±5,4**
* The disparity is accurate at P≤0.05 between the 1 st and the 2 nd groups. ** The disparity is accurate at P≤0.05 between the 1 st and the 3 rd groups.			

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145 In the I group the high output of cytokines IFN-γ, TNF-α and IL-6 were not observed. The
 146 increased output of IFN-γ was recorded at 10% of examined patients.

147 In the second group, there was an increase in output of IFN-γ at 15% of the patients, with
 148 adequately high content (in 4 times), recorded only at one patient (5%). The elevated level of
 149 TNF-α output was registered at 20% of the patients; herewith there was no any highest production
 150 (in 4 or more times). At the same time there was the increased output of IL-6 at 20%. It should
 151 be noted that high (in 5-13 times) output of IL-6 was at 15% (3/20). When comparing the rates of
 152 occurrence of high content with I group, the increased output of IFN-γ and IL-6 was more often

153 registered in the 2nd group. However, adequate cytokine explosion with the activation of
154 decreasing immunity was observed only in 1/6 cases.

155 In the III group the enhancement of IFN- γ output was observed at 23.8% of the surveyed.
156 Herewith, the adequately high content (in 4 times) was recorded at 3/42 (7.1%) of the surveyed.
157 An elevated output of TNF- α was at 21.4% of surveyed, with the highest (in 4 or more times)
158 was not a product of one. The increased production of IL-6 was observed at 14.3% of the
159 patients, while high production was recorded at 4.7% of the patients. In comparing values of
160 occurrence of high content with group I, increased production of IFN- γ and IL-6 was
161 significantly recorded in group III ($p \leq 0.05$).

162 Consequently, nearly 1.6 times more girls and adolescents with acute nonspecific
163 vulvovaginitis have the increased output of proinflammatory cytokine IFN- γ than with sub-acute,
164 indicating the direct dependence of the activity level of proinflammatory cytokine from the
165 clinical course of the inflammatory process. Although, its high output does not exist in both
166 variants of the disease.

167 The data obtained give the evidence of the oppression of anticontagious immunity both
168 cellular and humoral, as TNF- α has the co-stimulant function for T-cell activation and
169 activation of mononuclear phagocytes, also it helps the antibody formation by B-cells, and IL-6 is
170 responsible for the specificity and the adequacy of immune reactions. This fact is due, first of all,
171 with the presence of different pathological changes of immune system.

172 In the group I the absorptive function of the vaginal discharge complies with the similar
173 parameters of healthy women of reproductive age, while digestible function of the vaginal
174 discharge is more evident in adolescence.

175 The phagocytic function of vaginal discharge at the adolescent girls of the II group in
176 comparison with I group was observed in some improvement of the average absorption capacity
177 of vaginal discharge, but did not differ significantly. Herewith, the digestible ability of the
178 vaginal discharge was decreased, resulting in decreasing the NBTR-test value in the stimulated
179 version (NBTR-test stim. - $12.4 \pm 4.9\%$, $p \leq 0.05$). The spontaneous value of NBTR-test was
180 below the equivalent control values, but not significantly different, due to the wide scatter in
181 values.

182 In the III group all values both the absorption and digestive functions of vaginal discharge
183 did not differ significantly from those in the I group, however, there has been a tendency to
184 increase of the value both spontaneous and stimulated phagocyte index, and to the downstream of
185 the phagocyte number

186 By comparison of the sub-acute value there was no substantial difference in absorption
187 capacity, but was noted a significant reduction of bactericidal activity in sub-acute in comparison
188 with acute.

189 Thus, in the acute with nonspecific vulvovaginitis at young girls and adolescents
190 the phagocytic function of the local discharge is satisfactory; and during the sub-acute against
191 satisfactory absorption function there is observed the decrease of bactericidal activity of the
192 vaginal discharge, which demonstrates the need for the phagocyte correction of these patients.

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Dynamic of complaints and of clinical signs in the Group IIA (M ± m)

Analysed indices	until the treatment	on the 7 th day	on the 14 th day
Complaints			
Discharge	3(15.0±3.8%)	7(35±5.7%)*	3(15±3.8%)
Redness	11(55.0±7.0%)	8(40±6.1%)*	2(10±3.1%)
Vulvovaginal pruritus	7(35.0±5.7%)	5(25±4.9%)	2(10±3.1%)
Heat	3(15.0±3.8%)	1(5±2.2%)*	-
Vulvovaginal pain	3(15.0±3.8%)	2(10±3.1%)	-
Painfulurination	1(5.0±2.2%)	-*	-
Examination			
Hyperaemia of the labia majoria	6(30.0±5.3%)	3(15±3.8%)*	1(5±2.2%)**
Hyperaemia of the vestibule of vagina	11(55.0±7.0%)	5(25±4.9%)*	3(15±3.8%)
Hyperaemiaofthe urethral meatus	13(65.0±7.5%)	-*	-
Oedemataofthelabia majoria	4(20.0,0±4.4%)	-*	-
Oedemataof the vestibular mucous membrane	8(40.0±6.1%)	2(10±3.1%)*	-
Oedemataoftheurethral meatus	4(20.0±4.4%)	-*	-
Hyperaemiaofthe perineum	6(30.0±5.3%)	3(15±3.8%)*	2(10±3.1%)
Pathologicdischargefrom the genital tracts			
The value of discharge	14(70.0±7.8%)	11(55±7.0%)	5(25±4.9%)
Low	6(30.0±5.3%)	9(45±6.4%)*	15(75±8.0%)
Moderate	-	-	-
Plethorical			
Colour of discharge	-	7(35±5.7%)*	11(55±7.0%)
Off-white	13(65.0±7.5%)	9(45±6.4%)*	9(45±6.4%)
Off-white–xanthic	7(35.0±5.7%)	4(20±4.4%)*	-
Pyromucous (xanthic)	-	-	-
* The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 7 th day			
** The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 14 th day			

Dynamic of the immunological indices and of the indices of the local phagocytic system, Group II A(M ± m)

Indices	Control group (n=20)	Group IIA		
		until the treatment	on the 7 th day	on the 14 th day
CD3+ (%)	64.5 ± 0.3	59.32±1.9*	61.3 ± 1.6**	64.6 ± 1.4 #
CD4+ (%)	53.9 ± 1.0	49.4 ±5.1	49.6 ±1.6	49.5 ±2.0
CD8+ (%)	23.2 ± 0.7	20.8 ±2.7	21.4 ±1.2	21.6 ±1.1
CD20+ (%)	15.3 ± 0.7	12.4 ± 0.6*	13.2 ± 1.1	13.6 ±0.86
IL-6 (pg/ml)	22.6 ± 1.8	38.2±3.22	23.9 ± 0.87	22.3 ± 1.3
Interferon (pg/ml)	11.02 ± 0.74	12.5 ±6.5	12.04 ±0.37	12.1 ±0.23
TNF (pg/ml)	4.48 ± 0.6	3.9 ±0.7	4.57±0.36	4.4 ±0.2
* The disparity is accurate at P≤0.05 between the examination and until treatment				
** The disparity is accurate at P≤0.05 between the examination and on the 7 th day				
*** The disparity is accurate at P≤0.05 between the examination and on the 14 th day				
# The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 14 th day				
Phagocytic index, idiopathic, %	22.4±5.4	35.8±5.8	-	60.2±11.4**
Phagocytic number, idiopathic	8.0±0.2	5.9±1.4	-	7.9±1.6***
Phagocytic index, induced by pyrogenal, %	36.3±4.7	48.8±8.4	-	65.8±9.8**,***
Phagocytic number, induced by pyrogenal	10.0±0.3	6.6±1.2*	-	9.8±0.7***
Phagocytic index, induced by ronkoleukine, %	44.5±4.7	92.6±3.3*	-	99.3±0.4**
Phagocytic number, induced by ronkoleukine	10.0±0.3	14.0±1.9*	-	16.3±1.07**
HCT idiopathic, %	24.5±13.0	10.1±4.12	-	20.0±2.7
HCT stimulated by pyrogenal, %	44.1±5.7	12.4±4.9*	-	22.3±6.2**
HCT stimulated by ronkoleukine, %	44.1±5.7	35.5±7.25	-	52.0±4.6
* The disparity is accurate at P≤0.05 between the control and until the therapy.				
** The disparity is accurate at P≤0.05 between the control and after the therapy.				
*** The disparity is accurate at P≤0.05 between until and after the therapy.				

226 Before the treatment in cytokine composition the averages did not differ significantly from
227 the control group average.

228 The same pattern for mean values was observed on the 7th and 14th days after treatment. But
229 when comparing the percentage of occurrence of high levels before treatment there was observed
230 the high output of IFN- γ at 15% of the patients, output of TNF- α and IL-6 - at 20%, of the patients,
231 output of TNF- α at 25% of the patients increased on the 7th day after treatment, and increasing output
232 of cytokines was observed on the 14th day. These changes may indicate an inflammatory process
233 remitting, preceded by activation of exogenous cytokine.

234 Before the treatment in the group II (A) in comparison with the group I the average absorption
235 capacity of vaginal discharge was not different; the digestion ability of vaginal discharge was
236 reduced, that resulted in reduction of the NBTR-test value in the Pirogenal stimulated version. In
237 loading tests with Ronkoleukin in vitro, there was a sharp increase of phagocyte index, phagocyte
238 number and NBTR-test, which indicated a positive response of vaginal mucus.

239 Through 7 days after the vaginal irrigation by Ronkoleukin a local secret reaction was the
240 following: the value of spontaneous and induced by Pirogenal the phagocyte index, phagocyte
241 number, and NBTR-test sharply increased.

242 In comparison with the individual changes, it was observed a significant increase in
243 spontaneous phagocytic number to 139.2%, phagocyte number induced by Ronkoleukin to 113.5%,
244 spontaneous NBTR-test, induced by Pirogenal and Ronkoleukin to 353.3%, 220 percent and 161.9%
245 respectively.

246 Thus, in the sub-acute the local application of Roncoleukin leads to the increased absorption
247 and oxygen-dependent bactericidal ability of vaginal secretion, which contributes to the rapid
248 involution of clinical aspects.

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Table 6

Dynamic of complaints and of clinical signs in the Group IIB (M ± m)

Analysed indices	until the treatment	on the 7 th day	on the 14 th day
Complaints			
Discharge	12(48.0±6.5%)	7(28.0±5.1%)*	2(8.0±2.8%)**
Redness	19(76.0±7.8%)	8(32.0±5.4%)*	_***
Vulvovaginal pruritus	12(48.0±6.5%)	4(16.0±3.9%)*	2(8.0±2.8%)**
Heat	11(44.0±6.3%)	_*	_***
Vulvovaginal pain	8(32.0±5.4%)	_*	_***
Painfulurination	3(12.0±3.4%)	_*	_***
Examination			
Hyperaemia of the labia majoria	14(56.0±6.9%)	4(16.0±3.9%)*	1(4.0±2.0%)**
Hyperaemia of the vestibule of vagina	22(88.0±8.3%)	9(36.0±5.7%)*	_***
Hyperaemia of the urethral meatus	9(36.0±5.7%)	_*	_***
Oedemata of the labia majoria	8(32.0±5.4%)	_*	_***
Oedemata of the vestibular mucous membrane	17(68.0±7.5%)	2(8.0±2.8%)*	_***
Oedemata of the urethral meatus	6(24.0±4.7%)	_*	_***
Hyperaemia of the perineum	11(44.0±6.3%)	2(8.0±2.8%)*	2(8.0±2.8%)**
Pathologic discharge from the genital tracts			
The value of discharge	-	18(72.0±7.8%)*	9(36.0±5.7%)**
Low	14(56.0±6.9%)	7(28.0±5.1%)*	16(64.0±7.3%)**
Moderate	11(44.0±6.3%)	_*	_***
Plethorical			
Colour of discharge	-	6(24.0±4.7%)*	14(56.0±6.9%)
Off-white	8(32.0±5.4%)	11(44.0±6.3%)	11(44.0±6.3%)**
Off-white-xanthic	7(28.0±5.1%)	8(32.0±5.4%)	_***
Pyromucous (xanthic)	10(40.0±6.0%)	_*	_***
* The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 7 th day			
** The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 14 th day			

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Table 7

Dynamic of the lymphocytes subpopulations rates in the setting of the treatment, the Group II B(M ± m)

Indices	Group I A	Group II B		
		until the treatment		until the treatment
CD3+ (%)	64,5 ± 0,3	59,32±1,9*	61,3 ± 1,6**	64,6 ± 1,4 #
CD4+ (%)	53,9 ± 1,0	49,4 ± 5,1	49,6 ± 1,6	49,5 ± 2,0
CD8+ (%)	23,2 ± 0,7	20,8 ± 2,7	21,4 ± 1,2	21,6 ± 1,1
CD20+ (%)	15,3 ± 0,7	12,4 ± 0,6*	13,2 ± 1,1	13,6 ± 0,86
IL-6 (pg/ml)	22,6 ± 1,8	20,75±1,54	24,17±0,62	20,64±1,54
Interferon (pg/ml)	11,02 ± 0,74	10,44±0,57	11,50±0,28	12,01±0,23
TNF (pg/ml)	4,48 ± 0,6	4,88±0,29	5,07±0,22	4,64±0,20
* The disparity is accurate at $P \leq 0.05$ between the examination and until treatment				
** The disparity is accurate at $P \leq 0.05$ between the examination and on the 7 th day				
# The disparity is accurate at $P \leq 0.05$ between the beginning of the treatment and on the 14 th day				

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255 Before treatment in the group II B there was record of decrease of CD3 + and CD20 +
256 lymphocytes in the value of lymphocyte subpopulation composition. On the 7th day rate of CD3 +
257 increased slightly, but continued to be significantly reduced, and the content of CD20 + turned up to
258 stated value.

259 An increase of CD3+lymphocytes to the level of stated value was recorded on the 14th day,
260 and by comparing value before and after the treatment there was registered a significant
261 improvement of this indicator.

262 Thus, the appointment of Roncoleukin subcutaneously twice alternate days in the sub-acute
263 leads to the normalization of main value of lymphocyte subpopulation composition.

Dynamic of complaints and of clinical signs in the Group II C(M ± m)

Analysed indices	until the treatment	on the 7 th day	on the 14 th day
Complaints			
Discharge	7(41,2±3,9)	7(41,1±6,1)	-**
Redness	11(64,7±4,8)	3(17,6±2,6)*	-**
Vulvovaginal pruritus	12(70,5±5,0)	4(23,5±3,0)*	1 (5,9±1,5)**
Heat	11(64,7±4,8)	_*	-**
Vulvovaginal pain	10(58,8±4,6)	_*	-**
Painfulurination	2(11,8±2,2)	_*	-**
Examination			
Hyperaemia of the labia majoria	9(52,9±4,4)	_*	-**
Hyperaemia of the vestibule of vagina	17(100±5,8)	2(11,7±3,3)*	-**
Hyperaemia of the urethral meatus	7(41,2±3,9)	_*	-**
Oedemata of the labia majoria	5(29,4±3,4)	_*	-**
Oedemata of the vestibular mucous membrane	13(76,5±5,2)	_*	-**
Oedemata of the urethral meatus	5(29,4±3,4)	_*	-**
Hyperaemia of the perineum	5(29,4±3,4)	2(11,8±2,2)*	-**
Pathologic discharge from the genital tracts			
The value of discharge	-	9(52,9±4,4)*	7(41,2±3,9)**
Low	11(64,7±4,8)	8(47,1±4,2)*	10(58,8±4,6)
Moderate	6(35,3±3,7)	_*	-**
Plethorical			
Colour of discharge	-	7(41,2±3,9)*	15(88,2±5,5)**
Off-white	-	7(41,2±3,9)*	2(11,2±2,2)**
Off-white-xanthic	10(58,8±4,6)	3(17,6±2,6)*	-**
Pyromucous (xanthic)	7(41,2±3,9)	_*	-**
* The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 7 th day			
** The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 14 th day			

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Table 9

Dynamic of the immunological indices, Group II C(M ± m)

Indices	Group I	Group II C		
		until the treatment	on the 7 th day	on the 14 th day
CD3+(%)	64,5 ± 0,29	63,19±0,80	62.48±0,94	66,25±0,71
CD4+(%)	53,95 ± 0,9	47.13±2,43	50.25±1,25	46,18±1,06
CD8+(%)	23,2 ± 0,7	19.94±1.00	22.78±0,84	24,83±1,45
CD20+(%)	15,3 ± 0,7	11,52±0,55	16.61±1,04	16.06±0,79
IL-6 (pg/ml)	22,6 ± 1,8	19,77±1,46	26,25±0,65	24,22±0,49
Interferon(pg/ml)	11,02 ± 0,74	10.35±0,45	12.02±0,33	12.68±0,71
TNF(pg/ml)	4,48 ± 0,57	4.32±0,30	5.55±0,13	3.95±0,15

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Table 10

Dynamic of complaints and of clinical signs in theGroupIII(M ± m)

Analysed indices	until the treatment	on the 7 th day	on the 14 th day
Complaints			
Discharge	19(45.2±6.1%)	14(33.3±5.4%)	2(4.8±2.2%)**
Redness	30(71.4±7.1%)	11(26.2±4.8%)*	-**
Vulvovaginal pruritus	24(57.1±6.6%)	8(19.0±4.2%)*	3(7.1±2.6%)**
Heat	22(52.4±6.4%)	-*	-**
Vulvovaginal pain	18(42.9±5.9%)	-*	-**
Painfulurination	5 (11.9±3.4%)	-*	-**
Examination			
Hyperaemia of the labia majoria	23(54.8±6.5%)	4(9.5±3.0%)*	1(2.4±1.5%)**
Hyperaemia of the vestibule of vagina	39(92.9±7.5%)	11(26.2±4.8%)*	-**
Hyperaemiaofthe urethral meatus	16(38.1±5.7%)	-*	-**

Oedemataofthelabia majoria	13(31.0±5.3%)	_*	_**
Oedemataof the vestibular mucous membrane	30(71.4±7.1%)	2(4.8±2.2%)*	_**
Oedemataoftheurethral meatus	11(26.2±4.8%)	_*	_**
Hyperaemiaofthe perineum	16(38.1±5.7%)	4 (49.5±3.0%)*	2 (4.8±2.2%)**
Pathologicdischargefrom the genital tracts			
The value of discharge	-	27 (64.3±6.9%)*	16(38.1±5.7%)**
Low	25(59.5±6.7%)	15((35.7±5.5%)*	26(61.9±6.8%)
Moderate	17(40.5±5.8%)	-	_**
Plethorical			
Colour of discharge	-	13(31.0±5.2%)*	29(69.0±7.0%)**
Off-white	8 (19.0±4.2%)	18 (42.9±5.9%)*	13(31.0±5.2%)**
Off-white–xanthic	17(40.5±5.8%)	11 (26.2±4.8%)	-
Pyromucous (xanthic)	17(40.5±5.8%)	_*	_**
* The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 7 th day			
** The disparity is accurate at P≤0.05 between the beginning of the treatment and on the 14 th day			

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Table 11

Dynamic of the immunological indices, Group III(M ± m)

Indices	Group I	Group III		
		until treatment	on the 7 th day	on the 14 th day
CD3+(%)	64,5 ± 0,29	69,2 ± 6,4	62,5 ± 1,6	66,8 ± 1,1
CD4+(%)	53,95 ± 0,9	53,0 ± 8,1	50,6 ± 1,6	49,1 ± 2,7
CD8+(%)	23,2 ± 0,7	22,0 ± 5,3	22,3 ± 1,1	24,04 ± 1,2
CD20+(%)	15,3 ± 0,7	14,9 ± 4,1	14,6 ± 1,9	15,3 ± 0,8
IL-6 (pg/ml)	22,6 ± 1,8	36,5 ± 26,7	25,3 ± 1,1	22,5 ± 1,9
Interferon(pg/ml)	11,02 ± 0,74	17,32 ± 12,2	11,8 ± 0,4	12,3 ± 0,5
TNF(pg/ml)	4,48 ± 0,57	4,14 ± 1,7	5,3±0,28	4,3 ± 0,4

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285 On the 7th and 14th days the value of lymphocyte subpopulation composition did not undergo
286 any significant changes, but the dynamics tended to decrease (normalization) of CD3+
287 lymphocytes.

288 Thus, in the acute period twice alternate days subcutaneous injection of Roncoleukin has no
289 effect on the normal values of the immune status, i.e. it has a modulatory effect depending on the
290 initial status.

291 On the 7th day of the treatment there were no any complaints to dysuria and objectively there
292 were no swelling and hyperemia of external urethral opening and swelling of the labia majoria.

293 The profuse discharge were completely absent on the 7th treatment day, corresponding to the
294 data of objective examination.

295 According to the patient's words there was an increased vaginal discharge, but the nature of
296 this discharge has changed to ordinary fluor albus with a gradual transition to the normal vaginal
297 discharge to the 10th day of the treatment.

298 Thus, the major clinical criteria of inflammatory process remitting were the quantity reduction
299 of the pathologic discharge from the reproductive tract, appearance of light whitish discharge in
300 moderate and scarce quantities, disappearance of vulvovaginal pain, burning sensation in the
301 genitals, decurrence of dysuric syndrome, the blushed mucous membranes of the vagina with an
302 absence of any pathological changes

303 **Conclusion**

304 Thus, in the subacute the appointment of twice alternate days subcutaneous injection of
305 Roncoleukin leads to the normalization of main value of lymphocyte subpopulation.

306 In the acute Roncoleukin has no influence on the normal values of the immune status, i.e. it
307 has an effective immune protection depending on the initial status.

308 Prior to treatment with cytokine composition the mean value did not differ significantly from
309 the control group value in the sub-acute.

310 The high percentage of IFN- γ value before treatment was observed at 15% of the patients,
311 TNF and IL-6 (at 20%, after treatment for 7th day it was reported an increased production of TNF-at
312 25% of patients, and on the 14th day the increased production of cytokines were not observed. These
313 changes may indicate a decrement of inflammation, which was preceded by the activation of
314 cytokine.

315 In the acute period prior to treatment in the cytokine composition the value did not differ
316 significantly from the average of the control group.

317 Interest occurrence of increased production value IFN- γ was observed at 23.8% of patients,
318 TNF - at 21.4% of patients, IL-6 - at 14.3% of patients, which was significantly more frequently
319 than in the control group.

320 On the 7th day there was evaluated an increased production of IL-6 at 14.2% of the patients,
321 TNF at 78.6%, and on the 14th day the increasing of IL-6 was only 4.8%. In this case the
322 introduction of Roncoleukin has also helped to reduce the inflammatory response.

323 Changes of phagocytic activity of the local secretion were observed in the sub-acute.

324 After the local application of Ronkoleukin through 7 days reaction of the local secret sharply
325 increased performance spontaneous and induced by pyrogenal of phagocytic index, phagocytic
326 number, NBTR-test.

327 Thus, locally administering of preparation greatly increases the efficiency of patients'
328 treatment with a positive effect on the clinical course of the disease, providing immunomodulatory
329 effects.

330 All patients with the sub-acute and acute vulvovaginitis should be made an assessment, in
331 addition to clinical examination, for the state of the immune and cytokine status, as well as
332 phagocytic system of vaginal contents in order to address the issue of immune system correction.

333 Immune system correction by Ronkoleukin locally can be recommended to patients with sub-
334 acute disease.

335 Application of Ronkoleukin can reduce treatment costs and shorten the time of treatment.

336 These phenomena were leveled out independently at the 2nd day and were not accompanied
337 by abnormality of general well-being.

338 In view of the high tolerability and rare complications, the local application can be performed
339 on an outpatient basis.

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