The Influence of Propionate on the Spectrum of Short-Chain Fatty Acids in Cervical Cells During Dis- and Neoplastic Transformation

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Abstract: Introduction. There is some data of the short-chain fatty acids (SCFA) participation in cancerogenesis. The cervical cancer is an “ideal” model for determining the molecular mechanisms underlying the development of cancer. The aim of study was to estimate the influence of propionate on the spectrum of SCFA in cervical cells during dis- and neoplastic transformation. Materials and methods of research. Materials of the study - cervical biopsy verified morphologically. Study groups: IA - the focus of the pretumor lesion of cervix; IB - paradisplastic cells; IIA - locus of cervical cancer; IIB - paraneoplastic cells. The spectrum of fatty acids was analyzed before, after 24 hours incubation with 50 µmol/l propionic acid by the gas chromatography method. Methods of nonparametric statistics the Mann-Whitney test were used. Differences were considered statistically significant at p<0.05. Results of the study. The incubation of cervical cancer cells with propionate leads to an increase the level of propionate and butyrate. The incubation of “precancer”, paraneoplastic and paradisplastic cells leads to an increase the levels of isobutyric, valeric and caproic acids against the background of a significant drop in the concentration of propionate and butyrate. Conclusion. It can be concluded that the effect of propionate on the metabolism of fatty acids in the cervical epithelium of the studied samples is multidirectional and depends on the cell type. The dates indicate the modifications of SCFA in the cervical carcinogenesis.

Keywords: Cancerogenesis, Cervical Cancer, Short Chain Fatty Acid

1. Introduction.

Cervical cancer is a long and staged oncological disease based on malignant transformation of exocervix and endocervix cells. From the perspective of oncogenesis research malignization of cervical epithelium is viewed as an «ideal» model to study molecular mechanisms provoking the development of oncological diseases. It were the cervical cancer cells (HeLa-cells) obtained as early as 1951 that were the first human “immortal” cells still being used for the research of biology of many diseases [1]. Currently there is no unified theory of cervical oncogenesis. Literature sources describe several ethiopathogenetic variants of the development of this disease:
viral, bacterial, immunological, hormonal, and alteration.

However, regardless of the trigger nature, further life of the
cell is determined by the following processes: metabolic
reprogramming, proliferation, differentiation, apoptosis, cell
cycle kinetics, and necrosis. It is known that the state of
tumor cell biological membrane and its fatty acid
composition potentiate the main elements of carcinogenesis.

[2-5].

Recently data on participation of short-chain fatty acids
(SCFA) in tumor growth have appeared in literature; they are
viewed as metabolic by products of microorganisms. Only
some information concerning SCFA metabolism peculiarities
was found [6].

Earlier we determined that SCFA deficiency was observed
in tumors and dysplastic cervical epithelium cells. It was
the most obvious in nidus of dysplasia and neoplasia [7, 8],
since SCFA were used inter alia for synthesis of higher fatty
acids in the tumor tissue.

2. Aim of the Research

The Aim of the Research is to estimate the influence of
propionate, one of the SCFAs, in vitro, on the SCFA profile
cervical epithelium tissue lipids in premalignancies and
neoplasia of exocervix, in order to get more extended data on
interconversion of SCFAs in carcinogenesis.

3. Materials and Methods of Research

Nonrandomized controlled prospective study was
performed. It included 58 women who were examined and
trained in the Zabaikalye Territorial Oncology Center. The
average age of the patients was 38±8.26 years. All patients
were informed about the study and gave their written consent
to participate. The study was compliant with the principles of
WMA Declaration of Helsinki, 1964, rev. 2013 and was
confirmed morphologically.

Control group samples were collected from 18 apparently
healthy women aged from 28 to 48 years (34.5±6.5 years),
under dispensary observation because of non-tumor
exocervix pathology treated earlier. They were informed
about the study design, and gave informed consent to
participate. Integrity of the cervical epithelium was
confirmed morphologically.

Cervical tissue fragments received by target biopsy or
intraoperatively under morphological control served as
samples for the research. According to histological
examination, each tissue fragment was divided into two loci:
A – diseased area, B (B) – tissue showing no signs of
dysplastic and neoplastic transformation. The patients were
divided into three clinical groups: I – patients with cervical
premalignancies (20 – grade III cervical intraepithelial
neoplasia), II – cervical cancer (20 - histologically
squamous cell cancer, stage 0-1b), control group – healthy
women (18).

In order to get cell-rich fluid biopsy, samples were
pulverized and homogenized in Gentle MACS Dissociator
(Miltenyi Biotec GmbH, Germany) using C type tubes, and
Tumor Dissociation Kit reagents (Miltenyi Biotec GmbH,
Germany) in compliance with the procedure. The obtained
cell-rich fluid was filtered through capronic filter with the
mesh size of 30 µm (Miltenyi Biotec GmbH, Germany).
The cells were washed by autoMACS Rinsing Solution
(Miltenyi Biotec GmbH, Germany). The cells were counted
by FC500 flow cytometer (Beckman Coulter, USA)
using FLOW-COUNT Fluospheres (Beckman Coulter,
USA).

The amount of SCFAs in washed cell-rich fluid was
determined by procedure developed by Ardatskaya M.D.
(2004) and modified by us. It comprised two stages: sample
preparation, and gas-liquid chromatography [10].

SCFA extraction was performed as follows: diethyl ether
and cell suspension were mixed in equal amounts, then 0.50
ml of 50.0% sulfuric acid and internal standard (a,a-
dimethyl-butanoic acid) were added. The mixture was shaken
generously for 10 minutes and centrifuged at 3000 rpm for
the same time. The supernatant was evaporated to dryness,
dissolved by hexane and then analyzed by «Кристалл-
2000M» chromatograph (Crystal-2000M) chromatograph (Russia). The
following acids were detected: C$_{3}$ – propionic acid, C$_{4}$ –
butyric acid, isoC$_{4}$ – isobutyric acid, C$_{5}$ – valeric acid,
and C$_{6}$ – capronic acid.

Statistical data processing was performed by
«BIOSTAT» software. The results were presented as a
median, 25th and 75th percentile. Methods of
nonparametric statistics using the Wilcoxon signed-rank
test for paired measurements and Mann-Whitney U test for
the control group were used. The differences were
considered statistically significant at p<0.05.

4. Results of Research

Cervical epithelium cells regardless of pathology type
(neoplasia or precancer), their localization (tumor nidus or
distal cells), and incubation conditions (with propionic acid
or without propionic acid) showed the total deficiency of
SCFA, and the deficiency grew in the premalignancy nidus
of dysplasia and neoplasia [7, 8], since SCFA were used inter alia for synthesis of higher fatty
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and paraneoplastic cells under the influence of propionate
(Figure 1).

In the presence of propionic acid cervical cancer cells
showed increased share of short-chain analogs – 1.5 times
higher at the expense of both even-numbered and odd-
numbered fatty acids; however, their level never reached the
control value (p<0.001).

SCFA content in paradysplastic biopsy material after
incubation with C$_{3}$ increased only by 6%, mainly at the
expense of analogs with an even number of carbon atoms
(p<0.001).
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Cervical epithelium cells of both groups regardless incubation conditions (with propionic acid or without propionic acid) showed total deficiency of SCFA, while this deficiency grew in the premalignancy nidus and paraneoplastic cells under the influence of propionate. The detailed changes of SCFA pool in the cervical epithelium cells depending on conditions of incubation are shown in the table 1.

In the «premalignancy» group the level of C_3:0, C_4:0 statistically did not differ from localization of cells. After incubation with propionate, concentration of the latter both in premalignancy locus and in paraneoplastic cells decreased by 27.2% and 30% correspondingly (p<0.02). The share of the remaining SCFA (iC_4:0, C_5:0, C_6:0) in the cells of this group grew, but did not reach the control values (p<0.001).

In cervical cancer biopsy materials after incubation with C_3:0 changes of SCFA pool were caused only by increasing concentration of propionate itself, and butyrate by 3.2 and 2 times correspondingly (p<0.02).

In paraneoplastic tissue fragments, under the influence of propionic acid, their level decreased by 40.6% each (p<0.02), while the share of isobutyrate, valeriate and capronate increased by 54.5%, 50%, and 50% correspondingly (p<0.02).

Table 1. SCFA Content (ng/cell) in Cervical Cells of Different Localization at Dysplastic and Neoplastic Transformations Depending on the Conditions of Incubation.

| group | Control (n=25) | I group | II group |
|-------|----------------|---------|----------|
| SCFA  | incubation 24 hours | locus A (n=35) | locus B (n=35) | locus A (n=45) | locus A (n=45) |
| C_3:0 | 0.060 (0.049; 0.089) | 0.022 (0.022; 0.023) | 0.022 (0.016; 0.029) | 0.019 (0.09; 0.058) | 0.074 (0.035; 0.120) |
| C_4:0 | 0.026 (0.021; 0.039) | 0.010 (0.009; 0.010) | 0.010 (0.007; 0.012) | 0.08 (0.004; 0.025) | 0.032 (0.015; 0.053) |
| iC_4:0 | 0.059 (0.045; 0.067) | 0.005 (0.004; 0.006) | 0.013 (0.013; 0.014) | 0.013 (0.008; 0.018) | 0.011 (0.006; 0.020) |
| C_5:0 | 0.032 (0.025; 0.037) | 0.005 (0.004; 0.006) | 0.012 (0.011; 0.013) | 0.011 (0.007; 0.016) | 0.010 (0.006; 0.018) |
| C_6:0 | 0.112 (0.089; 0.150) | 0.003 (0.002; 0.004) | 0.007 (0.007; 0.008) | 0.007 (0.004; 0.010) | 0.006 (0.003; 0.011) |
| incubation with propionic acid | | | | | |
| C_3:0 | 0.044x (0.036; 0.063) | 0.016# (0.016; 0.017) | 0.016# (0.012; 0.020) | 0.062* (0.033; 0.106) | 0.044** (0.022; 0.064) |
| C_4:0 | 0.019x (0.016; 0.027) | 0.007# (0.007; 0.007) | 0.007# (0.005; 0.009) | 0.027* (0.014; 0.046) | 0.019** (0.010; 0.028) |
| iC_4:0 | 0.082x (0.062; 0.099) | 0.008# (0.006; 0.009) | 0.019# (0.018; 0.020) | 0.013 (0.008; 0.018) | 0.017** (0.011; 0.028) |
| C_5:0 | 0.072x (0.056; 0.087) | 0.007# (0.005; 0.008) | 0.017# (0.016; 0.018) | 0.011 (0.007; 0.016) | 0.015** (0.009; 0.025) |
| C_6:0 | 0.045x (0.034; 0.054) | 0.004# (0.003; 0.005) | 0.010# (0.010; 0.011) | 0.007 (0.004; 0.010) | 0.009** (0.006; 0.015) |

*, **, #, ##, x, xx – statistical significance of parameters from the corresponding subgroups incubated with propionic acid and without propionic acid (p<0.001)
In order to exclude possible influence of conditions of incubation and to estimate real influence of propionate on SCFA metabolism in cervical cells, we performed comparative study of their profile in the researched clinical groups before and after the incubation, in the presence of propionic acid and without propionic acid (Figure 2).

Under the influence of C\text{3:0} in cervical cancer biopsy materials the pool of odd-numbered SCFA increased by 80%, as opposed to precancer locus and paraneoplastic cells, where the content of short-chain analogs was reduced regardless of the amount of carbon atoms: by 21.3% and 20.6% - for odd-numbered SCFA and by 8.6% and 9.5% - for even-numbered SCFA in conditionally healthy cells of the “cancer” group and premalignancy nidus correspondingly (p<0.001).

In cells of paradyplastic localization after incubation with propionic acid the share of odd-numbered SCFA decreased by 11% (p<0.001).

The detailed changes of SCFA profile in cervical epithelium cells depending on the pathology type, distance from the tumor nidus and conditions of incubation are shown in the figure 3.

In cervical cancer locus during 24-hour incubation all SCFA showed only the tendency of decreased concentration. After incubation of cervical cancer cells with C\text{3:0} increase of propionic acid and butyric acid pool by more than 2 times (p<0.001) was registered.

It was observed in paraneoplastic cells during incubation without propionate addition that the levels of C\text{3:0}, C\text{4:0} increased by 1.9 and 1.8 times correspondingly (p<0.001) against the background of decreased share of iC\text{4:0}, C\text{5:0}, C\text{6:0}by 21.4%, 23.1% and 25% correspondingly (p<0.001). In conditions of incubation with propionate the opposite situation was observed: increased pool of isobutyric, valerianic, and capronic acid by 21.4%, 15.4% and 12.5% correspondingly against the background of decreased concentration of propionate and butyrate approximately by 30% each (p<0.001).

In the premalignancy nidus after incubation of cells with C\text{3:0} deficiency of propionate and butyrate by 30% was determined for each SCFA (p<0.001), the levels of iC\text{4:0}, C\text{5:0} increased by 14% and 40% correspondingly (p<0.001); also a constant level of C\text{6:0} was observed. Similar SCFA profile was observed in cells of paradyplastic localization with the exception of capronic acid, which increased by 42.9% after incubation with C\text{3:0} (p<0.001).

* * *, #, ##, х, хх – statistical significance of parameters from the corresponding subgroups incubated with propionic acid and without propionic acid (p<0.001)

**Figure 2. SCFA Structure (ng/cell) in Cervical Cells at Dysplastic and Neoplastic Transformations before and after Incubation with Propionic Acid and without Propionic Acid.**
The incubation of cervical cancer cells with propionate leads to an increase in the level of propionate and butyrate. The incubation of "precancer", paraneoplastic and paradysplastic cells leads to an increase in the levels of isobutyric, valeric and caproic acids against the background of a significant drop in the concentration of propionate and butyrate.

5. Discussion

The analyzed Russian and foreign literature did not contain any data concerning changes of SCFA metabolism in tumor tissue. We can assume that excess of C$_{3:0}$ and C$_{4:0}$ in cervical cancer lesion after incubation with propionate is caused not only by its exogenous inflow, but also by interconversions of SCFA, oxidation reaction of even-numbered (for butyric acid) and odd-numbered (for propionic acid) fatty acids, as well as by the possibility of preliminary formation of C$_{3:0}$ in tumor tissue from glucose. Such changes in malignant tissue are possibly caused by high energy status of tumor cells [11-15].

Moreover, in order to preserve liquid crystalline structure of its membranes against the background of deficiency of polyunsaturated fatty acids, cancer cells possibly initiate synthesis of α-methylstearinic acid from propionate, according to the theory of Khyshiktuev B.S. (1995). We also found this compound in cervical cancer tumor growth [8].

6. Conclusion

Thus, the influence of propionate on SCFA metabolism in cervical epithelium of the studied samples is various and depends on the type of the cell: increased levels of C$_{3:0}$ and C$_{4:0}$ are observed in cancer cells, while in paraneoplastic and paradysplastic areas share of isobutyric, valerianic, and capronic acid is increased with considerable decrease of propionate and butyrate concentrations.

Conflict of Interest

The authors of the article declare no conflict of interest.

The authors of the article certify the original work, he manuscript has not previously been published and is not currently being considered for publication elsewhere.

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Compliance with the Ethical Principles

The study was compliant with the principles of WMA Declaration of Helsinki, 1964, rev. 2013 and was performed under consent of the Local Ethics Committee of Chita State Medical Academy. All patients gave written consent to participate in the research.

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