Fecal short-chain fatty acids at different time points after ceftriaxone administration in rats

Yu. V. Holota¹, O. O. Holubenko², A. M. Ostapchuk²,
T. M. Serhiyuchuk³, L. V. Zakordonets³, G. M. Tolstanova¹

¹Educational and Scientific Center “Institute of Biology and Medicine”,
Taras Shevchenko National University of Kyiv, Ukraine;
²D.K. Zabolotny Institute of Microbiology and Virology, National
Academy of Sciences of Ukraine, Kyiv;
³Bogomolets National Medical University, Kyiv, Ukraine
e-mail: gtolstanova@gmail.com

Short-chain fatty acids (SCFAs) are major products of the microbial fermentation of dietary fiber in the colon. Recent studies suggest that these products of microbial metabolism in the gut act as signaling molecules, influence host energy homeostasis and play major immunological roles. In the present study, we defined the long-term effects of ceftriaxone administration on the fecal SCFAs concentration in Wistar rats. Ceftriaxone (300 mg/kg, i.m.) was administered daily for 14 days. Rats were euthanized in 1, 15 and 56 days after ceftriaxone withdrawal. Caecal weight and fecal concentration of SCFAs were measured by gas chromatography. Ceftriaxone administration induced time-dependent rats’ caecal enlargement through accumulation of undigestable substances. One day after ceftriaxone withdrawal, the concentrations of acetic, propionic, butyric acids and total SCFAs were decreased 2.9-, 13.8-, 8.5-, 4.8-fold (P < 0.05), respectively. Concentration of valeric, isovaleric and caproic acids was below the detectable level. That was accompanied by 4.3-fold decreased anaerobic index and increased relative amount of acetic acid (P < 0.05). In 56 days, concentration of SCFAs was still below control value but higher than in 1 day (except for propionic acid). Anaerobic index was 1.3-fold lower (P < 0.05) vs. control. Conclusion: antibiotic therapy induced long-term disturbance in colonic microbiota metabolic activity.

Keywords: antibiotics, ceftriaxone, colon, short-chain fatty acids (SCFAs).

The interchange of low molecular weight metabolites between gut microorganisms and macroorganism have attracted a lot of attention during last years [1-3]. The gut microbiota affects predominantly host physiology by the production of short-chain fatty acids (SCFAs). SCFAs are saturated aliphatic organic acids that consist of one to six carbons of which acetate (C2), propionate (C3), and butyrate (C4) are most abundant (≥95%). Acetate, propionate, and butyrate are present in an approximate molar ratio of 60:20:20 in the colon and stool [4]. Depending on the diet, the total maximum concentration of SCFAs decreases from 70 to 140 mM in the proximal colon from 20 to 70 mM in the distal colon [5].

These metabolites, especially butyrate, serve as an important source of energy for the intestinal epithelial cells, providing about 60-70% of their energy demand. Colonocytes from germ-free mice are in an energy-deprived state and exhibit decreased expression of enzymes that catalyze key steps in intermediary metabolism including the tricarboxylic acid cycle. Consequently, there is a marked decrease in NADH/NAD⁺, oxidative phosphorylation, and ATP levels, that results in AMP-activated protein kinase activation, cyclin-dependent kinase inhibitor 1B phosphorylation and autophagy. When butyrate is added to germ-free colonocytes, it rescues their deficit in mitochondrial respiration and prevents them from undergoing autophagy [6].
Moreover, they indeed play major immunological roles both systemically and locally in the gut. This pathway involves the activation of SCFA-binding G-protein-coupled receptors (GPCRs). In 2003, GPR41 and GPR43 were deorphanised as SCFA receptors [7-9]. Anti-inflammatory functions refer to the property of SCFAs modulate immune cell chemotaxis, adhesion, production of reactive oxygen species and cytokines [10, 11]. Butyrate elicits anti-inflammatory effects via inhibition of interleukin (IL)-12 and upregulation of IL-10 in human monocytes [12]; inhibition of the tumor necrosis factor (TNF)-α-induced nuclear translocation of the proinflammatory nuclear transcription factor-κB (NF-κB) in human colorectal adenocarcinoma (HT-29) cell line [13] induction of colonic regulatory T cells differentiation, which suppresses the inflammatory and allergic response [14]. However, in sites of anaerobic bacteria infection or after the loss of intestinal epithelial integrity, high concentrations of SCFAs may lead to neutrophil accumulation and amplification of the inflammatory process [11].

Significant violations in gut and the whole body microbiome often occur after using antibiotics and other chemotherapeutic agents that occupied a great place in modern medical practice [15]. There is the increasing concern that antibiotic exposure may have long-term consequences for immune tolerance of the gastrointestinal tract and susceptibility to pathogens, contributing to the development of inflammatory diseases [16]. The effect of antibiotic therapy on the fecal SCFAs level has been studied before [17-19], but most studies were conducted during or immediately after antibiotics administration. Therefore little is known about the duration of microbiota metabolic profile disorders caused by antibiotic therapy. The aim of this work was to determine the fecal SCFAs level at different time points after 14-day administration of ceftriaxone in Wistar rats.

**Materials and Methods**

The experiments were performed on Wistar male rats (140-160 g, n = 12), which were bred and housed in the conventional animal facility of the ESC “Institute of Biology and Medicine” of Taras Shevchenko National University of Kyiv (Ukraine) under standard environmental conditions (12-h light/dark cycle at a constant temperature of 22 °C). All animals had unlimited access to animal chow and tap water throughout the study. To normalize gut microbiota, rats from all groups were kept in the same room and maintained by the same personnel. The study was approved by the animal committee of ESC “Institute of Biology and Medicine” Taras Shevchenko National University of Kyiv (Protocol No 8 issued in Nov, 2015). The day 1 of the experiment is considered the first day of ceftriaxone or vehicle administration. Ceftriaxone (Ind. Stock Company Darnytsya, Ukraine) was administered for 14 days in a dose 300 mg/kg intramuscularly (i.m.). Control rats were treated with sterile water (0.1 ml/rat, i.m.). Caecal weight was measured directly one (15th day of experiment) and 15 days (29th day of experiment) after ceftriaxone withdrawal and calculated as % of body weight. The fecal concentration of SCFAs of rats was determined one (15th day of experiment) and 56 days (72nd day of experiment) after ceftriaxone withdrawal.

Fecal samples weighing 1 g were homogenized in 2 ml of 0.02N HCl and then kept at room temperature for extraction during 30 min in air-tight containers to prevent the loss of volatile SCFA. Then samples were centrifuged for 10 min (11 000 g). 300 µl of supernatants were transferred to an autosampler vial for GC-MS analysis and added 100 µl of 0.05% internal standard (4-methyl-valeric acid, Sigma-Aldrich, Germany).

The gas chromatographic (GC) analysis was carried out at the Center for collective usage at the D.K. Zabotolny Institute of Microbiology and Virology of NAS of Ukraine using an Agilent 6890N GC system (Agilent Technologies, USA) equipped with an automatic liquid sampler Agilent 7683B, (Agilent Technologies, USA). Separation was performed using a fused-silica capillary column with a free fatty acid phase DB_FFAP, 0.25 mm × 0.25 mm × 30 m (Agilent Technologies Inc., USA). Helium was supplied as the carrier gas at a flow rate of 1 ml/min. The initial oven temperature was 100 °C, maintained for 5 min, raised to 190 °C at 10 °C/min. The temperature of the injection port was 250 °C. The injected sample volume for GC analysis was 1 µl with split ratio 1 : 20. The run time for each sample was 16 min. A single quadrupole mass spectrometer Agilent, 5973 inert MSD (Agilent Technologies Inc., USA) was used for detection of SCFAs. Data handling was carried out with Chem Station Data Analysis D.01.02.16 software.

The SCFAs were identified on chromatograms by specific retention times of standard SCFA mixture of acetic, propionic, i-butyric, n-butyric,
n-valeric, i-valeric, n-caproic acids (Sigma-Aldrich, Germany) under the above GC conditions. To quantify the peak area in terms of concentration, the peak area was plotted against the peak area and concentration of the internal standard. Absolute quantities of SCFAs were normalized to sample mass and were expressed as μmol/g faeces. The anaerobic index was counted as the proportion the sum of total propionate and butyrate to acetate concentration (AI = (C₃ + C₄)/C₂), relative SCFA abundance as proportion of each acid to the sum of total acetate, propionate and butyrate (Cₙ/∑(C₂ + C₃ + C₄)).

Data were analyzed with the Statistica 6.0 software. Statistical significance was determined by the Student’s t-test or nonparametric Mann-Whitney criterion where appropriate. P-values less than 0.05 were considered statistically significant. Data are presented as mean±SEM.

Results and Discussion

Although antibiotics are effective medications for the treatment of infections, antibiotic therapy often has adverse consequences to the host. Consistent with our previous study [20, 21], ceftriaxone administration leads to profound changes in colonic microbiota in rats as well as in patients, immediately and long after antibiotic withdrawal. Moreover, the disturbance of the surface mucus barrier (level of glycoproteins) in the rat colon was observed long after the antibiotic therapy. We revealed also qualitative changes in the structure of glycan part of mucins, e.g. a decrease of hexoses and fucose content and increase of the level of sialation of mucus glycoproteins [20]. These changes are typical of inflammatory bowel diseases (IBD) [22]. However, the molecular basis of these alterations is not fully understood. It is important to find out the mechanisms by which antibiotic-induced microbial perturbations induce the disturbance of the intestinal barrier in remote terms after antibiotic therapy. We hypothesized that these disturbances involve the changes in gut microbiota metabolic profile.

As hindgut fermenters, rodents and rabbits comprise either a large cecum or colon, where food undergoing digestion is retained, mixed and fermented by microbes [23]. Undigested materials in the caecum are a potential source for intestinal bacteria to produce SCFAs. We observed caecal enlargement after ceftriaxone administration (Fig. 1). Thus one day after ceftriaxone withdrawal the caecal weight was 2.9-fold increased (P < 0.05) (Fig. 1, A), in 15 days – 1.5-fold (P < 0.05) (Fig. 1, B). Tulstrup et al. [24] have shown the increased caecum weight after amoxicillin and vancomycin administration. The development of an enlarged caecum is a typical feature in germ-free rodents [25]. It seems that caecal enlargement depends on the amount of accumulated substances that in the conventional animals should be degraded by microbial fermentation.

Further, we determined how ceftriaxone administration influences the fecal SCFAs concentrations. One day after ceftriaxone withdrawal the total concentration of the fecal SCFAs (Σ(C₂-C₆)) was decreased 4.8-fold (P < 0.05) (Table).

Concentrations of acetic, propionic and butyric acids were 2.9-, 13.8- and 8.5-fold decreased
Concentrations of valeric, isovaleric and caproic acids were below the detectable level of parameters of chromatographic separation. These data are consistent with the results of Meijer-Severs et al. [26], in which treatment of healthy volunteers with ceftriaxone for 5 days resulted in a significant decrease in the fecal concentration of SCFAs on the 3rd and 5th day. Our previous study [27] showed that next day after ceftriaxone withdrawal the composition of colon parietal microbiota was not significantly changed. Moreover, the number of *Propionibacterium* and *Bacteroides* were even increased 1.6- and 2.3-fold (*P* < 0.05), respectively. Thus the observed changes in fecal SCFAs concentration represent disturbance of microbiota metabolic activity rather than its composition. According to the fact that SCFAs are the main energy substrates for the enterocytes, SCFAs loss might lead to disturbance of mucosal cells homeostasis in the rat colon. Moreover, the disruption of biochemical signaling between microbiota and host can profoundly affect inflammatory response leading to unresolved inflammation in the models of colitis, arthritis and asthma [28]. It should be noted that the decrease of butyrate-producing bacteria and SCFAs level detected in feces of patients with ulcerative colitis, indicated the role of intestinal dysbiosis and decrease of butyrate during colitis development [29]. Similar changes were also observed in patients with nosocomial diarrhea and *Clostridium difficile*-associated colitis [30]. A reduction in fecal butyrate has been found in patients with colorectal adenocarcinoma [31].

Acetic, propionic and butyric acids are most important among SCFAs. They are metabolic end products of proteolytic and saccharolytic microbiota used for the integrated assessment of its state [32]. We observed the 1.6-fold increased relative amount of acetic acid and 3.3- and 1.9-fold reduced proportion of propionic and butyric acids respectively (*P* < 0.05) one day after ceftriaxone withdrawal (Fig. 2). That was accompanied by 4.3-fold decrease of the anaerobic index (Fig. 3). This index reflects the redox potential of the colon environment (balance between putatively anti-inflammatory properties of butyric and propionic acids and proinflammatory acetic acid) [32]. It was shown that butyrate and propionate were equipotent at inhibiting the transcription of NF-κB and had anti-inflammatory properties, while acetate revealed a significantly lower efficiency [33]. In addition, propionate and butyrate, but not acetate, induced intestinal hormone release, that is an important mechanism for the regulation of host metabolism by microbiota [34]. Acetate might increase macrophage cells histone acetylation *in vitro* by end-product inhibition of histone deacetylases and activation of histone acetyltransferases. These changes promote activation of proinflammatory genes transcription and cytokines synthesis [35]. Thus observed changes may indicate a trend toward a proinflammatory condition in the colon.

Allowing for the fact that dysbiotic disturbance of mucosal microbiota were observed for a long time after ceftriaxone administration [27], we tested how these changes affect the SCFAs level in rat’s feces.

### Concentration of the short-chain fatty acids (SCFAs) in rats in 1 and 56 days after ceftriaxone (300 mg/kg, i.m., 14 days) withdrawal

<table>
<thead>
<tr>
<th>SCFAs</th>
<th>Concentration (μmol/g wet faeces)</th>
<th>Control</th>
<th>Ceftriaxone, after withdrawal</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>in 1 day</td>
<td>in 56 days</td>
</tr>
<tr>
<td>Acetic</td>
<td>10.98 ± 1.82</td>
<td>3.79 ± 1.03*</td>
<td>8.27 ± 0.66^</td>
</tr>
<tr>
<td>Propionic</td>
<td>7.32 ± 1.41</td>
<td>0.53 ± 0.19*</td>
<td>2.41 ± 0.15^</td>
</tr>
<tr>
<td>Butyric</td>
<td>3.75 ± 0.72</td>
<td>0.44 ± 0.14*</td>
<td>3.91 ± 0.15^</td>
</tr>
<tr>
<td>Valeric</td>
<td>0.37 ± 0.11</td>
<td>–</td>
<td>0.27 ± 0.03</td>
</tr>
<tr>
<td>Isovaleric</td>
<td>0.32 ± 0.13</td>
<td>–</td>
<td>0.38 ± 0.02</td>
</tr>
<tr>
<td>Caproic</td>
<td>0.45 ± 0.13</td>
<td>–</td>
<td>0.17 ± 0.09</td>
</tr>
<tr>
<td>Total</td>
<td>22.65 ± 3.47</td>
<td>4.75 ± 1.36*</td>
<td>15.14 ± 0.95^</td>
</tr>
</tbody>
</table>

* *P* < 0.05 compare to control values; ^ P < 0.05 compared to values one day after ceftriaxone withdrawal; “—” acid concentration below detectable level of parameters of chromatographic separation.
The concentration of fecal SCFAs in rats increased significantly 56 days after ceftriaxone withdrawal vs. the values observed one day after ceftriaxone withdrawal, but did not reach the control values. Moreover, the concentration of propionic acid was still significantly lower vs. control \( (P < 0.05) \) (Table). Interestingly, the amount of butyric acid corresponded to control value 56 days after ceftriaxone withdrawal.

The relative amount of acetic acid was 1.5-fold decreased \( (P < 0.05) \) compared to the values observed one day after ceftriaxone withdrawal and corresponded to control values, while relative amount of propionic acid remained 2-fold reduced \( (P < 0.05) \) vs. control. The relative amount of butyric acid increased 1.6-fold \( (P < 0.05) \) vs. control. According to similar anti-inflammatory properties of these acids, an increased level of butyric acid might occur due to a compensatory metabolic activation of butyrate-producing bacteria in response to a significant reduction in propionate formation. Propionate and butyrate are produced from hexose sugars by different microbial species. Only two species of Lachnospiraceae \( (\) Coprococcus catus and Roseburia inulinivorans \( ) \) might switch from butyrate to propionate production on different substrates [36]. However, these changes do not lead to restoration of the redox status of colon lumen. The anaerobic index was 1.3-fold lower \( (P < 0.05) \) 56 days after ceftriaxone withdrawal vs. the control values.

Thus, our results indicate that administration of ceftriaxone leads to a severe decrease in the total concentration of the fecal SCFAs in rats immediately after antibiotic withdrawal. The most significant changes were observed in the concentration of propionic and butyric acids that led to the increase of relative amount of acetic acid. Reduction in the anaerobic index may indicate the inhibition of anaerobic and activation of conditionally pathogenic microorganisms after ceftriaxone withdrawal. These changes are stable over time. Even 56 days after ceftriaxone withdrawal the concentration of propionic acid and anaerobic index were still reduced significantly. Therefore antibiotic therapy leads to the stable long-term disturbances of colon microbiota metabolic activity in rats which underlie the pathogenesis of antibiotic-associated colon homeostasis.
disorders and might increase susceptibility to IBD development.

ФЕКАЛЬНІ КОРОТКОЦЕПОЧЕЧНІ ЖИРНІ КИСЛОТЫ В РІЗНИ СТРОКИ ПІСЛЯ ВВЕДЕННЯ ЦЕФТРИАКСОНА ЩУРАМ

Ю. В. Голота, О. О. Голубенко, А. М. Остапчук, Л. В. Закордонець, Г. М. Толстанова

1НИЦ «Інститут біології та медицини», Київський національний університет імені Тараса Шевченка, Україна;
2Інститут мікробіології і вірусології ім. Д. К. Заболотного НАН України, Київ;
3Національний медичний університет ім. О. О. Богомольца, Київ, Україна;
e-mail: gtolstanova@gmail.com

Короткоцепочечні жирні кислоти (КЦЖК) є основними продуктами бактеріальної ферментації харчових волокон у товстій кишці. Недавні дослідження показали, що ці продукти мікробного метаболізму в кишечнику діють як сигнальні молекули, впливають на енергетичний гомеостаз організму хазяїна та відіграють головну роль в імунологічній відповіді. У цьому дослідженні визначали вплив цефтриаксона на концентрацію фекальних КЦЖК щурів лінії Вістар. Цефтриаксон (300 мг/кг, в/м) вводили щоденно впродовж 14 діб. Щурів умертвляли через 1, 15 і 56 діб після відміни цефтриаксону. Определяли масу сліпої кишки та концентрацію КЦЖК в фекаліях методом газової хроматографії. Введення цефтриаксону індукувало залежне від часу збільшення маси сліпої кишки щурів у зв’язку з накопиченням неперетравлених залишків.

Через 1 добу після відміни цефтриаксону концентрація масляної кислоти знизилась в 2,9; 13,8; 8,5; 4,8 раза (P < 0,05) відповідно. Концентрація валеріанової, ізовалеріанової і капронової кислоти була нижче рівня детекції. Це супроводжувалося зниженням в 4,3 раза значення анаеробного індексу та збільшенням відносного вмісту оцтової кислоти (P < 0,05) в порівнянні з контролем. Дійшли новку, що антибактеріальна терапія спричиняє тривале порушення метаболічної активності мікробіоти товстої кишки.

Ключові слова: антибіотики, цефтриаксон, товста кишка, короткоцепочечні жирні кислоти (КЦЖК).
(changes относительного содержания уксусной кислоты (Р < 0,05). Через 56 суток концентрация КЦЖК была все еще ниже контрольных значений, но выше, чем через 1 сутки (за исключением пропионовой кислоты). Анаэробный индекс был снижен в 1,3 раза (Р < 0,05) по сравнению с контролем. Сделан вывод о том, что антибактериальная терапия вызывает длительное нарушение метаболической активности микрофлоры толстой кишки.

Ключевые слова: антибиотики, цефтриаксон, толстая кишка, короткоцепочечные жирные кислоты (КЦЖК).

References


Received 13.12.2016