

**IMMUNOLOGICAL ASPECTS OF VACCINATION IN HIV-INFECTED PATIENTS**

Zhestkov A. V. <sup>a</sup>,

Zolotov M. O. <sup>a</sup>,

Kadantseva E. V. <sup>a</sup>,

Nikitina T. R. <sup>a</sup>,

Protasov A. D. <sup>a</sup>

<sup>a</sup> Samara State Medical University.

**ИММУНОЛОГИЧЕСКИЕ АСПЕКТЫ ВАКЦИНАЦИИ ВИЧ-ИНФИЦИРОВАННЫХ ПАЦИЕНТОВ**

Жестков А. В. <sup>1</sup>,

Золотов М. О. <sup>1</sup>,

Каданцева Е. В. <sup>1</sup>,

Никитина Т. Р. <sup>1</sup>,

Протасов А. Д. <sup>1</sup>

<sup>1</sup> ФГБОУ ВО СамГМУ Минздрава России.

## Резюме

Проблема ВИЧ-инфекции не теряет своей актуальности до настоящего времени. В 2022 году 630 000 человек умерли и 1,3 миллиона человек заразились вирусом иммунодефицита человека (ВИЧ). У ВИЧ-инфицированных инфекционные заболевания развиваются чаще, чем у здоровых лиц, возбудителями которых являются в основном условно-патогенные микроорганизмы. Основным возбудителем пневмонии у ВИЧ-инфицированных является *Streptococcus pneumoniae*. Для предотвращения развития тяжелых форм пневмококковой пневмонии и преодоления антибиотикорезистентности разработаны вакцины, содержащие антигены из капсулы *S.pneumoniae*. Созданы и активно применяются в клинической практике пневмококковые полисахаридные (ППВ) и пневмококковые конъюгированные вакцины (ПКВ). Однако не обнаружено данных о влиянии ПКВ13 на клетки иммунологической памяти. Целью данного исследования является изучение влияния вакцинации ПКВ13 на иммунную систему у ВИЧ-инфицированных лиц. Материалы и методы. В исследование были включены 200 больных ВИЧ-инфекцией, которые были разделены на две группы: 1 - получившая дозу ПКВ13 (n=100) и контрольная (n=100). Во время первого визита были проведены иммунологические и микробиологические исследования. Во второй визит осуществлялось введение в дельтовидную мышцу ПКВ13. Третий визит был выполнен через 1 год, повторно проводились иммунологическое и микробиологическое исследования. Участники были разделены на 4 подгруппы в зависимости от уровня CD4+ клеток. Микробиологическое исследование проводилось в мазке с задней стенки глотки. Результаты. При проведении иммунологического обследования на визите 1 были выявлены отклонения во всех изученных популяциях и субпопуляциях клеток иммунной системы. Через 12 месяцев после вакцинации медиана уровня CD3+CD4+ и CD45RO+ лимфоцитов в

иммунизированной группе стала выше довакцинального уровня, в отличие от контрольной группы, в которой значения статистически не изменились. Наши данные подтверждают иммунологическую эффективность введения ПКВ13 у ВИЧ-инфицированных пациентов. У больных с дефицитом CD19+ лимфоцитов в периферической крови выявлено увеличение частоты выделения микроорганизма ( $p=0,003$ ). Заключение. Таким образом, в связи с высоким риском развития пневмококковой пневмонии ВИЧ-инфицированные больные должны быть вакцинированы 13-валентной пневмококковой конъюгированной вакциной.

**Ключевые слова:** ВИЧ-инфекция, ПКВ13, *Streptococcus pneumoniae*, вакцинация, пневмококковая пневмония.

### Abstract

Until recently, HIV infection does not lose its relevance. In 2022, 630,000 people died and 1, 3 million people became infected with the human immunodeficiency virus (HIV). HIV-positive persons develop more infectious diseases than healthy people do; the causative agents are mainly opportunistic microorganisms. *Streptococcus pneumoniae* is the main causative agent of infection in the lungs in HIV-infected persons. In order to prevent the development of severe pneumococcal infections and to overcome antibiotic resistance, vaccines have been developed. There are polysaccharide (PPV) and conjugate (PCV) vaccines. According to clinical recommendations, vaccination of previously unvaccinated HIV-infected patients is carried out regardless of T-helper cell level. However, no data were found on the effect of PCV13 on immunological memory cells. The purpose of this study is to assess an effect of PCV13 vaccination on the immune system in HIV-infected subjects. Materials and Methods. The study included 200 patients with HIV infection, which were divided into two groups: 1 - received a dose of PCV13 (n=100) and control group (n=100). During the first visit, immunological and microbiological studies were carried out. On the second visit, a PCV13 was injected into the deltoid muscle. The third visit was made a year later, where immunological and microbiological studies were repeated. Participants were divided into 4 subgroups depending on CD4+ T cell level. The microbial study was done using a swab collected from the back of the throat. Results. During the immunological examination at visit 1, abnormalities were detected in all examined populations and immune cell subsets. At 12 months post-vaccination, the median levels of CD3+CD4+ and CD45RO+ T lymphocytes in the immunized group were higher than pre-vaccination levels compared to control group, in which the values changed insignificantly. Our data confirm the immunological effectiveness of PCV13 administration in HIV-infected patients. In patients with peripheral blood CD19+ lymphocyte deficiency, had increased microbial detection rate (p=0.003).

**Conclusion.** As a result, due to the high risk of pneumococcal pneumonia, HIV-infected patients should be immunized with a 13-valent pneumococcal conjugate vaccine.

**Keywords:** HIV-infected; PCV13; Streptococcus pneumoniae; vaccination; pneumococcal pneumonia.

## 1 Introduction

### 1 Backgrounds

In 2022, 630,000 people died and 1.3 million people became infected with the human immunodeficiency virus (HIV), and there are 39 million people living with HIV (PLHIV) in the world [14]. In the Russian Federation in 2022, 63 150 new cases of infection were detected, 34 410 people died, which is 0.9% more than in 2021 [6].

The causative agents of infectious diseases in PLHIV are opportunistic microorganisms [11]. In 25-60% of HIV-infected patients, lung damage of infectious origin (tuberculosis, pneumocystis and bacterial pneumonia) is observed [3]. The most common causative agents of community-acquired pneumonia are *S. pneumoniae* (up to 20%), *H. influenzae* (10-15%) and *S. aureus* (5-10%). Pneumonia caused by *Mycoplasma pneumoniae* and *Clamydia pneumoniae* also occurs, but is quite rare [1, 2, 4, 5, 7, 9].

To prevent the development of severe forms of pneumococcal infections and to overcome antibiotic resistance, vaccines have been created. Polysaccharide (PPV) and conjugated (PCV) pneumococcal vaccines have been developed. The 13-valent conjugate pneumococcal vaccine contains antigens from thirteen serotypes of pneumococcus, and the 23-valent polysaccharide vaccine contains from twenty-three serotypes. [7]. PCV13 contains polysaccharide antigens conjugated to a carrier protein (diphtheria toxoid CRM197). This binding of molecules allows one to attract T-lymphocytes to the site of inflammation and increase the effectiveness of the immune response. PCV13 is approved for use in children from 6 months of age; it reduces the carriage of pneumococcus on the mucous membrane of the upper respiratory tract and does not require repeated administration in adults.

According to clinical recommendations, vaccination of HIV-infected against pneumococcus, who have not been previously vaccinated, is carried out regardless of the level of T-helpers. Previously unvaccinated people are given PCV13, no earlier than 8 weeks later PCV23 is given, after 5 years PCV23 is revaccinated [8].

29 Several studies have been conducted to evaluate the effectiveness of PCV13  
30 in HIV-infected adults. Research carried out at the University Hospital of Reims to  
31 determine the immunological efficacy of PCV13 in PLHIV. As a result was found  
32 that a single vaccination leads to a positive immunological response and protection  
33 1 month after vaccination in a group of patients with an initial CD4+ level of 200  
34 cells/ $\mu$ l and a CD4+ ratio /CD8+ more than 0.8. After 6 and 12 months, the  
35 percentage of protected patients decreases. In addition, after vaccination with  
36 PCV13, an increase in the opsonophagocytic reaction of anti-pneumococcal  
37 antibodies was noted [12].

38 A study from South Korea examined the effectiveness of PCV13 in PLHIV  
39 depending on baseline T-helper cell levels [13]. A lower immune response to the  
40 vaccine was found in individuals with a T-helper cell count of less than 350 cells/ $\mu$ L.  
41 In addition, lower levels of specific IgG were reported in this group.

42 A meta-analysis that included 28 cohort studies and 11 randomized clinical  
43 trials examined the effectiveness of different pneumococcal vaccination regimens  
44 [10]. The immunological effectiveness of PCV13 administration has been  
45 demonstrated. At the same time, multiple vaccination led to only a slight increase in  
46 IgG and was not economically effective.

47 Limitations of the different studies include the absence of information about  
48 the effect of PCV13 administration on immunological memory cells. In addition, the  
49 main attention was paid to the assessment of the T-cell component of immunity and  
50 the effect on B cells was not presented. However, it is B-lymphocytes produce  
51 antibodies, which are a key link in the fight against pneumococcus.

52 **The purpose of the study** is to evaluate the immunological efficacy of  
53 PCV13 vaccination in HIV-infected people.

## 54 **2 Materials and methods**

55 In the study were included 200 HIV-infected people, which were divided into  
56 two groups: 1 - received a dose of PCV13 (n=100) and control (n=100).



57 For the entire period of the study, for patients were planned 3 visits. At the  
58 first visit, the following studies were performed: microbiological examination of a  
59 smear from the posterior pharyngeal wall, immunological examination of peripheral  
60 blood (determination of CD3+CD4+, CD19+, CD45RO+ lymphocytes). The second  
61 visit was only for the immunized group, where PCV13 was injected into the deltoid  
62 muscle. The third visit was performed after 12 months, immunological and  
63 microbiological studies were repeated.

64 Determination of populations and subpopulations of lymphocytes was carried  
65 out by flow cytometry. Depending on the level of CD4+ cells (T-helpers), the  
66 participants were divided into 4 subgroups: 1st - from 50 to 199 cells/ $\mu$ l of blood  
67 (immunized group n=19, control group n=25); 2nd – 200-349 cells/ $\mu$ l (immunized  
68 group n=20, control group n=14); 3rd - 350-499 cells/ $\mu$ l (immunized group n=17,  
69 control group n=21); 4th - 500 or more cells / $\mu$ l. (immunized group n=44, control  
70 group n=40).

71 To conduct a microbiological study, a smear was taken from the posterior  
72 pharyngeal wall. The biomaterial was seeded on the following media: media: 5%  
73 blood agar, chocolate agar, Sabouraud's medium, universal chromogenic media. The  
74 isolated strains were identified by matrix-activated laser desorption/ionization time-  
75 of-flight (MALDI-ToF) mass spectrometry.

76 Statistical data processing was carried out using the Statistica 13.0 program  
77 (STATSOFT, USA; license 2883). All the studied samples were distributed  
78 abnormally, so non-parametric statistics methods were applied to their study. The  
79 results were considered statistically significant if the significance level (p) was less  
80 than 0.05.

### 81 **3 Results and its discussion**

82 At the first visit, the following median lymphocyte levels were obtained in  
83 vaccinated patients (Figure 1). The median level of T-helpers in those examined in  
84 the 1st subgroup was 115 cells/ $\mu$ l, in the 2nd subgroup - 254.5 cells/ $\mu$ l, in the 3rd

85 subgroup - 421 cells/ $\mu$ l, in the 4th subgroup - 663, 5 cells/ $\mu$ l. At the 3rd visit, there  
86 was an increase in the 1st subgroup up to 205 cells/ $\mu$ l ( $p=0.001$ ), in the 2nd one - up  
87 to 377 cells/ $\mu$ l ( $p=0.001$ ), in the 3rd one - up to 478 cells/ $\mu$ l ( $p=0.004$ ) and in the 4th  
88 - up to 735 cells/ $\mu$ l ( $p=0.096$ ).

89 The dynamics of changes in the number of B-lymphocytes in all the studied  
90 subgroups had no statistical significance. At the same time, a statistically significant  
91 increase in the number of cells involved in the mechanisms of immunological  
92 memory (CD45RO+ lymphocytes) was registered in vaccinated PLHIV. The value  
93 of the median level of CD45RO+ lymphocytes in the first subgroup increased from  
94 556 cells/ $\mu$ l at visit 1 to 623 cells/ $\mu$ l at visit 3 ( $p=0.030$ ); in the second subgroup  
95 from 778.5 cells/ $\mu$ l to 818 cells/ $\mu$ l ( $p=0.028$ ); in the third subgroup from 701 cells/ $\mu$ l  
96 to 749 cells/ $\mu$ l ( $p=0.022$ ); in the fourth subgroup, from 848 cells/ $\mu$ l at the 1st visit to  
97 924 cells/ $\mu$ l at the 3rd visit ( $p=0.01$ ).

98 Participants in the control group showed no statistically significant changes in  
99 the median levels of CD3+CD4+, CD19+, CD45RO+ lymphocytes in all four  
100 subgroups. The dynamics of changes in the content of T-helpers in peripheral blood  
101 was: in the examined in the 1st subgroup - from 102 cells/ $\mu$ l to 118 cells/ $\mu$ l ( $p =$   
102  $0.322$ ), in the 2nd - from 293.5 cells/ $\mu$ l to 372, 5 cells/ $\mu$ l ( $p=0.164$ ), in the 3rd  
103 subgroup - from 432 cells/ $\mu$ l to 421.5 ( $p=0.446$ ), in the 4th subgroup from 640  
104 cells/ $\mu$ l to 634 cells/ $\mu$ l ( $p=0.326$ ).

105 A similar trend was found when assessing the median of B cells: in the first  
106 subgroup from 126 cells/ $\mu$ l to 107 cells/ $\mu$ l ( $p=0.717$ ), in the second subgroup - from  
107 117 cells/ $\mu$ l to 135 cells/ $\mu$ l ( $p= 0.808$ ). In addition, there were no statistically  
108 significant differences in the change in the level of median immunological memory  
109 cells. In the 1st subgroup, their value decreased from 564 to 499.5 cells/ $\mu$ l ( $p=0.478$ ),  
110 in the 2nd, 3rd and 4th subgroups it increased from 706.5 to 743.5 cells/ $\mu$ l ( $p$   
111  $=0.520$ ), from 640 to 758.5 cells/ $\mu$ l ( $p=0.542$ ), from 864 to 884.5 cells/ $\mu$ l ( $p=0.412$ ),  
112 respectively.

113           When conducting a statistical analysis, a correlation was found between the  
114 level of B-cells in the peripheral blood and the detection of pneumococcus on the  
115 mucous membrane of the upper respiratory tract. In those examined with a  
116 deficiency of CD19+ lymphocytes in the peripheral blood, an increase in the  
117 frequency of isolation of the microorganism was found ( $p=0.003$ ).

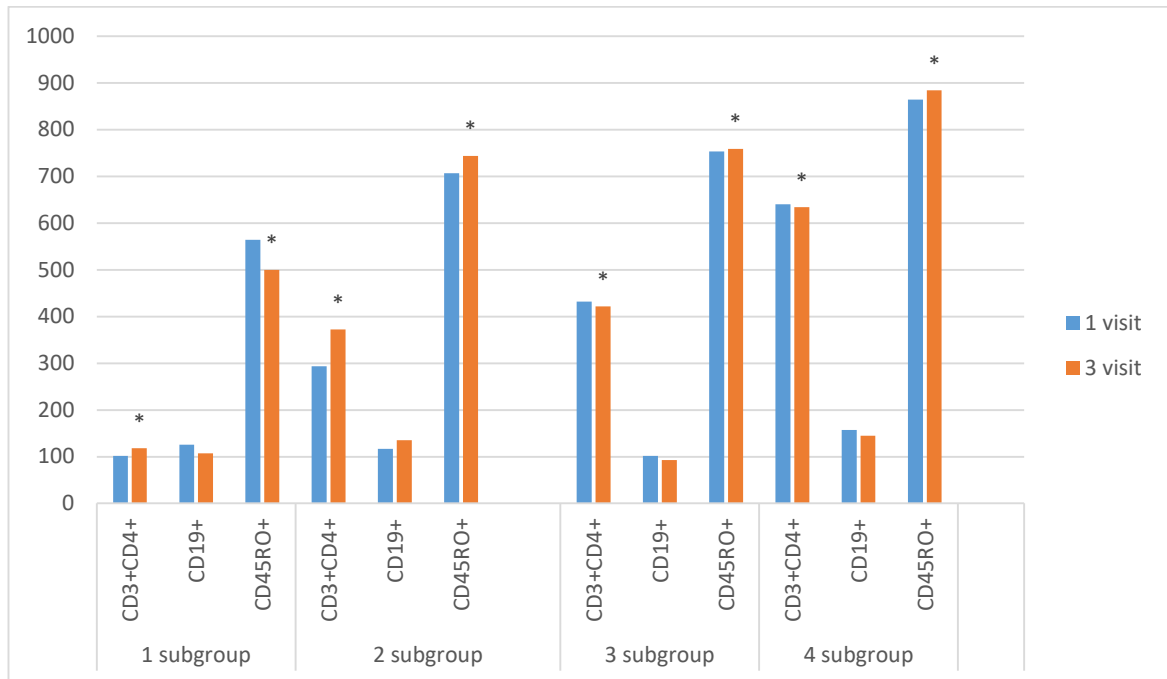
118           Thus, the obtained data correlate with literary sources and confirm the  
119 immunological effectiveness of PCV13 administration [10, 12, 13]. A pronounced  
120 stimulation of the cellular link of the immune system was revealed, which was  
121 expressed in a statistically significant increase in the level of T-helpers in those  
122 examined with a CD3+CD4+ deficiency at the time of the start of the study. It is  
123 important to note that immunization led to an increase in the number of  
124 immunological memory cells, which demonstrates the effectiveness of the  
125 vaccination. At the same time no statistically significant differences were found in  
126 the control group.

#### 127 **4 Conclusion**

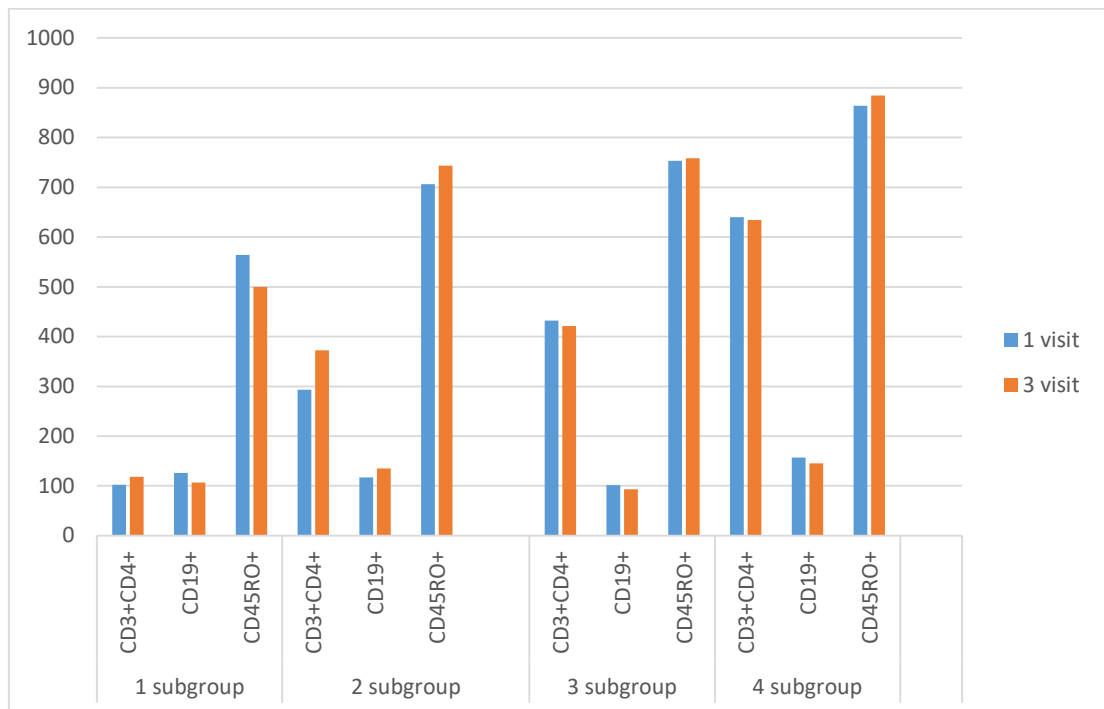
128           Thus, the introduction of a 13-valent pneumococcal conjugate vaccine to  
129 HIV-infected patients causes immunological effects and can effectively reduce the  
130 risk of infections caused by *S. pneumoniae*.

РИСУНКИ

**Figure 1.** The value of the median parameters of the immunogram in patients of the immunized group at visits 1 and 3.



**Figure 2.** The value of the median of immunogram parameters in patients of the control group at visits 1 and 3.



**Note:** \*p<0.05.

## ТИТУЛЬНЫЙ ЛИСТ\_МЕТАДААННЫЕ

### **Блок 1. Информация об авторе ответственном за переписку**

**Каданцева Елизавета Викторовна** – студентка 6 курса; Федеральное государственное бюджетное образовательное учреждение высшего образования "самарский государственный медицинский университет" министерства здравоохранения российской федерации; Кафедра общей и клинической микробиологии, иммунологии и аллергологии ФГБОУ ВО СамГМУ;

адрес: Самара, ул Гагарина, 18, 443079;

телефон: 8(987)437-45-19;

e-mail: lizandria1134@mail.ru

**Kadantseva Elizaveta Victorovna** – Samara State Medical University

Department of General and Clinical Microbiology, Immunology and Allergology of the Samara State Medical University;

address: Samara, Gagarina st., 18, 443079;

telephone: 8(987)437-45-19;

e-mail: lizandria1134@mail.ru

### **Блок 2. Информация об авторах**

**Золотов Максим Олегович** – кандидат медицинских наук, ассистент кафедры общей и клинической микробиологии, иммунологии и аллергологии ФГБОУ ВО СамГМУ.

**Zolotov Maxim Olegovich** – Candidate of Medical Sciences, Assistant of the Department of General and Clinical Microbiology, Immunology and Allergology of the Samara State Medical University.

**Жестков Александр Викторович** – доктор медицинских наук, профессор, заведующий кафедрой общей и клинической микробиологии, иммунологии и аллергологии ФГБОУ ВО СамГМУ.

**Zhestkov Alexander Viktorovich** – Doctor of Medical Sciences, Professor, Head of the Department of General and Clinical Microbiology, Immunology and Allergology of the Samara State Medical University.

**Никитина Татьяна Рудольфовна** – кандидат медицинских наук, доцент кафедры общей и клинической микробиологии, иммунологии и аллергологии ФГБОУ ВО СамГМУ.

**Nikitina Tatyana Rudolfovna** – candidate of Medical Sciences, docent of the Department of General and Clinical Microbiology, Immunology and Allergology of the Samara State Medical University.

**Протасов Андрей Дмитриевич** – доктор медицинских наук, профессор кафедры общей и клинической микробиологии, иммунологии и аллергологии ФГБОУ ВО СамГМУ, врач высшей квалификационной категории по специальности «Аллергология и иммунология».

**Protasov Andrey Dmitrievich** – Doctor of Medical Sciences, Professor of the Department of General and Clinical Microbiology, Immunology and Allergology of the Samara State Medical University, doctor of the highest qualification category in the specialty "Allergology and Immunology".

### **Блок 3. Метаданные статьи**

IMMUNOLOGICAL ASPECTS OF VACCINATION OF HIV-INFECTED PATIENTS

ИММУНОЛОГИЧЕСКИЕ АСПЕКТЫ ВАКЦИНАЦИИ ВИЧ-ИНФИЦИРОВАННЫХ ПАЦИЕНТОВ

**Сокращенное название статьи для верхнего колонтитула:**

IMMUNOLOGICAL ASPECTS OF VACCINATION OF HIV-INFECTED PATIENTS

ИММУНОЛОГИЧЕСКИЕ АСПЕКТЫ ВАКЦИНАЦИИ ВИЧ-ИНФИЦИРОВАННЫХ ПАЦИЕНТОВ

**Ключевые слова:** ВИЧ-инфекция, ПКВ13, Streptococcus pneumoniae, вакцинация, пневмококковая пневмония.

**Keywords:** HIV-infected; PCV13; Streptococcus pneumoniae; vaccination; pneumococcal pneumonia.

Краткое сообщение.

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## СПИСОК ЛИТЕРАТУРЫ

<b>Порядковый номер ссылки</b>	<b>Авторы, название публикации и источника, где она опубликована, выходные данные</b>	<b>ФИО, название публикации и источника на английском</b>	<b>Полный интернет-адрес (URL) цитируемой статьи и/или DOI</b>
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