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The virulence of mixed infection with *Streptococcus constellatus* and *Fusobacterium nucleatum* in a murine orofacial infection model

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1 **ABSTRACT** - Orofacial infections are usually polymicrobial, and it is the microbial $\mathbf{2}$ interactions of pathogenic species that cause tissue destruction. In this study, the 3 microbial interaction between Streptococcus constellatus and Fusobacterium 4 nucleatum was characterized using a murine orofacial infection model. A mixture of viable S. constellatus and F. nucleatum cells (both $2 \ge 10^8$ cfu/mice) was injected into $\mathbf{5}$ 6 the submandible; as a result, all of the test mice died. In contrast, none of the 7 experimental animals monoinjected with either S. constellatus or F. nucleatum died 8 (p<0.001), indicating that the synergism between the two resulted in the virulence. 9 When a mixture of viable S. constellatus cells and a culture filtrate of F. nucleatum 10 was tested, lethality and the bacterial cell count per lesion were significantly enhanced 11 as compared with monoinjections (p < 0.02). However, the virulence of F. nucleatum 12was not enhanced by infection of a culture filtrate of S. constellatus. The 13enhancement of virulence was observed even when viable S. constellatus cells and the 14culture filtrate of F. nucleatum were injected at separate sites. Heat-treatment of the 15culture filtrate of *F. nucleatum* did not affect the enhancement. These results indicate 16 that a heat-stable substance(s) produced by F. nucleatum contributes to the microbial 17synergy of S. constellatus and F. nucleatum in orofacial infections.

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virulence synergy / Streptococcus constellatus / Fusobacterium nucleatum /
 orofacial infection / murine model

21

1 **1. Introduction**

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3 Most human orofacial infections involve bacteria that reside in the oral cavity, 4 particularly viridans streptococci, Peptostreptococcus species, Prevotella species, and $\mathbf{5}$ *Fusobacterium* species [1-8]. Moreover, such orofacial infections are usually 6 Previous studies have shown that facultative streptococci and polymicrobial. Fusobacterium nucleatum along with other oral bacterial species are often isolated 7 8 simultaneously from odontogenic infections [7,8], and that the Streptococcus milleri 9 group and F. nucleatum are isolated more frequently from severe orofacial infections 10 than from mild infections [6]. Several investigators have examined the pathogenicity 11 of Streptococcus constellatus, a member of the S. milleri group, and F. nucleatum 12using animal infection models and have demonstrated that these two bacteria have the 13potential to produce abscesses [9-14]. These studies indicate that bacterial interaction 14is one of the most important factors in the occurrence and progress of orofacial 15infections [10-14]. In this study, we attempted to elucidate the nature of the bacterial 16interaction between S. constellatus and F. nucleatum in a murine orofacial infection 17model [15] using the mouse submandible as the injection site.

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20 **2. Materials and methods**

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- 22 **2.1.** Bacterial strains and preparation of bacterial inocula
- 23

24 2.1.1. Bacteria

25 S. constellatus ATCC 27823 and F. nucleatum ATCC 25586 [15] were used in the

present study. The colony forming unit(s) (cfu) of the bacterial inocula was
 determined by counting the number of bacterial colonies grown under the same
 conditions as those described below.

- 4
- 5 2.1.2. Cell suspension

6 Colonies of S. constellatus were cultured on Brucella HK agar (Kyokuto 7Pharmaceutical Industrial Co., Tokyo, Japan) with 5 % (v/v) sheep blood in an 8 atmosphere of 10 % CO₂ (v/v), 20% H₂ (v/v), and 70 % N₂ (v/v), at 37 °C for 48 h. F. 9 nucleatum was cultured on Brucella HK agar with 5 % sheep blood in an atmosphere 10 of 5 % CO₂ (v/v), 10 % H₂ (v/v), and 85 % N₂ (v/v), at 37 °C for 78 h. The resultant 11 colonies were collected and suspended in a Peptone-Yeast-Glucose (PYG) broth (1 % 12(w/v) Bacto-peptone (Becton Dickinson, Cockeysville, MD, USA), 1 % (w/v) yeast 13extract (Becton Dickinson), 1 % (w/v) glucose, and 0.01 % (w/v) vitamin K (Wako Pure Chemical Industries, Osaka, Japan), pH 7.0) at concentrations between 1.0 x 10⁹ 14and $4.0 \ge 10^9$ cfu/ml. 15

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17 2.1.3. Broth culture

Fluid cultures of *S. constellatus* and *F. nucleatum* were prepared in PYG broth under the following atmospheric conditions: 10 % CO₂ (v/v), 20 % H₂ (v/v), and 70% N₂ (v/v) at 37 °C for 48 h for *S. constellatus*, and 5 % CO₂ (v/v), 10% H₂ (v/v), and 85 % N₂ (v/v) at 37 °C for 72 h for *F. nucleatum*. Each PYG broth culture was used as the culture for the examination of the synergistic effect of these strains on virulence. The bacterial concentrations of *S. constellatus* and *F. nucleatum* in broth cultures were 4.0×10^9 cfu/ml and 7.0 x 10^9 cfu/ml, respectively.

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1 2.1.4. Culture filtrate

For the preparation of bacteria-free culture filtrates, fluid cultures were centrifuged at 600 x g for 10 min and the supernate was sterilized by 0.22 μ m pore size membrane filtration (Millipore, Bedford, MA, USA). The pH values of these filtrates were in the range 6.5 - 6.7. A heat-treated culture filtrate was prepared with an autoclave at 121 °C for 20 min.

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8 **2.2. Animals and inoculation in mice**

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ICR Crj CD-1 mice (Charles River Japan Inc., Yokohama, Japan) raised under
conventional conditions were used. The mice were six-week-old females (25 - 28g).
A murine orofacial infection model [15] was employed for the experiments.

13

14 2.2.1. The synergistic effect of living cells on virulence

An aliquot of 50 µl of a cell suspension or a broth culture of one strain was mixed
with an equal volume of a cell suspension or a broth culture of the other strain (Table
I). In the control group, uncultured sterile PYG broth was used instead of the cell
suspension or the broth culture. The pH value of the mixture was in the range 6.8 7.0.

The mixture was injected into the submandible of each mouse, as described previously [15]; the mice were anesthetized with diethyl ether (Wako Pure Chemical Industries) and the skin at the submandible was disinfected with 70 % (v/v) ethanol. The skin was pricked with a 26 gauge needle along the midline of the submandible and an aliquot of 100 μ l of the mixture was injected into the space between the skin and smooth muscular layers at the center of the oral floor. 1

2 2.2.2. The synergistic effect on virulence of a culture filtrate and a cell suspension
3 when injected at the same site

An aliquot of 50 µl of a cell suspension of one strain was mixed with an equal
volume of a culture filtrate of the other strain. In the control group, uncultured sterile
PYG broth was used instead of the culture filtrate. The pH value of the mixture was
in the range 6.8 - 7.0. The mixture was injected into the submandible of each mouse.

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9 2.2.3. The synergistic effect on virulence of a culture filtrate and a cell suspension
10 when injected at separate sites

An aliquot of 50 µl of a cell suspension of one test strain mixed with an equal
amount of uncultured sterile PYG broth was injected into the submandible of the mice.
At the same time, an aliquot of 50 µl of a culture filtrate of the other test strain was
injected into the back of the same mice (Table IV). In the control group, uncultured
sterile PYG broth was injected into the back of mice instead of the culture filtrate.

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17 **2.3.** Assessment

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Bacterial virulence was assessed by (1) the presence or absence of abscesses, (2) lethality, (3) the bacterial count per lesion. The mice were sacrificed by cervical dislocation. The formation of a submandibular abscess was determined by visual inspection on the seventh day after the injection. The bacterial count of the abscesses was determined as follows: the collected submandibular abscesses were homogenated with 1 ml of sterile PYG broth, and the homogenated abscesses were diluted at serial 10-folds. Each diluted material was cultured on Brucella HK agar with 5 % (v/v)

1	sheep blood under the conditions stated before. The number of bacterial colonies
2	grown on the blood-agars were counted. In addition, the above culture confirmed the
3	presence of the test bacterial species and the absence of contamination.
4	
5	2.4. Statistical analysis
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7	Statistical comparisons of lethality and the abscess formation rate were performed
8	by a x^2 test. The statistical differences in the bacterial cell count per murine lesion
9	were analyzed by Student's-t test.
10	
11	
12	3. Results
13	
14	3.1. Virulence of living cells
15	
16	In the present animal model, viable cells of S. constellatus ATCC 27823 produced a
17	single abscess, and those of F. nucleatum ATCC 25586 produced a single abscess or
18	multiple abscesses at the injection site.
19	Although injection of pure cell suspensions of S. constellatus and F. nucleatum
20	produced abscesses in the submandible, all of the mice in this group survived (Table I).
21	The injection of pure PYG broth cultures of S. constellatus (2.0 x 10^8 cfu) or F.
22	nucleatum (3.5 x 10^8 cfu) failed to produce lesions. When a mixture of cell
23	suspensions of both S. constellatus and F. nucleatum, a mixture of a cell suspension of
24	S. constellatus and a broth culture of F. nucleatum, or a mixture of a cell suspension of
25	F. nucleatum and a broth culture of S. constellatus was injected, all of the mice died

- 1 (Table I).
- $\mathbf{2}$

3 3.2. Virulence of a culture filtrate and a cell suspension when injected at the same 4 site

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6 When a cell suspension of F. nucleatum mixed with a culture filtrate of S. 7 constellatus was injected into the submandible, the rate of abscess formation, the 8 lethality, and the bacterial cell count per lesion were the same as those produced by F. *nucleatum* alone, even at a concentration of 2.0×10^8 cfu/mouse (Table II). However, 9 10 when a bacterial cell suspension of S. constellatus was injected with a culture filtrate 11 of F. nucleatum, enhanced virulence was observed (Table III). When a concentration 12of 5.0 x 10^7 cfu of S. constellatus was injected with a F. nucleatum culture filtrate, 13three mice died and the remaining seven mice developed abscesses (Table III). In 14contrast, all of the mice in the control group survived and only six mice formed 15The bacterial cell count of S. constellatus isolated from lesions in the abscesses. experimental group was also significantly larger than that of the control group 16(p<0.02). When a higher number of S. constellatus cells (2.0 x 10^8 cfu/mouse) was 17injected in the same manner, virulence again significantly increased (Table III). In 18 19 this group, eight mice died and the remaining two mice developed abscesses with a 20bacterial cell count of $7.35 \pm 0.04 \text{ Log}_{10}$ cfu/lesion. This cell count was significantly 21higher than that of the control group (p<0.02). Furthermore, the heat-treated culture 22filtrate of *F. nucleatum* also enhanced the virulence of *S. constellatus*.

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3.3. Virulence of a culture filtrate and a cell suspension when injected at separatesites

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 $\mathbf{2}$ When a cell suspension of F. nucleatum was injected into the submandible and a 3 culture filtrate of S. constellatus was injected into the back of the mice, the rate of 4 abscess formation, lethality, and the bacterial cell count did not increase as compared with those of the control group (data not shown). In contrast, when a cell suspension $\mathbf{5}$ 6 of S. constellatus was injected into the submandible of the mice along with a 7 subcutaneous injection of a culture filtrate of F. nucleatum into the back, the virulence of S. constellatus was enhanced (Table IV). When 5.0 x 10^7 cfu of S. constellatus 8 9 cells were injected with F. nucleatum culture filtrate at separate sites, the bacterial cell 10 count was significantly greater (p < 0.05) than that of the control group. When 2.0 x 10⁸ cfu of S. constellatus cells were injected in the same manner, four mice died and 11 12the remaining six mice developed abscesses. The bacterial cell count was 13significantly greater (p<0.05) than that of the control group (Table IV). In cases 14which received injections of heat-treated F. nucleatum culture filtrate, the bacterial cell 15count was significantly greater (p<0.05) than that of the control group (Table IV).

16We further investigated the effect of the F. nucleatum culture filtrate upon the 17virulence of S. constellatus by injecting a standardized S. constellatus cell suspension $(5.0 \times 10^7 \text{ cfu})$ with a 10-fold dilution of F. nucleatum culture filtrate at the same site 18 19 and also at separate sites. In the group with same site injection, abscesses formed in 20all of the 10 mice used (Table V). In the group with the separate site injection, the 21effect was weaker than that observed in the same site injection group; however, 22abscesses formed in response to the 10^1 -fold dilution in all of the 10 mice used. The 23number of S. constellatus cells per lesion was significantly larger (p<0.05) than that of 24the control group when a 10^3 -fold dilution of the filtrate was used, irrespective of 25whether the mice received same site or different site injections (Table V).

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3 4. Discussion

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 $\mathbf{5}$ In a previous study, we determined the lesion type formed by clinical isolates of S. 6 constellatus and F. nucleatum in the orofacial infection model: S. constellatus 7 produced a single abscess, and F. nucleatum produced a single abscess or multiple 8 abscesses at the submandible in mice [15]. Moreover, we obtained the same 9 experimental results regarding lesion type in the present study by using not only 10 ATCC type strains but also clinical isolates of S. constellatus and F. nucleatum, 11 although the clinically isolated strains differed from the ATCC strains in terms of 12bacterial number in the abscesses formed (data not shown).

13Microbial interactions in polymicrobial infections have been emphasized in the 14literature [16], and the microbial synergy of pathogens involved in odontogenic 15infections has also been studied [10-14]. The microbial synergy of S. constellatus 16 and F. nucleatum has been described in a previous study using animal subcutaneous 17abscess models, in which the groin was used as the injection sites [12]. Similarly, in 18 this study, the synergistic effect of S. constellatus and F. nucleatum was observed with 19 the murine orofacial infection model using the submandible as the injection site. 20Furthermore, it was shown that bacterial virulence depended on the preparation of the 21bacterial inocula. Pure PYG broth cultures of S. constellatus and F. nucleatum did 22not induce any lesions, although pure cell suspensions of both bacteria did induce 23lesions (Table I). The virulence of some anaerobes is demonstrated to be influenced 24greatly by the type and amount of nutrients contained in the medium in which the 25bacteria are cultured [17,18]. The findings observed in this study may be partly due

1 to the different expression of virulence factors that are directly and indirectly regulated $\mathbf{2}$ by different nutrients contained in the media. The virulence factors of S. constellatus 3 and F. nucleatum have been previously reported [9,19]. In particular, it has been 4 suggested that the possession of a bacterial capsule relates closely to the induction of The virulence factors, especially the bacterial $\mathbf{5}$ abscess formation [9,10,20,21]. 6 capsule, may contribute to the varied pathogenic potentials of different strains, 7 although the present study did not assess these factors in the test strains.

8 When a cell suspension of F. nucleatum was injected into mice with a culture 9 filtrate of S. constellatus at the same site, enhancement of F. nucleatum virulence was 10 not observed (Table II). Virulence was not enhanced even when the culture filtrate of 11 S. constellatus was injected two or three times a day for seven days (data not shown). 12Interestingly, when a cell suspension of F. nucleatum was injected with a broth culture 13of S. constellatus, a synergistic effect on virulence was observed (Table I), resulting in 14the death of all of the mice used. These results indicate that viable S. constellatus 15cells, and not S. constellatus products in culture filtrate, enhanced F. nucleatum 16 virulence. As regards the enhancement of F. nucleatum virulence, the cell-to-cell 17interaction between F. nucleatum and S. constellatus may be important.

18 A culture filtrate of F. nucleatum enhanced S. constellatus virulence when the 19 filtrate and a cell suspension of S. constellatus were injected at the same site (the 20submandible) (Table III). These findings indicate that a product(s) of F. nucleatum 21in the culture filtrate enhanced S. constellatus virulence. Even when a cell 22suspension of S. constellatus and a culture filtrate of F. nucleatum were injected at 23separate sites, the enhancement of S. constellatus virulence was observed (Table IV). 24Four principal mechanisms have been identified by which bacteria may interact to 25increase the net pathogenicity of the infection: (1) the effects on host defenses, in

1 particular, inhibition of phagocytosis, (2) the provision of essential nutrients, (3) the $\mathbf{2}$ improvement of the local environment, and (4) the increased virulence of the 3 organisms [22]. We presume that the mechanism of the synergistic effect is as 4 follows: (1) the culture filtrate of F. nucleatum affected the host cell's function, and $\mathbf{5}$ this alteration of host defense against S. constellatus, e.g., the change in host immunity, 6 enhanced S. constellatus virulence; (2) the culture filtrate of F. nucleatum injected into 7 the backs of mice was transmitted to the submandible through hematogenous pathways, 8 and some substance(s) in the filtrate enhanced S. constellatus virulence directly at the 9 submandible. Some investigators have demonstrated that F. nucleatum inhibits the 10 normal function of host cells; certain molecular candidates such as proteins and 11 volatile fatty acids are possibly associated with this phenomenon [23-25]. The 12present results indicate that a heat-stable product(s) of F. nucleatum plays an important 13role in the enhancement of S. constellatus virulence. Further experimental study is 14required to characterize this heat-stable substance and to clarify the mechanism by 15which this substance affects host defenses. The enhancement of S. constellatus virulence by the F. nucleatum culture filtrate was observed even at a 10^3 -fold dilution 16 17This finding suggests that a culture filtrate of F. nucleatum of the filtrate (Table V). 18 possesses great potential to elicit a synergistic effect with S. constellatus as regards 19 virulence. In this respect, even when S. constellatus is accompanied by only a small 20number of F. nucleatum, it may express great virulence in an oral lesion involving 21various pathogens. It appears, therefore, that the microbial synergism between S. 22constellatus and F. nucleatum contributes to a large extent to the occurrence and 23progression of orofacial infections.

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Abbreviations

PYG broth: Peptone-Yeast-Glucose broth

cfu: colony forming unit(s)

Injection combination		No.of	No. of	Bacterial cell count	
S. constellatus	F. nucleatum	dead mice	abscess-forming mice	per lesion $(\pm SD, Log_{10} cfu/mouse)$	
Cell suspension	None	0	8	5.45 ± 0.93^{c}	
Cell suspension	Cell suspension	10 ^{<i>a</i>, <i>b</i>}	NA	NA	
Cell suspension	Broth culture	10 ^{<i>a</i>, <i>b</i>}	NA	NA	
Broth culture	None	0	0	0.00	
Broth culture	Cell suspension	10 ^{<i>a</i>, <i>b</i>}	NA	NA	
None	Cell suspension	0	10	7.39 ± 0.29^{d}	
None	Broth culture	0	0	0.00	

Ten mice were used in each group.

- ^{*a*} p<0.001 vs. the cell suspension or the broth culture of *S. constellatus* alone.
- ^{*b*} p<0.001 vs. the cell suspension or broth culture of *F. nucleatum* alone.
- ^c Bacterial cell count of *S. constellatus* per lesion
- ^d Bacterial cell count of *F. nucleatum* per lesion
- NA, not applicable because all mice died before abscess formation.

Injection cor	No. of	No. of	Bacterial cell count of	
<i>F. nucleatum</i> ^{<i>a</i>} (cfu/mouse)	S. constellatus	dead mice	abscess-forming mice	<i>F. nucleatum</i> per lesion $(\pm SD, \text{Log}_{10} \text{ cfu/mouse})$
5.0 x 10 ⁷	None	0	9	7.25 ± 0.69
5.0 x 10 ⁷	Culture filtrate	0	10	7.20 ± 0.38
5.0 x 10 ⁷	Heat-treated culture filtrate	0	8	7.01 ± 0.46
2.0 x 10 ⁸	None	0	10	7.39 ± 0.29
2.0 x 10 ⁸	Culture filtrate	0	10	7.37 ± 0.62
2.0 x 10 ⁸	Heat-treated culture filtrate	0	9	7.15 ± 0.33

Table II. Effect of *S. constellatus* culture filtrate on the virulence of *F. nucleatum* when the mixture of a cell suspension of *F. nucleatum* and a culture filtrate of *S. constellatus* was challenged at the same site (submandible)

Ten mice were used in each group.

^a cell suspension.

Injection combination		No. of	No. of	Bacterial cell count of	
S. constellatus ^a (cfu/mouse)	F. nucleatum	dead mice	abscess-forming mice	S. constellatus per lesion (\pm SD, Log ₁₀ cfu/mouse)	
5.0 x 10 ⁷	None	0	6	4.40 ± 0.20	
5.0 x 10 ⁷	Culture filtrate	3	7	6.34 ± 0.14 ^c	
5.0 x 10 ⁷	Heat-treated culture filtrate	1	9	5.53±0.42 ^c	
2.0 x 10 ⁸	None	0	8	5.45 ± 0.93	
2.0 x 10 ⁸	Culture filtrate	8 ^b	2	7.35 ± 0.04 ^c	
2.0 x 10 ⁸	Heat-treated culture filtrate	6 ^{<i>b</i>}	4	6.82 ± 0.42 ^c	

Table III. Effect of *F. nucleatum* culture filtrate on the virulence of *S. constellatus* when the mixture of a cell suspension of *S. constellatus* and a culture filtrate of *F. nucleatum* was challenged at the same site (submandible)

Ten mice were used in each group.

^{*a*} cell suspension.

 b,c p<0.02 vs. the cell suspension of *S.constellatus* alone.

Table IV. Effect of *F. nucleatum* culture filtrate on the virulence of *S. constellatus* which were injected at different sites when a bacterial cell suspension of *S. constellatus* and a culture filtrate of *F. nucleatum* were injected into the submandible and the subcutaneous tissue of the back of mice, respectively

	Injection combination		No. of		No. of	Bac	Bacterial cell count of	
	onstellatus ^a /mouse)	F. nucleatum	dead mice		abscess-formin mice	0	S. constellatus per lesion $(\pm SD, Log_{10} cfu/mouse)$	
5.0	0 x 10 ⁷	None		0	6		4.71 ± 0.17	
5.0) x 10 ⁷	Culture filtrate		0	10		6.49 ± 0.48 ^b	
5.0) x 10 ⁷	Heat-treated culture filtrate		0	9		$5.50 \pm 0.43 \ ^{b}$	
2.0) x 10 ⁸	None		0	8		5.54 ± 0.90	
2.0) x 10 ⁸	Culture filtrate		4	6		$6.77 \pm 0.31 \ ^{b}$	
2.0) x 10 ⁸	Heat-treated culture filtrate		1	9		6.43 ± 0.43^{b}	

Ten mice were used in each group.

^{*a*} cell suspension.

^{*b*} p<0.05 vs. the cell suspension of *S. constellatus* alone.

Injection combination		Injecti	on at same site	Injection at separate sites		
S. constellatus	F. nucleatum	No. of abscess-	Bacterial count of <i>S. constellatus</i>	No. of abscess-	Bacterial count of <i>S. constellatus</i>	
Cells suspension Culture filt (cfu/mouse) dilution		forming mice (per lesion \pm SD, Log ₁₀ cfu/mouse)	forming mice	per lesion $(\pm SD, Log_{10} cfu/mouse)$	
5.0 x 10 ⁷	None	6	4.40 ± 0.20	6	4.71 ± 0.17	
5.0 x 10 ⁷	10 ⁰	10	6.34 ± 0.14 ^{<i>a</i>}	10	$6.49 \pm 0.48 \ ^{b}$	
5.0 x 10 ⁷	10 ¹	10	5.58±0.45 ^a	10	5.44 ± 0.64 ^b	
5.0 x 10 ⁷	10 ²	10	5.90±0.68 ^a	7	$5.50 \pm 0.10^{\ b}$	
5.0 x 10 ⁷	10 ³	8	5.36 ± 0.12^{a}	7	5.53±0.07 ^b	
5.0 x 10 ⁷	10 4	6	5.26 ± 0.90	6	4.94 ± 0.60	

Table V. Effect of various concentrations of *F. nucleatum* culture filtrate on the virulence of *S. constellatus* when the cell suspension of *S. constellatuas* and a culture filtrate of *F. nucleatum* were challenged at the same site or separate sites

Ten mice were used in each group.

^{*a*, *b*} p<0.05 vs. the cell suspension of *S*. *constellatus* alone.