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Abstract	It was previously demonstrated that brief ( $\leq 1$ h) exposures to echinocandins are as effective to kill <i>Candida albicans</i> cells as continuous 24-h exposure. However, killing rates after continuous and short (1 h) echinocandin exposures to <i>C. albicans</i> have not yet been evaluated in RPMI-1640 with and without 50 % serum. We evaluated four echinocandin susceptible <i>C. albicans</i> bloodstream isolates, ATCC 10231 type strain and an echinocandin-resistant isolate (DPL20, FKS F645P). Caspofungin MICs, time-kill and postantifungal effect (PAFE) tests were performed in RPMI-1640 with and without 50 % serum. Killing rates ( <i>k</i> values) in time-kill and PAFE experiments were determined for each strain and concentration. In time-kill experiments, colony count decreases were isolate- and concentration-dependent at 0.25, 1, 4, 8, 16 and 32 mg/L in RPMI-1640, but concentration-independent at 1, 4, 8, 16 and 32 mg/L in 50 % serum. One-hour caspofungin exposure at 4, 16 and 32 mg/L resulted in CFU decreases comparable with the results obtained in time-kill experiments in RPMI-1640, but 50 % serum at 4, 16 and 32 mg/L allowed growth of all isolates ( <i>k</i> values were negative) ( <i>P</i> < 0.05–0.001). PAFE in 50 % serum decreased markedly at 4, 16 and 32 mg/L. Killing rates remained high and concentration-independent in 50 % serum in case of continuous but not in case of brief caspofungin exposure. As only a short growth inhibition without killing was observed in 50 % serum, clinical relevance of caspofungin PAFE in vivo is questionable.					
Keywords (separated by '-')	Echinocandins - Serum-ba	ased susceptibility testing - Postantifungal effect				
Footnote Information						

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# Killing Rates for Caspofungin Against *Candida albicans* After Brief and Continuous Caspofungin Exposure in the Presence and Absence of Serum

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11 Abstract It was previously demonstrated that brief 12  $(\leq 1 h)$  exposures to echinocandins are as effective to 13 kill Candida albicans cells as continuous 24-h expo-14 sure. However, killing rates after continuous and short 15 (1 h) echinocandin exposures to C. albicans have not 16 yet been evaluated in RPMI-1640 with and without 17 50 % serum. We evaluated four echinocandin suscep-18 tible C. albicans bloodstream isolates. ATCC 10231 19 type strain and an echinocandin-resistant isolate 20 (DPL20, FKS F645P). Caspofungin MICs, time-kill 21 and postantifungal effect (PAFE) tests were performed 22 in RPMI-1640 with and without 50 % serum. Killing 23 rates (k values) in time-kill and PAFE experiments 24 were determined for each strain and concentration. In 25 time-kill experiments, colony count decreases were isolate- and concentration-dependent at 0.25, 1, 4, 8, 26 27 16 and 32 mg/L in RPMI-1640, but concentrationindependent at 1, 4, 8, 16 and 32 mg/L in 50 % serum. 28 29 One-hour caspofungin exposure at 4, 16 and 32 mg/L

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resulted in CFU decreases comparable with the results 30 obtained in time-kill experiments in RPMI-1640, but 31 50 % serum at 4, 16 and 32 mg/L allowed growth of 32 all isolates (k values were negative) (P < 0.05-0.001). 33 PAFE in 50 % serum decreased markedly at 4, 16 and 34 32 mg/L. Killing rates remained high and concentra-35 tion-independent in 50 % serum in case of continuous 36 but not in case of brief caspofungin exposure. As only 37 a short growth inhibition without killing was observed 38 in 50 % serum, clinical relevance of caspofungin 39 PAFE in vivo is questionable. 40

KeywordsEchinocandins · Serum-based41susceptibility testing · Postantifungal effect42

#### Introduction

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Echinocandins (caspofungin, micafungin and anidu-44 lafungin) are reported to exhibit concentration-depen-45 dent fungicidal or fungistatic activity against the 46 majority of Candida species. Additionally, echino-47 candins exert prolonged postantifungal effect (PAFE) 48 after short (1 h) echinocandin exposure against many 49 *Candida* species, including *C. parapsilosis* [1–4]. 50 Recent in vitro findings with RPMI-1640 suggest that 51 a very short ( $\leq 1$  h) exposure to caspofungin kills 52 Candida cells as effectively as a continuous 24-h 53 exposure [5]. As echinocandins are highly protein-54 bound (≥96.5 %) agents (i.e., serum fundamentally 55

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56 influences killing activities of echinocandins) [6-8], 57 we compared killing rates produced by short (1 h) and 58 continuous (24 h) caspofungin exposure as well as 59 PAFE in RPMI-1640 and 50 % serum against C. 60 albicans.

#### 61 **Materials and Methods**

#### 62 Isolates

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All C. albicans originated from blood samples (Table 1) and were identified as described previously [9]. ATCC 10231 and an echinocandin-resistant isolate (DPL20, FKS F645P) were also included in the study.

68 MIC Determination

69 Caspofungin (Sigma, Budapest, Hungary) MICs in 70 RPMI-1640 with and without 50 % human serum 71 (from a human male, type AB, Sigma, Budapest, 72 Hungary) were determined using the CLSI broth macrodilution method [6-8, 10]. Caspofungin final 73 74 concentration ranged between 0.015 and 32 mg/L. 75 MICs were read after 24 h using the partial inhibition 76 criterion [10].

77 Postantifungal Effect and Time-Kill Curves

78 PAFE was measured in both media simultaneously. As in our preliminary experiments 5-, 10- and 30-min 79 80 exposures to 0.5, 1 or 2 mg/L caspofungin did not produce measurable PAFEs in 50 % serum 81

Table 1 Candida albians isolatos MICs of asspotungin

Fig. 1 Time-kill plots of caspofungin against four Candida albicans isolates (averages  $\pm$  standard deviation) in RPMI-1640 (a) and 50 % serum (d) against ATCC 10231 type strain in RPMI-1640 (b) and 50 % serum (e), and against the echinocandin-resistant C. albicans DPL20 in RPMI-1640 (c) and 50 % serum (f). Clinical isolates and type strain were exposed to 0.25, 1, 4, 8, 16 and 32 mg/L, while DPL20 isolate was exposed to 4, 8, 16 and 32 mg/L of caspofungin for 24 h (continuous caspofungin exposure)

 $(1-16 \times \text{MIC in 50 \% serum})$ , we used caspofungin 82 at 4, 16 and 32 mg/L concentrations with a 60-min 83 exposure time (brief caspofungin exposure) [3–5]. As 84 the maximum administrable daily 150-200 mg ca-85 spofungin doses produce 30.4-40.6 mg/L geometric 86 mean of peak concentrations in humans [11], the 87 highest caspofungin concentration used in this study 88 was 32 mg/L [6]. 89

The starting inocula in PAFE experiments were 90  $1-5 \times 10^5$  cells/ml [3-5]. After 1 h, the cells were 91 collected by centrifugation at  $1.500 \times g$  for 10 min and 92 were washed three times with sterile saline, resus-93 pended in 10 ml drug-free warm RPMI-1640 with and 94 without 50 % human serum. Samples (100 µl) were 95 removed at 0, 4, 8, 12 and 24 h, serially diluted 96 tenfold, plated  $(4 \times 30 \ \mu l)$  onto Sabouraud dextrose 97 agar and incubated at 35 °C for 48 h [3–5]. 98

For time-kill assays (continuous 24-h caspofungin 99 exposure), test solutions were not centrifuged or 100 washed. We used 0.25, 1, 4, 8, 16 and 32 mg/L 101 caspofungin. In case of isolate DPL20, we used 4, 8, 102 16 and 32 mg/L caspofungin. Otherwise, the method 103 was the same as described for PAFE [3-5]. All 104 experiments were performed at least twice, and means 105 of data are presented. 106

and the offect cosposition in time kill studies in **PDMI** 1640 (**PDMI**) and

	Cununuu uibicui	is isolates	s, which of	casporungin ai	iu uie ciree	i casporung	III III UIIIC-K	in studie	cs m	KI WII-10-	+0 (IXI IVII	<i>i)</i> and
RPMI-16	540 supplemented	1 with 50	% serum	(50 % serum)								
	11			· /								
r 1.		MIC	/T \	E CC	1.11					(T.)		

Isolates	MIC (mg/L)	Effect in time-kill s	Effect in time-kill studies at concentrations shown (mg/L)						
	RPMI	50 % Serum	RPMI	50 % Serum					
183	0.03	0.25	≥0.03 fungistatic	≥0.25 fungistatic					
3666	0.03	0.125	≥0.03 fungistatic	$\geq 0.25$ fungistatic					
12132	0.015	0.25	≥0.03 fungistatic	$\geq 0.25$ fungistatic					
10920	0.03	0.125	≥0.03 fungistatic	$\geq 0.25$ fungistatic					
ATCC 10231	0.03	0.5	≥0.03 fungistatic	$\geq 0.5$ fungistatic					
DPL20	4	>32	$\geq 16$ fungistatic	No effect					



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32 mg/L

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32 mg/L

#### 107 Data Analysis

108 Killing kinetics was analyzed mathematically, as 109 described previously [6, 12]. An exponential equation 110 was fitted to the mean data at each time point: 111  $N_t = N_0 \times e^{-kt}$ , where  $N_t$  is the number of viable 112 yeasts at time t,  $N_0$  is the number of viable yeasts in the initial inoculum, k is the killing rate and t is the 113 incubation time. Negative k values indicate growth, 114 and positive k values indicate killing. The goodness of 115 fit for each isolate was assessed by the  $r^2$  value (>0.8) 116 [6, 12]. PAFE was defined as the difference between 117 the time required for control and test isolates to grow 1 118  $\log_{10}$  following drug removal [3–5]. 119



**Fig. 2** Postantifungal effect curves of caspofungin against *Candida albicans* isolate 183 in RPMI-1640 (**a**) and 50 % serum (**c**), and against *C. albicans* isolate 3666 in RPMI-1640 (**b**) and

50 % serum (**d**). Isolates were exposed to 4, 16 and 32 mg/L of caspofungin for 1 h (brief caspofungin exposure)

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One-way ANOVA with Tukey's posttesting was
used to analyze differences in killing rates by
concentrations in either RPMI-1640 or 50 % serum
[6]. Paired *T* test was used to compare the effect of the
medium as well as the killing and growth rates in timekill and PAFE experiments at the same drug
concentration.

Results

## Susceptibility 128

MIC values are presented in Table 1. Paradoxical129growth was not observed. Clinical isolates and 10231130ATCC strain were susceptible to caspofungin131

 Table 2
 Postantifungal effect (PAFE) of caspofungin against Candida albicans isolates at 4, 16 and 32 mg/L in RPMI-1640 (RPMI) and RPMI-1640-50 % serum (50 % serum)

Isolate number	Medium	PAFE in hours					
		4 (mg/L)	16 (mg/L)	32 (mg/L)			
183	RPMI-1640	>19.34	>19.34	14.81			
	50 % Serum	2.27	10.14	>18.79			
3666	RPMI-1640	7.40	>19.23	>19.23			
	50 % Serum	0	0.79	1.01			
10920	RPMI-1640	>19.05	>19.05	>19.05			
	50 % Serum	0	0.24	0.24			
12132	RPMI-1640	14.21	13.84	7.65			
	50 % Serum	0.60	0.95	3.46			
ATCC 10231	RPMI-1640	4.89	>19.88	>19.88			
	50 % Serum	0.09	0.34	1.39			
DPL20	RPMI-1640	0	0	0			
	50 % Serum	0	0	0			

Table 3	Maximum log	changes	in log C	FU/mL	compared	to starting	inoculum	in ti	ime-kill	and	postantifungal	(PAFE)	studies in
RPMI-16	40 and RPMI-	1640-50 %	% serum	(50 % :	serum)								

Isolate number	Media	Maximum log decreases in CFU in time-killing and PAFE experiments at the indicated caspofunging concentration								
		4 mg/L		16 mg/L		32 mg/L				
		Time kill	PAFE	Time kill	PAFE	Time kill	PAFE			
183	RPMI-1640	-1.1	-0.53	-1.18	-0.42	-0.75	-0.48			
	50 % serum	-1.06	+0.37	-0.9	-0.08	-0.9	-0.04			
3666	RPMI-1640	-1.38	-0.66	-0.54	-1.6	-0.34	-2.08			
	50 % serum	-1.40	+0.13	-1.12	+0.12	-2.68	-0.12			
10920	RPMI-1640	-1.23	-1.23	-1.22	-1.48	-0.69	-1.48			
	50 % serum	-0.69	+0.49	-1.46	+0.38	-1.43	+0.64			
12132	RPMI-1640	-0.44	-0.30	-0.19	-0.27	-0.23	-0.62			
	50 % serum	-1.56	+0.67	-1.71	+0.66	-1.86	+0.27			
ATCC 10231	RPMI-1640	-1.56	-1.48	-0.55	-2.08	-1.65	-1.78			
	50 % serum	-1.18	+0.26	-1.38	+0.09	-1.95	+0.08			
DPL20	RPMI-1640	-0.39	+0.61	-1.19	+0.78	-1.00	+0.37			
	50 % serum	+0.81	+0.84	+0.69	+0.57	-0.05	+0.71			

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◄ Fig. 3 Killing rates of caspofungin and the corresponding fitted regression lines (dashed lines) in time-kill experiments against four C. albicans isolates (averages  $\pm$  standard deviation) in RPMI-1640 (a) and 50 % serum (d), against C. albicans DPL20 in RPMI-1640 (b) and 50 % serum (e), and against ATCC 10231 type strain in RPMI-1640 (c) and 50 % serum (f). Positive and negative k values indicate the decrease and increase, respectively, in viable cell numbers

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according to the revised CLSI break points in RPMI-1640 [13]. As expected, the DPL20 isolate with a 134 prominent fks mutation was resistant to caspofungin 135 [13]. MIC values were 4- to 16-fold higher in the 136 presence of 50 % serum.

137 **Time-Kill Experiments** 

138 In time-kill experiments, caspofungin was fungistatic 139 at  $>1-2\times$  MIC in both media against the clinical 140 isolates as well as against the 10231 ATCC strain 141 (<99.9 % reduction in viable cell count compared to 142 the starting inoculum) (Table 1; Fig. 1a, b, d, e).

143 Against the resistant strain DPL20, caspofungin 144 produced a weak fungistatic effect in RPMI-1640 (Fig. 1C). In 50 % serum, the killing curves were 145 146 generally similar to control; at 32 mg/L, a negligible 147 reduction was observed after 4 h, but later the killing 148 curves again became similar to control (Fig. 1f).

#### 149 Postantifungal Effect

150 The time required for control (drug-free) isolates to 151 grow  $1 \log_{10}$  was similar in both media (4.12–4.95 h in 152 RPMI-1640 and 4.69–5.17 h in 50 % serum). PAFE 153 plots for isolates 183 and 3666 in RPMI-1640 and in 154 50 % serum are shown in Fig. 2. In RPMI-1640, 155 clinical isolates and the ATCC 10231 strain showed the inhibition of re-growth at 4, 16 and 32 mg/L for 156 4.89 to >19.34, 13.84 to >19.88 and 7.65 to >19.88 h, 157 respectively (Table 2). PAFE in 50 % serum 158 159 decreased markedly at 4, 16 and 32 mg/L concentra-160 tions (Table 2; Fig. 2). Most isolates showed growth 161 in 50 % serum; colony count decreases occurred only 162 in cases of isolates 183 at 16 and 32 mg/L (Fig. 2c) 163 and 3666 at 32 mg/L (Fig. 2d) and only after 4 h and 164 were negligible (Table 3). In case of DPL20, isolate 165 PAFE was never observed regardless of media (Table 2). 166

Comparison of Colony Count Changes in Time-167 Kill and Postantifungal Effect Experiments 168

Maximum colony count changes compared to the 169 starting inocula in time-kill and PAFE experiments at 170 4, 16 and 32 mg/L are presented in Table 3. 171

Comparing the different media at the same con-172 centrations in killing experiments, the CFU decrease 173 was generally higher in 50 % serum than in RPMI-174 1640 (Table 3). Contrastingly, the CFU decrease in 175 PAFE experiments was significantly higher in RPMI-176 1640 than in 50 % serum with all isolates and 177 concentrations (P < 0.05-0.001) (Table 3). 178

Comparing the colony counts reductions at the 179 same concentrations in PAFE and time-kill experi-180 ments, we noticed comparable or sometimes higher 181 reductions (in cases of 10920 and 3666 isolates) in 182 PAFEs than that seen with continuous 24-h exposure 183 in RPMI-1640 (Table 3). However, 50 % serum 184 significantly decreased the PAFE killing for all tested 185 isolates (P < 0.01-0.001) (Table 3; Fig. 4a–e). 186

Killing Rates in Time-Kill Experiments

In time-kill experiments, killing activity of caspofun-188 gin was significantly weaker at 16-32 mg/L than at 189 0.25, 1, 4 and 8 mg/L (P < 0.05-0.001) in RPMI-190 1640 (mini-paradoxical effect) (Fig. 3a). However, 191 killing rates at 1-32 mg/L were concentration-inde-192 pendent in 50 % serum against the susceptible isolates 193 (Fig. 3d). Similar effect was noticed in case of the 194 strain ATCC 10231 (Fig. 3c, f). In case of isolate, 195 196 DPL20 k values were negative (indicating the growth instead of killing) regardless of media (Fig. 3b, e). 197

Killing Rates in Postantifungal Effect Experiments 198

In PAFE experiments, killing rates for clinical isolates 199 and the ATCC strain in RPMI-1640 were isolate- and 200 concentration-dependent (k values from -0.111 to 201 +1.019 1/h), while in 50 % serum, the k values 202 showed markedly narrower range (from -0.017 to -203 0.185 1/h) (Fig. 4a-e). 204

#### Discussion

This study is the first in which killing rates in short and 206 continuous caspofungin exposures to C. albicans were 207

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**Fig. 4** Killing rates of caspofungin and the corresponding fitted regression lines (*dashed lines*) in PAFE experiments against 183 (a), 10920 (b), 3666 (d) and 12132 (e) *C. albicans* isolates in RPMI-1640 and 50 % serum, and against *C. albicans* ATCC 10231 type strain (c) in RPMI-1640 and 50 % serum. Positive and negative k values indicate the decrease and increase, respectively, in viable cell numbers. For panels C and D, the scale of the y axis was broken for better visualization of regression lines

208 compared head to head in RPMI-1640 with and 209 without 50 % serum. In agreement with previous results [3-5], the CFU decreases in time-kill and 210 PAFE experiments were similar in RPMI-1640; 211 212 however, at 16 and 32 mg/L, killing rates decreased. 213 Decreased killing rates in time-kill experiments at 16 and 32 mg/L can be explained by the adaptive and 214 215 compensatory response to high caspofungin concen-216 trations in the fungal cells that limit killing effect and 217 allow for the growth [14]. Addition of 50 % serum 218 significantly decreased killing rates at 4, 16 and 219 32 mg/L in the PAFE experiments as compared to 220 RPMI-1640, while the killing rates in time-kill 221 experiments (continuous exposure) at  $\geq 1$  mg/L con-222 centrations remained high and concentration-indepen-223 dent (i.e., killing rate reached its maximum at 1 mg/L). 224 These findings are in accordance with our previous 225 results, where killing rates of C. krusei and C. 226 inconspicua did not differ significantly in 50 % serum 227 at effective concentrations [6]. Moreover, 1-h expo-228 sure of C. albicans to 0.5, 1 and 2 mg/L of caspofungin 229 in 50 % serum is not long enough to produce any 230 growth inhibition, as opposed to what found with RPMI-1640 alone. 231

232 Louie et al. [15] demonstrated that tissues serve as 233 drug reservoirs from which the drug is released slowly, 234 explaining that serum caspofungin half-life tripled 235 when both serum and tissues half-life were taken into 236 account in the terminal half-life calculation. They 237 concluded that the primary tissue reservoir rather than 238 PAFE was responsible for the excellent in vivo 239 activity of caspofungin [15]. Their results are in line 240 with the present study, as 1-h exposure to caspofungin produced negative k values (growth) and significantly 241 242 decreased PAFEs in 50 % serum.

PAFE is frequently regarded as a contributor to
clinical efficacy of echinocandins [1–4, 13]. It is
defined as prolonged growth inhibition following
limited (generally 1 h) in vitro drug exposure; however, it must be noted that this is not equal to killing, as

slower growth may also be regarded as prominent 248 PAFE. While killing may directly lead to eradication, 249 growth, even slower growth, of fungi still carries a risk 250 of persistent infection and fungal re-growth. These 251 facts should be taken into consideration when trans-252 lating PAFE as a contributor to clinical efficacy. 253 Present results strongly suggest that PAFE is lost in the 254 presence of 50 % serum, even though marked PAFE is 255 detected in RPMI-1640 after 5-min exposure [5]. The 256 negligible PAFEs found in 50 % serum indicate that 257 prolonged in vitro PAFEs (in RPMI-1640), frequently 258 interpreted as a contributor to better clinical efficacy 259 [1–4], may be less important in vivo (better mimicked 260 by the serum-containing medium) at least against C. 261 albicans. Whether these also apply for non-albicans 262 species is to be answered by further studies. 263

In summary, continuous but not brief caspofungin 264 exposure produced measurable killing rates against C. 265 *albicans* clinical isolates in killing studies in the 266 presence of 50 % serum. PAFE after brief exposure to 267 caspofungin (and probably to other echinocandins), 268 even when marked, may play a limited role in the 269 excellent clinical efficacy of echinocandins. 270

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Conflict of interestL. Majoros received conference travel276grants from MSD, Astellas and Pfizer.277278

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![](_page_12_Picture_16.jpeg)

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