Echinocandins Accelerate Particle Transport Velocity in the Murine Tracheal Epithelium: Dependency on Intracellular Ca\(^{2+}\) Stores

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ABSTRACT

The mucociliary clearance of lower airways is modulated by different physiologic stimuli and also by pathophysiologic agents like polluting substances or pharmaceutical molecules. In the present investigation, we measured the particle transport velocity (PTV) of mouse tracheae as a surrogate for mucociliary clearance. In mouse tracheal preparations, we detected a sustained increase in the PTV under the application of the echinocandins caspofungin, anidulafungin, and micafungin. In further experiments, we observed the effects of echinocandins on the PTV were dependent on intracellular Ca\(^{2+}\) homeostasis. In Ca\(^{2+}\)-free buffer solutions, the amplitude of the echinocandin-evoked rise in the PTV was significantly reduced relative to that in the experiments in Ca\(^{2+}\)-containing solutions. Depletion of intracellular Ca\(^{2+}\) stores of the endoplasmic reticulum (ER) by caffeine completely prevented an increase in the PTV with subsequent caspofungin applications. Mitochondrial Ca\(^{2+}\) stores seemed to be unaffected by echinocandin treatment. We also observed no altered generation of reactive oxygen species under the application of echinocandins as probable mediators of the PTV. Consequently, the observed echinocandin effects on the PTV depend upon the Ca\(^{2+}\) influx and Ca\(^{2+}\) contents of the ER. We assume that all three echinocandins act intracellularly on ER Ca\(^{2+}\) stores to activate Ca\(^{2+}\)-dependent signal transduction cascades, enhancing the PTV.

KEYWORDS

particle transport velocity, echinocandins, mucociliary clearance, intracellular calcium, antifungal agents

Mucociliary clearance of the lower airways is a pivotal mechanism to prevent deposition or colonization of the lungs by potential pathogens (1). This clearance function depends upon the metachronal continuous beating of cilia located on the apical membranes of tracheal epithelial cells (2, 3). The cilia immerse into the soluble phase of the mucus layer, and their continuous movement propels the mucus forward toward the gastrointestinal tract (4, 5), thereby removing particles trapped in the upper mucus layer from the lower airways (6). This particle transportation can be accelerated by increasing the ciliary beat frequency (CBF), which leads to more effective mucus flow (7–10). Here, the importance of proper cilia coordination of ciliary beating has to be accentuated, while in patients suffering from ciliopathies, like primary cilia dykinesia (PCD), the ciliary dysfunction leads to an ineffective mucociliary clearance, resulting in recurrent or chronic bacterial infections (11, 12). Mucociliary clearance is influenced by...
multiple physiological signal cascades, including humoral, neural, mechanical, and immunologic processes (13–17). The invocation of many control mechanisms leads to enhanced CBF and, consequently, increased particle transport velocity (PTV), with improved rates of lower airway clearance. However, the CBF can also be inhibited by several factors, such as a drop in temperature, acidosis, reactive oxygen species (ROS), or polluting agents. All these determinants lead to a reduced PTV and subsequent ineffective mucociliary clearance (5, 18–21).

Many pharmacological agents have been described to date as direct or indirect modulators of mucociliary clearance (22–24). These pharmacological agents reach ciliary-bearing tracheal epithelial cells via diffusion from blood vessels or by inhalation in order to interfere with them (25, 26). They may attain concentrations that are sufficient enough to alter CBF and, furthermore, for many pharmacological agents, e.g., betasympathomimetics or purines, research has elucidated how they influence cilia-bearing cells (27, 28). Most pharmacological substances act primarily on membrane-bound receptors or ion channels (29–32). However, they are also able to directly alter intracellular signal cascades or modulate cellular energy metabolism, leading to an altered turnover of ATP, which is a main activator and substrate of the cilia motion apparatus (32). Especially in critically ill patients or immunocompromised individuals, the mucociliary function plays a key role in reducing the risk of bacterial colonization of the lower airways, which would result in severe infections (33–35). Besides the risk for bacterial infection, severe pulmonary fungal infections, e.g., by Candida or Aspergillus, represent significant hazards that trigger increased mortality in critically ill or in immunocompromised patients (36, 37). In these patients, the use of effective antifungal therapies is crucial to reduce the fungal load and lessen the risk for organ failure and mortality (38–40).

In recent years, new strategies for the therapy of systemic fungal infections have been developed, including the administration of echinocandins, a new class of antymycotic drugs that are effective against invasive fungal infections, especially of diverse Candida strains (41–43). This class of antymycotic drugs mainly includes caspofungin, anidulafungin, and micafungin, which can reach high concentrations in the airways, since they are also suitable for direct bronchial administration via a nebulizer (44). However, so far, it is not known whether echinocandins can influence the mucociliary clearance or intracellular signal cascades of ciliary-bearing epithelial cells. In the present study, we investigated the PTV as an indicator of the mucociliary function of the lower airways under exposure to all three clinical established echinocandins in organ preparations of mouse tracheae. We elucidated the possible signal cascades involved in how echinocandins influence mouse tracheal epithelial function by pharmacological intervention. We separately inhibited receptors, kinases, and proteins of the concerned main signal cascades known to stimulate the PTV by administering pharmacological inhibitors. We also investigated whether echinocandins induce the generation of ROS in tracheal preparations as a hazard of many cell functions. Classically, ROS generation mainly occurs as a by-product of mitochondrial oxidative phosphorylation. ROS are highly reactive molecules that can oxidize other molecules of various cell signaling pathways, while the liberation of ROS acts like an alarm system of the cell during hypoxia, inflammation, or contact with pathogens (45–47).

The amount of ROS liberation plays a crucial role in cell homeostasis. Moderate ROS liberation leads to cell adaption to the stressor, while larger amounts of ROS could cause cell damage or even induce cell death (48). We performed ROS measurements to investigate if caspofungin induces ROS liberation, indicating cell stress or cell damage.

RESULTS

The application of either caspofungin, anidulafungin, or micafungin significantly increased the PTV of the mouse tracheal epithelium in a concentration-dependent manner (Fig. 1A to F). Under control conditions without the administration of echinocandins, the PTV experienced no alterations during the observation period (Fig. 1A, C,
Echinocandins elicted an increase in the PTV in mice trachea. (A, C, and E) Caspofungin (60 µM), anidulafungin (44 µM), or micafungin (50 µM) rapidly induced a sustained elevation in the PTV that reached up to 200% of the basal PTV of control tracheae. Only experiments responding to ATP (100 µM) at the end of the exposure period were included for further statistical evaluation. (B) A dose-response curve presents the effects of caspofungin on the PTV. Using the Hill equation, EC<sub>50</sub> was estimated at 13.4 µM. (C) Anidulafungin rapidly triggered an increase in the PTV, reaching more than 200% of the basal PTV under control conditions. (D) A dose-response curve presents the concentration-dependent effect of anidulafungin on the PTV following a Hill equation EC<sub>50</sub> value of 6.5 µM. (E) Micafungin induced an increase in the PTV that seemed rather transient compared to that of caspofungin or anidulafungin; the maximum response reached more than 170% of the initial basal PTV. (F) The micafungin response occurred in a concentration-dependent manner. The Hill equation EC<sub>50</sub> value for micafungin was estimated as 7.4 µM. In all panels, the PTV of the initial recording time point was normalized to 100%. Recordings show mean ± standard error of the mean values for individual time points. α, control; Δ, caspofungin; ■, anidulafungin; ●, micafungin; horizontal bars in experimental recordings present exposure periods of defined pharmacological agents; time scale bars are present in all individual panels.

and E). All three tested echinocandins evoked a prolonged elevation of the PTV that declined only slightly with long-term exposure; still, the tracheae responded to ATP as a vitality control at the end of all individual experiments, which is presented exemplarily for all tests in Fig. 1A. An acceleration of the PTV following drug application became evident after a brief delay, which then rapidly reached a sustained plateau. The response to echinocandins occurred in a dose-dependent manner, following a Hill
equation with 50% effective concentration (EC50) values of 13.4 μM for caspofungin (Fig. 1B), 6.5 μM for anidulafungin (Fig. 1D), and 7.4 μM for micafungin (Fig. 1F), respectively.

Different signal pathways are described to alter the mouse tracheal PTV, also including membrane-bound receptors. Due to similar characteristics for all three echinocandins, we only show data for caspofungin concerning the investigation of membrane-bound receptors. In subsequent experiments, we added atropine (10 μM) to inhibit muscarinic receptors prior to caspofungin application. We observed no intrinsic effect of atropine on the PTV (Fig. 2A). The following application of caspofungin, still in the presence of atropine, evoked a significant increase of the PTV to 163% ± 12% (n = 7; P < 0.001) relative to control therapy. Atropine did not reduce the effect of caspofungin on the PTV (Fig. 2B).

ICI-118,511 (10 μM), a nonselective inhibitor of adrenergic beta-receptors, induced a significant increase in the PTV, and application of caspofungin in the presence of ICI further evoked a sustained rise in the PTV that was higher than that triggered by caspofungin alone (Fig. 2C and D). Exposure to caspofungin in the presence of ICI induced a significant increase in the PTV that was also different from that measured under caspofungin application alone. (atrop., atropine; caspo., caspofungin; horizontal bars in experimental recordings present exposure periods of defined pharmacological agents; n, number of individual experiments; ns, not significant; time scale bars in all individual panels; *, P < 0.05; ***, P < 0.001; Mann-Whitney U test).
Since serotonin is involved in the modulation of murine tracheal mucociliary clearance via different serotonin receptors, we conducted a further series of experiments to prove the involvement of this signal pathway. Application of methysergide (100 μM), an unselective inhibitor of 5-hydroxytryptamine (5-HT) receptors 1 and 2 (5-HT1,2), did not alter the PTV. Caspofungin was applied in the presence of methysergide and immediately induced a rise in the PTV to a stable plateau above the PTV level that was evoked by caspofungin alone (●, control; ▲, caspofungin; x, methysergide and caspofungin). (B) Methysergide did not influence the PTV relative to control conditions (P = 0.03, Student’s paired t test). Caspofungin induced a significant increase in the PTV in the presence of methysergide. This increased PTV was also higher than the rise in the PTV evoked by caspofungin alone. (C) Recording of the PTV during exposure to the inhibitor of serotonin 5-HT1A receptor WAY-100635 (20 μM). WAY-100635 had no effect on the PTV. Caspofungin effects on the PTV were not inhibited by WAY-100635. (●, control; ▲, caspofungin; ○, WAY and caspofungin). (D) The application of WAY did not alter the PTV; however, caspofungin application following treatment with WAY-100635 induced a significant rise in the PTV. This rise was not different from that in the PTV seen under caspofungin application alone. (E) A selective inhibitor of 5-HT2, cyproheptadine, led to a prolonged increase in the PTV. Subsequent caspofungin application, still in the presence of cyproheptadine, induced a further increase in the PTV (●, control; ▲, caspofungin; △, cyproheptadine and caspofungin). (F) Cyproheptadine induced a significant rise in the PTV, while the application of caspofungin in the presence of cyproheptadine induced a further significant rise in the PTV (casp., caspofungin; cyroph., cyproheptadine; methys., methysergide; WAY, WAY-100635; horizontal bars in experimental recordings present exposure periods of defined pharmacological agents; n, number of individual experiments; ns, not significant; time scale bars presented in all individual panels; *, P < 0.05; **, P < 0.01; ***, P < 0.001; Mann-Whitney U test).

Since serotonin is involved in the modulation of murine tracheal mucociliary clearance via different serotonin receptors, we conducted a further series of experiments to prove the involvement of this signal pathway. Application of methysergide (100 μM), an unselective inhibitor of 5-hydroxytryptamine (5-HT) receptors 1 and 2 (5-HT1,2), did
not alter the basal PTV \( (P = 0.93) \) (Fig. 3A and B). The application of caspofungin (60 \( \mu \)M) in the presence of methysergide evoked a significant rise in the PTV to 181\% \( \pm \) 9\% \( (n = 7; P < 0.001) \) relative to methysergide alone (Fig. 3A and B). This rise in the PTV was also different from that observed under caspofungin application alone \( (P < 0.01) \); still, we observed a sustained rise in the PTV that remained at a plateau throughout the observation period.

WAY-100635 (50 \( \mu \)M), an inhibitor of 5-HT\(_{1A}\) receptors, also did not influence the PTV itself. When caspofungin was administered in the presence of WAY-100635, we observed a significant increase in the PTV to 160\% \( \pm \) 5.7\% \( (P < 0.01 \text{ versus control}) \) that was not different from that measured under caspofungin application alone \( (P = 0.135) \) (Fig. 3C and D).

In a further series of experiments, we inhibited 5-HT\(_{3}\) receptors prior to caspofungin application using the inhibitor cyproheptadine (2.5 \( \mu \)g/ml). Application of caspofungin in the presence of cyproheptadine induced a significant sustained increase in the PTV that was significantly higher than the increase in the PTV observed under cyproheptadine administration alone. This rise in the PTV was not different from the rise in the PTV that was evoked by caspofungin alone (Fig. 3E and F).

We found no evidence that caspofungin influenced the PTV via known membrane-bound receptors. Subsequently, we investigated whether caspofungin was able to induce the PTV by interfering with known intracellular signal cascades.

Inhibition of the cyclic adenosine 3',5'-monophosphate (cAMP)-dependent protein kinase signal pathway was achieved by the application of H-89 (10 \( \mu \)M) and reduced the baseline PTV to 59\% \( (n = 5; P < 0.01) \) relative to control therapy. Meanwhile, caspofungin applied in the presence of H-89 significantly enhanced the PTV to a maximum of 168\% \( \pm \) 25\% \( (n = 5; P < 0.001) \). This increase in the PTV evoked by caspofungin while inhibiting the cyclic AMP (cAMP)-dependent protein kinase signal pathway by H-89 was not different from the increase in the PTV triggered by caspofungin alone (154\% \( \pm \) 32\%) \( (n = 6; P = 0.86) \) (Fig. 4A and B). However, it showed a slow decrease in the PTV at the end of the observation period.

We further examined whether caspofungin acts via a nitric oxide (NO) signal pathway to enhance the PTV. The possible interference of this signal pathway was verified by the inhibition of soluble guanylate cyclase (sGC) using the selective inhibitor 1H-[1,2,4]oxadiazolo[4,3-a]quinazolin-1-one (ODQ) (10 \( \mu \)M). ODQ alone did not alter the basal PTV (89\% \( \pm \) 2.8\%) \( (n = 5) \) after a 15-min incubation period relative to control therapy with resting for 15 min (87\% \( \pm \) 3.6\%) \( (n = 7; P = 0.093) \). Caspofungin in the presence of ODQ induced a sustained rise in the PTV to 187\% \( \pm \) 15\% \( (n = 5) \) that was significantly higher than the rise in the PTV induced by caspofungin alone (146\% \( \pm \) 8\%) \( (n = 6; P < 0.001) \) (Fig. 4C and D). Thus, we found no evidence that caspofungin enhances the PTV via an sGC pathway in these cilia-bearing cells.

**Interference of ROS generation by caspofungin.** Since ROS generation changes cilia function and, here, interferes with signal pathway cascades, we elucidated whether caspofungin changes ROS generation in isolated mouse tracheae. We measured ROS generation using the aforementioned Amplex red assay kit. Fluorescence was recorded for 30 min under each condition, i.e., control therapy or echinocandin exposure. Under control conditions, the fluorescence showed only a trend of increase after an exposure period of 30 min to 126\% \( \pm \) 27\% relative to the initial value \( (n = 4; P = 0.38) \). Tracheae exposed to caspofungin for 30 min also showed no significant alterations in ROS generation, 114\% \( \pm \) 12\% \( (n = 5; P = 0.13) \), compared with the initial fluorescence value. The fluorescence under the application of anidulafungin or micafungin did not trigger ROS generation (anidulafungin 100.5\% \( \pm \) 3\% \( [n = 4; P = 0.56]\); micafungin 98.4\% \( \pm \) 1\% \( [n = 4; P = 0.12]\)). There was no difference in ROS generation between control and echinocandin-treated organs at the end of the exposure period (caspofungin, \( P = 0.29 \)); anidulafungin, \( P = 0.34 \); micafungin, \( P = 0.17 \) (Fig. 5).

**Ca\(^{2+}\)-dependent effects of echinocandins on PTV. (I) Caspofungin.** We further investigated whether the effects of echinocandins depend upon the cytosolic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]). Tracheae resting in Ca\(^{2+}\)-free conditions showed no changes in
the basal PTV, remaining constant during the entire observation period. However, the application of echinocandins in Ca\(^{2+}\)-free buffer immediately triggered a significant rise in the PTV, reaching a peak of 153\% ± 6\% (\(n = 5\); \(P < 0.001\) versus control) (Fig. 6A and B), although the response to caspofungin in Ca\(^{2+}\)-free buffer was significantly lower than that in Ca\(^{2+}\)-containing buffer solution (182\% ± 20\%) (\(n = 6\); \(P < 0.01\)).

The depletion of internal endoplasmic reticulum (ER) Ca\(^{2+}\) stores prior to caspofungin application was achieved by caffeine (30 mM). Caffeine exposure alone did not change the PTV (78\% ± 1.2\%) relative to Ca\(^{2+}\)-free controls (82\% ± 13\%) (\(P = 0.23\)) (Fig. 6C). Exposure to caspofungin (60 \(\mu\)M) after the depletion of caffeine-sensitive Ca\(^{2+}\) stores did not increase the PTV (65\% ± 13\%), which was significantly lower than the PTV evoked by caspofungin alone in Ca\(^{2+}\)-free solution (164\% ± 20\%) (\(P < 0.001\)) (Fig. 6C and D). Hence, the depletion of caffeine-sensitive Ca\(^{2+}\) stores completely inhibited the caspofungin effect on the PTV.

(ii) Anidulafungin. The application of anidulafungin (44 \(\mu\)M) in Ca\(^{2+}\)-free buffer solution still evoked a significant rise in the PTV to 140\% ± 16\% (\(n = 5\); \(P < 0.001\)); however, this accelerated PTV was lower than that in Ca\(^{2+}\)-containing buffer solution (210\% ± 19\%) (\(n = 6\); \(P < 0.05\)) (Fig. 7). Still, during the inhibition of the PKA signal pathway by H-89 (10 \(\mu\)M), the application of anidulafungin (44 \(\mu\)M) evoked a significant rise in the PTV to 136\% ± 5\% (\(n = 4\); \(P < 0.01\)) (Fig. 7D). A high concentration of ryanodine (40 \(\mu\)M) was used to inhibit ryanodine receptors of the ER. Subsequent
prolonged exposure to anidulafungin (44 μM) evoked no change in the PTV (119% ± 8%) (n = 3; P = 0.34) compared with ryanodine (116% ± 7%) (n = 3) (Fig. 7C and D).

(iii) Micafungin. Micafungin (50 μM) induced a rise in the PTV up to 165% ± 16% (n = 6) in Ca^{2+}-containing solution, which was reduced in Ca^{2+}-free solution to 126% ± 10% (n = 6; P < 0.01) (Fig. 8A, B, and D). The evoked PTV slowly decelerated during the exposure period (Fig. 8A). In tracheae treated with BAPTA-AM [1,2-bis(2-aminophenoxy)ethane-N,N,N',N'-tetraacetic acid tetrakis(acetoxymethyl ester)], the application of micafungin (50 μM) still evoked a transient rise in the PTV to a maximum of 145% ± 24% (n = 4; P < 0.05) (Fig. 8D). The inhibition of the PKA signal pathway by H-89 did not prevent a micafungin-induced rise in the PTV of 162% ± 16% (n = 5; **, P < 0.01) (Fig. 8C and D). The depletion of mitochondrial Ca^{2+} stores by the application of DNP (125 μM), an uncoupler of the mitochondrial respiratory chain, reduced the PTV to 74% ± 2% (n = 6). The subsequent application of micafungin (50 μM) significantly enhanced the PTV to 133% ± 25% (n = 6; P < 0.01) (Fig. 8D) relative to DNP alone.

Further, we investigated whether Ca^{2+} stores in the ER contribute to micafungin-induced PTV. We depleted ER Ca^{2+} stores using caffeine. Under low-dose caffeine (15 mM) application, we then measured an increase in the PTV that was not further enhanced under micafungin exposure (Fig. 9A). A higher caffeine concentration (30 mM) did not induce the PTV, which also was not induced by subsequent micafungin application (Fig. 9B). The inhibition of the ryanodine receptors by ryanodine (120 μM) almost completely prevented the PTV response to micafungin (Fig. 9C and D).

DISCUSSION

In the present study, we gained evidence that echinocandins increase the PTV in intact murine tracheal epithelia. Furthermore, we investigated the possible signal cascades of echinocandin-induced alteration of mice tracheal epithelial function and revealed that they alter the PTV via intracellular Ca^{2+}-dependent signal cascades in a dose-dependent manner. The mucociliary clearing system is an important mechanism that functions to disburden the airway surface area of the lung from debris and pathogen particles (49, 50). Besides bacterial and viral pathogens as well as the fungal cell wall, the polysaccharides galactomannan and zymosan can inhibit the PTV via reversible ROS generation in the tracheal tissue (51). Echinocandins are antifungal agents.

![Figure 5](https://example.com/fig5.png)  
**FIG 5** ROS generation in tracheal epithelial cells during the application of echinocandins. ROS were detected using the Amplex Red reagent for whole mouse trachea. Fluorescence was measured at 580 nm for 30 min and expressed as a percentage of the initial fluorescence value. In untreated organs, only a slight change in the fluorescence intensity was recorded, suggesting a slight change in ROS generation. In tracheae exposed to either caspofungin, anidulafungin, or micafungin, we recorded only minute changes in ROS generation. Each organ was assessed at three consecutive time points, and statistical testing compared fluorescence recordings taken at 15 min and 30 min with the initial fluorescence (control, n = 4; caspofungin, n = 5; anidulafungin, n = 4; micafungin, n = 4; ●, control; ○, caspofungin; □, anidulafungin; X, micafungin; Wilcoxon rank-sum test).
recommended for the empirical and specific treatment of invasive *Candida* and *Aspergillus* infections and are widely used in clinical practice (52, 53).

First of all, we analyzed the impact of the three existing echinocandins (caspofungin, anidulafungin, and micafungin) on the PTV in murine tracheal epithelia and found that application of caspofungin, anidulafungin, or micafungin significantly increased the PTV (Fig. 1A). The effects of all three echinocandins appeared to be similar in a dose-dependent manner. Thus, we elucidated according to the signal pathway how echinocandins enhance the PTV; specifically, different signal pathways are described to alter the mouse tracheal PTV, including membrane-bound receptors, intracellular signal cascades, ROS generation, and intracellular Ca²⁺ homeostasis (13, 51). Subsequently, we used pharmacological inhibitors to identify the underlying mechanism of a caspofungin-induced increase of the PTV and whether membrane-bound receptors are involved in echinocandin-induced PTV. Since all three echinocandins triggered a similar response in the PTV when known plasma membrane-bound receptors are pharmacologically inhibited, here, we only present data for caspofungin. First, the potential effect on muscarinic receptors was investigated via the application of atropine, prior to caspofungin application, but

![Graph A: Caspofungin effects on PTV](image)

![Graph B: Caspofungin effects on PTV](image)

![Graph C: Caspofungin effects on PTV](image)

![Graph D: Caspofungin effects on PTV](image)

**FIG 6** Caspofungin effects depend upon Ca²⁺ derived from caffeine-sensitive stores. (A) In Ca²⁺-free buffer medium, caspofungin induced a significant rise in the PTV that slowly decelerated during the exposure period. (B) The basal PTV was not affected by Ca²⁺-free buffer solutions (Wilcoxon rank-sum test). The significant rise in the PTV evoked by caspofungin in Ca²⁺-free buffer solution was significantly lower than that in Ca²⁺-containing buffer solution (Mann-Whitney U test). (C) Mouse tracheae exposed to caffeine (30 mM) in Ca²⁺-free buffer solution did not demonstrate alterations in the PTV. The application of caspofungin to tracheae primarily exposed to caffeine in Ca²⁺-free solution also did not evoke an increase in the PTV. Instead, the PTV slowly decelerated below the basal PTV value. (D) Caffeine (30 mM) in Ca²⁺-free medium did not change the PTV relative to the control in Ca²⁺-free solution (Mann-Whitney U test). The application of caspofungin in the presence of caffeine (30 mM) did not affect the basal PTV. In Ca²⁺-free buffer medium, the PTV level during caspofungin exposure in the presence of caffeine was significantly lower than that measured during sole caspofungin exposure. (B, Ca²⁺-free control buffer medium; , Ca²⁺-free buffer medium with caspofungin; , Ca²⁺-free buffer medium with caffeine; , Ca²⁺-free buffer medium with caffeine and caspofungin; horizontal bars in experimental recordings present exposure periods of defined pharmacological agents; n, number of individual experiments; ns, not significant; time scale bars are present in all individual panels; **, P < 0.01; ***, P < 0.001; Mann-Whitney U test).
our data revealed that caspofungin did not alter the PTV via muscarinic receptor activation. Second, the inhibition of beta-receptors did not alter the caspofungin-induced increase in the PTV. Third, neither the inhibition of 5-HT1 nor the inhibition of 5-HT2 receptors affected the rise of the PTV evoked by caspofungin. Accordingly, we assumed that caspofungin did not influence the PTV via known membrane-bound receptors and subsequently investigated whether caspofungin could induce the PTV by interfering with intracellular signal cascades. Further inhibition of the cAMP-dependent protein kinase signal pathway, which is a cornerstone of signal transduction in tracheal epithelial cells, did not affect echinocandin-evoked activation in the PTV.

We further investigated whether echinocandins act via a NO signal pathway to enhance the PTV. It is known that NO spontaneously generated by airway epithelial cells plays a modulatory role in beta-adrenoreceptor-mediated stimulation of ciliary motility (54, 55). In the present study, we found no evidence that endogenously generated NO contributed to echinocandin-induced acceleration of the PTV. To elucidate this, we pharmacologically inhibited the NO signal transduction pathway prior to echinocandin application. This was achieved by the inhibition of sGC, the known target...
molecule of NO in the tracheal epithelium (56, 57), although we observed no alteration in the PTV under the subsequent echinocandin exposure. Although intrinsic generated NO plays a key role in the modulation of beta-adrenergic stimulation of ciliary motility, the observed echinocandin effect was not influenced or amplified by NO.

Since ROS generation impairs airway cilia function, we furthermore investigated whether echinocandins alter ROS generation in isolated mouse tracheae to accomplish an enhanced ciliary motility (18, 58). In experiments using whole mouse tracheae, we found no alterations in ROS generation during echinocandin exposure. We revealed that there was no difference in ROS generation at the end of the exposure period between control and echinocandin-treated organs. Consequently, we assume that echinocandins do not act via the inhibition of an ROS-sensitive pathway to provoke the PTV acceleration. Many of the established signal transduction cascades in ciliary cells investigated here depend upon an increased [Ca\(^{2+}\)] (15, 30, 59, 60). Therefore, we investigated whether the echinocandin effect also depends upon homeostasis of [Ca\(^{2+}\)]. However, the increase in the PTV was only partly dependent on extracellular calcium. We assume that Ca\(^{2+}\) channels of the plasma membrane were activated directly or indirectly by echinocandins. Store-operated Ca\(^{2+}\) entry (SOCE) is a common mechanism in many cells to regulate cell function (61). Calcium influx via SOCE is

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**FIG 8** Micafungin evokes a prolonged rise in the PTV partly dependent on extracellular Ca\(^{2+}\). (A) In Ca\(^{2+}\)-containing solution, micafungin induced a prolonged rise in the PTV, up to 165% of the basal PTV. The PTV slowly declined under long-term micafungin exposure. (B) The micafungin-induced rise in the PTV was partly dependent on the extracellular Ca\(^{2+}\) concentration, since the observed rise in the PTV was reduced in Ca\(^{2+}\)-free buffer solution. Still, under prolonged micafungin exposure, the PTV declined. (C) The inhibition of PKA using H-89 reduced the basal PTV, while subsequent application of micafungin induced a significant rise in the PTV, which declined after reaching a maximum velocity. (D) Micafungin induced a significant rise in the PTV either in Ca\(^{2+}\)-containing or Ca\(^{2+}\)-free medium. However, the PTV in Ca\(^{2+}\)-free buffer solution was significantly lower than that in Ca\(^{2+}\)-containing buffer solution. In tracheae treated with BAPTA-AM to reduce [Ca\(^{2+}\)], micafungin induced a significant transient rise in the PTV. H-89, the inhibitor of the PKA signal transduction pathway, did not prevent the micafungin-induced rise in the PTV. The depletion of mitochondrial Ca\(^{2+}\) stores and simultaneous interruption of mitochondrial ATP synthesis using DNP did not prevent the micafungin-evoked rise in the PTV (horizontal bars in experimental recordings present exposure periods of defined pharmacological agents; n, number of individual experiments; time scale bars are present in all individual panels; *, P < 0.05; **, P < 0.01; ***, P < 0.001; Mann-Whitney U test).
mediated by STIM-operated ORAI channels (62–64) and may also be involved in Ca\textsuperscript{2+} influx to activate the PTV and replenish Ca\textsuperscript{2+} stores. Another important role could concern the involvement of the transient receptor potential vanilloid 4 cation channel (TRPV4) (65–67), a Ca\textsuperscript{2+}-permeable nonselective cation channel in airway epithelial cells, which can also interact with calmodulin or other calcium-sensing molecules (68). Without the presence of extracellular Ca\textsuperscript{2+}, TRPV4 currents are diminished (69), which may support our PTV measurement findings. In Ca\textsuperscript{2+}-free buffer, the echinocandin-initiated rise in the PTV was as rapid as that in calcium-containing buffer solutions; however, it was still significantly reduced, leading to the assumption of the participation of TRPV4. These kinetics were similar to those observed in previous investigations (30). The subsequent rundown of the PTV in Ca\textsuperscript{2+}-free buffer, still in the presence of echinocandins, could also be caused by a slow efflux of Ca\textsuperscript{2+} ions from the cytosol, and the reduced [Ca\textsuperscript{2+}], may lead to an inactivation of the signal transduction cascade. The response to caspofungin and micafungin was completely prevented when ER Ca\textsuperscript{2+} stores were depleted by caffeine prior to their application. In experiments using ryanodine to inhibit ER-bound ryanodine receptors, the response in the PTV to micafungin or anidulafungin was prevented. Therefore, we assume that echinocandins release Ca\textsuperscript{2+} ions from the ER, which further activates signal transduction cascades to increase the ciliary beat frequency, propelling the PTV. We gained no evidence of the contribution of mitochondrial Ca\textsuperscript{2+} stores. Still, while we cannot say whether the increase in the PTV evoked by echinocandins is the consequence of direct liberation of Ca\textsuperscript{2+} ions from the ER and activation of store-operated channels of the plasma membrane, we

![FIG 9 Micafungin induced the PTV in a manner dependent on ER Ca\textsuperscript{2+} stores. (A) The exposure of mouse tracheae to low-dose caffeine (15 mM) induced a sustained rise in the PTV that was not further increased by micafungin application. (B) A high caffeine concentration (30 mM) did not trigger a PTV increase, and subsequent application of micafungin also did not provoke a rise in the PTV. (C) Ryanodine at a high concentration (40 μM) prevented the micafungin-induced rise in the PTV. (D) A low caffeine concentration (15 mM) evoked a rise in the PTV that was not further enhanced in the presence of micafungin. A high caffeine concentration (30 mM) did not accelerate the PTV, and subsequent application of micafungin did not change the PTV. A high ryanodine concentration did not evoke an increase in the PTV, and successive micafungin application did not increase the PTV (horizontal bars in experimental recordings present exposure periods of defined pharmacological agents; n, number of individual experiments; ns, not significant; time scale bars in all individual panels; Mann-Whitney U test).](image-url)
can conclude that the activation mechanism is located downstream of the cAMP-dependent PKA (Fig. 4A, 7, and 8). The PKA is involved in the liberation of Ca\(^{2+}\) ions from the ER (30). However, in a study on isolated human tracheal epithelial cells, we previously demonstrated that caspofungin liberates Ca\(^{2+}\) from ER stores via ryanodine receptors (70). To elucidate further the underlying mechanisms, we will focus on store-operated Ca\(^{2+}\) channels, assessing how echinocandins activate ryanodine receptors or how they are involved in alternative Ca\(^{2+}\) liberation from the ER. Additionally, it has to be investigated if the echinocandin-induced PTV alteration also occurs in human tracheal epithelium. We know that the cilia function in mammals is similar, but it is difficult to establish an equivalent human model to examine the mucociliary clearance in intact human tracheal epithelium. To approach clinical in vivo conditions, an initial way is to examine cultured human ciliated tracheal epithelial cells that were harvested during bronchoalveolar lavage. However, this model discounts cell-to-cell signaling and the coordinated ciliary beat function of adjacent epithelial cells. Thus, further clinical studies are necessary to investigate whether patients receiving echinocandins for antifungal therapy reveal pulmonary benefits like lower rates of or faster recovery from pulmonary infections.

**Conclusions.** The echinocandins caspofungin, anidulafungin, and micafungin provoked a sustained elevation in the PTV that strongly depended upon extracellular Ca\(^{2+}\) and the content of internal caffeine- and ryanodine-sensitive ER stores. We can rule out any echinocandin interference with membrane-bound receptors or with further downstream-located signal transduction cascades. In mouse tracheae, echinocandins did not alter ROS generation, which is assumed to inhibit ciliary function. The mechanism of how echinocandins increase the PTV via Ca\(^{2+}\) influx or liberation ought to be further investigated to understand its interactions with mammalian cell organelles. Finally, experimental investigations on human tracheal epithelium and clinical studies are needed to clarify if these findings in echinocandin administration lead to a noticeable benefit for patients in the ICU.

**MATERIALS AND METHODS**

**Mouse tracheal segment preparation and imaging.** For the present study, we used male C57BL/6J mice aged 12 to 15 weeks (weighing 25 to 35 g) delivered by Charles Rivers (Sulzbach, Germany). All experiments were performed according to the German guidelines for the care and use of laboratory animals. The study protocol was approved by the local committee for animal care of Justus Liebig-University Giessen (Giessen, Germany) (permit no. 443_M 932). Study animals were sacrificed by inhalation of an overdose of isoflurane (Baxter, Unterschleissheim, Germany) in a closed chamber. All of the following preparation steps were performed within 30 min after euthanasia. Immediately after confirmation of death, we removed the trachea with a parasternal incision of the thorax and a median incision of the throat. Still in situ, the trachea was gently disconnected by slicing cranial to its bifurcation and directly downward. Immediately afterward, the organ was transferred into a Delta T culture dish (Biotech, Butl, PA, USA) containing 2 ml of preheated (30°C) 4-(2-hydroxyethyl)-1-piperazine ethanesulfonic acid (Hepes) buffer (pH 7.4). For direct visualization of the respiratory epithelium, Hepes buffer was then replaced by 2 ml of fresh Sylgard polymer (Dow Corning, Wiesbaden, Germany) to attain precise positioning of the trachea. Here, we fixed the submerged organs with two fine minuta (Fiebig Lehrlmittel, Berlin, Germany) so that the cartilage arches faced the Sylgard polymer and the musculus trachealis faced upwards; this position enabled the following fine preparation. Connective tissue and surrounding blood vessels were gently removed by employing spring scissors (Vannas-Tübingen; FST, Heidelberg, Germany). Finally, the pars membranacea, including the musculus trachealis, was cut open in the longitudinal direction, allowing for direct visualization of the respiratory epithelium. Hepes buffer was then replaced by 2 ml of fresh Hepes buffer (pH 7.4; 30°C). The dish containing the trachea was transferred to the stage-holder of an upright transmission light microscope (BX50 WI; Olympus, Hamburg, Germany). The stage-holder of the microscope was equipped with a temperature control unit that allowed the operator to maintain a constant temperature of 30°C in the center of the buffer solution throughout each experiment. In our experiments the temperature was set to 30°C after measuring the CBF and PTV during different temperatures, reaching from 22 to 42°C (data not published). Here, the best measuring conditions and imaging, depending on the camera resolution, could be achieved at 30°C. Therefore, we have to note that the measured PTV in our experiments might be slightly slower than real time, while the PTV is temperature dependent and accelerates with increasing temperatures. Imaging was performed using the TiliVisION imaging software program (Till Photonics, Gräfeling, Germany).

**Measurement of PTV.** For the measurement of the PTV, we used 4 μl of Dynabeads (mean diameter, 2.8 to 4.5 μm; Dynal Biotech GmbH, Hamburg, Germany) added prior to each experiment into the bath solution. Dynabeads directly floating on the tracheal epithelium were focused in a bright-field mode between two cartilages using a 20× water immersion lens (BW50 WI; Olympus, Hamburg, Germany). At certain time points, we recorded short movie sequences with high sampling rates.
Subsequently, the movie sequences were analyzed using the Image Pro Plus software (Media Cybernetics, Rockville, MD, USA). The viability of the tracheal ciliated cells was confirmed at the end of each experiment by the application of ATP (100 μM), leading to a maximal increase in the PTV. Only experiments responding to ATP with an increase in the PTV were included for further data processing. For each individual experiment, we evaluated approximately 200 to 400 particle tracks at certain predefined time points. At these time points, we acquired 200 images during a period of 16.726 s. Further, nonmoving objects were excluded and background subtraction was performed pixel by pixel (200 images/16.726 s = 1 image/83.63 ms). Following the background subtraction, the images were converted and the formerly dark polystyrene Dynabeads now appeared as bright images. Only particles that displayed a lateral deviation of less than 15% were selected for further data acquisition and evaluation. These images were transformed into a binary system with a threshold allowing for the discrimination between bright bead images and the dark background. Finally, on a greyscale, we reduced the 12-bit film to an eight-bit film so that we could monitor individual particle movements using the TILLvision software (Till Photonics, Gräfeling, Germany). From these measurements, we calculated an average PTV.

Estimation of ROS. To gain evidence for ROS generation from the trachea, we employed the Amplex red hydrogen peroxide/peroxidase assay kit (Life Technologies, Carlsbad, CA, USA) for the detection of H2O2. Freshly harvested tracheae were prepared as described above and equilibrated in fresh HEPES buffer (2 ml, 30°C, pH 7.4) for 60 min. After this equilibration period, we added 20 μl of H2O for control experiments (n = 4) or 120 μM caspofungin, 44 μM anidulafungin, 50 μM micafungin (each diluted in 20 μl of H2O, n = 5) to the bath solution. The fluorescence sampling was started directly following the administration of echinocandins (60 min after tissue preparation), after 75 min (15 min after echinocandin administration), and after 90 min (30 min after echinocandin administration). The tracheal tissue specimens were incubated with 10 μM Amplex red reagent (10-acetyl-3,7-dihydroxyphenoxazine) to detect released H2O2. The emitted fluorescence was measured with a microplate reader equipped for excitation wavelengths in the range of 530 to 560 nm and the detection of fluorescence emission wavelengths at ~590 nm (Infinite M200; Tecan, Männedorf, Switzerland). For data processing, fluorescence was normalized and initial fluorescence was arbitrarily set as 100%.

Drugs and buffer solutions. The following drugs were administered: ATP (100 μM, Sigma-Aldrich, St. Louis, MO, USA), caspofungin (60 μM or 120 μM diluted in 20 μl of H2O; MSD, Kenilworth, NJ, USA), anidulafungin (44 μl diluted in 20 μl of H2O; Pfizer, New York, NY, USA), micafungin (50 μM diluted in 20 μl of H2O; Astellas Pharma GmbH, Munich, Germany), atropine (10 μM; B.Braun, Melsungen, Germany), ICI-118,551 (10 μM; Sigma-Aldrich, St. Louis, MO, USA), caffeine (30 mM; Roth, Karlsruhe, Germany), WAY-100635 (20 μM, diluted in 20 μl of H2O; Sigma-Aldrich, St. Louis, MO, USA), ryanodine (40 μM, diluted in 25 μl of H2O; Bio-Technne GmbH, Wiesbaden-Nordenstadt, Germany), H-89 (10 μM; Sigma-Aldrich, St. Louis, MO, USA), methysergide (100 μM Sigma-Aldrich, St. Louis, MO, USA), cyproheptadine (2.5 μg/mI; Sigma-Aldrich, St. Louis, MO, USA), 1H-(1,2)-4oxadiazolol[4,3-a]quinazolin-1-one (ODQ) (10 μM; Sigma-Aldrich, St. Louis, MO, USA), or 2,4-dinitrophenol (DNP) (125 μM; Sigma-Aldrich, St. Louis, MO, USA) was diluted in ethanol for stock solution. H-89, methysergide, and ODQ were diluted in dimethyl sulfoxide. End concentrations were achieved by applying the stock solution directly to the buffer solution in the recording chamber. Dimethyl sulfoxide did not exceed volumes of 20 μl in 2 ml of HEPES to prevent the induction of unspecific disturbances of cilia-bearing cells. Cyproheptadine was diluted in ethanol for the stock solution. All preparations and experiments were carried out in HEPES solution consisting of 20 mM HEPES, 4.5 mM KCl, 2.5 mM CaCl2, 11 mM glucose, 140 mM NaCl, and 1 mM MgCl2; the pH was adjusted using NaOH (4 M) to 7.4 at 30°C for tissue preparation. For Ca2+-free solutions, CaCl2 was replaced by 1 mM ethylene glycol tetraacetic acid. As a balance in buffer solutions with KC1 concentrations of 30 mM, the NaCl concentration was reduced by the same amount. Solutions containing 30 mM KC1 also contained 2.5 mM CaCl2.

Statistical analysis. Only tracheal preparations were included in further statistical evaluation that responded to ATP application. The end of each individual recording. The Mann-Whitney U test was used to compare equivalent measuring points from different experiments. The Wilcoxon rank-sum test was used to compare paired variables. EC50 (median effective concentration) values were calculated using the Hill equation. Statistical data evaluation and testing were performed using the GraphPad Prism software (version 5.04; GraphPad Software, La Jolla, CA, USA).

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.

SUPPLEMENTAL FILE 1, PDF file, 0.3 MB.

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