Lanostane Triterpenes from *Gloeophyllum odoratum* and Their Anti-Influenza Effects

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ABSTRACT

In an in vitro screening for anti-influenza agents from European polypores, the fruit body extract of Gloeophyllum odoratum dose-dependently inhibited the cytopathic effect of the H3N2 influenza virus A/Hong Kong/68 (HK/68) in Madin Darby canine kidney cells with a 50% inhibitory concentration (IC₅₀) of 15 µg/mL, a noncytotoxic concentration. After a chromatographic work-up, eight lanostane triterpenes (1-8) were isolated and their structures were elucidated based on high-resolution electrospray ionization mass spectrometry analyses, and one- and two-dimensional nuclear magnetic resonance experiments. Constituents 1 (gloeophyllin K) and 2 (gloeophyllin L) are reported here for the first time, and compounds 5, 7, and 8 have not been described for the investigated fungal material so far. The highest activity was determined for trametenolic acid B (3) against HK/68 and the 2009 pandemic H1N1 strain A/Jena/8178/09 with IC₅₀ values of 14 and 11 µM, respectively. In a plaque reduction assay, this compound was able to bind to cell-free viruses and to neutralize their infectivity.

Introduction

Polypores, also known as conks, shelf fungi, or bracket fungi, are a morphologically distinct but polyphyletic group of Agaricomycetes (Basidiomycota) [1,2]. They are mostly growing on tree trunks, where they feed on wood by exuding enzymes to break down cellulose or lignin, thus causing wood rot. In central European folk medicines, polypore species (e.g., *Laetiporus sulphureus, Fomes fomentarius, Fomitopsis pinicola, Piptoporus betulinus*, and *Laricifomes officinalis*) have been widely used for the treatment of various diseases [3].

Gloeophyllum odoratum (Wulfen) Imazeki (syn. *Osmoporus odoratus* [Wulf. ex Fr.] or *Trametes odorata* [Wulf. ex Fr.]) is a bracket fungus from the order Gloeophyllales. This perennial wood-rotting fungus grows mainly on stumps of spruce and pine trees in Central Europe, Asia, and North America [4]. Due to its pleasant odor similar to anise or fennel it is commonly called "anise mazegill" [5].

Fruit bodies of *G. odoratum* have been mycochemically investigated concerning their volatile constituents (i.e., aliphatic alcohols, ketones, aldehydes, and monoterpenes) and nonvolatile constituents (i.e., aromatic compounds and tetracyclic lano-

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ABBREVIATIONS						
CC ₅₀	50% cytotoxic concentration					
CPE	cytopathic effect					
ELSD	evaporative light scattering detector					
FTIR	fourier-transform infrared spectroscopy					
HK/68	influenza virus A/Hong Kong/68 (H3N2)					
HSQC	heteronuclear single quantum coherence					
IAV	influenza A virus					
MDCK	Madin Darby canine kidney					
PDA	photo diode array					
SFC	supercritical fluid chromatography					
SI	selectivity index					

stane-type triterpenes) [4–12]. Regarding bioactivity on the extract level, immunomodulatory activity and cytotoxicity have been tested *in vitro* [13]. On the compound level, bioassay-guided fractionation led to the discovery of three *G. odoratum* lanostanes as thrombin inhibitors [9]. To date, no studies are available investigating the anti-influenza or any kind of antiviral properties of this species. However, lanostane triterpenes isolated from other polypores (e.g., *Ganoderma sp.* [Polyporales]) are reported to have an activity against IAV [14–16].

The limited portfolio of currently available and efficient agents against influenza prompted us to investigate the fruit body extracts of ten European polypore species in the search for highly needed novel anti-influenza scaffolds. In this study, we report on the isolation, structure elucidation, and the *in vitro* anti-influenza properties of two so far unknown triterpenes named gloeophyllin K (1) and gloeophyllin L (2) and six known triterpenes (3–8) from the EtOH fruit body extract of *G. odoratum*.

Results and Discussion

To identify anti-influenza natural leads from polypores, EtOH extracts of fruit bodies of 10 species (i.e., *F. fomentarius, F. pinicola, Ganoderma lucidum, Ganoderma applanatum, G. odoratum, Ischnoderma benzoinum, L. sulphureus, Phellinus robustus, P. betulinus, and Trametes gibbosa*) were screened in a CPE inhibition assay with the H3N2 HK/68 in MDCK cells [17]. The EtOH fruit body extract of *G. odoratum* strain 0054 (GO54E) was identified as pronounced inhibitor of the CPE with an IC₅₀ of 15 µg/mL [17]. Moreover, this extract was not cytotoxic to MDCK cells (measured up to 100 µg/mL).

GO54E (31.2 g) was suspended in water and partitioned sequentially between petroleum ether (GO54EP), dichloromethane (GO54ED), and water (GO54EW). By using TLC analysis, GO54ED was found to be enriched with lanostane triterpenes. This fraction was then subjected to several chromatographic steps including size exclusion, SFC, and silica flash chromatography, resulting in the isolation of eight triterpenes (1–8), namely (24*E*)-(3 β ,15 α)-15-acetyloxy-3,26-dihydroxy-lanosta-8,24-dien-21-oic acid, referred to as gloeophyllin K (1), (3*S*,4*S*,4a*R*,6b*R*,9*R*,10a*R*,11b*S*)-3hydroxy-4,11b-dimethyl-10-methylene-9-((*R*)-6-methylhept-5en-2-yl)-2,3,4,4a,5,6,6b,7,8,9,10,10a,11,11b-tetradecahydro1H-benzo[a]fluorine-4-carboxylic acid, referred to as gloeophyllin L (2), trametenolic acid B (3), gloeophyllin B (4), gloeophyllin A (5), ergosterol peroxide (6), eburicodiol (7), and 21-hydroxylanosterol (8) (**> Fig. 1**).

The constituents were identified by means of HRESIMS and 1D and 2D NMR, and comparison of their spectroscopic data with published values. Constituents 1 and 2 are new triterpenes never described in the literature so far. The isolates 3, 4, and 6 have been reported previously from the fruit body of *G. odoratum*. The constituents 5, 7, and 8 are known natural compounds but new for this species.

The full assignments of the ¹³C and ¹H NMR spectroscopic data of isolated new compounds 1 and 2 are listed in ► **Table 1**. Their key HMBC, ¹H-¹H COSY, and NOESY correlations are presented in ► **Fig. 2**. The respective NMR and HRESIMS spectra are provided in the Supporting Information.

Compound 1 was isolated as white, amorphous powder with the molecular formula C₃₂H₅₀O₆, obtained by HRESIMS from the $[M - H_2O + H]^+$ ion at m/z 513.3575 (calcd for $C_{32}H_{49}O_5^+$, 513.3575, $\Delta = -0.1$ ppm) and the [M + Na]⁺ ion at m/z 553.3499 (calcd for $C_{32}H_{50}NaO_6^+$, 553.3500, $\Delta = 0.1$ ppm). The ¹H NMR spectrum exhibits resonances of seven methyl singlets at $\delta_{\rm H}$ 0.80, 0.87, 0.98, 1.00, 1.03, 1.62, and 2.03. ¹³C NMR and APT spectra show resonances for one carboxylic acid carbon (δ_c 180.5, C-21), one ester carbon (δ_{C} 172.9), two oxygenated methines (δ_C 79.5, C-3; δ_C 77.2, C-15), and seven methyl moieties $(\delta_{C}$ 16.7, 19.5, 13.6, 28.6, 16.0, 18.7, 21.2). The HSQC spectrum was used to classify the remaining carbons as four methines, ten methylenes, and eight quaternary carbons. For the determination of the orientation of the acetoxy residue in position C-15, the Karplus equation describing the relationship between the dihedral angle and the vicinal coupling constant ³/ was used. Dihedral angles for both possible orientations of the acetoxy residue were calculated using Chem3D. The values obtained for α orientation clearly correspond to the experimentally obtained coupling constants in the ¹H NMR spectrum. Moreover, the interpretation of the NOE correlations allowed for the determination of the relative stereochemistry of compound 1. In particular, the correlation between H-19/H-29 proved the β orientation of both methyl groups. Moreover, the correlation between H-3/H-28 (both are α oriented) proved the β orientation of the hydroxyl group in C-3. The relative configurations at C-15, C-17 and C-20 were determined by the NOEs of H-15/H-16a (both are β) and H-15/H-18 (both are β), H-17/H-30 (both are α) and H-18/H-20 (C-20 is R) and H-16a/ H-22 (C-20 is R). The assignment of H-16a and H-16b was evaluated by the interpretation of the correlations between H-18/H-16b (H-16b is α oriented) and H-15/H-16a (H-16a is β oriented). The stereochemistry of the double bond between C-24/C-25 was deduced from the correlations between H-24/H-26 (E) and H-23/ H-27 (E).

Based on these data, the new natural compound 1 was identified as (24E)- $(3\beta, 15\alpha)$ -15-acetyloxy-3,26-dihydroxy-lanosta-8,24-dien-21-oic acid and named gloeophyllin K.

Constituent **2**, having a molecular formula of $C_{29}H_{44}O_3$ (*m/z* 439.3225 [M – H]⁻; calcd for $C_{29}H_{43}O_3^-$, 439.3218, $\Delta = -1.6$ ppm and *m/z* 463.3183 [M + Na]⁺; calcd for $C_{29}H_{44}O_3Na^+$, 463.3183, $\Delta = -0.2$ ppm), was also obtained as a white, amorphous powder.



Fig. 1 Chemical structures of isolated and identified G. odoratum triterpenes.

The ¹H NMR spectrum exhibits resonances of one secondary methyl at $\delta_{\rm H}$ 0.89 (d, J = 6.7 Hz), singlets of four tertiary methyl groups at $\delta_{\rm H}$ 0.98, 1.12, 1.61, and 1.66 and two vinyl proton signals at $\delta_{\rm H}$ 4.66 and $\delta_{\rm H}$ 4.82 (H-18a/b). Additionally, the ¹³C NMR and APT spectra of **2** show one oxygenated methine at $\delta_{\rm C}$ 76.5 (C-3) and one carboxylic moiety at $\delta_{\rm C}$ 181.7 (C-28) in its structure. As reported for gloeophyllin A (5) [18], the interpretation of the ¹H-¹H COSY and the HMBC data confirmed the presence of a rare 6/6/5/6 ring system, a so called C-nor-D-homoergosteroid skeleton.

The configurations of the methyl group at C-29 and the hydroxyl group in C-3 were found to be β oriented by NOEs of H-19/H-29 and H-3/H-5, respectively. The configurations at C-12, C-14, and C-17 were assigned via NOE correlations between H-12/H-14 and H-12/H-17, proving an α orientation of H-12, H-14, and H-17. Thus, the structure of the new compound **2** can be assigned to (3*S*,4*S*,4*aR*,6*bR*,9*R*,10*aR*,11b*S*)-3-hydroxy-4,11b-dimethyl-10-methylene-9-((*R*)-6-methylhept-5-en-2-yl)-2,3,4,4a,5, 6,6b,7,8,9,10,10a,11,11b-tetradecahydro-1H-benzo[a]fluorine-4-carboxylic acid (atom numbering according to systematic name generation), for which we gave the trivial name gloeophyllin L.

Triterpenes with this rare and unusual backbone have only been reported once from a fungal source (i.e., *Gloeophyllum abie-tinum* grown in Tibet) [18]. Biosynthetically, it is assumed that C-nor-D-homoergosteroids arise via a Wagner-Meerwein rearrangement reaction from a classical lanostane triterpene precursor. Here, the cleavage of carbon bonds between C-13/C-14 and C-8/C-14 are key steps.

The six known lanostane triterpenoids were identified by comparing their spectroscopic and spectrometric data (1D and 2D NMR and HRESIMS) with those in the literature: trametenolic acid B (3) [19], gloeophyllin B (4) [18], gloeophyllin A (5) [18], ergosterol peroxide (6) [20], eburicodiol (7) [21], and 21-hydroxylanosterol (8) [22]. Copies of the original spectra for known compounds are obtainable from the corresponding author. All *G. odoratum* isolates were tested in an *in vitro* CPE inhibition assay using the IAV H3N2 strain HK/68 and the 2009 pandemic H1N1 strain A/Jena/8178/09 (Jena/8178; **Table 2**). Additionally, their cytotoxicity was evaluated in MDCK cells. The SI of each compound was calculated (CC_{50}/IC_{50}) to evaluate the specificity of antiviral activity. The higher the SI, the lower is the risk of false-positive results.

Compounds 1, 2, and 4–6 showed no or weak anti-influenza virus activities, whereby compounds 2 and 4–6 exerted strong cytotoxicity with CC_{50} values between 7.3 and 45 µM. Whereas compounds 7 and 8 could only moderately protect MDCK cells against HK/68 infection, they significantly reduced the CPE caused by Jena/8178 to 50% at low micro molar concentrations (15.4 µM and 9.0 µM, respectively).

Compound **3** was identified as novel anti-influenza virus agent against both HK/68 and Jena/8178. Its IC_{50} values are in the low micro molar range with 14.1 and 11.3 μ M, respectively.

Based on the available data, it seems that an OH group in the form of an alcohol (compounds 7 and 8) or carboxylic acid (compounds 1 and 3) at position C-21 is favorable for the observed anti-influenza activity. However, more conclusive structure activity relationships between the features of the investigated triterpenes and the observed antiviral effects could not be established. Regarding the anti-influenza mechanism of action of the most effective compound 3, results from a fluorescence-based neuraminidase inhibition assay showed that this target protein on the viral surface can be excluded (data not shown). To evaluate the effects of compound 3 on the cell-free virus and different stages of the viral replication cycle, a modified plaque-reduction assay was performed. Here, 3 was able to bind to cell-free viruses and to neutralize their infectivity (> Fig. 3). Virus neutralization can be based on inhibitor binding to the viral envelope glycoproteins (i.e., hemagglutinin or neuraminidase, which are crucial for viral attachment and release, respectively) [23–25]. Hence, our data give first hints about possible targets of compound 3, which warrants

Table 1 ¹H and ¹³C NMR spectroscopic data of 1 and 2 (¹H: 500 MHz, CD₃OD, δ in ppm, *J* in Hz; ¹³C: 125 MHz, CD₃OD, δ in ppm, multiplicities determined by APT experiments).

Position	1		2		
	δ _H	δ _C	δ _H	δ _C	
1	1.75, m 1.22, m	33.9, CH ₂	1.70, m 1.37, m	36.8, CH ₂	
2	1.63, m	28.4, CH ₂	1.70, m 1.68, m	28.1, CH ₂	
3	3.15, dd (9.1, 7.0)	79.5, CH	4.01, dd (10.8, 5.1)	76.5, CH	
4	-	39.9, C	-	54.7, C	
5	1.03, m	51.7, CH	1.87, dd (12.4, 1.5)	48.6, CH	
6	2.07, m 2.02, m	21.7, CH ₂	1.66, m 1.37, m	22.5, CH ₂	
7	2.10, m 1.89, m	27.7, CH ₂	2.05, m 1.91, m	25.7, CH ₂	
8	-	134.0, C	-	138.0, C	
9	-	137.1, C	-	143.9, C	
10	-	38.3, C	-	36.1, C	
11	1.71, m 1.54, m	19.3, CH ₂	2.27, d (9.2)	36.2, CH ₂	
12	1.81, m 1.48, m	30.2, CH ₂	2.86, dt (9.2, 9.2, 8.5)	46.6, CH	
13	-	45.8, C	-	152.7, C	
14	-	51.9, C	2.36, dt (8.5, 7.5, 7.5)	49.6, CH	
15	5.03, dd (9.6, 5.6)	77.2, CH	1.54, m 1.16, m	24.8, CH ₂	
16	2.08, m 1.72, m	36.4, CH ₂	1.68, m 1.45, m	27.3, CH ₂	
17	2.22, overlapped	47.2, CH	1.94, m	50.2, CH	
18	0.87, s	16.8, CH ₃	4.82, d (2.4) 4.66, d (2.4)	112.4, CH ₂	
19	1.01, s	19.5, CH ₃	0.98, s	20.4, CH ₃	
20	2.22, overlapped	49.2, CH	1.60, m	33.9, CH	
21	-	180.5, C	0.89, d (6.7)	18.9, CH ₃	
22	1.51, m	33.4, CH ₂	1.59, m 1.00, m	35.3, CH ₂	
23	2.03, m	26.5, CH ₂	2.05, m 1.96, m	26.2, CH ₂	
24	5.38, dt (7.2, 7.2, 1.1)	125.8, CH	5.08, m	125.9, CH	
25	-	136.7, C	-	132.1, C	
26	3.91, s	68.9, CH ₂	1.67, s	26.0, CH ₃	
27	1.62, s	13.7, CH ₃	1.60, s	17.8, CH ₃	
28	0.98, s	28.6, CH ₃	-	181.7, C	
29	0.80, s	16.2, CH ₃	1.12, s	11.5, CH ₃	
30	1.04, s	18.7, CH ₃	-	-	
Ac <u>C</u> OO	-	172.9, C	-	-	
Ac <u>C</u> H ₃	2.03, s	21.2, CH ₃	-	-	

further investigation through more specific tests in a follow-up study.

In sum, *G. odoratum* fruit bodies have shown to be a rich source of anti-influenza triterpenes (**3**, **7**, and **8**). In particular, the most active ingredient–that is, trametenolic acid B (**3**)–is one of the major triterpenes contained in the EtOH extract (~ 3%).

Materials and Methods

General experimental procedures

Optical rotations were measured at 20 °C in MeOH on a Perkin-Elmer 341 polarimeter. FTIR spectra were acquired on a Bruker Tensor 27 spectrometer. 1D and 2D NMR experiments were performed by using a Bruker Avance 500 NMR spectrometer (Ultra-Shield) with a 5-mm switchable probe (TCI Prodigy CryoProbe, 5 mm, triple resonance inverse detection probe head) as described before [26]. The samples (1–2 mg) were measured at 298 K in fully deuterated MeOH referenced to the residual nondeuterated solvent signals. HRMS analyses were performed on a Bruker Daltonics maXis HD ESI-Qq-TOF mass spectrometer equipped with ESI in the positive and negative modes: capillary voltage, 1.5-4.5 kV; nebulizer, 0.4 bar (N₂); dry gas flow, 4 L/min (N₂); dry temperature, 200°C. The sum formulas of the ions were determined using Bruker Compass DataAnalysis 4.2 based on the mass accuracy (Δ $m/z \le 2$ ppm) and isotopic pattern matching (SmartFormula algorithm). Flash chromatography was performed on an Interchim puriFlash 4250 system, equipped with an ELSD and a PDA detector. Semi-preparative SFC was performed on a Waters Prep-15 System as described previously [27]. A Waters Viridis BEH column (5 µm; 10 × 250 mm) served as stationary phase. The mobile phase consisted of a supercritical CO₂/organic modifier (EtOH) gradient (T, 40 °C; flow rate, 15 mL/min). The fractions obtained from all chromatographic steps were analyzed by TLC (mobile phase: dichloromethane-MeOH-water [10:1.0:0.25]); stationary phase: Merck silica gel 60 PF254, detected after derivatization with vanillin/ H₂SO₄ (5% in MeOH) under visible light, UV₂₅₄, and UV₃₆₆. HPLC analysis was performed on a Dionex Ultimate 3000 RS system, equipped with a PDA detector and a charged aerosol detector on an Agilent Zorbax SB-C₁₈ rapid-resolution column (3.5 µm, 4.6 × 150 mm) with a MeCN/water gradient as mobile phase (0 min 45%/55%, 42 min 80%/20%, 43 min 95%/5%, 53 min 95%/ 5%, 54 min 45%/55%, 64 min 45%/55%). Conditions: T, 35°C; flow rate, 1 mL/min; injection volume, 10 µL; detection wavelength, 210, 254 nm. UPLC analysis was performed on a Waters Acquity UPLC system (H-class) equipped with a PDA detector and an ELSD. The stationary phase was a Waters Acquity UPLC BEH Phenyl column (1.7 µm, 2.1 × 100 mm). The mobile phase consisted of a MeCN/water gradient (0 min 84%/16%, 1.5 min 86%/14%, 1.6 min 98%/2%, 2.9 min 98%/2%, 3 min 84%/16%). Conditions: T, 40 °C; flow rate, 0.6 mL/min; injection volume, 5 µL; detection wavelength, 210 nm, and full range scan 190-400 nm.

(Ultrahigh-)gradient grade solvents and deuterated solvents from Merck were used.



▶ Fig. 2 Key HMBC, ¹H⁻¹H COSY, and NOESY correlations of compounds 1 and 2.

Fungal material

The fruit bodies of investigated polypores were collected in 2013 in Tyrol, Austria, and South Tyrol, Italy. They were dried at 40 °C for 5 d. Voucher specimens are deposited in the Herbarium of the Department of Pharmacognosy, University of Vienna, Austria. Exact collection locations/source and voucher ID numbers are the following: F. fomentarius: Neue Magdeburger Hütte (1300 m), Zirl, Austria (grown on beech), ID: FomfomE0019; F. pinicola: Viggartal, Ellbögen, Austria (grown on dead spruce trunk), ID: FompinE0010; G. lucidum: Plantasia GmbH, Oberndorf bei Salzburg, Austria, ID: 680898; G. applanatum: Oberperfuss, Austria (grown on spruce), ID: GanappE0012; G. odoratum: Oberperfuss, Austria (grown on spruce), ID: GloodoE0054, JR-20140310-A1; I. benzoinum: Geigerhof, Innichen/Vierschach in South Tyrol (Pustertal), Italy (grown on conifer wood), ID: IscbenE0038, L. sulphureus: Oberperfuss, Austria (grown on deciduous tree), ID: LaesulE0043; P. robustus: Hötting, Austria (grown on dead beech), ID: PherobE002; P. betulinus: Vill, Innsbruck, Austria (grown on birch); ID: PipbetE0029; T. gibbosa: Gnadenwald, Austria (grown on beech), ID: TragibE0052 [17]. Phylogenetic analyses were used to confirm the morphological identification and partial sequences of ITS rRNA were used to characterize the strains [28].

Extraction and isolation

The dried and powdered fruit bodies of *G. odoratum* strain 54 (389.0 g) were macerated at r. t. with EtOH 96% (3×5 L, 5 d each). After removal of the solvent under vacuum, the EtOH extract (31.2 g) was suspended in water and partitioned sequentially between petroleum ether (GO54EP; 5.6 g), dichloromethane (GO54ED; 18.4 g), and water (GO54EW; 2.8 g).

Of the GO54ED fraction, 12.8 g were subjected to silica gel column chromatography (Merck silica gel 60 PF₂₅₄, 517 g; 55 × 5 cm) using a gradient system of dichloromethane-ethyl acetate-MeOH (7.5 L) to give 17 fractions (A1–A17). Fraction A10 (2958–3247 mL) was recrystallized from acetone and identified as compound **3** (980 mg). Fractions A14–A15 (4896–6154 mL, 1.2 g) were combined and further separated via flash chromatography (PuriFlash C₁₈ HQ column [15 µm, 55 g]; flow rate: 26 mL/min) applying a gradient system of MeCN/water as mobile phase (0 min 10%/90%, 40 min 95%/5%, 70 min 95%/5%) to yield 10 fractions (B1–B10). Fraction B3 (574–602 mL, 48.0 mg) was purified via

Code	CC ₅₀ (CI95) [µM] in MDCK cells	IC_{50} (CI95) [μM] of CPE in MDCK cells		SI IAV HK/68	SI IAV Jena/8178
		IAV HK/68	IAV Jena/8178	[CC ₅₀ /IC ₅₀]	[CC ₅₀ /IC ₅₀]
Gloeophyllin K (1)	>100	n.a.	46.4 (40.14–54.45)	-	>2
Gloeophyllin L (2)	45.5 (37.35–54.26)	n.a.	n.a.	-	-
Trametenolic acid B (3)	>100	14.1 (9.76–19.16)	11.3 (9.74–14.41)	>7	>9
Gloeophyllin B (4)	35.5 (27.88–43.94)	n.a.	n.a.	-	-
Gloeophyllin A (5)	17.8 (15.81–19.81)	n.a.	n.a.	-	-
Ergosterol peroxide (6)	7.31 (4.96–11.47)	n.a.	n.a.	-	-
Eburicodiol (7)	>100	31.2 (17.66–48.01)	15.4 (8.13–25.07)	>3	>6
21-Hydroxylanosterol (8)	>100	34.5 (23.84–47.14)	9.00 (7.72–10.42)	>3	>11
Oseltamivir	-	0.004 (0.0004–0.01)	0.076 (0.06–0.10)	-	-

Table 2 Cytotoxicity and anti-influenza virus activity of compounds 1–8. CC₅₀ in MDCK cells and IC₅₀ determined against HK/68 and Jena/8178 in the CPE inhibition assay in MDCK cells are presented (n = 3; n. a.: not active).



▶ Fig. 3 Inhibition of different stages of the life cycle of IAV HK/68 by compound 3 at 100 μ M (plaque reduction assay performed in MDCK cells). Bars represent the mean values and SD of at least three experiments, each with two parallels per concentration.

another flash chromatography (PuriFlash C₁₈ HQ column [15 μ m, 35 g]; flow rate: 15 mL/min) applying a gradient system of MeCN/ water as mobile phase (0 min 35%/65%, 50 min 50%/50%, 52 min 95%/5%) to yield three fractions (C1–C3). Fraction C3 (432–448 mL) was identified as compound 1 (16.2 mg).

Fraction A12 (1.3 g) was suspended in water and partitioned sequentially between n-hexane (D1), ethyl acetate (D2), and water (D3). Fraction D2 (1.1 g) was further separated via flash chromatography (PuriFlash C₁₈ HQ column [15 μ m, 55 g]; flow rate: 26 mL/min) applying a gradient system of MeCN/water as mobile phase (0 min 45%/55%, 30 min 80%/20%, 55 min 98%/2%, 65 min 98%/2%) to yield nine fractions (E1–E9). Fraction E6

(1050–1155 mL, 130 mg) was subjected to semi-preparative SFC (Viridis BEH column [5 µm, 10 × 250 mm]; flow rate: 15 mL/min) applying a gradient system of supercritical CO₂/EtOH as mobile phase (0 min 90%/10%, 8 min 90%/10%, 9.5 min 50%/50%, 11.5 min 50%/50%, 12 min 90%/10%, 13 min 90%/10%) to yield three fractions (F1-F3). Fraction F2 was identified as compound 4 (45.0 mg). Fraction E7 (1156–1215 mL, 34 mg) was subjected to semi-preparative SFC (Viridis BEH column [5 µm, 10 × 250 mm]; flow rate: 15 mL/min) applying a gradient system of supercritical CO₂/EtOH as mobile phase (0 min 92%/8%, 9.5 min 92%/8%, 10 min 50%/50%, 11 min 50%/50%, 11.5 min 92%/8%, 12 min 92%/8%) to yield three fractions (G1–G3). Fraction G2 was identified as compound 2 (5.2 mg). Fraction E8 (1216-1395 mL, 63 mg) was subjected to semi-preparative SFC (Viridis BEH column [5 µm, 10 × 250 mm]; flow rate: 15 mL/min) applying a gradient system of supercritical CO2/EtOH as mobile phase (0 min 92%/8%, 9.5 min 92%/8%, 10 min 50%/50%, 11 min 50%/ 50%, 11.5 min 92%/8%, 12 min 92%/8%) to yield three fractions (H1–H3). Fraction H3 was identified as compound 5 (16.7 mg).

Fraction A8 (2.6–2.8 L, 350 mg) was separated via flash chromatography (PuriFlash C₁₈ HQ column [15 μ m, 35 g]; flow rate: 15 mL/min) applying a gradient system of MeCN/water as mobile phase (0 min 45%/55%, 35 min 80%/20%, 60 min 98%/2%, 90 min 98%/2%) to yield 19 fractions (I1–I19). Fraction I17 (1068– 1176 mL) was identified as compound **6** (11.9 mg).

The MeOH soluble part of fraction A9 (2788–2941 mL, 260.4 mg) was purified via Sephadex LH-20 column chromatography (mobile phase: MeOH) yielding seven fractions (J1–J7). Fractions J2–J3 (66–114 mL, 191 mg) were combined and further separated via silica gel column chromatography (Merck silica gel 60 PF₂₅₄, 110 g; 95 × 2 cm) eluting with a gradient solvent system of dichloromethane-MeOH, yielding nine fractions (K1–K9). Fraction K2 (856–912 mL, 69.5 mg) was further separated via silica gel column chromatography (Merck silica gel 60 PF₂₅₄, 14 g; 40 × 1 cm) eluting with an isocratic solvent system of dichloromethane-MeOH (10:0.25) to yield eight fractions (L1–L8). Fraction L3 (48–60 mL, 35 mg) was purified via silica gel column chromatography (Merck silica gel 60 PF₂₅₄, 11 g; 26 × 1 cm) eluting with a gradient solvent system of dichloromethane-MeOH, yielding 10 fractions (M1–M10). Fraction M4 (282–300 mL, 2.2 mg) was finally purified via UPLC including a fraction manager giving 1.0 mg of compound **7** and 1.0 mg of compound **8**.

The purity of the isolated compounds was determined by HPLC-PDA-MS to be > 98%.

Gloeophyllin K (**1**): white, amorphous powder; $[α]_D^{20} + 43.6^\circ$ (c 0.10, MeOH); ¹H and ¹³C NMR, see **Table 1**; HRESIMS *m*/*z* 553.3499 [M+Na]⁺ (calcd for C₃₂H₅₀NaO₆⁺, 553.3500, Δ = 0.1 ppm).

Gloeophyllin L (**2**): white, amorphous powder; $[α]_D^{20} + 21.9^\circ$ (c 0.33, MeOH); ¹H and ¹³C NMR, see **Table 1**; HRESIMS *m/z* 439.3225 [M – H]⁻ (calcd for C₂₉H₄₃O₃⁻, 439.3218, Δ = – 1.6 ppm).

Cell culture and viruses

Jena/8178 isolated from nasal swabs of an influenza patient during the pandemic of 2009 [29] and HK/68 (strain collection of the Institute of Medical Microbiology, Section Experimental Virology, Jena University Hospital Jena, Germany) were used in antiviral studies. They were propagated in MDCK cells (Friedrich-Loeffler Institute, Riems, Germany) in serum-free Eagle's minimum essential medium, $2 \mu g/mL$ trypsin, and 1.2 mM bicarbonate [30]. Titers of virus stocks were determined according to [31] in MDCK cells.

Determination of cytotoxicity and CPE inhibition

The CC₅₀ as well as IC₅₀ (inhibition of virus-induced CPE) was determined on 2-d-old confluent MDCK cell monolayers grown in 96-well plates as described previously (maximum tested concentration: 100 µg/mL for extracts/fractions and 100 µM for pure compounds) [32]. The CC₅₀ value represents the inhibitor concentration that reduces the viability of the treated cells in comparison to untreated control cells (no inhibitor) by half. For the calculation of these values, the mean optical density of six crystal violet stained cell controls was set to 100% viability [32]. In the CPE inhibition assay, six wells of untreated, noninfected (100% viability; no CPE) and six wells of untreated, infected cells (nearly complete CPE and no viability), without the test compound served as cell and virus control, respectively, on each plate. The percentage of antiviral activity of the tests compounds was calculated according to Pauwels et al. (1988) [33] using the following equation: antiviral activity = [(mean optical density of 6 cell controls - mean optical density of 6 virus controls)/(optical density of test - mean optical density of 6 virus controls)] × 100%. A 100% CPE inhibition means that 100% of virus-infected, inhibitor-treated cells were viable. The IC_{50} is the concentration of a virus inhibitor, where the response (CPE caused by the virus) is reduced by half (resulting in 50% cell viability = 50% of the mean optical density of the six cell controls).

All compounds were solved in DMSO. The maximum applied solvent concentration was 0.5% (v/v). Cytotoxicity was analyzed 72 h after adding the samples. CPE inhibition was measured 48 h after infection. A multiplicity of infection of 0.0008 and 0.003 TCID₅₀/cell of Jena/8178 and HK/68, respectively, resulted in a complete CPE at this time point. Each concentration was tested twice. At least three experiments were performed. Data leading to the determination of CC_{50} and IC_{50} values are provided as dose-response curves in **Fig. 24S**, Supporting Information. CC_{50} and IC_{50} values were calculated from mean dose-response curves of at least three independent experiments. Linear regression analysis using Microsoft Excel was applied in the linear scaled dose-dependent sample concentrations. To evaluate the specificity of antiviral activity, the SI was calculated as CC_{50}/IC_{50} ratio. Oseltamivir served as a positive control.

Plaque reduction assay

The assay time and temperature conditions of the plaque reduction assay were described recently [25, 34] and slightly modified. Here, confluent MDCK cell monolayers grown in 12-well plates were inoculated with ~ 30 pfu HK/68 in 500- μ L test medium. Cells and/or virus were exposed to **3** (100 μ M) before infection, during the viral adsorption phase, after adsorption phase, or during the whole replication cycle. Nontreated virus controls were included in all assays. Three independent assays were performed.

Supporting Information

The NMR, HRESIMS, and IR spectra of compounds 1 and 2 and graphs showing dose-dependencies for the investigated compounds (Fig. 24S) are provided as Supporting Information (Fig. 1S–23S).

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Conflict of Interest

The authors declare no conflicts of interest.

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