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Mass Spectrometry Guided Discovery and Design of Novel Asperphenamate Analogs From *Penicillium astrolabium* Reveals an Extraordinary NRPS Flexibility

Karolina Subko¹, Xinhui Wang¹, Frederik H. Nielsen^{1†}, Thomas Isbrandt¹, Charlotte H. Gotfredsen², Carmen Ramos³, Thomas Mackenzie³, Francisca Vicente³, Olga Genilloud³, Jens C. Frisvad¹ and Thomas O. Larsen^{1*}

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Subko K, Wang X, Nielsen FH, Isbrandt T, Gotfredsen CH, Ramos C, Mackenzie T, Vicente F, Genilloud O, Frisvad JC and Larsen TO (2020) Mass Spectrometry Guided Discovery and Design of Novel Asperphenamate Analogs From Penicillium astrolabium Reveals an Extraordinary NRPS Flexibility. Front. Microbiol. 11:618730. doi: 10.3389/fmicb.2020.618730 Asperphenamate is a small peptide natural product that has gained much interest due to its antitumor activity. In the recent years numerous bioactive synthetic asperphenamate analogs have been reported, whereas only a handful of natural analogs either of microbial or plant origin has been discovered. Herein we describe a UHPLC-HRMS/MS and amino acid supplement approach for discovery and design novel asperphenamate analogs. Chemical analysis of Penicillium astrolabium, a prolific producer of asperphenamate, revealed three previously described and two novel asperphenamate analogs produced in significant amounts, suggesting a potential for biosynthesis of further asperphenamate analogs by varying the amino acid availability. Subsequent growth on proteogenic and non-proteogenic amino acid enriched media, revealed a series of novel asperphenamate analogs, including single or double amino acid exchange, as well as benzoic acid exchange for nicotinic acid, with the latter observed from a natural source for the first time. In total, 22 new asperphenamate analogs were characterized by HRMS/MS, with one additionally confirmed by isolation and NMR structure elucidation. This study indicates an extraordinary nonribosomal peptide synthetase (NRPS) flexibility based on substrate availability, and therefore the potential for manipulating and designing novel peptide natural products in filamentous fungi.

Keywords: natural product discovery, mass spectrometry, filamentous fungi, asperphenamate, amino acid incorporation, biological activity, NRPS flexibility

INTRODUCTION

Asperphenamate (1) is a linear amino acid (AA) ester, comprised of N-benzoylphenylalanine 111 (2) and N-benzoylphenylalaninol (3). Asperphenamate, first discovered from *Aspergillus flavipes*in 1977 (Clark et al., 1977), was since found to be produced by a wide range of *Aspergillus*(Samson et al., 2011; Zheng et al., 2013; Ratnaweera et al., 2016; Hou et al., 2017) and *Penicillium*

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(Frisvad et al., 2004, 2013) species. Additionally, the compound 115 has also been isolated in trace amounts from a number of 116 unrelated plant species (Wu et al., 2004; Dang et al., 2014; 117 Zhou et al., 2017; Bunteang et al., 2018; Caridade et al., 2018), 118 suggesting endophytic fungi being the actual producers, rather 119 than the plants. Although asperphenamate is mainly known 120 for its antitumour activity and immense synthetic chemists 121 interest in asperphenamate backbone modification (Li et al., 122 2012; Yuan et al., 2012, 2018, 2019, 2020; Liu et al., 2016), 123 recent studies have also shown asperphenamate to be a potential 124 neuroinflamatory inhibitor (Zhou et al., 2017), and to possess 125 anti-HIV (Bunteang et al., 2018) and antidiabetic (Del Valle 126 127 et al., 2016) properties. In recent years, a handful of new natural 128 analogs have been isolated, namely Asperphenamates B (4) and 129 C (5) (Liu et al., 2018), and 4-OMe-asperphenamate (Zheng 130 et al., 2013; Ratnaweera et al., 2016) (6) from filamentous fungi. Other analogs containing partial structural similarities include: 131 patriscabratine (7), a N-benzoylphenylalanine phenylalanynol 132 acetate ester, aurantiamide (8) and aurantiamide acetate (9) 133 (Zhou et al., 2017), N-benzoylphenylalanine phenylalanynol 134 135 and phenylalanynol acetate amides, all isolated from plant material; cordyceamides A (10) and B (11) (Jia et al., 136 2009), a N-benzoyl-L-tyrosinyl-L-phenylalaninol and N-benzoyl-137 L-tyrosinyl-L-p-hydroxyphenylalaninol acetates, from an insect 138 pathogen fungus (Figure 1); along with a number of tentatively 139 identified related metabolites (Kildgaard et al., 2014; Sica et al., 140 2016). 141

Biosynthesis of asperphenamate was first described in the 142 filamentous fungus, P. brevicompactum (Li et al., 2018). Here, 143 a two module NRPS system was described, where the first 144 module, ApmA, is responsible for amide bond catalysis between 145 146 the phenylalanine and benzoic acid moieties, and subsequent 147 reduction to afford N-benzoylphenylalaninol (3), while the second module, ApmB, utilizes the same substrates to produce 148 N-benzoylphenylalanine (2), as well as catalyses the ester bond 149 formation between the two intermediates to release the final 150 product asperphenamate (1). Assuming, that other filamentous 151 fungi may follow the same or a similar biosynthetic pattern, the 152 production of 4-5 in Penicillium sp. and 6 in Aspergillus sp., 153 involving tyrosine, 4-OMe-phenylalanine and 4-hydroxybenzoic 154 acid instead of phenylalanine and benzoic acid as substrate 155 molecules, indicates promiscuity of either one or both NRPS 156 modules and provides new insights for production of novel 157 asperphenamate analogs and laying the grounds for molecular 158 biology work to achieve higher production of asperphenamate 159 and related analogs. 160

To contribute to a better understanding of the diversity 161 of asperphenamate biosynthesis and address the increasing 162 163 resistance toward anticancer drugs (Vasan et al., 2019), 164 P. astrolabium IBT 28865, a distant relative of P. brevicompactum from section Brevicompacta (Serra and Peterson, 2007), was 165 investigated for production of asperphenamate and related 166 analogs. In this study, we employ an ultra-high performance 167 liquid chromatography diode array detection quadrupole time 168 of flight high-resolution tandem mass spectrometry (UHPLC-169 DAD-QTOF-HRMS/MS) to dereplicate known and novel 170 asperphenamate analogs. As a result, 22 novel asperphenamate 171

analogs were characterized by HRMS/MS, of which 21 were 172 designed using proteogenic and non-proteogenic AAs as a 173 supplement to the growth media. This study has further revealed 174 a rare promiscuity of a fungal NRPS, laying the grounds for 175 future NRPS research in filamentous fungi. Altogether, this 176 study demonstrates the HRMS/MS based dereplication and 177 characterization of novel analogs of a known bioactive peptide 178 scaffold to be a powerful strategy in natural product discovery. 179

MATERIALS AND METHODS

Reagents, Strains, and Media

All solvents and reagents were purchased from Sigma-Aldrich185(St. Louis, MO, United States), for the exception of para-186substituted phenylalanines, which were acquired from Bachema187(Bubendorf, Switzerland); ultra-pure water used throughout the188study was filtered with a Milli-Q system (Millipore, Burlington,189MA, United States).190

Penicillium astrolabium (IBT 28865), Penicillium olsonii (IBT28864), Penicillium bialowiezense (IBT 28294), and Penicillium192brevicompactum (IBT 30524) are filamentous fungi from the193IBT culture collection at the Department of Biotechnology andBiomedicine, Technical University of Denmark.

For the chemical profile analysis, P. astrolabium was cultivated 196 with 3-point inoculation on Czapek yeast agar (CYA), yeast 197 extract sucrose agar (YES) and malt extract agar (MEA; Oxoid) 198 for 7, 10, and 14 days at 20 and 25°C in the dark. For large scale 199 cultivation, the fungus was cultivated with 3-point inoculation on 200 200 YES agar plates, and incubated for 10 days at 25°C in the 201 dark. For a proteogenic AA incorporation study, the fungus was 202 cultivated with 3-point inoculation on Czapek (CZ) agar plates 203 (10 cm) for 14 days at 25°C in the dark. Here, triplicates of 24 204 sets of supplemented media were used: 20 with all proteogenic 205 AAs at 5 g/L; two for anthranilic acid and 4-hydroxybenzoic 206 acid at 2.5 g/L, and two with additional inorganic nitrogen 207 supplement of NaNO₃ at 5 g/L and 10 g/L. For a non-proteogenic 208 AA incorporation study, the fungus was cultivated with 1-point 209 inoculation on CZ agar plates (6 cm) for 14 days at 25°C in the 210 dark. Here, triplicates of four sets of 4-chloro-L-phenylalanine, 4-211 bromo-L-phenylalanine, 4-amino-L-phenylalanine, and 4-nitro-212 L-phenylalanine supplemented media were used at 2.5 g/L. 213

For the chemical profile analysis and comparison, all four 214 fungi were cultivated with 3-point inoculation on minimal 215 media (MM), CZ, CYA, and YES 10 cm agar plates for 7 days 216 25°C in the dark. 217

Extraction and Isolation

For chemical profiling and the asperphenamate analog design 220 study, five 6 mm diameter plugs taken in triplicates and extracted 221 with acidic (1% formic acid; FA) isopropanol (iPr) – ethyl acetate 222 (EtOAc) (1:3 v/v) as described by Smedsgaard (1997). All samples 223 were re-dissolved ultrasonically for 10 min in 100 μ L methanol 224 (MeOH) and centrifuged prior to analysis by LC-MS. 225

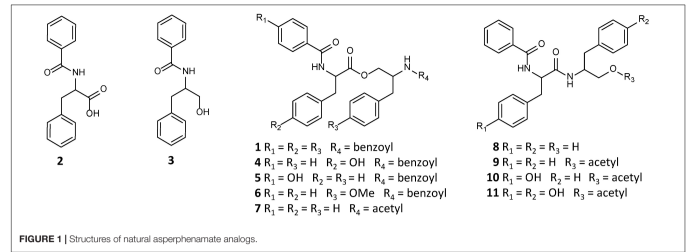
For large-scale extraction, the agar plates were extracted twice 226 with acidic (1% FA) EtOAc. The liquid-liquid partitioning was 227 then performed on the crude extract, by dissolving it with 90% 228

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data, see Table 1 and Supplementary Figure S1; HRESIMS m/z $507.2279 [M+H]^+$ (calculated for $C_{32}H_{30}N_2O_4$, m/z 507.2278). Asperphenamate Y (4): white powder; $[\alpha]^{20}D - 25.8^{\circ}$ (c 0.12, CHCl₃); UV (MeCN) λ_{max} 239 and 275 sh nm; ¹H and ¹³C NMR 286

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data, see Table 1 and Supplementary Figure S2; HRESIMS m/z $523.2227 [M+H]^+$ (calculated for $C_{32}H_{30}N_2O_5$, m/z 523.2227). Asperphenamate W (12): white powder; $[\alpha]^{20}$ _D -43.6° (c 0.14, CHCl₃); UV (MeCN) λ_{max} 236 and 278 nm; ¹H and ¹³C NMR data, see Table 1 and Supplementary Figure S3; HRESIMS m/z 546.2389 $[M+H]^+$ (calculated for C₃₄H₃₁N₃O₄, m/z 546.2387) Asperphenamate L (13): white powder; UV (MeCN) λ_{max}

314 237 and 272 sh nm; ¹H and ¹³C NMR data, see Table 1 and 315 Supplementary Figure S4; HRESIMS m/z 473.2436 [M+H]⁺ 316 (calculated for $C_{29}H_{32}N_2O_4$, m/z 473.2435).

Asperphenidine F1 (1a): white powder; UV (MeCN) λ_{max} 238 318 and 272sh nm; ¹H NMR data, see Supplementary Table S5 and 319 Supplementary Figure S5; HRESIMS m/z 508.2230 [M+H]⁺ (calculated for C₃₁H₂₉N₃O₄, m/z 508.2231). 321

UHPLC-DAD-QTOF-MS Analysis

All samples were analyzed on an Agilent Infinity 1290 UHPLC 324 system (Agilent Technologies, Santa Clara, CA, United States) 325 equipped with a diode array detector (DAD), monitoring between 326 190 and 640 nm. Separation was achieved on an Agilent Poroshell 327 120 phenyl-hexyl column (150 mm \times 2.1 mm, 1.9 μ m particles) 328 with a flow rate of 0.35 mL/min at 40°C, using a linear acetonitrile 329 (MeCN)/water (both buffered with 20 mM FA) gradient of 330 10 to 100% MeCN in 10 min, followed by 2 min flush at 331 100% MeCN, return to starting conditions in 0.1 min and 332 equilibration at 10% for 2 min before the following run. It 333 was coupled to an Agilent 6,545 QTOF MS equipped with 334 Dual Jet Stream ESI source with the drying gas temperature 335 of 250°C and gas flow of 8 L/min and sheath gas temperature 336 of 300°C and flow of 12 L/min, capillary voltage 4,000 V and 337 nozzle voltage of 500 V. The mass spectrometer was operated 338 in positive polarity, recording centroid data in m/z range 100 339 to 1,700 for MS mode, and 30-1,700 for MS/MS mode, with 340 acquisition rate of 10 spectra/s. Automated HRMS/MS was 341 done for ions detected in the full scan above 50,000 counts, 342

246 MeOH:water and treating it with the same amounts of heptane, 247 resulting in two phases. After separating the heptane phase, the 248 90% MeOH:water fraction was then diluted with water to get 249 50% MeOH:water, and further treated with dichloromethane 250 (DCM), resulting in three phases overall. The DCM phase 251 was dried before loading onto a 50 g SNAP column (Biotage, 252 Uppsala, Sweden) with diol material (Isolute diol, Biotage). Crude 253 fractionation was performed using an Isolera One automated 254 flash system (Biotage) with stepwise increments of 25% at 255 50 mL/min in heptane-DCM-EtOAc-MeOH system, starting at 256 100% heptane, finishing at 100% MeOH, resulting in 13 fractions 257 (i.e., heptane, heptane 3:1 DCM, heptane:DCM, heptane 1:3 258 DCM, DCM, DCM 3:1 EtOAc, DCM:EtOAc, DCM 1:3 EtOAc, 259 EtOAc, EtOAc 3:1 MeOH, EtOAc:MeOH, EtOAc 1:3 MeOH, 260 and MeOH), with 300 mL each. Selected resulting fractions 261 were further fractionated on a 25 g SNAP column with RP C18 262 material (Grace, 15 μ m/100 Å) at a flow rate of 30 mL/min 263 using a stepwise 30-100% MeOH:water (both buffered with 264 50 ppm TFA) gradient as follows: in 10% increments at 30-265 50, 5% increments of 50-80, and 10% increments of 80-266 100%, resulting in 11 fractions (i.e., 30, 40, 50, 55, 60, 65, 267 70, 75, 80, 90, and 100%). Further separation was achieved 268 on an Agilent Infinity 1290 HPLC-DAD (Agilent Technologies, 269 Santa Clara, CA, United States) system, with UV monitoring 270 at 230 and 280 nm, a flow rate of 4 mL/min and column 271 temperature at 40°C as follows: Asperphenamate (1) and 272 Asperphenamate L (13) were purified on a Gemini C₆-Phenyl 273 column (5 μ m, 110 Å, 250 \times 10 mm, Phenomenex) using a 274 linear gradient of 57 to 64% acetonitrile (MeCN)/water over 275 30 min; Asperphenamate W (12) on a Kinetex RP C18 column 276 (5 μ m, 100 Å, 250 \times 10 mm, Phenomenex) using a linear 277 gradient of 55 to 65% MeCN/water over 20 min at a flow rate 278 of 4 mL/min; Asperphenamate Y (4) and Asperphenidine F1 (1a) 279 280 on a Kinetex RP C18 column (5 μ m, 100 Å, 250 mm \times 10 mm, Phenomenex) using a linear gradient of 55 to 67% MeOH/water 281 over 20 min at a flow rate of 4 mL/min. All solvents were buffered 282 with 50 ppm TFA. 283

Asperphenamate (1): white powder; $[\alpha]^{20}_{D} - 25.5^{\circ}$ (c 0.11, 284 CHCl₃); UV (MeCN) λ_{max} 238 and 272 sh nm; ¹H and ¹³C NMR 285

	1		4		12		13	
Position	δ c *	δ _H (J in Hz)	δc	δ _H (J in Hz)	δc	δ _H (J in Hz)	δc	δ _H (J in Hz)
1			167.7		167.6		167.8	
2			133.4		133.5		133.3	
3	127.0	7.70 dd (8.3, 1.1)	127.2	7.65 m	127.3	7.63 m	127.1	7.72 m
4	128.6	7.39 m	128.8	7.39 m	128.6	7.29 m	128.6	7.41 m
5	131.9	7.50 tt (7.5, 1.1)	132.2	7.50 tt (7.4, 1.2)	132.1	7.48 t (7.4)	132.2	7.52 m
1′			172.2		172.5		173.2	
2′	54.4	4.92, q (6.6)	54.8	4.87 d (6.7)	54.3	5.04 q (6.5)	52.1	4.71 m
3' NH		6.58 d (6.6)		6.59 d (6.6)		6.69 d (6.4)		6.46 d (6.8)
4′	37.5	3.29 dd (14.0, 6.6)	36.9	3.20 dd (14.0, 6.5)	27.7	3.43 d (6.0)	40.8	1.79 m
		3.21 dd (14.0, 7.0)		3.14 dd (14.0, 6.9)				1.69 m
5′			127.6		110.1		25.1	1.75 m
6′	129.1	7.21 m	130.5	7.05 m	127.5		22.2	0.99 d (6.5)
7′	128.8	7.29 m	116	6.76 m	118.7	7.64 m	22.8	1.02 d (6.5)
8′	126.7	7.24 m	155.3		120.1	7.12 t (7.4)		
9′					122.7	7.20 m		
10′					111.6	7.33 d (7.9)		
11′					136.4			
12' NH						8.06 s		
13′					123.1	7.06 d (2.2)		
1″	65.3	4.54 dd (11.4, 3.4)	65.5	4.50 dd (11.4, 3.6)	65.4	4.46 dd (11.6, 3.6)	65.1	4.59 dd (11.5, 3.3
		4.04 dd (11.4, 4.4)		4.04 dd (11.4, 4.5)		4.06 dd (11.6, 4.6)		4.08 dd (11.5, 4.6
2″	50.2	4.62 m	50.6	4.60 m	50.6	4.55 m	50.5	4.69 m
3″ NH		6.67 d (8.4)		6.71 d (8.4)		6.59 d (8.4)		6.73 d (8.2)
4″	37.2	3.00 dd (13.7, 6.4)	37.4	3.01 dd (13.9, 6.5)	37.3	2.94 dd (13.6, 6.7)	37.3	3.10 dd (13.6, 6.5
		2.89 dd (13.8, 8.5)		2.91 dd (13.9, 8.2)		2.81 dd (13.6, 8.4)		3.01 dd (13.8, 8.2
5″			137.1		137.4		137.3	
6″	129.2	7.23 m	129.5	7.22 m	129.5	7.18 s (7.6)	129.3	7.29 m
7″	128.3	7.32 m	128.9	7.31 m	128.7	7.36 t (7.8)	128.7	7.32 m
8‴′	127.3	7.25 m	127	7.24 m	126.9	7.23 m	126.8	7.25 m
1‴′			167.8		167.4		167.3	
2‴′			134.1		134.4		134.3	
3‴′	126.9	7.65 dd (8.3, 1.1)	127.3	7.68 m	127.2	7.63 m	127.1	7.70 m
4″	128.6	7.31 m	128.6	7.31 m	128.8	7.29 m	128.4	7.28 m
5‴	131.3	7.43 tt (7.5, 1.1)	131.7	7.43 tt (7.4, 1.1)	131.5	7.43 t (7.4)	131.3	7.42 m

*13C NMR data available only from HSQC experiment.

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with a cycle time of 0.5 s, quadrupole width of m/z \pm 0.65 384 using fixed CID energies of 10, 20, and 40 eV with maximum 385 three precursor ions per cycle. A lock mass solution of 70% 386 MeOH was infused in the second sprayer, with an extra LC 387 pump at a flow of 15 µL/min using a 1:100 splitter. The 388 solution contained 1 µM tributylamine (Sigma-Aldrich) and 389 10 µM hexakis(2,2,3,3-tetrafluoropropoxy)phosphazene (Apollo 390 Scientific Ltd., Cheshire, United Kingdom) as lock masses. The 391 $[M + H]^+$ ions of both compounds (m/z 186.2216 and 922.0098, 392 respectively) were used. 393

In-house fungal metabolite library search was done as described by Kildgaard et al. (2014). Data files were processed in MassHunter workstation B.07.00 with "Find by Auto MS/MS function" with a processing limit to 200 largest peaks and mass match tolerance m/z 0.05. HRMS/MS library search using parent and fragment ion accuracy of 20 ppm + 2 mDa, with minimal forward score of 50 and reverse score of 80.

Targeted analysis for the asperphenamate analog design study 443 was performed using expected masses of individual AA and 444 benzoic acid analogs, for all potential precursors, intermediates 445 and final products, see Supplementary Table S1. Relative 446 amounts of asperphenamate analogs were quantified by direct 447 integration of peak area of target compounds, normalized to 448 xanthoepocin water loss adduct peak area ([M-H₂O+H]⁺ m/z 449 589.0975) in control samples. All analyses were performed in 450 triplicates. All MS/MS spectra reported were at 20 eV, unless 451 stated otherwise. 452

General Experimental Procedures

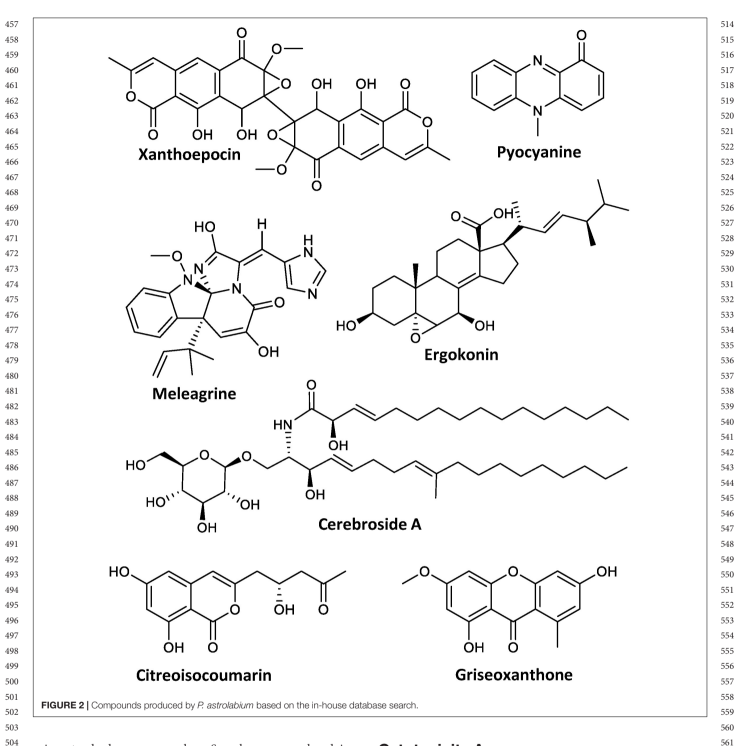
1D and 2D NMR analyses were performed on a Bruker Avance 455 800 MHz spectrometer (Bruker, Billerica, MA, United States), 456

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using standard sequence pulses. Samples were analyzed in a 3 mm TCl cryoprobe using deuterated chloroform (CDCl₃) and referenced to the residual solvent signals $\delta_H = 7.26$ ppm and $\delta_C = 77.16$ ppm. J-couplings are reported in hertz (Hz) and chemical shifts (δ) in ppm. For 1D and 2D NMR data, see **Supplementary Tables 2–4** and **Supplementary Figures 1–5**.

Optical rotations were measured in chloroform (CHCl₃) on a PerkinElmer 341 Polarimeter (PerkinElmer, Waltham, MA, United States).

Cytotoxicity Assay

Compounds 1, 1a, 4, 12, and 13 were tested in triplicates 563 against five cancer cell lines, i.e., human lung carcinoma 564 A549 ATCC CCL-185, breast adenocarcinoma MCF7 565 ATCC HTB-22, human skin melanoma A2058 ATCC CRL-566 11147, hepatocyte carcinoma HepG2 ATCC HB-8065 and 567 pancreas carcinoma MiaPaca-2 ATCC CRL-1420 following 568 previously described methodology (Audoin et al., 2013; 569 Lauritano et al., 2020). 570

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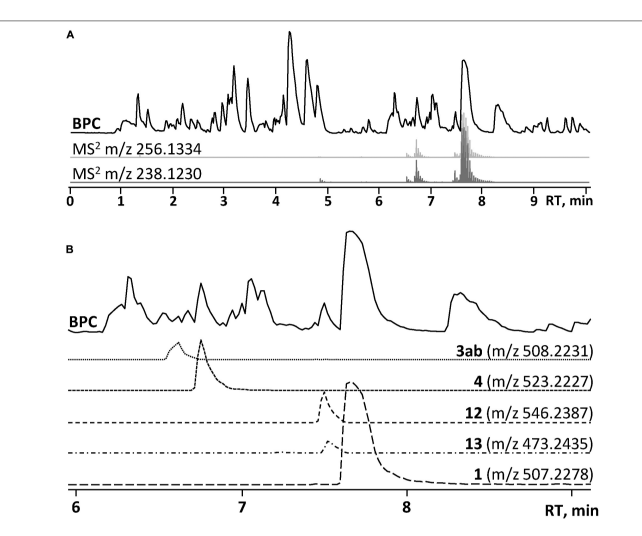


FIGURE 3 | HPLC-MS analysis for asperphenamate analog production on YES media. (A) Base peak chromatogram (BPC) of crude *P. astrolabium* extract, with extracted ion chromatogram (EIC) showing "bait" ions m/z 256.1334 and m/z 238.1230. (B) Zoomed BPC with EIC for asperphenamate analogs at full scan.

RESULTS

⁶¹⁰ Chemical Profile of Penicillium ⁶¹² astrolabium

To investigate the secondary metabolite profile from P. astrolabium, the fungus was inoculated on three media (MEA, CYA, and YES) and incubated at 20 and 25°C for 7, 10, and 14 days. The resulting 18 extracts were analyzed by UHPLC-DAD-QTOF-MS and used for automated in-house library search of fungal secondary metabolites (Kildgaard et al., 2014). In addition to previously reported asperphenamate (1), N-benzoylphenylalanine (2) and xantoepocin (Perrone et al., 2015), all 18 extracts also revealed the presence of meleagrine and its biosynthetic intermediates neoxaline, glandicoline B, roquefortine C and histidyltryptophanyldiketopiperazine (Ali et al., 2013), as well as cerebroside A. Other notable secondary metabolites include ergokonin B and pyocyanine detected on YES and CYA media extracts; citreoisocoumarin detected only in YES extracts; griseoxanthone C mainly seen

on CYA 20°C (**Figure 2**). Additionally, a series of di- and tetracyclopeptides with varying AA composition, depending on growth conditions, were produced.

Targetted Asperphenamate Daughter Ion Search Reveals Novel Analogs

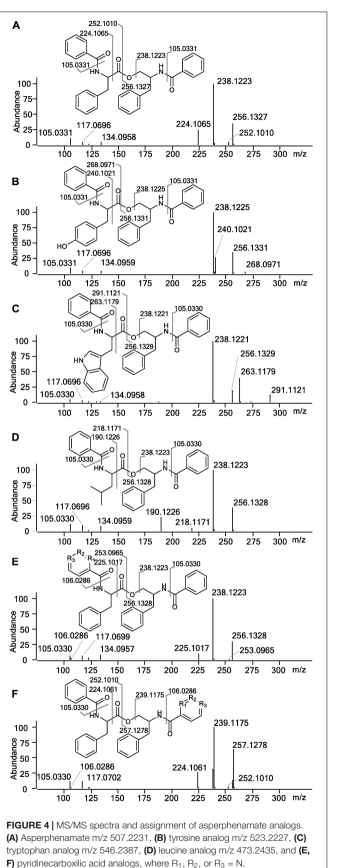
In the HRMS/MS analysis, asperphenamate (1) and reported fungal analogs (4-6) share two major fragment ions m/z 256.1334 and m/z 238.1230, corresponding to ester bond cleavage to result in a N-benzovlphenvlalaninol protonated ion $[C_{16}H_{18}NO_2]^+$, followed by water loss on the same moiety to get $[C_{16}H_{16}NO]^+$. In addition, a minor fragment of m/z 105.0335 [C₇H₅O]⁺ corresponding to a benzoyl loss was also observed. To screen for potential asperphenamate analogs, the two major fragment ions were used as "bait" (Figure 3A), resulting in five major peaks (Figure 3B): asperphenamate $([M+H]^+ m/z 507.2277, C_{32}H_{30}N_2O_4)$, the most abundant analog with an extra oxygen atom ([M+H]⁺ m/z 523.2227, C₃₂H₃₀N₂O₅), an analog indicating a single carbon-nitrogen

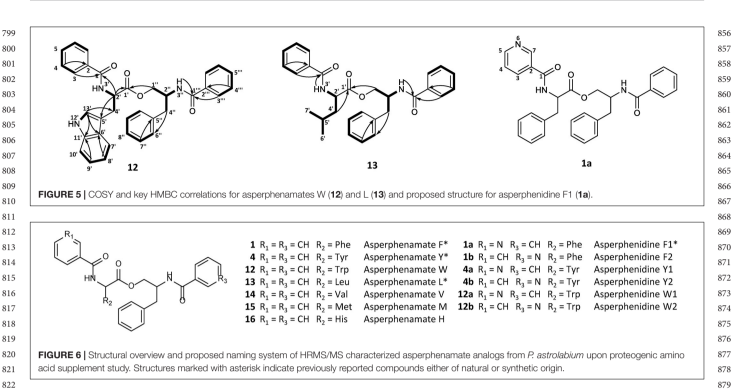
exchange ([M+H]+ m/z 508.2232, C31H29N3O4), and two other analogs with a significant mass differences, one 34Da lower $([M+H]^+ m/z 473.2436, C_{29}H_{32}N_2O_4)$ and the other 39Da higher ([M+H]⁺ m/z 546.2385, C₃₄H₃₁N₃O₄) to that of asperphenamate. Subsequent MS/MS analysis revealed, that two asperphenamate-specific fragments m/z 252.1010 and m/z 224.1064, corresponding to ester cleavage and subsequent CO loss of an N-benzoylphenylalanine moiety (Figure 4A), have been exchanged for fragments 16Da higher, namely m/z 268.0971 and m/z 240.1021 (Figure 4B), for compound with m/z 523.2227. Therefore, with the fragment of m/z 105.0331 being present in both of compounds, and no other major differences in fragmentation patterns observed, a phenylalanine exchange for tyrosine in a non-reduced N-benzoyl AA moiety could be proposed, resulting in 4. The asperphenamate analog with m/z 546.2385 produced unique fragments of m/z 291.1121 $[C_{18}H_{15}N_2O_2]^+$ and m/z 263.1179 $[C_{17}H_{15}N_2O]^+$ (Figure 4C). Taking into account the presence of a benzoyl ion $[C_7H_5O]^+$, the rest of m/z 291.1121 fragment suggest molecular formula $C_{11}H_{10}N_2O$, corresponding to phenylalanine exchange for tryptophan in a non-reduced N-benzoyl AA moiety. Similarly, the differences between the unique fragments in m/z 473.2436 and the benzoyl ion led to proposal of leucine containing analog (Figure 4D). Finally, a compound with m/z 508.2232 showed similar fragmentation patterns to those of asperphenamate, however, fragments corresponding to fragmentation of nonreduced AA moiety showed a fragment mass increase by 1Da, with fragment ion m/z 106.0286 $[C_6H_4NO]^+$ indicating a pyridinecarboxylic acid incorporation (Figure 4E). Moreover, MS/MS data revealed trace amounts of a coeluting isomer, with the two major ion fragments weighing 1 Da higher, namely m/z 257.1278 and m/z 239.1175, hence indicating that the pyridinecarboxylic acid can also be incorporated into the reduced AA part of the molecule (**Figure 4F**).

NMR Confirms Phenylalanine Exchange for Other Amino Acids in the Non-reduced N-Benzoyl Amino Acid Moiety

To confirm the structures proposed by HRMS/MS fragmentation patterns, a large scale of 200 agar plates was grown for targeted isolation of asperphenamate (1), and tyrosine (4), tryptophan (12) and leucine (13) analogs, as well as one of the pyridinecarboxylic acid analogs (1a). ¹H and ¹³C NMR data shown in Table 1, with full assignment table and spectra available in supplementary material (Supplementary Tables 2-4 and Supplementary Figures 1-5). Data for asperphenamate (1) and the tyrosine analog (4) fit with previously published data (Catalán et al., 2003; Liu et al., 2018). ¹H and ¹³C NMR shifts of N-benzoylphenylalaninol and the N-benzoyl part of non-reduced AA moiety was in agreement within all four compounds, further supported by COSY and HMBC correlations for tryptophan (12) and leucine (13) analogs (Figure 5 and Supplementary Tables 3-4).

The rest of the shifts corresponding to **12** showed three spin systems, with the first comprised of an amino group at NH-3'





823 ($\delta_{\rm H}$ 6.69), a methine at H-2' ($\delta_{\rm H}$ 5.04) and a methylene at H-824 4' ($\delta_{\rm H}$ 3.43), the second consisted of four aromatic methines at 825 H-7' ($\delta_{\rm H}$ 7.64), H-8' ($\delta_{\rm H}$ 7.12), H-9' ($\delta_{\rm H}$ 7.20) and H-10' ($\delta_{\rm H}$ 826 7.33), and the third one included aromatic amino and methine 827 groups, NH-12' ($\delta_{\rm H}$ 8.06) and H-13' ($\delta_{\rm H}$ 7.06), respectively. 828 The HMBC correlations of the last two spin systems from H-829 7' and H-13' to C-5' (δ_C 110.1), H-8' and H-13' to C-6' (δ_C 830 127.5), and H-9' and H-13' to C-11' (δ_C 136.4), revealed the 831 presence of indole, which was connected to the first spin system 832 by H-2' to C-5' ($\delta_{\rm C}$ 110.1) and H-4' to C-13' ($\delta_{\rm C}$ 123.1), to 833 confirm presence of tryptophan. The HMBC correlations from 834 H-3 ($\delta_{\rm H}$ 7.63) and H-2' to C-1 ($\delta_{\rm C}$ 167.6) and H-2' and H-835 1'' ($\delta_{\rm H}$ 4.46/4.06) to C-1' ($\delta_{\rm C}$ 172.5), connected tryptophan 836 moiety to the benzoyl and N-benzoylphenylalaninol parts of the 837 molecule (Figure 5). 838

For 13, the rest of the shifts comprised a single spin system of 839 amino group NH-3' ($\delta_{\rm H}$ 6.46), two methines at H-2'($\delta_{\rm H}$ 4.71) and 840 H-5'($\delta_{\rm H}$ 1.75), a diastereotopic methylene at H-4'($\delta_{\rm H}$ 1.79/1.69), 841 and two methyl groups at H-6'($\delta_{\rm H}$ 0.99) and H-7'($\delta_{\rm H}$ 1.02), to 842 give a leucine backbone. The spin system was connected to the 843 rest of the structure by H-3 (δ_H 7.72) and H-2' to C-1 (δ_C 167.8) 844 and H-2' and H-1' ($\delta_{\rm H}$ 4.59/4.08) to C-1' ($\delta_{\rm C}$ 173.2) (Figure 5). 845 The NMR data was eventually found to be in agreement with the 846 commonly overlooked lichen secondary metabolite hypothallin 847 (Huneck et al., 1992). 848

Compound **1a** was purified in trace amounts (0.2 mg) and only 1H NMR was acquired (**Supplementary Figure S5**). In comparison to asperphenamate, all the proton shifts in the aliphatic range for **3a** were of same multiplicity and similar shift values, as well as shifts for both amino groups. In the aromatic range, three asperphenamate shifts at H-3 ($\delta_{\rm H}$ 7.70), H-4 ($\delta_{\rm H}$ 7.39) and H-5 ($\delta_{\rm H}$ 7.50) were swapped for more downfield shifts at $\delta_{\rm H}$ 7.94 (m), $\delta_{\rm H}$ 8.72 (dd), and $\delta_{\rm H}$ 8.87 (d). Based on the two latter shifts and their multiplets, they were assigned as H-5 and H-7, respectively, with $\delta_{\rm H}$ 7.94 (m) assigned at H-3, and the H-4 shift assigned to the general aromatic region at 7.18–7.35, led to confirmation of pyridinecarboxylic acid moiety as nicotinic acid. This fits with the published NMR data for nicotinic acid (Chen et al., 1999) and the corresponding synthetic asperphenamate analog (Liu et al., 2016).

Herein we propose a new asperphenamate analog naming system using one letter AA abbreviation to denote a specific AA incorporation, based on similar azaphilone pigment naming system proposed by Isbrandt et al. (2020). Compounds **12** was named asperphenamate W, whereas compounds **4** and **13** will be referred to as asperphenamate Y and L, respectively. Compound **1a** was named asperphenidine F1, to signify it being an asperphenamate analog, with phenylalanine incorporation and benzoic acid exchange for nicotinic acid at the non-reduced part of the molecule.

Amino acid Enriched Media Induces Phenylalanine Exchange in the N-Benzoylphenylalanine Moiety

To investigate if higher AA availability can induce AA exchange 904 in asperphenamate biosynthesis, the fungus was incubated 905 on CZ media supplemented with one of each of the 20 906 proteogenic AAs. Subsequently, targeted MS analysis was 907 performed by search of masses corresponding to phenylalanine 908 exchange for one AA moiety within the asperphenamate 909 backbone (Supplementary Table S1). In addition tyrosine (4), 910 tryptophan (12), and leucine (13) analogs, the novel valine 911 (14), methionine (15), histidine (16), alanine and isoleucine 912

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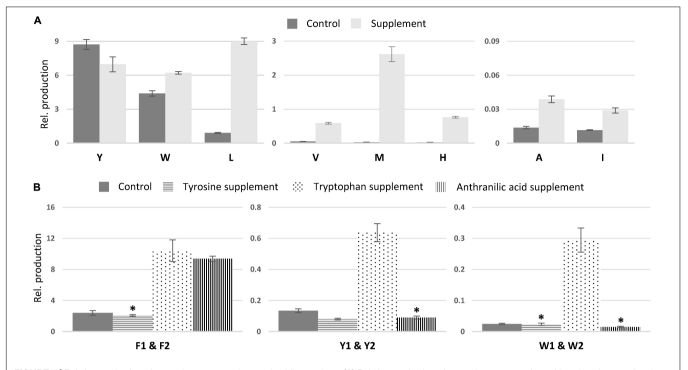


FIGURE 7 | Relative production of asperphenamate and asperphenidine analogs. (**A**) Relative production of asperphenamate amino acid analogs in control and corresponding amino acid supplement media. (**B**) Production of asperphenidines F, Y, and W in control, tyrosine, tryptophan and anthranilic acid supplemented media. Error bars show standard deviation. All data, except the ones marked with asterisk, indicate statistically significant differences compared to the control (*ρ* < 0.05).

analogs could also be observed, and AA exchange in the nonreduced AA part of the molecule was confirmed for compounds
14–16 by HRMS/MS fragmentation patterns (Figure 6 and
Supplementary Figure S6). The new analogs were accordingly
named as asperphenamates V, M, and H.

In comparison to non-fed control cultures of previously characterized compounds, only small changes in production were observed for asperphenamate Y and W production, a slight decrease and increase, respectively (Figure 7A). However, upon leucine supplement, asperphenamate L production increased 10-fold in comparison to the control. A similar increase pattern was also observed in the valine supplement experiment, whereas the production of histidine and methionine analogs was drastically boosted upon respective AA supplement, with 33-fold increase for asperphenamate H and more than a 100-fold increase for asperphenamate M. Although, the production of alanine and isoleucine analogs increased by three-fold for each, the relative amounts were still marginally lower in comparison to other uptake experiments, and were not sufficient for MS/MS data acquisition and assignment of structures (Figure 7A).

Additionally, MS analysis targeting reduced and non-reduced AA N-benzoyl precursors, and single or double AA exchange in asperphenamate, patriscabratine, aurantiamide and aurantiamide acetate backbones, was performed to result in the discovery of an additional double leucine asperphenamate analog characterized by HRMS/MS (Supplementary Figure S6). No other AA analogs or analogs

for benzoic acid exchange for 4-hydroxybenzoic or anthralinic acids were observed.

Tryptophan Induces Nicotinic Acid Incorporation

In contrast to the expected asperphenamate W (12) being one of the major metabolites upon growth on tryptophan supplemented media, both asperphenidines F1 and F2 (1ab), with nicotinic acid exchange on either the non-reduced or reduced part of the molecule, respectively, showed the most drastic increase in the relative amount in comparison to the control. With the similar behavior observed in anthranilic acid supplement, targeted MS analysis of all extracts was performed with masses corresponding to the benzoic acid exchange for nicotinic acid for novel AA analogs described above (Supplementary Table S1). The MS profile indicated the potential for nicotinic acid incorporation in all eight AA analogs, however, only tyrosine (4ab) and tryptophan (12ab) analogs, with nicotinic acid incorporation in either of two possible positions, could be confirmed by MS/MS (Figure 6 and Supplementary Figure S7). With individual analog peaks strongly overlapping in the MS profile, MS/MS of two coeluting analogs as well as combined peak area were used for further structure assignment and relative quantification, respectively.

The relative amounts of the most common AA and 1023 nicotinic acid analogs, asperphenidines Y1-Y2 (**4ab**), and 1024 asperphenidines W1-W2 (**12ab**), were further compared to 1025 those of the asperphenidines F1-F2 (**1ab**) (Figure 7B). 1026

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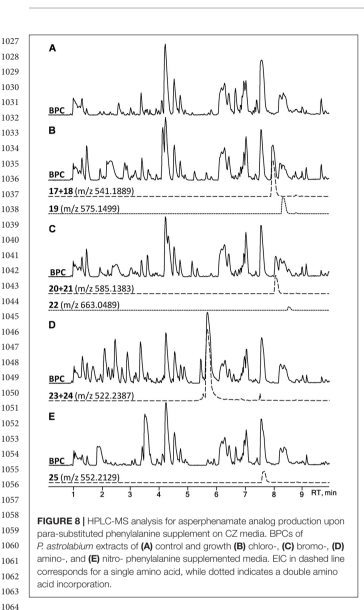
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1066 The relative production of asperphenidines followed the 1067 same pattern as observed in asperphenamate production: 1068 asperphenidines F1-F2 being the major nicotinic acid analogs, 1069 followed by asperhenidines Y1-Y2 and W1-W2. Moreover, 1070 production of all three upon growth on tryptophan supplemented 1071

media was drastically higher in comparison to control, with 1084 four- and five-fold increase in aspherphenidines F1-F2 and 1085 Y1-Y2, and 12-fold increase for asperphenidine W1-W2 1086 production. However, upon anthranilic acid supplement, a 1087 significant increase was observed only in asperphenidine F1-1088 F2 production. Additionally, production of all asperhenamates 1089 and asperhenidines was slightly lower to that of the control. 1090 Additional analysis of other AA supplement cases or additional 1091 inorganic nitrogen supplement experiments did not trigger 1092 similar nicotinic acid incorporation response. 1093

Para-Substituted Phenylalanines Are Incorporated in Either of N-Benzoyl Amino Acid Moieties

A set of four para-substituted, namely chloro-, bromo-, 1099 amino- and nitro-, phenylalanines were used to investigate 1100 the uptake of non-natural phenylalanines in asperphenamate 1101 1102 biosynthesis. The targeted MS analysis was performed by search 1103 of masses corresponding to a single or double AA exchange both asperphenamate and asperphenidine backbones 1104 in (Supplementary Table S1). This revealed, that incorporation 1105 of single para-substituted AA was successful in all four 1106 1107 supplement cases, however, a double para-substituted AA 1108 exchange was also observed in the halogenated phenylalanine 1109 supplement experiments (Figure 8). Moreover, small amounts of asperphenidine derivatives for single amino- and nitro-1110 prenylalanine incorporation analogs were also detected. 1111 Subsequent MS/MS analysis revealed, that halogenated para-1112 substituted phenylalanine can be incorporated at either or both 1113 1114 reduced or non-reduced parts of the molecule, resulting in three analogs each for chloro- (17-19) and bromo- (20-22) 1115 1116 asperphenamates (Figure 9 and Supplementary Figure S8). In 1117 case of amino- and nitro- phenylalanine exchange, the single AA incorporation was clearly preferred at the non-reduced part 1118 1119 of the molecule, with only trace amounts of substituted AA 1120 incorporation at the reduced part of the molecule detected (23-1121 25). As a result, asperphenidine analogs were detected only with AA exchange at the non-reduced part of the molecule, resulting 1122 in two analogs each for amino- (23ab) and nitro - (25ab) 1123 1124 asperphenidines (Figure 9 and Supplementary Figure S9). 1125 Additionally, non-reduced and reduced pathway intermediates, 1126 containing para-substituted phenylalanines, were also detected in each of the supplement study cases. 1127 1128

$$\begin{array}{c} 1129\\ 1130\\ 1074\\ 1075\\ 1076\\ 1077\\ 1078\\ 1088$$

1141 **TABLE 2** | Cytotoxic activities of 1, 1a, 4, 12, and 13.

Compound					
	A549	MCF7	A2058	HepG2	MiaPaca
1	>46	>46	1.1	28.5	>46
4	>46	23	24.8	21.6	>46
12	>46	>46	16.6	>46	>46
13	>46	>46	>46	>46	>46
1a	>46	>46	13.3	>46	13.3

A549, lung adenocarcinoma; MCF7, breast carcinoma; A2058, skin melanoma;
 HepG2, hepatocellular carcinoma; MiaPaca, pancreas carcinoma.

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¹¹⁵⁴ Asperphenamate Amino Acid Exchange ¹¹⁵⁵ Is Also Observed in Other Section Brevicompacta strains

1157 To compare the production of asperphenamate analogs among 1158 section Brevicompacta, the chemical profile of P. astrolabium 1159 was compared to three other section species, namely P. olsonii, 1160 P. bialowiezense, and P. brevicompactum (Houbraken et al., 1161 2020). Targeted MS search based on 5 readily observed 1162 asperphenamates in P. astrolabium (asperphenamates F, Y, W, 1163 L, and asperphenidine F1) revealed, that all the other strains 1164 were also able to exchange phenylalanine in the non-reduced 1165 AA moiety (Supplementary Figure S10). Additionally, all three 1166 strains were producing isoleucine analog in similar or higher 1167 amount in comparison to asperphenamate L and were able to 1168 produce compound 5, which was not detected in P. astrolabium. 1169 Moreover, P. brevicompactum revealed two peaks corresponding 1170 to the value of Asperphenamate W protonated adduct (m/z 1171 546.2385), both with the same fragmentation patterns, suggesting 1172 them being structural isomers. 1173

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Amino Acid Exchange Affects the Asperphenamate Analog Cytotoxicity

1177 Asperphenamates F, Y, W, and L (1, 4, 12 and 13) as 1178 well as aspephenidine F1 (1a) were tested for their cytotoxic 1179 activities against five cancer cell lines, i.e., lung carcinoma 1180 A549, breast adenocarcinoma MCF7, skin melanoma A2058, 1181 hepatocyte carcinoma HepG2 and pancreas carcinoma MiaPaca (Table 2). Asperphenamate Y and Asperphenidine F1 were the 1182 1183 only compounds exhibiting moderate cytotoxic activities against 1184 MCF7 and MiaPaca cell lines, respectively. Asperphenamates F and Y exhibited moderate activity against HepG2 cell lines, 1185 1186 whereas all but asperphenamate L showed activity toward A2058 1187 cell line, with asperphenamate F exhibiting the strongest activity. 1188 Asperphenamate L did not show activity against any of the 1189 cell lines at the tested concentrations. None of the compounds 1190 exhibited activity against A549 cell line at tested concentrations. 1191

1193 DISCUSSION

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Only a handful of asperphenamate analogs from fungal sources, including endophytic and parasitic fungi, have been isolated to date (Jia et al., 2009; Frisvad et al., 2013; Houbraken et al., 2020). In this study we have demonstrated yet another powerful 1198 application of a HRMS/MS guided discovery approach, for 1199 detection and structural characterization of novel peptide natural 1200 products via HRMS/MS fragmentation pattern analysis, which 1201 is in line with current HRMS/MS based peptide detections and 1202 characterization approaches (Mohimani et al., 2017; Jarmusch 1203 et al., 2020; Ricart et al., 2020). Moreover, we report that the 1204 choice of complex growth medium and/or a simple growth media 1205 supplement with selected building blocks, such as proteogenic 1206 and non-proteogenic AAs, P. astrolabium and related species can 1207 produce a series of novel asperphenamate analogs, which can be 1208 readily characterized by HRMS/MS. 1209

Incubation of P. astrolabium IBT 28865 on complex 1210 media revealed the strain being readily capable of exchanging 1211 phenylalanine for tyrosine, leucine or tryptophan in 1212 N-benzoylphenylalanine moiety, suggesting that the preferred 1213 AA substrate should be either aromatic or aliphatic. A subsequent 1214 proteogenic AA supplement study supported the hypothesis, with 1215 all but one AA incorporated being either aromatic or aliphatic. 1216 It can be speculated, that the incorporation of AA is dependent 1217 on the side chain size and conformational similarity to that of 1218 phenylalanine, since tyrosine, leucine and methionine have the 1219 highest production rate in the AA supplement studies, whereas 1220 the smaller alanine and isoleucine analogs are produced at the 1221 lower rate. Lastly, histidine was the only non-hydrophobic AA to 1222 be incorporated into the asperphenamate backbone, something 1223 that can be attributed to its similarity to the other aromatic AAs, 1224 thereby likely interacting via similar $\pi - \pi$ interactions. 1225

Although the relative production upon tyrosine supplement 1226 decreased in comparison to the control, it might be attributed 1227 to tyrosine being preferentially taken up by other pathways, 1228 such as di- or tetra-peptide biosynthesis. Nevertheless, upon 1229 growth on non-supplemented media, it was observed that 1230 tyrosine incorporation in general was preferred over any other 1231 AA incorporation. Subsequent supplement study with other 1232 synthetic para-substituted phenylalanines confirmed previous 1233 observation, with all four selected substrates being incorporated 1234 into the asperphenamate backbone irrespective of the size of the 1235 para-moiety. In comparison to proteogenic AAs being mainly 1236 incorporated into the non-reduced part of asperphenamate 1237 backbone, para-substituted AAs were readily incorporated into 1238 either or both the reduced and the non-reduced part of 1239 the molecule. Additionally, no other pathway intermediates 1240 rather than N-benzoylphenylalanine, N-benzoylphenylalaninol 1241 and respective para-substituted phenylalanine analogs were 1242 observed. This suggests, that only intermediates with the highest 1243 similarity to phenylalanine intermediates can be recognized and 1244 released from the asperphenamate biosynthetic machinery. 1245

The production of nicotinic acid containing analogs, 1246 asperphenidines, was strongly correlated with production 1247 of the corresponding asperphenamates upon growth on 1248 complex media. However, upon proteogenic AA supplement 1249 study, asperphenidines F1-F2 and Y1-Y2 were produced 1250 in higher amounts when tryptophan was supplemented, 1251 and in case of asperphinidines F1-F2 also in anthranilic 1252 acid supplemented media. This suggest, that biosynthesis 1253 of asperphenamate directly intercepts primary metabolism, 1254

since both anthranilic and nicotinic acids are intermediates of 1255 tryptophan catabolism in nicotinamide adenine dinucleotide 1256 (NAD) biosynthesis (Foster and Moat, 1980). This hypothesis 1257 can be further substantiated by the fact that no other organic 1258 and inorganic nitrogen source resulted in similar nicotinic 1259 acid production and incorporation response. However, no 1260 incorporation of directly supplemented benzoic acid derivatives, 1261 namely 4-hydroxybenzoic and anthranilic acids, was observed. 1262 This indicated, that incorporation of nicotinic acid is most 1263 likely driven by the availability of the substrate, rather than 1264 promiscuity of either of NRPS domains, since there are only a 1265 minute structural difference among benzoic and nicotinic acids, 1266 1267 as well as no clear discrepancies among preference of nicotinic 1268 acid over benzoic acid by either of the NRPS domains.

In general natural product biosynthesis of small peptides 1269 1270 involves a very strict uptake of AAs controlled by the NRPS adenylation domains leading to a conserved sequence of AAs 1271 present in the final product (Fischbach and Walsh, 2006). 1272 However, certain cyanobacteria have been reported to possess 1273 adenylation domains capable of activation of two or more 1274 chemically distant AA (Kaljunen et al., 2015; Meyer et al., 2016). 1275 In contrast, our study revealed an unusually high flexibility, 1276 rather than specificity of fungal adenylation domain toward the 1277 uptake of structurally related natural AAs, as well as synthetic 1278 para-substituted phenylalanine analogs. Such unusual NRPS 1279 flexibility is rather uncommon, with only one recent similar case 1280 observed in filamentous fungi (Hai et al., 2020) . Recently, the 1281 Tang lab demonstrated that the hybrid NRPS-NRPKS involved in 1282 biosynthesis of α-pyrones in Aspergillus niger is also promiscuous 1283 toward the uptake of tyrosine, leucine and a number of para-1284 substituted phenylalanines with small substitution groups (Hai 1285 1286 et al., 2020). However, with the higher variety of natural AA being 1287 tolerated in asperphenamate biosynthesis, our results altogether suggest an even more relaxed substrate specificity in comparison 1288 to that of α-pyrone biosynthesis. 1289

Interestingly, three related species other from 1290 1291 section Brevicompacta, P. olsonii, P. bialowiezense, and P. brevicompactum, were also found to be producers of the 1292 same analogs as observed in P. astrolabium when grown 1293 on complex media. In addition, detection of several other 1294 asperphenamates, such as a 4-hydroxybenzoic acid containing 1295 analog (5), indicates an even more relaxed substrate specificity 1296 in comparison to that of P. astrolabium. Nonetheless, it might 1297 be speculated that the aforementioned analogs are not observed 1298 in P. astrolabium due to a lower growth rate in comparison 1299 to the other three strains (Serra and Peterson, 2007; Perrone 1300 et al., 2015). Moreover, the presence of two Asperphenamate 1301 W stereoisomers in P. brevicompactum suggest the presence 1302 1303 of a biosynthetically unrelated enzymatic activity responsible 1304 for epimerization of tryptophan. Similar enzymatic activity was previously characterized in a single-module NRPS responsible 1305 for specific stereoconversion of L-tryptophan to D-tryptophan in 1306 A. niger (Hai et al., 2019). 1307

Asperphenamates F, Y, L, and W, as well as asperphenidine F1 were tested against five cancer cell lines. Although asperphenamate L did not exhibit activity against any of the cell lines, the four other compounds revealed moderate activities against breast, skin, liver or pancreas cell lines. In particular, 1312 asperphenamate Y was the only active compound against the 1313 breast cell line, suggesting that the presence of tyrosine at the 1314 non-reduced AA moiety might be essential for the observed 1315 activity. Therefore, further investigations of asperphenamates 1316 harboring a para-substituted phenylalanine could be of interest 1317 for future cytotoxicity studies. Moreover, asperphenidine F1, was 1318 the only active candidate against the pancreas cell line, suggesting 1319 the nicotinic acid analogs being more active than the benzoic acid 1320 analogs. Although our cytotoxicity results for asperphenamates F 1321 and Y, as well as asperphenidine F1 are comparable to previously 1322 published data, the natural analogs, none of them show improved 1323 bioactivity compared to synthetic asperphenamate derivatives (Li 1324 et al., 2012; Yuan et al., 2012, 2018, 2019, 2020; Liu et al., 2016, 1325 2018). 1326

In conclusion, HRMS/MS based analysis and the use 1327 of a targeted media supplement approach demonstrated 1328 an extraordinary relaxed substrate specificity in the double 1329 NRPS system responsible for asperphenamate production. 1330 The proteogenic and non-proteogenic para-substituted 1331 L-phenylalanine analog supplements led to biosynthesis of 1332 22 new analogs, all of which could readily be characterized by 1333 HRMS/MS. Here we proposed a standardized naming system for 1334 asperphenamate and asperphenidine analogs denoting specific 1335 amino acid incorporation. This strategy illustrates the potential 1336 for future combinatorial biosynthesis of asperphenamate and 1337 similar small NRPS products. 1338

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DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/**Supplementary Materials**, further inquiries can be directed to the corresponding author/s.

AUTHOR CONTRIBUTIONS

KS and TL designed the experiments. KS and JF performed HRMS/MS library search. KS, XW, and FN performed purification and structure elucidation, with the assistance by CG. KS performed asperphenamate design study and subsequent HRMS/MS the data analysis compounds, assisted by TI. OG, FV, CR, and TM designed and performed the bioassay. KS wrote the manuscript with contribution from all authors.

FUNDING

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2020. 618730/full#supplementary-material

uai for this article ersin.org/article y-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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