

1 *Review*

## 2 **A Decade of Antifungal Leads from Natural Products**

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22 **Abstract:** In this review, we discuss novel natural products discovered within the last decade that  
23 are reported to have antifungal activity against pathogenic species. Nearly a hundred natural  
24 products were identified that originate from bacteria, alga, fungi, sponges and plants. Fungi were  
25 the most prolific source of antifungal compounds discovered during the period of review. The  
26 structural diversity of these antifungal leads encompasses all the major classes of natural products  
27 including polyketides, shikimate metabolites, terpenoids, alkaloids and peptides.

28 **Keywords:** fungal pathogens; antifungal agents; natural products

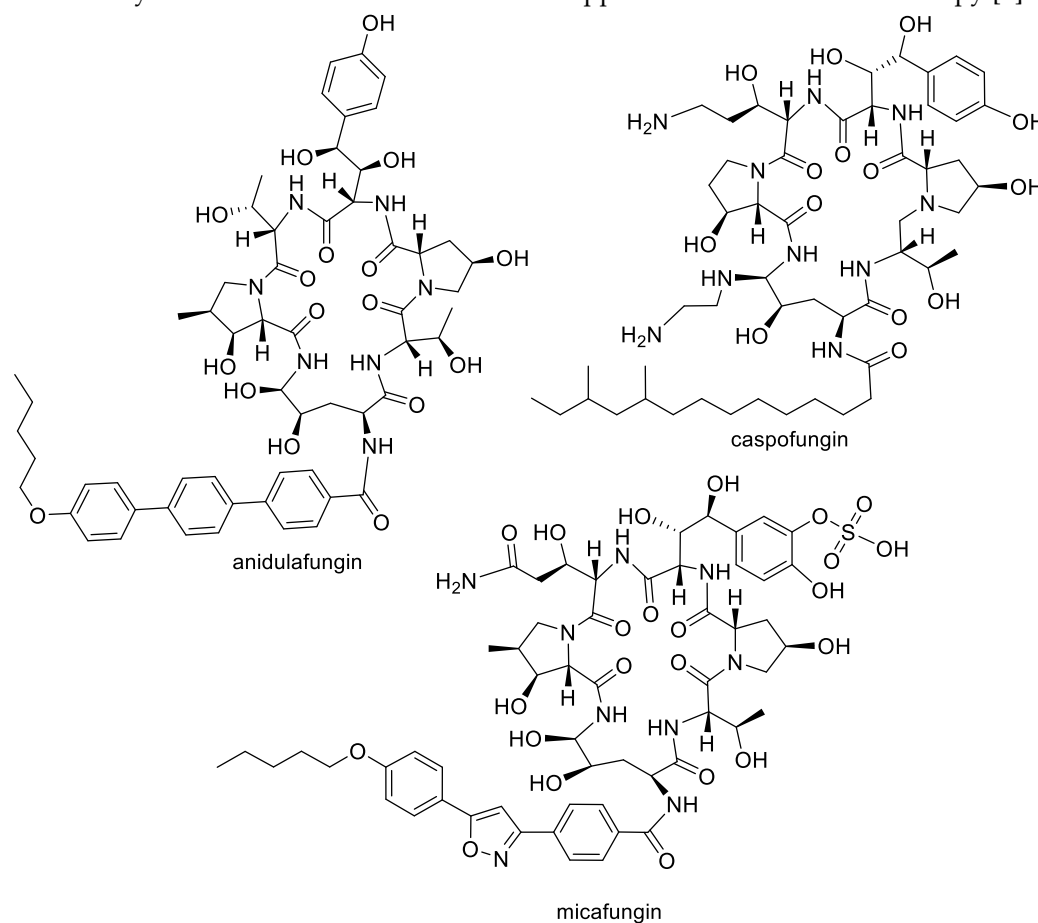
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### 30 **1. Introduction**

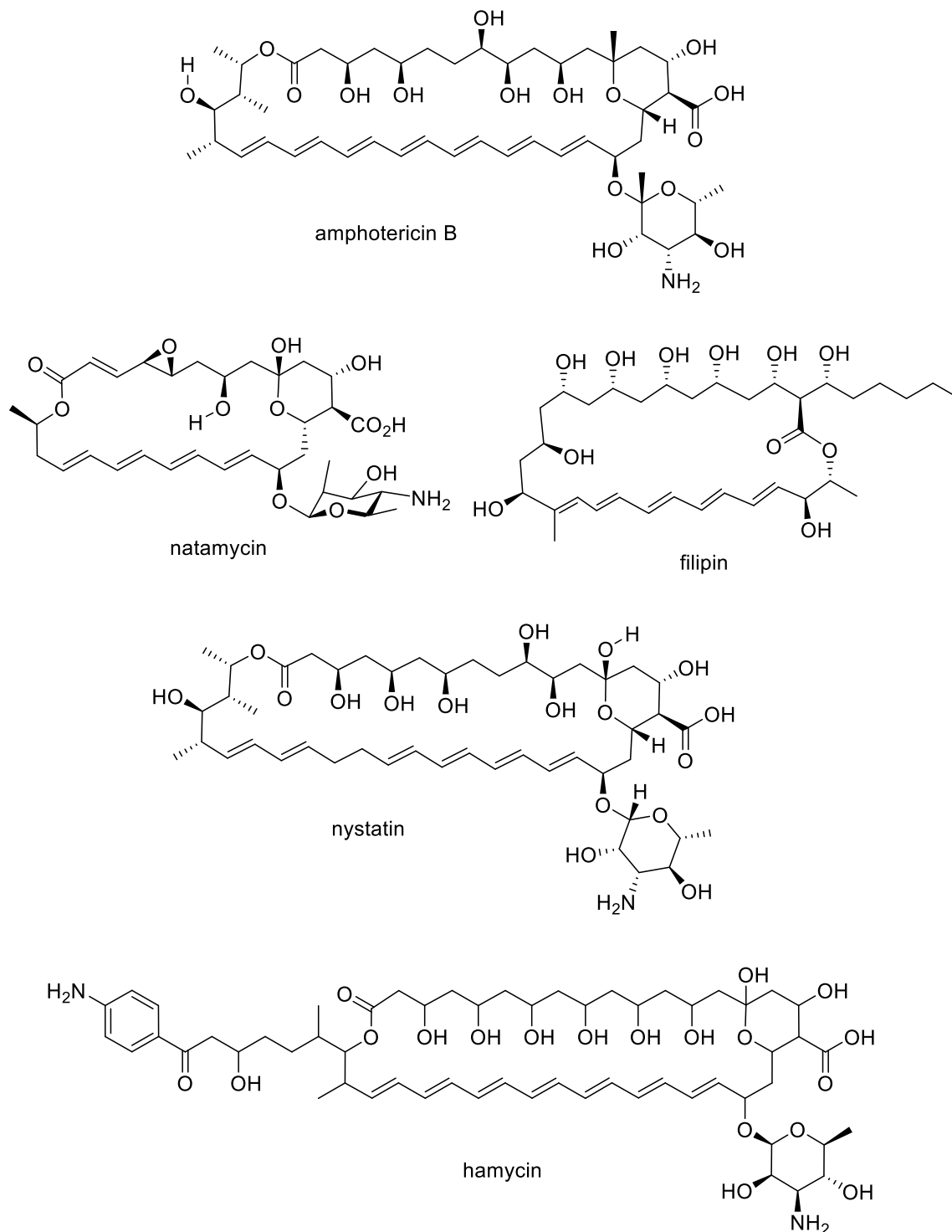
31 The global increase in antimicrobial resistance among pathogenic bacteria, viruses, fungi and  
32 parasites is a serious concern for human healthcare. In the case of fungi, more than one billion  
33 individuals worldwide are affected by fungal infections and the associated mortality, over 1.5 million  
34 deaths each year, is equivalent to that caused by tuberculosis and more than triple that of malaria [1].  
35 Although relatively rare in healthy individuals, the incidence of superficial and invasive fungal  
36 infections has dramatically risen in recent years. This is due to a growing 'at-risk' population with  
37 impairments in their immune system, breaches in physical barriers to fungal entry or an altered  
38 microbiome. Skin mycoses are predominantly caused by *Trichophyton*, *Microsporum* and  
39 *Epidermophyton* genera while *Candida*, *Cryptococcus*, *Aspergillus* and *Pneumocystis* genera, and  
40 *Mucorales* are the most common invasive fungal pathogens [2]. Meanwhile, emerging pathogenic  
41 fungi that are either new species such as the recently described *Candida auris* [3] or well-known  
42 species spreading in their ecological distribution represent additional threats to human health.

43 The growing challenges posed by fungal diseases are further heightened as antifungal treatment  
44 is mainly limited to the azoles and echinocandins. The azoles are the most widely used antifungals

45 and are synthetic compounds that reversibly inhibit cytochrome P450-dependent lanosterol or  
46 eburicol 14 $\alpha$ -demethylase with moderate specificity for the fungal enzyme over the human  
47 counterpart [4]. Nevertheless, they suffer from off-target toxicity as well as issues with fungistatic  
48 rather than fungicidal activity in yeast that promotes the development of resistance. The  
49 echinocandins are fungal lipopeptide natural products (Figure 1) that are non-competitive inhibitors  
50 of 1,3- $\beta$ -glucan synthase, an enzyme involved in fungal cell wall biosynthesis. While the natural  
51 products are not optimal in terms of pharmacokinetics, three semi-synthetic derivatives are approved  
52 for clinical use: anidulafungin prepared from echinocandin B, caspofungin prepared from  
53 pneumocandin B<sub>0</sub> and micafungin prepared from FR901379 [5]. Although the selectivity of the  
54 echinocandin target for fungi provides a good safety profile, these compounds are large peptides,  
55 requiring intravenous administration. In addition to the azoles and echinocandins, the polyenes and  
56 pyrimidines are two other classes approved for antifungal therapy. The natural product polyenes  
57 (Figure 2) are macrolides isolated from various *Streptomyces* strains. The prototypical amphotericin B  
58 has been in clinical use for the treatment of systemic fungal infections since the 1950s and is still an  
59 important option in critical cases. Several additional polyenes -nystatin, natamycin, hamycin and  
60 filipin- have received regulatory approval. As a class, the polyenes have significant nephrotoxicity  
61 due to their relatively nonselective mechanisms of ergosterol binding and pore formation within the  
62 cell membrane [6,7]. Finally, synthetic pyrimidine antimetabolites such as flucytosine interfere with  
63 nucleic acid biosynthesis but resistance restricts their application to combination therapy [8].



64 **Figure 1.** Semi-synthetic derivatives of the echinocandin family of natural products approved for  
65 antifungal therapy.  
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**Figure 2.** Polyene natural products approved for antifungal therapy.

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In summary, the current drugs have numerous limitations including toxicity, drug-drug interactions, poor pharmacokinetics, narrow spectrum of activity and fungistatic versus fungicidal action. These inherent liabilities are exacerbated in immunocompromised patients since their immune system cannot effectively assist in the eradication of the infection, thus requiring complex and prolonged treatment regimens [9]. A further alarming trend is the rising incidence of fungal

75 clinical isolates that are resistant to the currently used antifungals [10,11]. The scale of the problem is  
76 highlighted by the fact that the newest class of approved antifungals, the echinocandins, were  
77 actually discovered fifty years ago. The American Food and Drug Administration (FDA) has  
78 recognized the need for new antifungals by placing *Candida* and *Aspergillus* on their list of qualifying  
79 pathogens [12]. Therapies directed against these species will benefit from incentives including an  
80 additional five-year marketing exclusivity besides eligibility for designation as a fast-track drug.

## 81 2. A pipeline of antifungal natural product leads

82 While antifungal agents with novel mechanisms of action are in various stages of clinical  
83 development, their number is relatively small compared to other therapeutic indications [13]. A  
84 pipeline of additional preclinical leads is clearly needed, and natural product screening is an  
85 important contributor in this regard. One unique feature of natural products is their high structural  
86 diversity, sampling areas of chemical space that are difficult to access through purely synthetic  
87 compounds [14,15]. Natural products are also well validated to possess biological activity, with many  
88 examples approved as therapeutic agents either in their native form or as semi-synthetic derivatives  
89 [16]. For this review, we searched *Natural Product Updates* for publications that reported novel natural  
90 products with antifungal activity within the last decade January 2010 - November 2019. From the  
91 publications, we selected novel natural products that were active against human pathogenic fungi  
92 with a MIC < 10 µg/mL or IC<sub>50</sub> < 10 µM. In the discussion, we include any information on additional  
93 biological activity observed or mechanistic studies on the mode of action. The compounds are  
94 classified below according to the type of producing organism.

### 95 2.1 Natural product antifungal leads from bacteria and algae

96 Actinomycetes are the most prolific source of bacterial natural products, and this remains the  
97 case for recently discovered antifungal leads (Figures 3-6, 1-29). In addition, there were three  
98 examples isolated from non-actinomycete species (Figure 7, 30-35) and two from algae (Figure 8, 36-  
99 37). A strain of *Streptomyces albolongus* YIM 101047 isolated from elephant dung produced a number  
100 of bafilomycins in laboratory fermentation. The new example 21-deoxybafilomycin A1 (**1**) and the  
101 sesquiterpene (1β,4β,4aβ,8aα)-4,8a-dimethyloctahydronaphthalene-1,4a(2H)-diol (**2**) displayed  
102 antifungal activity against *Candida parapsilosis* with a MIC of 3.2 µg/mL while being inactive against  
103 other species [17]. Genome sequencing of the strain suggested the presence of forty-six putative  
104 biosynthetic gene clusters [18]. In the course of biosynthetic labelling experiments, it was discovered  
105 that supplementation by acetate produced new metabolites in a *Streptomyces hyaluromycini* MB-PO13  
106 strain. Among these, rubromycin CA1 (**3**) was active against Gram-positive bacteria and *Candida*  
107 *albicans* NBRC 1594 with a MIC of 6.3 µg/mL whereas an analogue with an additional alcohol was  
108 inactive [19]. A strain of *Actinoalloteichus* isolated from marine sediment was the source of  
109 neomaclafungins A-I (**4-12**), a series of macrolides of the oligomycin family of antibiotics. The  
110 neomaclafungins were active against *Trichophyton mentagrophytes* with MIC values between 1 and 3  
111 µg/mL, compared to 10 µg/mL for oligomycin A [20].

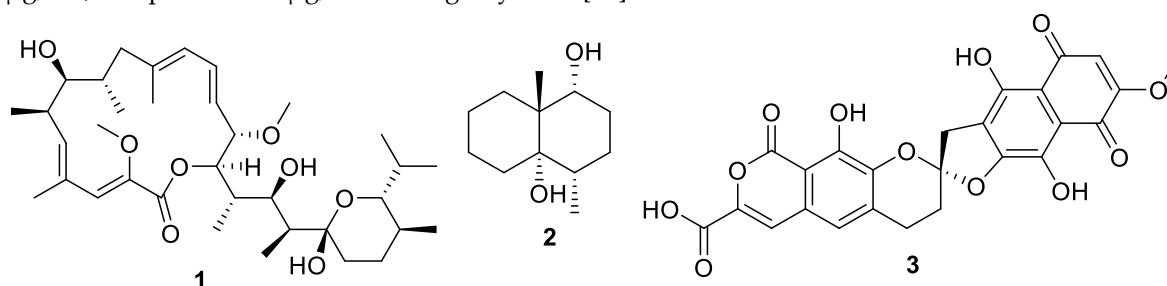


Figure 3. Structures of natural products 1-3

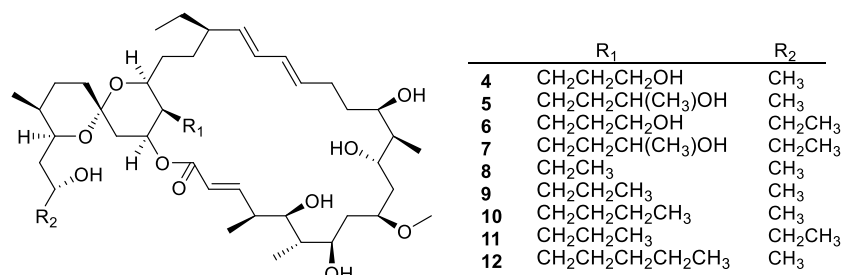
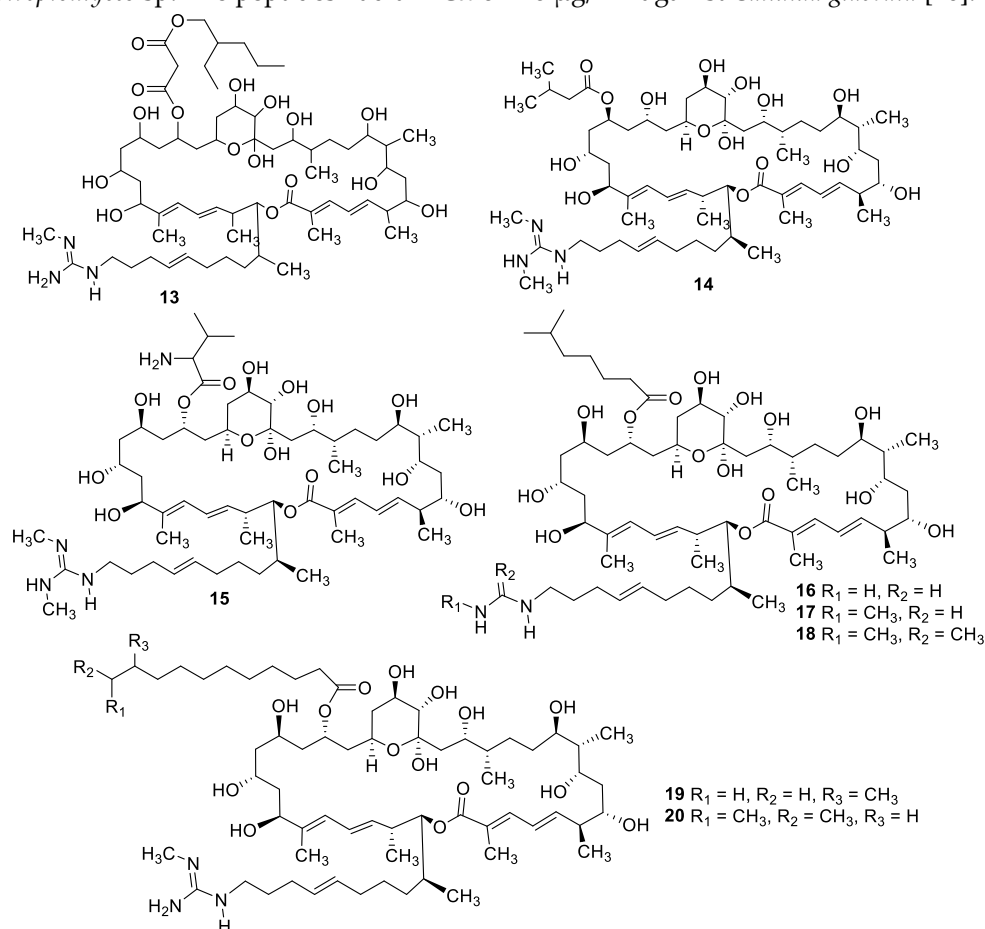
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Figure 4. Structures of neomaclafungins A-I 4-12

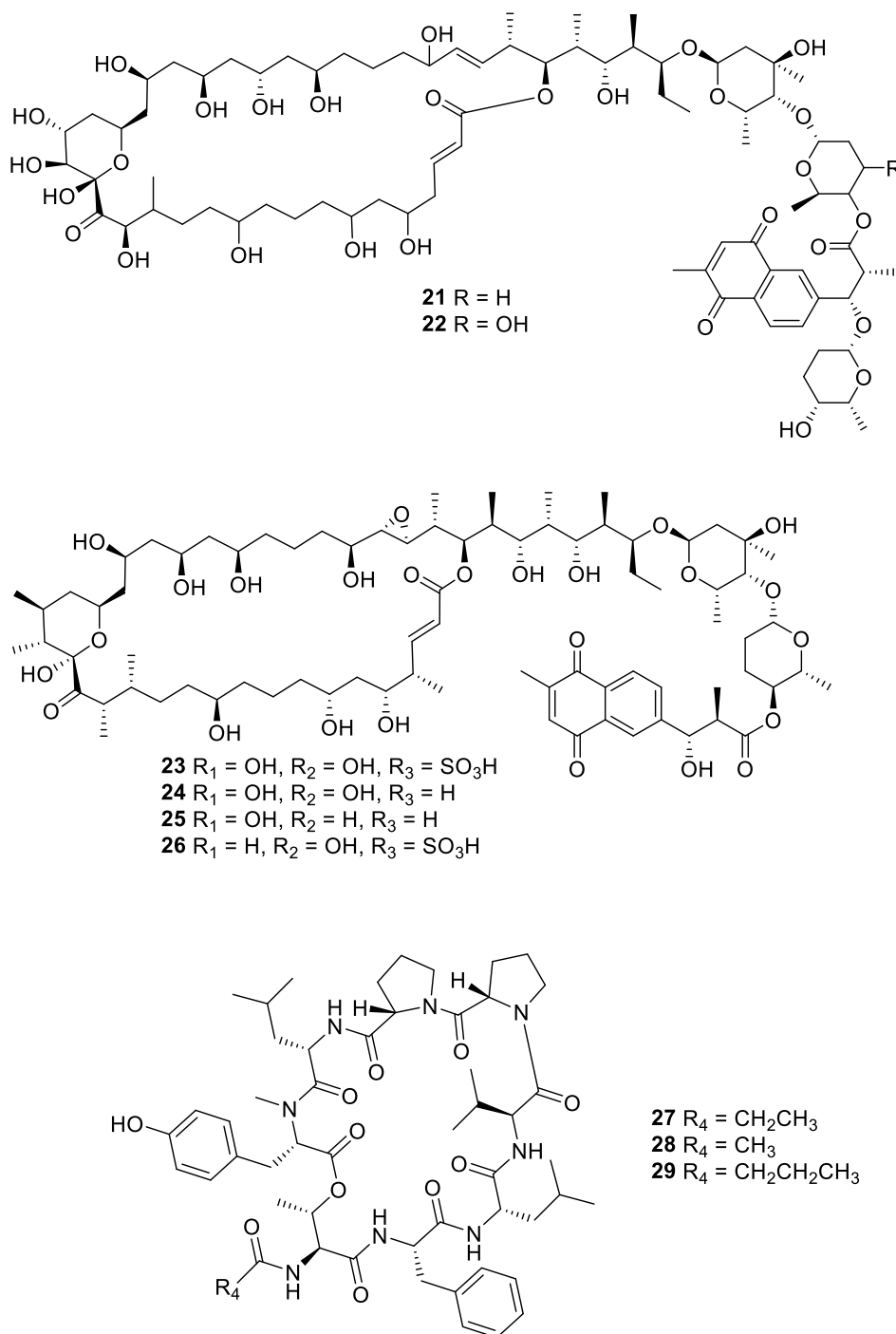
116 Fermentation of a *Streptomyces* sp. isolated from mangrove rhizosphere soil led to the isolation  
 117 of a series of azalomycin F natural products (13-20) with MIC values of 1.6-6.3 µg/mL against *Candida*  
 118 *albicans* as well as having antibacterial and cytotoxic activity [21,22]. Astolides A (21) and B (22) are  
 119 polyol macrolides isolated from *Streptomyces hygrosopicus* collected from alkaline soil [23]. The  
 120 compounds have MICs of 1-2 µg/mL against *Candida albicans*, *Candida tropicalis* and *Aspergillus niger*.  
 121 Related macrolides caniferolides A-D (23-26) were isolated from the marine-derived *Streptomyces*  
 122 *caniferus* CA-271066 [24]. Like the astolides, the caniferolides displayed potent antifungal activity  
 123 with MICs of 0.5-2 µg/mL against *Candida albicans* and 2-8 µg/mL against *Aspergillus fumigatus*, as  
 124 well as similar levels of cytotoxicity against human tumor cell lines. Caniferolide A was also shown  
 125 to have in vitro activity against targets relevant to Alzheimer's disease [25]. Enduspeptides A-C (27-  
 126 29) are depsipeptides that differ in the acyl chain attached to the threonine residue and were isolated  
 127 from a *Streptomyces* sp. The peptides had an IC<sub>50</sub> of 2-8 µg/mL against *Candida glabrata* [26].



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Figure 5. Structures of azalomycin F macrolides 13-20



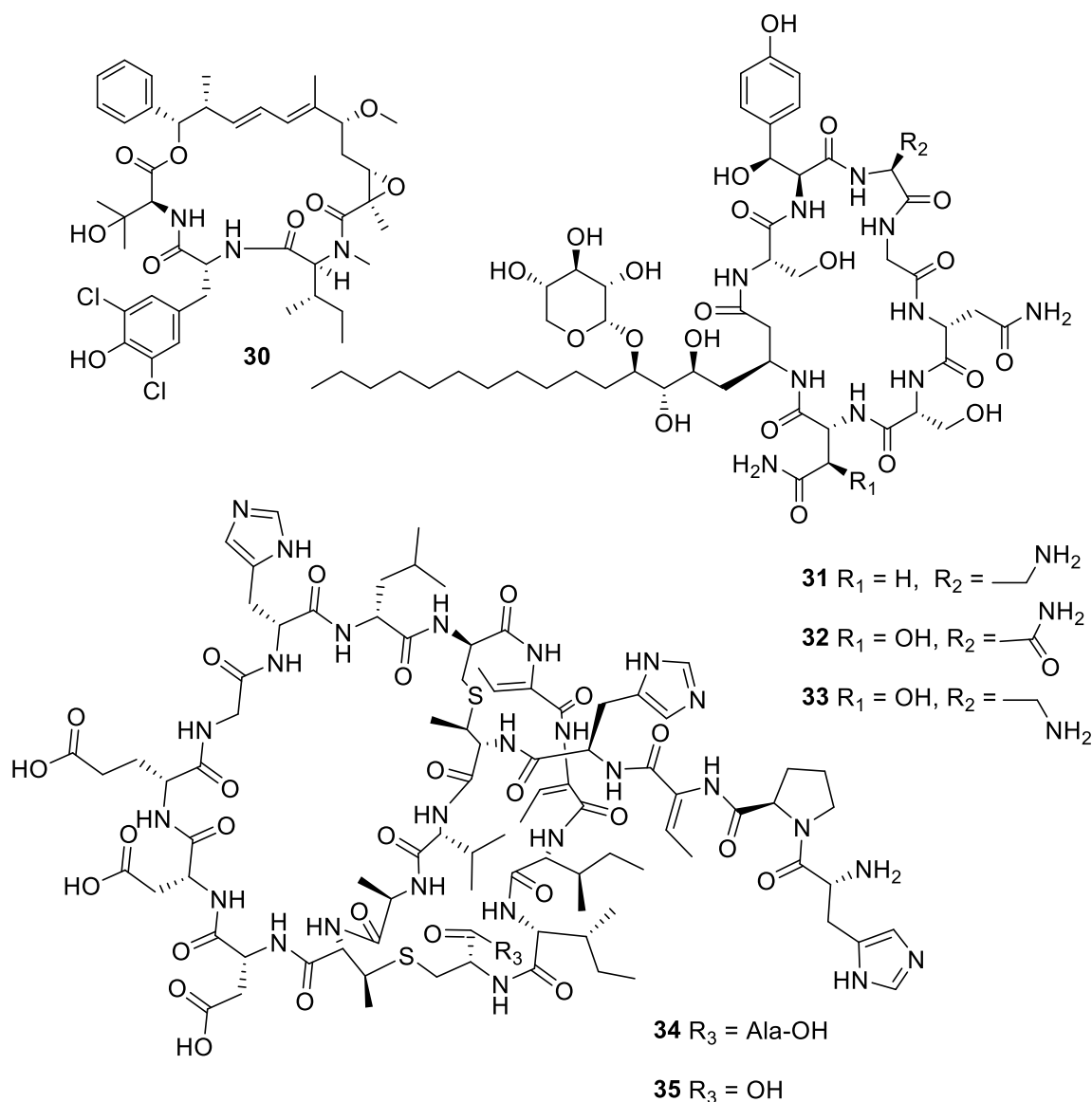
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**Figure 6.** Structures of natural products 21-29

132 Within the period under review, three antifungal leads were isolated from non-actinomycete  
 133 bacterial strains. Fermentation of a myxobacterial *Nannocystis* sp. led to the isolation of nannocystin  
 134 A (**30**) with a novel macrocyclic scaffold. While the compound inhibited *Candida albicans* with a MIC<sub>50</sub>  
 135 of 73 nM, it also inhibited human cancer cell lines at a nanomolar level [27]. The mechanism of action  
 136 involves binding to the eukaryotic translation elongation factor 1 $\alpha$  and SAR has been established  
 137 through the total synthesis of analogues [28]. The burkholdines are lipopeptide antifungal agents  
 138 previously isolated from *Burkholderia ambifaria* 2.2N, with three new examples Bk-1119, Bk-1213, and

139 Bk-1215 (**31-33**) displaying potent activity against *Candida albicans* and *Aspergillus niger* [29]. Among  
 140 the burkholdines, Bk-1119 was the most active against *Aspergillus niger* with a MIC of 0.1  $\mu\text{g/mL}$  and  
 141 also had the best antifungal:hemolytic ratio. Additional analogues were prepared by total synthesis  
 142 [30]. The Gram-negative bacteria *Chitinophaga pinensis* DSM 28390 produces the novel lantibiotics  
 143 pinensins A and B (**34, 35**). Although lantibiotics are typically antibacterial, the pinensins were only  
 144 weakly so while having MICs of 2-4  $\mu\text{g/mL}$  against yeasts and filamentous fungi [31].



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**Figure 7.** Structures of natural products **30-35**

147 The marine alga *Laurencia* is a prolific producer of secondary metabolites. The sesquiterpene  
 148 eudesma-4(15),7-diene-5,11-diol (**36**) isolated from a Red Sea sample of *Laurencia obtusa* was  
 149 antifungal with MIC values of 2-7  $\mu\text{M}$  against *Candida* and *Aspergillus* strains [32]. The prenylated  
 150 xylene caulerprenylol B (**37**) was isolated from the green alga *Caulerpa racemosa* and had MIC<sub>80</sub> values  
 151 of 4  $\mu\text{g/mL}$  against *Candida glabrata* and *Cryptococcus neoformans* while being inactive against  
 152 *Aspergillus fumigatus* [33].

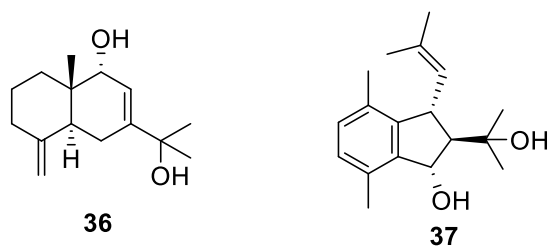


Figure 8. Structures of natural products 36 and 37

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## 156 2.2 Natural product antifungal leads from sponges

157 Marine sponges are an important source of novel natural products, and more than ten examples  
158 with antifungal activity were described in this period (Figures 9 and 10, 38-55). Extracts from the  
159 symbiotic two-sponge association *Plakortis halichondroides*-*Xestospongia deweerdtiae* yielded a number  
160 of peroxide natural products, of which plakinic acids I, J, K and L (38-41) were potent against *Candida*  
161 and *Cryptococcus* species with MIC  $\leq$  0.5  $\mu\text{g/mL}$  [34]. Plakinic acid M (42) was active against  
162 *Cryptococcus gattii*, *Cryptococcus grubii* and *Candida krusei* with MIC<sub>90</sub> values of 2.4-3.4  $\mu\text{g/mL}$  but less  
163 active against *Candida albicans* [33]. Extraction from the South China Sea sponge *Hippospongia lachne*  
164 was the source for hippolachnin A (43), a polyketide with an unprecedented scaffold [36]. The  
165 compound was potently antifungal with a MIC of 0.4  $\mu\text{g/mL}$  against *Cryptococcus neoformans*,  
166 *Trichophyton rubrum* and *Microsporium gypseum*. However, the natural product and analogues  
167 obtained by total synthesis were inactive, suggesting the initial report was in error [37]. Bioassay-  
168 guided fractionation of the same extract led to isolation of a racemic sesterterpene hippolide J (44)  
169 [38]. The natural product was resolved into its two enantiomers, and both were highly potent  
170 antifungals with MIC<sub>50</sub> of 0.13-0.25  $\mu\text{g/mL}$  against *Candida* and *Trichophyton* while weakly cytotoxic  
171 to the human embryonic kidney HEK293 cell line.

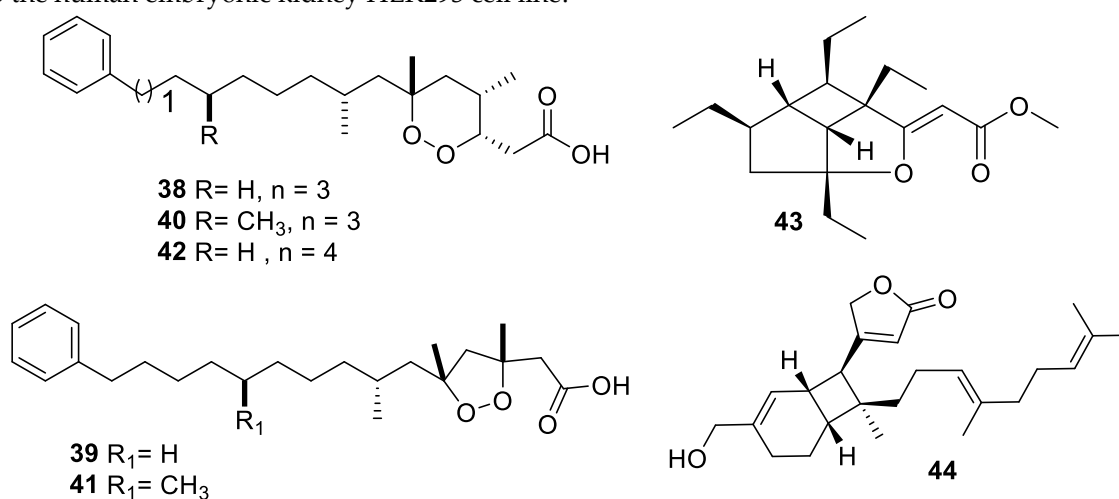


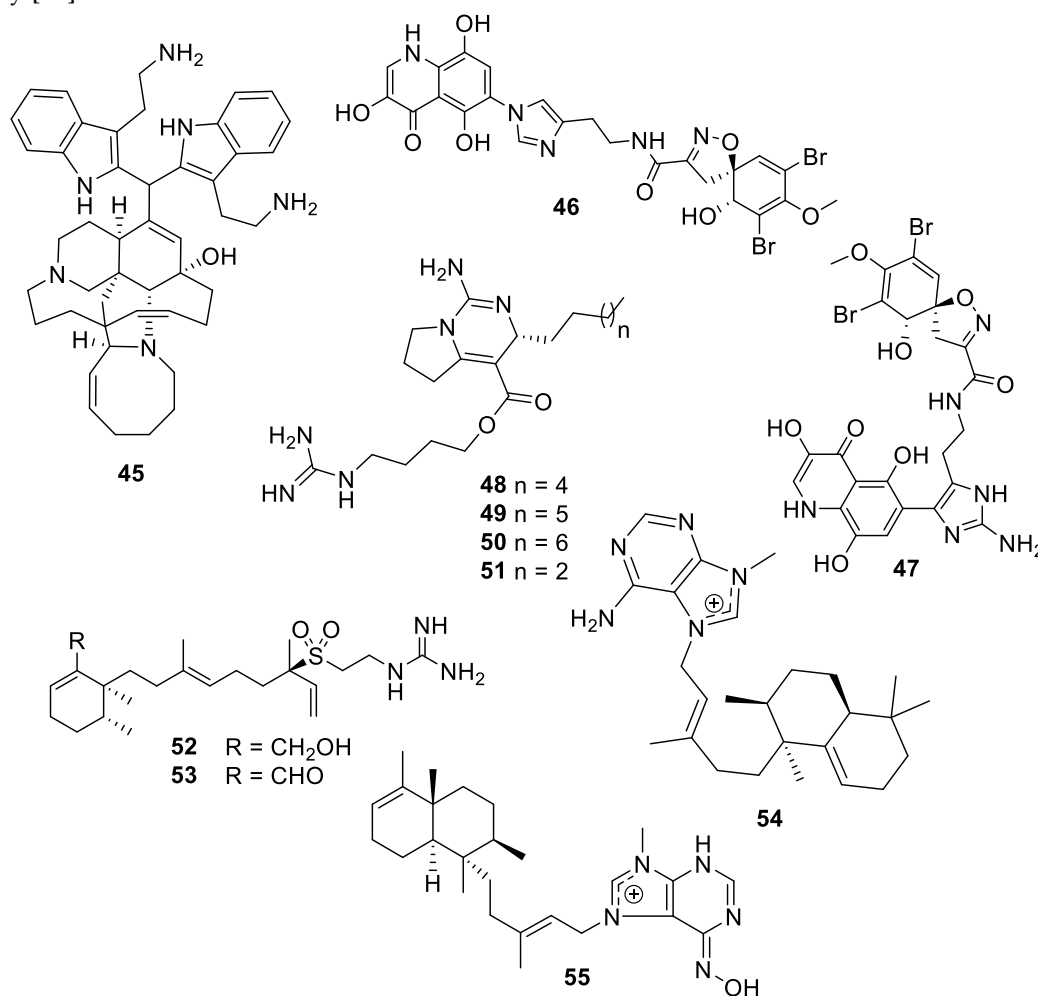
Figure 9. Structures of natural products 38-44

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174 A new member of the manzamine alkaloids, zamamidine D (45), was isolated from an Okinawan  
175 marine sponge *Amphimedon* sp. Zamamidine D had an IC<sub>50</sub> of 2  $\mu\text{g/mL}$  against *Cryptococcus*  
176 *neoformans* but was weakly active against other fungal and bacterial strains tested [39]. From another  
177 Okinawan marine sponge *Pseudoceratina* sp., ceratinadin A and B (46, 47) were isolated with MIC  
178 values of 4 and 8  $\mu\text{g/mL}$  respectively against *Cryptococcus neoformans* and 2 and 4  $\mu\text{g/mL}$  respectively  
179 against *Candida albicans* [40]. From an extract of the sponge *Pseudaxinella reticulata*, several crambescine  
180 guanidine containing alkaloids were isolated. Crambescine A2 392 and 406 (48, 49) inhibited



181 *Cryptococcus neoformans* with a MIC<sub>50</sub> of 1.2 and 0.9 µg/mL respectively while being relatively inactive  
 182 against *Candida albicans* [41]. The enantiomers of two known crambescins, crambescin A2 420 (**50**) and  
 183 Sch 575948 (**51**) were also isolated with a MIC<sub>50</sub> of 1.1 and 2.5 µg/mL respectively against *Cryptococcus*  
 184 *neoformans*. Among metabolites isolated from the marine sponge *Agelas*, two new diterpene alkaloids  
 185 from *Agelas citrina*, agelasidine E and F (**52**, **53**), were reported to have MIC values of 8 and 4 µg/mL  
 186 respectively against *Candida albicans* [42]. Isoagelasine C (**54**), isolated from *Agelas nakamura*, had a  
 187 MIC value of 4.7 µg/mL against *Candida albicans* [43]. Ageloxime B (**55**), isolated from *Agelas*  
 188 *mauritiana*, had an IC<sub>50</sub> value of 5.0 µg/mL against *Cryptococcus neoformans* as well as antibacterial  
 189 activity [44].

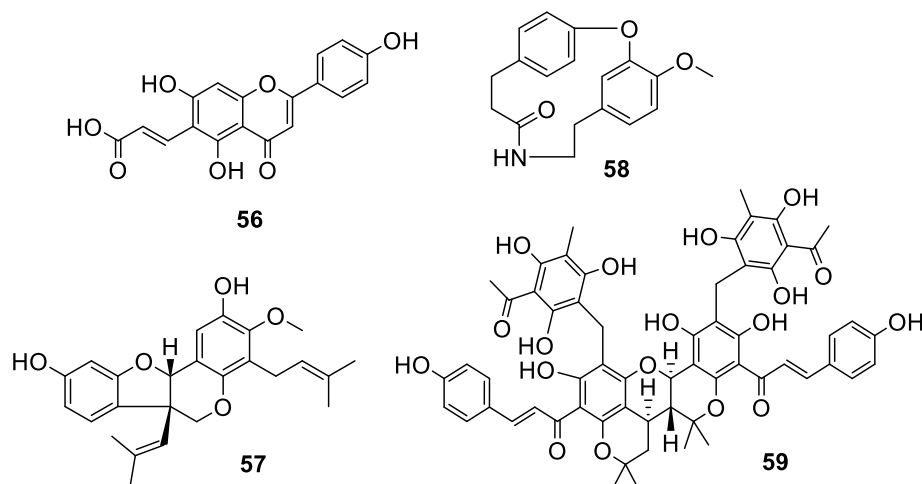


190  
 191 **Figure 10.** Structures of natural products 45-55

192 **2.3 Natural product antifungal leads from plants**

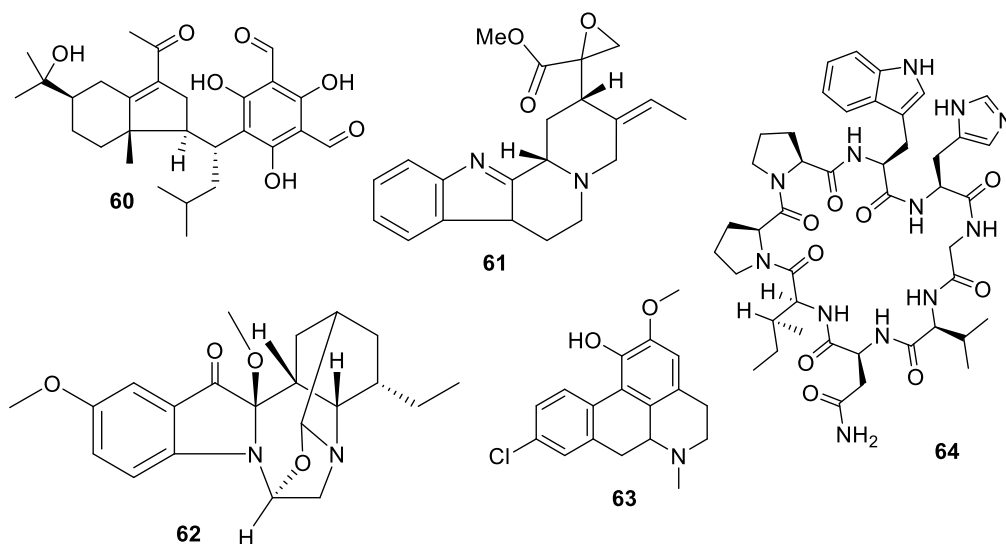
193 Plants accounted for nearly ten antifungal leads within the last decade (Figures 11 and 12, **56-**  
 194 **64**). The flavonoid (*E*)-6-(2-carboxyethenyl)apigenin (**56**) was isolated from an extract of *Mimosa*  
 195 *caesalpiniiifolia* Benth., a Brazilian medicinal plant commonly known as “sabiá” or “sansão-do-campo”  
 196 [45]. The compound inhibits *Candida krusei* with an IC<sub>50</sub> of 44 nM, although it was inactive against  
 197 *Candida glabrata*. The isoflavonoid vatacarpan (**57**) with a MIC of 1 µg/mL against *Candida albicans*  
 198 was isolated by bioassay-guided fractionation from the roots of *Vatairea macrocarpa* (Benth.) Ducke  
 199 [46]. The biaryl ether laevicarpin (**58**) was isolated from leaves of *Piper laevicarpu*, known as “falsa-  
 200 pimenteira” in Brazil [47]. Interestingly, the compound was previously prepared synthetically prior  
 201 to this isolation. Laevicarpin had an IC<sub>50</sub> of 7.9 µM against *Cryptococcus gattii*, in addition to an IC<sub>50</sub> of

202 50  $\mu\text{M}$  against the trypomastigote form of *Trypanosoma cruzi*. The dimeric chalcone kamalachalcone  
 203 E (59) was isolated from the red dye extracted from whole uncrushed fruits of *Mallotus philippinensis*  
 204 [48]. The chalcone exhibited an  $\text{IC}_{50}$  of 4-8  $\mu\text{g}/\text{mL}$  against two strains of *Cryptococcus neoformans*.



205  
 206 **Figure 11.** Structures of natural products 56-59

207 Investigation of the juvenile leaves of *Eucalyptus maideni* F. Muell led to the discovery of a  
 208 number of phloroglucinol derivatives, among which eucalmaidial A (60) showed antifungal activity  
 209 against *Candida glabrata* with an  $\text{IC}_{50}$  of 0.8  $\mu\text{g}/\text{mL}$  [49]. A monoterpene indole alkaloid, 16,17-  
 210 epoxyisositirikine (61), isolated from the evergreen shrub *Rhazya stricta* Decne. had an  $\text{IC}_{50}$  of 6.3  
 211  $\mu\text{g}/\text{mL}$  against *Candida glabrata* but was less active against other *Candida* species tested [50]. Erchinine  
 212 B (62), a monoterpene indole alkaloid with an unusual 1,4-diazepine ring embedded was isolated  
 213 from roots of *Ervatamia chinensis* and had a MIC of 6.3  $\mu\text{g}/\text{mL}$  against *Trichophyton rubrum*, with a  
 214 lower MIC of 0.8  $\mu\text{g}/\text{mL}$  against the Gram-positive bacteria *Bacillus subtilis* [51]. An aporphine  
 215 alkaloid (63) was isolated from the bark of a Costa Rican sample of *Beilschmiedia alloiophylla* [52]. The  
 216 alkaloid had a MIC of 8  $\mu\text{g}/\text{mL}$  against *Candida albicans*, as well as antileishmanial activity and  
 217 inhibition of acetylcholinesterase. The cyclic peptide tunicyclin D (64) was isolated from roots of the  
 218 medicinal herb *Psammosilene tunicoides* W. C. Wu *et. C. Y. Wu* [53]. The peptide exhibited MIC<sub>80</sub> values  
 219 of 0.3-16  $\mu\text{g}/\text{mL}$  against *Candida* species and 1.0  $\mu\text{g}/\text{mL}$  against *Cryptococcus neoformans*.

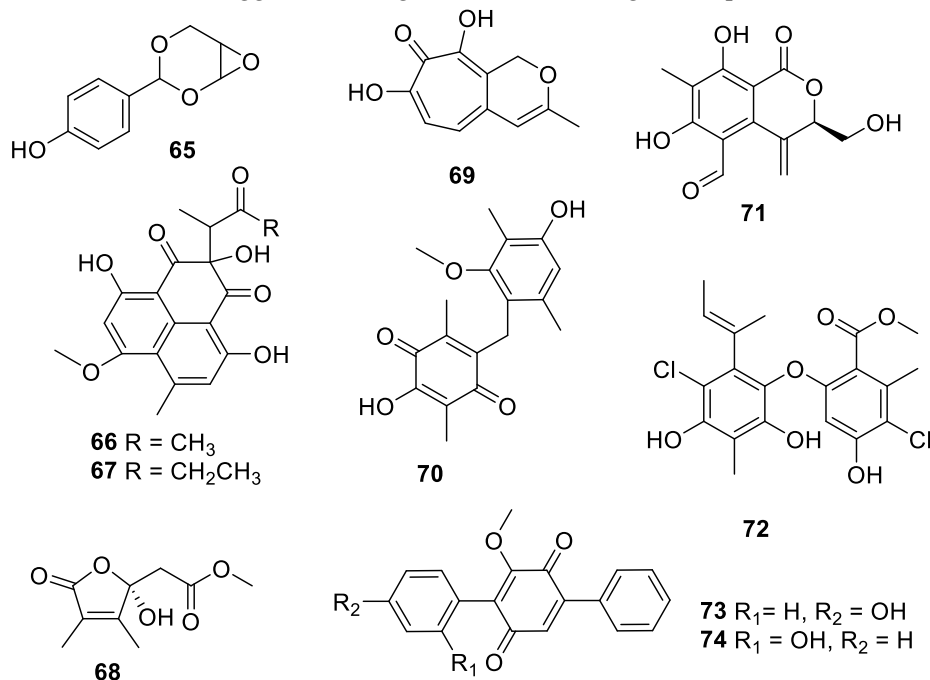


220  
 221 **Figure 12.** Structures of natural products 60-64

## 222 2.4 Natural product antifungal leads from fungi

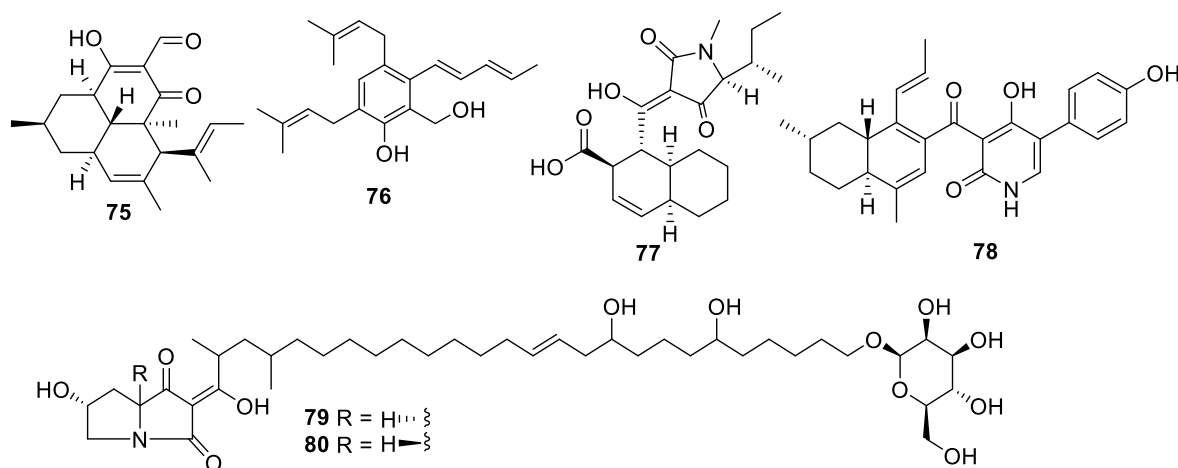
223 Within the last decade, fungi were the most prolific source of novel antifungal leads (Figures 13-  
 224 17, **65-98**). An extract of the endophytic species *Pestalotiopsis mangiferae* obtained from the leaves of  
 225 the plant *Mangifera indica* Linn. yielded an unprecedented epoxyacetal 4-(2,4,7-trioxa-  
 226 bicyclo[4.1.0]heptan-3-yl) phenol (**65**) with a MIC of 0.04  $\mu\text{g/mL}$  against *Candida albicans* strains and  
 227 1.3  $\mu\text{g/mL}$  against the bacterium *Micrococcus luteus* [54]. Two phenalenones, auxarthron A and D (**66**,  
 228 **67**) were obtained from fermentation extracts of an *Auxarthron pseudauxarthron* strain isolated from  
 229 rabbit dung [55]. The compounds have MIC values of 3.2 and 6.4  $\mu\text{g/mL}$  against *Cryptococcus*  
 230 *neoformans* and *Candida albicans* respectively. Further investigation into these compounds  
 231 demonstrated that they are unnatural artifacts, arising from reaction of natural products with ketone  
 232 solvents employed during the extraction. Grifolaone A (**68**) was isolated from the edible mushroom  
 233 *Grifola frondosa*. Interestingly, the hemiketal lactone was obtained in an optically active form and  
 234 assigned as the *S* enantiomer [56]. The furanone was a potent inhibitor, MIC of 0.15  $\mu\text{g/mL}$ , of the  
 235 opportunistic human pathogen *Pseudallescheria boydii* and also had a MIC of 10  $\mu\text{g/mL}$  against  
 236 *Aspergillus fumigatus*.

237 The tropolone nemanolone B (**69**) was isolated from fermentation of a *Nemania* sp. fungus and  
 238 displayed antifungal activity with an  $\text{IC}_{50}$  of 4.5  $\mu\text{g/mL}$  against *Candida albicans*, and similar levels of  
 239 activity against the parasite *Plasmodium falciparum* and human tumor cell lines [57]. The quinone  
 240 pleosporalin E (**70**), isolated from a marine-derived *Pleosporales* sp., inhibited *Candida albicans* with a  
 241 MIC of 7.4  $\mu\text{g/mL}$  [58]. Five new isocoumarins were isolated from fermentation of an endophytic  
 242 *Pestalotiopsis* sp. obtained from *Photinia fraseri*. Among these, pestalactone C (**71**) inhibited *Candida*  
 243 *glabrata* with a  $\text{MIC}_{50}$  value of 3.5  $\mu\text{g/mL}$  [59]. Aspergillusether D (**72**), isolated from fermentation of  
 244 *Aspergillus unguis* PSU-RSPG204, inhibited *Cryptococcus neoformans* with a MIC value of 8  $\mu\text{g/mL}$ , and  
 245 inhibited *Candida albicans* at a lower level [60]. A series of *p*-terphenyl natural products was isolated  
 246 from a strain of *Floricola striata* inhabiting the lichen *Umbilicaria* sp., among which the quinones  
 247 floricolin B and C (**73**, **74**) displayed  $\text{MIC}_{80}$  values of 8  $\mu\text{g/mL}$  against *Candida albicans* [61]. Further  
 248 investigation of floricolin C suggested a fungicidal action through disruption of mitochondria [62].



**Figure 13.** Structures of natural products 65-74

251 Extended fermentation (365 days) of a marine-derived strain of *Aioliomyces pyridodomos* led to  
 252 the appearance of new metabolites, of which onydecalin C (**75**) had a MIC of 2  $\mu\text{g}/\text{mL}$  against  
 253 *Histoplasma capsulatum* [63]. The same strain, in a more conventional fermentation period (25 days),  
 254 produced aintennol A (**76**) with an  $\text{IC}_{50}$  of 8  $\mu\text{g}/\text{mL}$  against *Histoplasma capsulatum* [64]. Genome  
 255 mining for potential Diels-Alderase enzymes identified a potential candidate in the sequence of  
 256 *Penicillium variable*. The putative biosynthetic gene cluster was engineered into an *Aspergillus nidulans*  
 257 expression host, enabling the isolation of varicidin A (**77**) with a  $\text{MIC}_{50}$  value of 8  $\mu\text{g}/\text{mL}$  against  
 258 *Candida albicans* [65]. The *N*-demethylated analogue, varicidin B, was two-fold less active. In the same  
 259 manner, the ilicicolin H biosynthetic gene cluster including a putative Diels-Alderase from a  
 260 producing strain, *Neonectria* sp. DH2, was heterologously expressed in *Aspergillus nidulans*. In  
 261 addition to ilicicolin H, a shunt metabolite ilicicolin J (**78**) was isolated with a MIC of 6.3  $\mu\text{g}/\text{mL}$   
 262 against *Candida albicans* [66]. Heterologous expression was also employed to confirm the biosynthetic  
 263 gene cluster involved in the production of the burnettramic acids A and B (**79** and **80**) in *Aspergillus*  
 264 *burnettii* FRR 5400 [67]. Burnettramic acid A had a MIC value  $< 1 \mu\text{g}/\text{mL}$  against *Candida albicans* and  
 265 *Saccharomyces cerevisiae* while burnettramic acid B was slightly less active with values of 1-2  $\mu\text{g}/\text{mL}$ .



**Figure 14.** Structures of natural products 75-80

269 Co-culture of two extremophilic fungal strains of *Penicillium fuscum* (Sopp) Raper & Thom and  
 270 *Penicillium camembertii/clavigerum* Thom isolated from a single sample of surface water from Berkeley  
 271 Pit Lake led to the production of novel metabolites. Berkeleylactone A (**81**) displayed modest  
 272 antifungal activity with an  $\text{IC}_{50}$  of 6  $\mu\text{g}/\text{mL}$  against *Candida glabrata* and higher antibacterial activity  
 273 [68]. Fermentation of a Saudi strain of *Petriella setifera* led to the identification of the triterpene  
 274 glycoside amnomopin (**82**) with MIC values of 0.5-2  $\mu\text{g}/\text{mL}$  against *Candida* species [69]. Sclerodol B  
 275 (**83**), a triterpene from extracts of the endophyte *Scleroderma* UFSM Sc1(Persoon) Fries obtained from  
 276 *Eucalyptus grandis* had a MIC of 6.3  $\mu\text{g}/\text{mL}$  against *Candida krusei* with weaker activity against other  
 277 species [70]. A strain of the marine-derived fungus *Stachybotrys chartarum* produced several novel  
 278 diterpenoids, of which atranone Q (**84**) had a MIC of 8  $\mu\text{g}/\text{mL}$  against *Candida albicans* and weaker  
 279 antibacterial activity [71].

280 An endophytic *Penicillium* sp. isolated from grass produced picolinic acid derivatives in  
 281 fermentation. Penicolinate B and C (**85**, **86**) had MIC values of 1.5 and 3.7  $\mu\text{g}/\text{mL}$ , respectively, against  
 282 *Candida albicans* [72]. The didymellamide series of pyridone alkaloids was isolated from cultures of  
 283 the marine-derived fungus *Stagonosporopsis cucurbitacearum* and *Coniochaeta cephalothecoides* [73,74].  
 284 Didymellamide A, F and G (**87-89**) were antifungal with MIC values of 3  $\mu\text{g}/\text{mL}$  against *Candida*  
 285 species. The fermentation also yielded (+)-*N*-hydroxyapiosporamide (**90**), the enantiomer of the  
 286 previously isolated natural product, with a MIC value of 6.3  $\mu\text{g}/\text{mL}$  against *Candida albicans*.

287 Fermentation of a *Cyathus* cf. *striatus* basidiomycete led to the isolation of the alkaloid pyristriatin A  
 288 (91) with a MIC of 8.3  $\mu\text{g}/\text{mL}$  against *Rhodotorula glutinis* and similar levels of activity against Gram-  
 289 positive bacteria and human tumor cell lines [75].

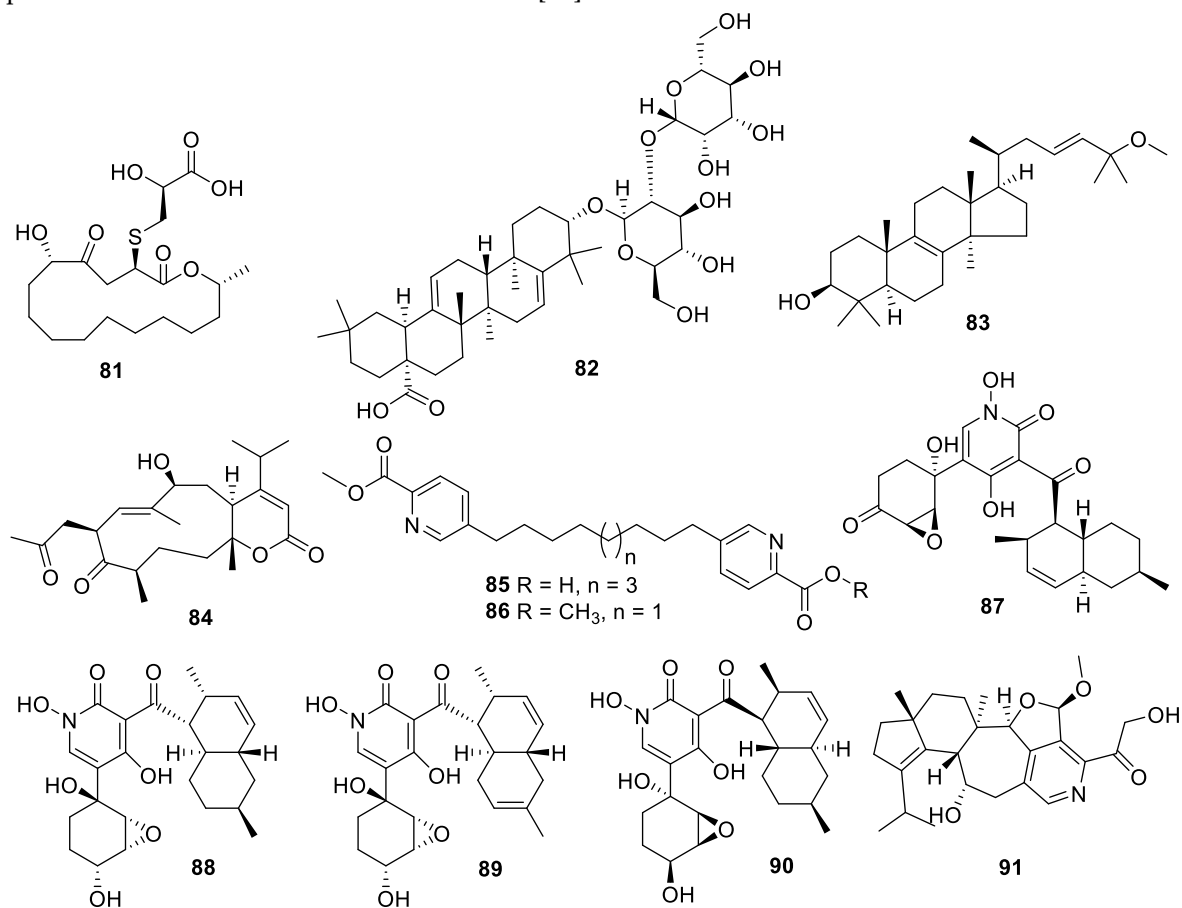
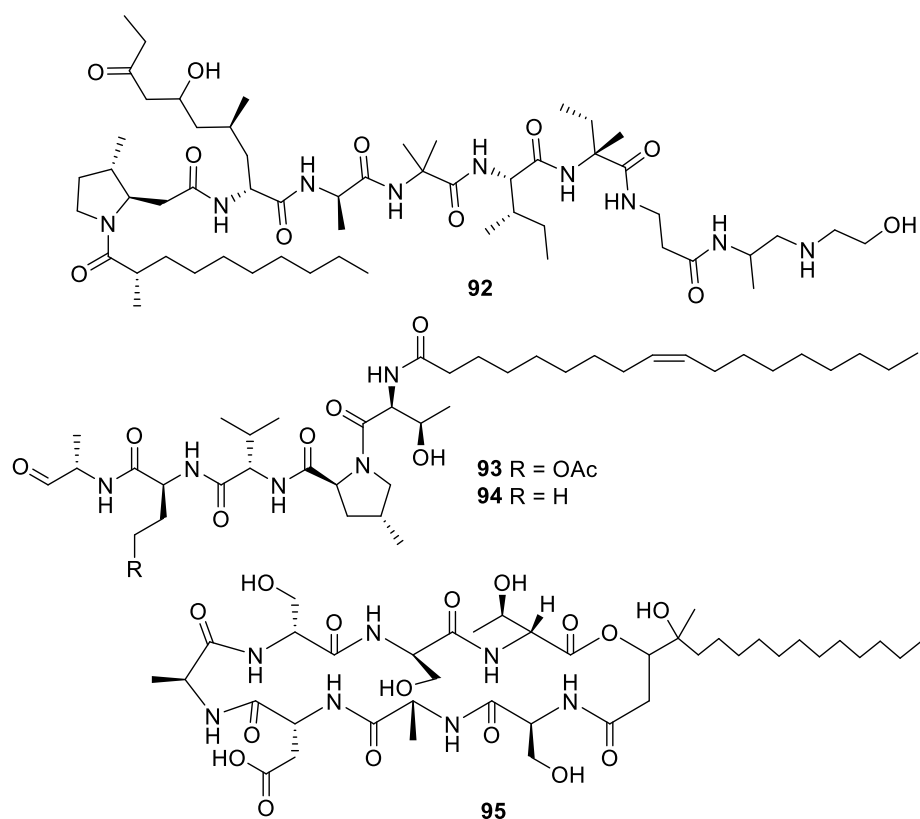


Figure 15. Structures of natural products 81-91

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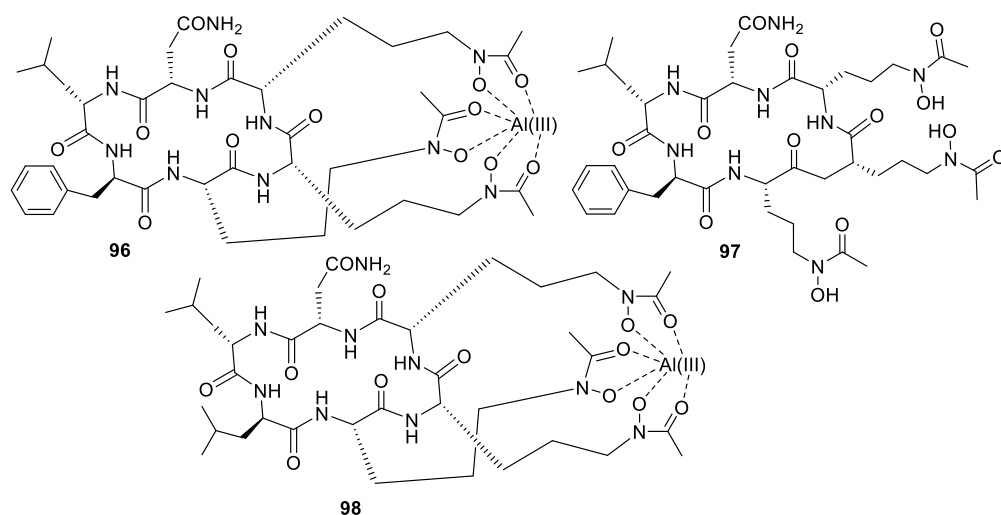
292 The alkalophilic extremophile fungus *Emericellopsis alkalina* VKPM F-1428 was the source of the  
 293 peptaibol emericellipsin A (92), which exhibited antifungal MIC values of 2-4  $\mu\text{g}/\text{mL}$  against *Candida*  
 294 and *Aspergillus* species as well as activity against Gram-positive bacteria. Bioassay-guided  
 295 fractionation of extracts of *Colispora cavincola* isolated from plant litter led to the discovery of the  
 296 linear peptides cavinafungin A and B (93, 94) [76]. The cavinafungins inhibited *Candida* species with  
 297 a MIC of 0.5-4  $\mu\text{g}/\text{mL}$  and *Aspergillus fumigatus* at 8  $\mu\text{g}/\text{mL}$ . However, the antifungal effects were lost  
 298 in the presence of mouse serum. Cavinafungin A also potently inhibits the Zika and dengue virus,  
 299 with the mechanism of action attributed to inhibition of the host signal peptidase [77]. The antifungal  
 300 activity of *Phaeosphaeria* sp. F-167,953 was ascribed to the lipodepsipeptide phaeofungin (95) with  
 301 some structural similarity to the previously known phomafungin [78]. Phaeofungin had a MIC of 4  
 302  $\mu\text{g}/\text{mL}$  against *Trichophyton mentagrophytes* and lower activity against other fungi tested.



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**Figure 16.** Structures of natural products 92-95

305 High-throughput screening by Astellas Pharmaceuticals against a silkworm model of *Aspergillus*  
 306 *fumigatus* infection led to bioassay-guided fractionation activity of an extract of *Acremonium*  
 307 *persicinum* MF-347833. The siderophore hexapeptide ASP2397 (**96**) was discovered as an aluminum  
 308 chelate with exceptional potency against *Aspergillus fumigatus*, with a MIC of 0.2  $\mu\text{g}/\text{mL}$  and efficacy  
 309 at 3.2 mg/kg in a mouse in vivo model [79]. The metal-free form AS2488059 (**97**) as well as the  
 310 congener AS2524371 (**98**) were also isolated, and the target was identified as a fungal siderophore  
 311 transporter [80,81]. The compound was out-licensed to Vical and renamed VL-2397, reaching Phase  
 312 II clinical trials that were recently discontinued.



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**Figure 17.** Structures of natural products 96-98

315 **3. Discussion**

316 Between 2010-2019, we identified nearly a hundred novel natural products reported with  
 317 antifungal activity against human pathogens. The compounds originate from a variety of sources  
 318 comprising bacteria, alga, fungi, sponges and plants with fungi being the most prolific source of  
 319 antifungal compounds. The techniques employed range from classical phytochemical studies with  
 320 plants to high-throughput screening of extract collections and modern microbiological strategies such  
 321 as co-cultivation and heterologous expression of biosynthetic gene clusters. All the major classes of  
 322 natural products including polyketides, shikimate metabolites, terpenoids, alkaloids and peptides  
 323 are represented. As the majority of examples in this review involve the initial disclosure of activity,  
 324 further investigations are needed to assess the therapeutic potential of highly active compounds as  
 325 well as their selectivity as antifungal agents. Meanwhile, it is interesting to observe the  
 326 physicochemical space occupied by these natural product leads (Table 1). Although the compounds  
 327 are diverse in their structural features, they are largely compliant with the typical guidelines for small  
 328 molecule drug-like chemical space. While many of the natural products are large in molecular weight,  
 329 resulting in an average of 569, other properties like hydrogen bonding potential, molecular flexibility  
 330 and polarity often remain within the recommended limits.  
 331

Compound	MW	clogP	HBD	HBA	nrot	TPSA
1	607	4.8	3	8	10	115
2	198	2.1	2	2	2	41
3	508	2.9	4	12	5	186
8	751	7.0	5	10	10	155
17	1123	7.6	13	18	26	312
21	1580	0.9	15	29	33	472
28	987	4.7	6	19	11	253
30	817	4.3	4	12	9	167
31	1200	-5.7	23	32	36	546
35	2144	-0.4	26	55	30	876
36	236	2.4	2	2	3	41
37	274	3.0	2	2	4	41
42	419	7.7	1	4	14	56
44	385	6.2	1	3	9	47
45	713	8.1	7	7	10	110
46	667	0.7	6	13	10	188
48	393	3.1	6	8	13	130
53	438	5.0	4	6	12	121
54	423	2.0	2	5	5	61
55	439	2.3	2	6	5	70
56	340	1.9	4	7	6	124
57	423	6.0	2	5	7	68
58	297	2.7	1	4	1	48
59	1065	8.0	11	18	23	319

60	487	6.5	4	7	12	132
61	352	3.0	0	5	3	64
62	370	0.5	0	6	3	51
63	281	2.5	1	3	2	33
64	901	-0.1	10	21	9	303
65	194	1.3	1	4	2	51
66	358	3.0	3	7	6	121
68	200	-1.6	1	5	4	73
69	206	-1.0	2	4	2	67
70	316	3.5	2	5	5	84
71	264	0.8	3	6	5	104
72	427	7.9	3	6	8	96
74	306	4.7	1	4	4	64
75	329	6.8	1	3	3	54
76	327	6.7	2	2	9	41
77	376	3.3	2	6	5	95
78	432	5.4	3	5	4	87
79	770	3.2	8	13	35	218
81	405	2.3	3	7	6	146
82	779	6.4	8	13	14	216
83	457	9.5	1	2	6	30
84	391	2.7	1	5	3	81
85	399	5.1	1	6	14	89
87	444	1.4	3	8	6	128
90	446	3.5	4	8	7	131
91	442	3.5	2	6	6	89
92	1064	4.9	10	20	38	294
93	792	6.6	5	14	31	200
95	904	-2.0	13	23	23	368
97	891	-2.5	11	23	21	339
<b>Average</b>	569	3.4	5	10	11	155

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**Table 1.** Physicochemical properties of antifungal natural products. MW = molecular weight, clogP = calculated log P, HBD = hydrogen bond donors, HBA = hydrogen bond acceptors, nrot = number of rotated bonds, TPSA = total polar surface area in Å<sup>2</sup>. The values were taken from SciFinder (<https://scifinder-n.cas.org>), based on calculations using Advanced Chemistry Development (ACD/Labs) Software V11.02. In certain cases where the data was absent in SciFinder, values were calculated using the Molinspiration website (<https://www.molinspiration.com/>).

For natural products where a series of related compounds was reported, one representative example was selected. Shaded cells indicate values above the recommended guidelines for small molecule drug-like chemical space (MW ≤ 500, Clog P ≤ 5, HBD ≤ 5, HBA ≤ 10, nrot ≤ 10, TPSA ≤ 140).



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