Metabolic Potential of Epichloë Endophytes for Host Grass Fungal Disease Resistance

Krishni Fernando 1,2, Priyanka Reddy 1, German C. Spangenberg 1,2, Simone J. Rochfort 1,2, and Kathryn M. Guthridge 1,*

1 Agriculture Victoria, AgriBio, Centre for AgriBioscience, Bundoora, VIC 3083, Australia; krishni.fernando@agriculture.vic.gov.au (K.F.); priyanka.reddy@agriculture.vic.gov.au (P.R.); german.spangenberg@agriculture.vic.gov.au (G.C.S.); simone.rochfort@agriculture.vic.gov.au (S.J.R.)
2 School of Applied Systems Biology, La Trobe University, Bundoora, VIC 3083, Australia
* Correspondence: kathryn.guthridge@agriculture.vic.gov.au

Abstract: Asexual species of the genus Epichloë (Clavicipitaceae, Ascomycota) form endosymbiotic associations with Pooidae grasses. This association is important both ecologically and to the pasture and turf industries, as the endophytic fungi confer a multitude of benefits to their host plant that improve competitive ability and performance such as growth promotion, abiotic stress tolerance, pest deterrence and increased host disease resistance. Biotic stress tolerance conferred by the production of bioprotective metabolites has a critical role in an industry context. While the known antimammalian and insecticidal toxins are well characterized due to their impact on livestock welfare, antimicrobial metabolites are less studied. Both pasture and turf grasses are challenged by many phytopathogenic diseases that result in significant economic losses and impact livestock health. Further investigations of Epichloë endophytes as natural biocontrol agents can be conducted on strains that are safe for animals. With the additional benefits of possessing host disease resistance, these strains would increase their commercial importance. Field reports have indicated that pasture grasses associated with Epichloë endophytes are superior in resisting fungal pathogens. However, only a few antifungal compounds have been identified and chemically characterized, and these from sexual (pathogenic) Epichloë species, rather than those utilized to enhance performance in turf and pasture industries. This review provides insight into the various strategies reported in identifying antifungal activity from Epichloë endophytes and, where described, the associated antifungal metabolites responsible for the activity.

Keywords: antifungal metabolites; fungitoxic; pasture and turf protection; metabolite annotation; bioprospecting endophytes

1. Introduction

Pasture is one of the main food sources for livestock throughout the world. While pure pasture-based farming systems rely solely on pasture, mixed farming systems supplement with commodities such as cereals and grains. More consumers are shifting towards “grass-fed” livestock produce, as it caters for both increasing global food demands and social concerns for animal-welfare [1,2]. Novel solutions to sustainably manage food supply systems also address increasing demands for nutritious food [3,4]. There is a wide variety of pasture species used in agriculture including annual grasses, perennial grasses, legumes, and herbs. Forage grasses that are adaptable to a wide range of climatic conditions are extensively used in farming systems [5,6].

With changing climates and evolving pest and pathogen pressures, pasture and turf grasses are threatened by abiotic and biotic stresses with increasing severity [5,7–9]. It has long been known that Epichloë fungi of the family Clavicipitaceae, associated with Pooidae grasses such as Lolium spp. (e.g., perennial ryegrass, short-term ryegrass and
tall fescue), improve host plant abiotic and biotic stress tolerance by producing bioactive metabolites [9–12]. Though *Epichloë* endophytes are historically well-characterised for their antimammalian and insecticidal alkaloid toxins, recent studies confirm that their bioactivities are not limited to these compounds alone [11,13–15]. Therefore, there is a requirement for an increase in applied research into *Epichloë* endophyte-derived bioactive metabolites to improve host grass performance and stress tolerance. This review focuses on prospects for *Epichloë* endophytes providing host plant disease resistance, and discusses modern experimental data analysis techniques for bioprospecting of antifungal metabolites in novel *Epichloë* strains.

2. *Epichloë* Endophytes

*Epichloë* fungi infecting grasses are categorized as sexual or asexual species based on their transmission mechanism. Sexually transmitting *Epichloë* spp. are known to cause choke disease in grasses during flowering, leading to reduced seed yield and aesthetic value of turf grasses [16,17]. In this process the fungus forms a stroma with spores and horizontal transmission occurs [16,17]. As they cause disease, sexual *Epichloë* spp. are considered pathogenic even though they live asymptomatically in the host plant during vegetative growth stages. Asexual *Epichloë* spp. are clonal, transmit vertically by infecting the seeds, and do not cause diseases to the host grass in any growth stage of the host plant [16]. Therefore, as they provide significant performance benefits, the pasture and turf grass industries utilize asexual (endophytic) strains of *Epichloë* species.

Asexual *Epichloë* spp. form endophytic associations with cool-season grasses by colonizing leaves, pseudostems, seeds and seedlings. In this mutualistic relationship, the endophyte relies on the plant for vertical dissemination while also gaining shelter and nutrients throughout its life [13]. The production of functional metabolites triggered by the association greatly benefits the host plant by conferring abiotic and biotic stress tolerances such as improved seedling vigour, persistence, and enhanced growth [13,18]. While most of these benefits are advantageous in an agricultural scenario, *Epichloë* endophytes are also well known for producing toxic antimammalian alkaloids (lolitrem B—ryegrass staggers, ergovaline—fescue toxicosis). These alkaloid toxins are extremely harmful to grazing animals and cause significant economic losses to the livestock industry [19–21]. Consequently, studies related to their biological and chemical properties, genetics, biosynthetic pathway and modes of action have been the focus from as early as 1980s. The biosynthetic intermediaries of these compounds have also been characterized for biological activity and many have been selected as candidates for insecticides [22]. Previous reports on *Epichloë* endophytes have demonstrated that the alkaloids lolitrem B, ergovaline, *n*-acetylloline, *n*-formylloline and peramine possess insecticidal and invertebrate pest deterrence properties [20,23,24]. However, the broad chemical diversity and metabolic capacity of *Epichloë* endophytes relating to antimicrobial activity remain largely unknown.

Field, glasshouse and lab-based studies have established that *Epichloë* endophytes improve host plant disease resistance, and a few studies have identified the production of metabolites with novel antimicrobial properties that may play a role in host plant protection from phytopathogens [11,12,25]. However, in most of these studies wild-type strains such as SE (Standard Endophyte; also referred to as wild type or common endophyte, CE) or Ky31 (Kentucky31) were investigated rather than the animal friendly strains utilized in pastoral agriculture. Thus, to better exploit endophyte-mediated disease resistance, improve pasture and turf quality, and reduce the impact of phytopathogen disease on animal welfare, *Epichloë* strains that are safe for animals, should be investigated.

3. Novel Endophytes

The search for endophytes that do not produce metabolites toxic to mammalian grazers led to the discovery of novel endophytes and their use commercially, as shown by the many novel strains for which Plant Breeders’ Rights has been granted worldwide, and those marketed with registered trademarks (Table 1) [26–28]. The significant economic
benefits of animal-safe endophyte strains that also improve pasture persistence by reducing the impact of invertebrate pests underpinned the investigation and exploration of high-performing endophyte infected pasture grasses [29–31]. Novel endophytes are typically screened and selected based on alkaloid profiles in planta, in particular low/no lolitrem B and low/no ergovaline combined with bioprotective (insect deterring) lolines and/or peramine. However, more recently, screening methods have been extended to biosynthetic pathways and gene clusters associated with these known alkaloids [27,29,32,33].

The search for animal-safe endophyte strains is important to the agricultural industry, with new endophytes being developed continuously (Table 1). Therefore, high-throughput methods and commercial standards are used regularly to detect the presence of alkaloids in pasture [34]. Those compounds that are not commercially available can also be isolated and purified using published methodologies [19,35]. The screening methods are, however, limited to the “known known” alkaloids—the well described *Epichloë*-derived compounds referred to in Table 1. Of the novel endophytes described on Table 1 only two, Nea 12 and Nea 23, have been investigated for production of antifungal metabolites in vitro and in planta [36,37]. Thus, further research is required to investigate the ‘known unknown’ metabolites, such as those responsible for disease resistance, that are beneficial to the pasture and turf grass related industries [38].
Table 1. Novel *Epichloë* endophyte strains for which Plant Breeders’ Rights (PBR) have been granted worldwide.\(^1\)

| Country | PBR Grant Date | Endophyte Strain (Market Name) | *Epichloë* Species | Host Common Name | Known Alkaloid Profile | Applicant |
|---------|---------------|---------------------------------|---------------------|------------------|------------------------|-----------|
| New Zealand | 23 April 1996 (expired) 20 October 2003 26 October 2004 | AR1 | *E. festucae* var. *lolii* (*Lp*TG-1) | Perennial ryegrass | P | Grasslanz Technology Ltd. (GTL; Palmerston North, New Zealand) |
| New Zealand | 21 April 2015 | AR1006 | *E. uncinata* MEADOW FESCUE | Meadow fescue | L | GTL |
| New Zealand | 5 October 2016 | AR1017 | *E. uncinata* MEADOW FESCUE | Meadow fescue | L | GTL |
| New Zealand | 29 November 2018 | AR127 | *E. festucae* var. *lolii* (*Lp*TG-1) | Perennial ryegrass | | GTL |
| New Zealand | 25 July 2008 | AR37 | *E. festucae* var. *lolii* (*Lp*TG-1) | Perennial ryegrass | J | Fischer Fleurquin Gustavo (FFG; Montevideo, Uruguay) |
| New Zealand | 25 July 2008 30 March 2010 | AR501 | *E. coenophiala* (*Fa*TG-1) | Tall fescue | LP | GTL |
| New Zealand | 1 February 1999 (expired) 26 October 2004 6 December 2004 | AR542 | *E. coenophiala* (*Fa*TG-1) | Tall fescue | LP | GTL |
| New Zealand | 25 July 2008 29 September 2010 2 August 2013 4 September 2014 | AR584 | *E. coenophiala* (*Fa*TG-1) | Tall fescue | LP | GTL |
| New Zealand | 12 May 2010 19 August 2013 22 May 2017 | AR601 (Avanex\(^®\)) | *E. coenophiala* (*Fa*TG-1) | Tall fescue | EL | GTL |
| New Zealand | 12 May 2010 22 May 2017 | AR604 | *E. coenophiala* (*Fa*TG-1) | Tall fescue | EL | GTL |
| New Zealand | 28 August 2014 22 May 2017 3 April 2018 21 November 2014 | AR95 (Avanex\(^®\)) | *E. festucae* var. *lolii* (*Lp*TG-1) | Perennial ryegrass | E | GTL |

\(^1\) For a list of approved *Epichloë* species with Host Common Name, known alkaloid profile and host plant, see https://www.grasslanz.com/epichloespecies.html.
Table 1. Cont.

| Country        | PBR Grant Date          | Endophyte Strain (Market Name) | Epichloë Species                          | Host Common Name | Known Alkaloid Profile | Applicant                                      |
|----------------|-------------------------|--------------------------------|-------------------------------------------|------------------|------------------------|------------------------------------------------|
| New Zealand    | 17 January 2019         | CM142                          | E. festucae var. lolii (Lp TG-1)          | Perennial ryegrass | J                      | Cropmark Seeds Ltd. (CSL; Rolleston, New Zealand) |
|                |                         |                                |                                           |                  |                        | Cropmark Seeds Australia Pty Ltd. (CSA; South Melbourne, Australia) |
| Australia      | 17 August 2020          |                                |                                           |                  |                        |                                                 |
| New Zealand    | 27 August 2014          | E815 (Edge)                    | E. festucae var. lolii (Lp TG-1)          | Perennial ryegrass | LtmEP                  | DLF Seeds A/S (DLF; Roskilde, Denmark) DLF      |
| Australia      | 23 October 2017         |                                |                                           |                  |                        |                                                 |
| New Zealand    | 23 June 2010            | Happe                          | E. siegelii                               | Perennial ryegrass | L                      | Barenbrug New Zealand Ltd. (BBNZ; Christchurch, New Zealand) DLF |
| Australia      | 23 October 2017         |                                |                                           |                  |                        |                                                 |
| New Zealand    | 25 July 2008            | Nea 2 (NEA/NEA2/NEA4) ³       | E. festucae var. lolii (Lp TG-1)          | Perennial ryegrass | LtmEP                  | Barenbrug New Zealand Ltd. (BBNZ; Christchurch, New Zealand) BBNZ |
|                | 30 June 2009            | Nea 3 (NEA4 ³)                 | E. festucae var. lolii (Lp TG-1)          | Perennial ryegrass | EP                     | BBNZ                                           |
|                | 25 July 2008            | Nea 6 (NEA2 ³)                 | E. festucae var. lolii (Lp TG-1)          | Perennial ryegrass | EP                     | BBNZ                                           |
|                | 29 August 2014          | Nea 10                         | E. festucae var. lolii (Lp TG-1)          | Perennial ryegrass | EP                     | BBNZ                                           |
|                | 13 August 2014          | Nea 11                         | E. festucae var. lolii (Lp TG-1)          | Perennial ryegrass | EP                     | BBNZ                                           |
|                | 19 March 2021           | Nea 12                         | Epichloë sp. (Lp TG-3)                    | Perennial ryegrass | J                      | Agriculture Victoria Services Pty Ltd. (Bundoora, Australia) |
|                | 29 August 2014          | Nea 21                         | Epichloë sp. (FaTG-3)                     | Tall fescue       | LP                     | BBNZ                                           |
|                | 29 August 2014          | Nea 23                         | Epichloë sp. (FaTG-3)                     | Tall fescue       | LP                     | BBNZ                                           |
|                | 10 July 2019            | Nea 47 (NEA2 ³)                | E. festucae var. lolii (Lp TG-1)          | Perennial ryegrass | EP                     | BBNZ                                           |
|                | 18 August 2014          | PTK647 (Protek®)               | E. coenophiala (FaTG-1)                   | Tall fescue       | EL                     | DLF                                           |
| Australia      | 23 October 2017         |                                |                                           |                  |                        |                                                 |
| Country           | PBR Grant Date  | Endophyte Strain (Market Name) | Epichloë Species | Host Common Name | Known Alkaloid Profile | Applicant |
|-------------------|-----------------|--------------------------------|------------------|------------------|------------------------|-----------|
| New Zealand       | 28 August 2014  | U12                            | *E. uncinata*    | Meadow fescue    | L                      | CSL, CSA |
| Australia         | 12 August 2021  |                                |                  |                  |                        |           |
| New Zealand       | 2 September 2016| U13                            | *E. uncinata*    | Meadow fescue    | L                      | CSL      |
| New Zealand       | 2 September 2016| U12 (GrubOUT®)                 | *E. uncinata*    | Meadow fescue    | L                      | CSL, CSA |
| Australia         | 30 January 2021 |                                |                  |                  |                        |           |
| Argentina         | 11 March 2015   |                                |                  |                  |                        | Gentos SA |
| European Union    | 22 May 2017     |                                |                  |                  |                        | CSL      |
| New Zealand       | 14 October 2008 | UNCl                           | *E. uncinata*    | Meadow fescue    | L                      | CSL      |
| Australia         | Not applicable  | AR5 (Endo5)                    | *E. festucae var. lolii (LpTG-1)* | Perennial ryegrass | EP                     | GTL      |
| USA               | Not applicable  | E34®                           | *E. coenophiala (FaTG-1)* | Tall fescue      | LP                     | Barenbrug USA (Tangent, OR, USA) |
| USA               | Not applicable  | KY31 (Kentucky31)              | *E. coenophiala (FaTG-1)* | Tall fescue      | EPL                    | Not applicable |
| worldwide         | Not applicable  | SE (Standard endophyte)        | *E. festucae var. lolii (LpTG-1)* | Perennial ryegrass | LtmEP                  | Not applicable |

1 UPOV Pluto database search for Plant Breeders’ Rights granted to 26 November 2021 [https://pluto.upov.int/search](https://pluto.upov.int/search) (Search: Botanical name - *Epichloë*). 2 Alkaloid profile, P = peramine, L = lolines, E = ergovaline, Ltm = lolitrem B, J = epoxy-janthitrems. 3 Strains are sometimes marketed as a combination e.g., NEA2 = Nea 2, Nea 6, Nea 47. 4 Commercial endophytes that were not listed in UPOV Pluto database search results. 5 Wildtype, toxic strains.
4. Disease Stress to Pasture Grasses

Pasture grasses are threatened by many pathogenic diseases, causing devastating losses to pasture yield and quality. These diseases include rusts, leaf spot diseases, blights, blotches, moulds, and wilts caused by pathogenic fungi, bacteria or viruses (Table 2).

In perennial ryegrass, crown rust (Puccinia coronata) is one of the most severe fungal disease-causing pathogens that affects foliage, causing substantial losses in pasture and turf grass industries. Severe rust infections can cause up to 37% loss in dry matter (DM) when infected plants are harvested and dried, and a 94% loss in fresh matter (FM) yield [39–42]. Water soluble carbohydrate content in rust-infected grasses is significantly lower compared to uninfected grasses, which in turn affects digestibility and palatability and can lead to low milk yields if used as feed [43–45]. Root growth is also reduced due to depleted carbohydrate reserves [46]. Further, the resultant increase in dead herbage tissue makes grass susceptible to Pithomyces chartarum, which causes serious disease of facial eczema in cattle and sheep [46,47].

Other foliage pathogens include Drechslera sissans, which causes significant production losses; disease incidence is significantly increased with high nitrogen (N) use conditions [44,48,49]. Though not prominent in Australia, snow mould (Microdochium nivale) and grey leaf spot (Pyricularia grisea) are another two serious diseases causing yield losses in perennial ryegrass and tall fescue in the northern hemisphere [50,51].

As well as the effect on foliage, several fungal pathogens also directly infect the inflorescence, thus altering seed yield. The most significant of these are stem rust (Puccinia graminis) and blind seed disease (Gloeotinia granigena), which both occur on seed heads on a wide range of pasture grasses. Stem rust infections can result in seed yield losses of up to 93% in turf-type perennial ryegrass cultivars, forcing seed producers, especially those growing late maturing cultivars, to use fungicides to prevent losses. Incidence of blind seed disease depends on environmental conditions. Infection usually results in seed death from heavily infected stands [44].

Pathogens are not limited to fungi. Bacterial pathogens include Xanthomonas translucens pv graminis, which causes bacterial wilt disease resulting in yield losses in ryegrass [52]. Pseudomonas syringae pv atropurpurea causes chlorosis in ryegrass [53]. Viral diseases can also infect perennial grasses, and occur in high incidences causing serious damage. These include barley yellow dwarf virus (BYDV), cereal yellow dwarf virus (CYDV), and ryegrass mosaic virus (RGMV). Upon infection, BYDV and CYDV cause leaf yellowing, stunting and tillering, and these symptoms have an effect on plant performance, productivity, quality and yield [7,54]. Mosaic streaking necrosis by RGMV leads to herbage yield loss [7]. Table 2 further describes common diseases in perennial ryegrass and tall fescue used for pasture and turf, detailing causative organism, symptoms, damage and current control measures.

Phytopathogenic fungi causing disease in grasses may also produce mycotoxins that pose a threat to animal health and wellbeing. Fusarium sp. derived toxins—T-2/HT-2, zearalenone, deoxynivalenol—affect food intake and animal performance [55,56]. Though actual toxins are not well characterized, Drechslera biseptata has been associated with acute bovine liver disease [57,58]. Rusts caused by Puccinia sp., pathogens reduce the palatability of infected grasses by altering total carbohydrate contents [41,59].

Disease outbreaks cause significant yield loss and degrade forage quality, and hence proper control measures are required. The most widely used method for disease control is chemical treatment, which includes spraying fungicides and insecticides for disease vectors [60–62]. Fungicides include dimethylation inhibitors, nickel salts and dithiocarbamates [63,64]. Another approach is to apply suitable fertiliser in desirable rates of application to enrich the growth medium with macro and micro nutrients necessary for plant growth and performance [63,65]. Cultural and mechanical control measures include grazing and irrigation management practices [49,60]. All these curative methods are costly and labour demanding. Continuous long term application of fungicides may have a negative impact on livestock health, as some fungicides are toxic when accumulated in large quantities [66,67]. Furthermore, fungicides have a negative impact on Epichloë
endophytes in perennial ryegrass and tall fescue and may lead to loss of benefits from the symbiotic association [64,68]. Breeding of resistant varieties is time consuming and costly. Thus, introducing beneficial microorganisms as biocontrol agents may be a more cost and time-efficient response.

Bacteria have been reported to be effective biocontrol agents when applied to the plant or seeds. *Pseudomonas aeruginosa* reduces disease incidence and disease severity of grey leaf spot in perennial ryegrass when applied to seeds or the plant in controlled environment pot trials and field trials [69]. *Paenibacillus elgii* SD17 reduces disease severity of brown patch disease and pythium blight in turf grasses in both controlled chamber and field trials [70]. Inoculating novel *Epichloë* strains to grass populations would have relatively minimal ecological impacts as the symbiota are naturally occurring [71].

The asexual *Epichloë* sp. utilized in pastures and turf are known to produce antimicrobial metabolites and inhibit pathogen growth under both in vitro and in planta conditions [25,37,72]. With the availability of new high throughput technology, rigorous analytical methods are available to identify and quantify antimicrobial activity and detect the responsible molecules [73–75]. Early studies have shown the potential of *Epichloë* endophytes to inhibit pathogen infections, while more recent studies have focused on isolating and characterising responsible antimicrobial molecules [25,36,72].
| Disease/ Common Name          | Causative Organism    | Symptoms                                                                 | Damage/Loss                      | Control Measures                                      | References   |
|------------------------------|-----------------------|--------------------------------------------------------------------------|---------------------------------|--------------------------------------------------------|--------------|
| Crown rust                   | *Puccinia coronata*   | Reddish brown spores on leaf                                            | Dry matter yield loss 30–40% and stock thrift | Fungicide; Judicious grazing management; Resistant varieties. | [41,45,76]  |
| Grey leaf spot               | *Pyricularia grisea*  | Small water-soaked lesions on leaf blades gradually turning to dark necrotic spots and to grey spots | Up to 90% pasture loss          | Fungicide; Controlled release of N fertiliser; Biocontrol, Resistant varieties. | [50,63,77,78]|
| Brown blight and net blotch  | *Drechslera sp.*      | Net lesions with small dark brown bars amphigenous lesions with dark brown margins, light brown center | Dry matter and herbage loss     | Fungicide; Managed grazing before it spreads.           | [49,60,79]  |
| Stem end rust                | *Puccinia graminis*   | Reddish brown spores on sheath and stem                                 | Seed yield loss, Dry matter loss | Fungicide; Judicious grazing management; Resistant varieties. | [49,80]      |
| Blind seed disease           | *Gloeotinia temulenta*| Fungal mycelia on seeds under microscopic observation                    | Seed yield loss, reduce seed germination 50–90% | Fungicide; Increased rate of N application.           | [64,65,81]  |
| Snow mould                   | *Microdochium nivele*  | Dark brown lesions and pink sporodochia rows parallel to veins           | Seedling damage leading to yield loss, seed loss and yield loss | Fungicide; Biological control; Compost application. | [51,60,82]  |
| Yellow mould                 | *Ceratobasidium cereale* | Root pathogen                                                          | Yield loss                      | Fungicide                                              | [83,84]      |
| seedling pathogen and leaf spot | *Fusarium solani*     | Wilting of seedlings and necrotic lesions in mature plants               | Yield loss, dry matter loss     | Fungicide                                              | [85,86]      |
| Bacterial wilt               | *Xanthomonas translucens* | Water-soaked lesions and turning to bluish purple colour                 | Forage yield loss 20–40%        | Biological control; Resistant varieties.               | [52,87–89]  |
| Ryegrass mosaic virus        | RGMV                  | Light green-yellow streaky mosaic or brown necrosis on leaves            | Dry matter yield loss 21–30%    | Resistant varieties; Mixed pasture species.           | [7,90]       |
5. Epichloë sp.-Derived Antifungal Activity and Host Disease Resistance Studies

There are limited reports of Epichloë endophyte-derived antifungal activity and host disease resistance. For this review we used the scientific search engine Scopus (www.scopus.com (accessed on 5 November 2021)) to identify peer reviewed scientific reports with the terms “Epichloë” or “Neotyphodium” and “fungitoxic” or “antifungal” in the title, abstract, or keywords. Only publications in English and published in the last 25 years (1996–2021) were considered. Data integrity and collation were performed by either individually checking and filtering nonrelated articles or preserving and adding related articles cited in reviews and journal publications. Relevant secondary documents listed in Scopus but not indexed in the database were also included. Subsequently, only 46 journal articles, three letters, and one short survey (Supplementary Table S1) were found. During the same time period, the number of English journal manuscripts found using the terms “Epichloë” or “Neotyphodium” was 1313. Although only 3.5% of Epichloë fungi-related studies concerned antifungal activity and host disease resistance against fungal pathogens, this number has been increasing (Figure 1). This demonstrates that researchers are recognizing the importance of Epichloë endophyte derived antifungal activity as an important driver for better performing pasture and turf.

![Figure 1](chart.png)

Figure 1. Number of documents (journal articles, letters and short surveys) published in last 25 years on Epichloë-derived antifungal activity and/or disease resistance.

6. Bioprospecting Antifungal Metabolites from Epichloë Endophyte Strains

Despite the paucity of publications on Epichloë-derived antifungal activity and host disease resistance, there have been papers describing various methodologies to demonstrate the presence of biological activity of Epichloë endophyte strains [12,25,91–94]. However, the methods utilized historically are subjective, qualitative and do not allow for accurate comparisons between studies. To address the current research gaps in Epichloë-mediated fungal disease resistance, it is necessary to establish rigorous analytical processes to characterize the bioactivity of Epichloë strains quantitatively and accurately to determine the effect of responsible compounds. Recent advances in analytical methods and software tools enable development of standardized protocols and processes to analyze antifungal activity. In this section we review current knowledge in the field, highlighting the methods and tools available, to create a schematic process for identification, characterization and application of antifungal compounds produced by Epichloë endophytes (Figure 2).
the methods and tools available, to create a schematic process for identification, characterisation and application of antifungal compounds produced by *Epichloë* endophytes (Figure 2).

**Figure 2.** An experimental workflow proposed for bioprospecting antifungal metabolites from *Epichloë* endophytes using a stepwise process. (a) *Epichloë* strain identification, (b) Identification of bioactive strains using in vitro antifungal activity assays, (c) antifungal metabolite isolation and characterisation, (d) untargeted metabolite annotation for antifungal compound detection, (e) qualitative and quantitative confirmation of antifungal metabolites in planta.
6.1. Epichloë Endophyte Strain Identification

Epichloë strains are commonly tested in vitro using different plate-based assays to identify their antifungal activity [92–95]. Endophyte strains must be isolated, purified, and transferred to in vitro cultures prior to being tested (Figure 2a). As they are naturally found in symbiotic association with host plants, it is ideal to freshly isolate the Epichloë strain from an infected plant. [27]. Strain identity should be confirmed in advance of any testing, which generally involves DNA-based PCR analysis, such as an SNP diagnostic test, to enable identification of known Epichloë strains [28,96]. Ideally, novel endophyte strains would be defined by whole genome sequencing prior to testing bioactivity. Alkaloid profiling endophyte strains for currently known Epichloë-derived alkaloids will provide additional information on animal safety concerns [26,27].

6.2. Identification of Bioactive Strains Using In Vitro and in Planta Assays

Antifungal activities of endophyte strains are commonly detected using in vitro dual culture assays [37,92–95,97]. In dual culture assays, phytopathogens and endophytes are grown in close proximity to study their antagonistic reactions (Figure 2b). Growth parameters (growth area, mycelial density, growth direction) of phytopathogens are observed over a period of time to detect inhibitory activity. Earlier studies qualitatively analysed the results based purely on observation. However, with the availability and accessibility of imaging and image analysis software, growth parameter data such as pathogen growth area can easily be converted to quantitative data. Quantitative data can be analysed for precision and errors, and subjected to statistical analysis to detect the significance of antifungal activity [94]. Dual culture assays are suitable for detecting antifungal activity as they are quick, easily replicated and can be used to test against a range of pathogens. Importantly, the in vitro antifungal phenotypes observed are consistent and observed in independent isolates of the same strain and across duplicate assays [37,98,99].

In addition, dual culture assays provide information on direct interaction of the endophyte with phytopathogens, for example, Epichloë festucae strain E437 was shown to reduce hyphal tip growth of the pathogens Drechslera erythropila and Colletotrichum graminicola [95]. Fernando et al. (2020) observed differential bioactivity between three asexual Epichloë strains (Nea 12, Nea 21 and Nea 23) evaluated against three phytopathogens (Ceratobasidium sp., Drechslera sp. and Fusarium sp.), indicating that there is variation in the production of bioactive metabolites and their composition [37]. In vitro liquid culture extracts exhibited differential antifungal activity consistent with dual culture assays [37], establishing that the Epichloë strains produce and secrete antifungal metabolites.

However, it should be noted that virulence of a pathogen may be different when it is infecting a plant; therefore, dual culture assay results may not directly relate to pathogen inhibition in planta. Dual culture assays are also unsuitable for biotrophic pathogens (e.g., rusts such as Puccinia coronata) that are unable to be grown in pure cultures. Spore germination assays are less common but have also been used to study the antifungal activity of Epichloë strains and understand the physiological mechanisms [93,100–102]. It is noteworthy that Christensen and Latch’s study in 1991 is the only available in vitro study of Epichloë coenophiala (previously known as Acremonium/Neotyphodium coenophialum) demonstrating antifungal activity against spores of the rust pathogen Puccinia graminis by inhibition of urediniospore germination [101]. Spore germination assays are also a useful tool for testing antifungal compounds isolated from Epichloë sp. [102].

Detached leaf assays are another type of semi-in vitro assay used to overcome some of the limitations associated with dual culture and spore germination assays. In detached leaf assays, leaves from endophyte infected host plants are inoculated with phytopathogens and disease symptom development parameters (leaf spot number, area/size of leaf spot) are used to characterise the antifungal activity [25,95]. These assays are complex, do not account for direct interaction of the pathogen with the endophyte [11] and depend on tiller age, endophyte incidence, concentration of pathogen inoculum, and environmental
conditions, which may lead to inconsistent results [92]. Nonetheless, detached leaf assays can be informative when performed in addition to dual culture assays.

There are a few instances where pot or field trials were conducted to detect endophyte mediated disease resistance in grasses [91,95,103–109]. Most studies investigated wild type endophyte strains, and others did not provide information on strain details. These in planta assays have confirmed that *Epichloë* endophytes improve host plant disease resistance. While some studies focused on measuring disease severity parameters (lesion numbers, lesion size) [91,110] other studies included leaf senescence characteristics (chlorophyll content, superoxide dismutase, peroxidase, catalase, and ascorbate peroxidase) to further understand the mechanism of disease resistance [95,103]. Conducting field and pot trials in quarantine is expensive and increases risk of outbreaks.

6.3. *Antifungal Metabolite Isolation and Characterisation*

While most studies have focused on detecting antifungal activity, a few characterised the mode of action and identify compounds responsible. Some studies provide evidence that *Epichloë* strains produce antifungal molecules in vitro liquid cultures and secrete to the culture media [37,97,111,112]. In these studies, the filtrate containing the fungal secretome can be extracted and bioassays performed to identify antifungal activity [37,111,112]. Three studies have extracted the secretome/culture filtrate using different solvent systems and tested their activity against selected grass pathogens [37,112,113]. These confirmed the ability of *Epichloë* strains to produce antifungal molecules in vitro even when not in contact with the pathogens. This characteristic is important to identify and isolate bioactive compounds (Figure 2c).

Historically, antifungal metabolites have been isolated from a few sexual *Epichloë* strains (Table 3), while antifungal metabolites produced by strains of asexual (endophytic) *Epichloë* species that are utilised by pasture and turf industries remain to be discovered. While most metabolites are isolated from in vitro cultures, others have been isolated from infected plants (Table 3 and Figure 3). Early studies identified a series of antifungal compounds from *Epichloë typhina* isolated from timothy grass (*Phleum pratense*) [114,115]. Yue et al. (2000) tested antifungal activity of three fractions of aqueous extracts from a range of *Epichloë* species and confirmed their antifungal activity against *Cryphonectria parasitica*, the causal phytopathogen of chestnut blight, using thin layer chromatography assay [112]. Subsequently, they used *E. festucae* (BM7, M. D. Richardson) from *Festuca rubra* to isolate six antifungal metabolites and confirmed their activity against *C. parasitica, Lactisaria fusiformis, Magnaporthe poae*, and *Rhizoctonia solani* using Potato Dextrose Agar (PDA) plate based assays [112]. Nuclear Magnetic Resonance (NMR) and Gas Chromatography-Mass Spectrometry (GC-MS) data were acquired to fully characterise the isolated compounds. Tian et al. (2017) isolated antifungal protein *Efe-AfpA*, active against *Sclerotinia homoeocarpa* [105]. They used *E. festucae* Rose City isolate (*E. festucae* RC) in association with *Festuca rubra* subsp. *rubra* (strong creeping red fescue) to isolate apoplastic proteins. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was used to isolate the proteins. The peptide sequence was determined by using Liquid Chromatography tandem Mass Spectrometry (LC/MS/MS2). Bioactivity of the isolated protein was confirmed using a PDA plate-based assay [105]. Purev et al. (2020) isolated antifungal ε-poly-L-lysines encoded by the *VibA* gene from *E. festucae* strain E437. They used NMR and MALDI-TOF MS to identify the molecule and determine the structure. Disk diffusion assays confirmed the antifungal activity against *Drechslera erythrospila* and *Phytophthora capsica* [116]. Structures of some of these antifungal metabolites are shown in Figure 3. A recent study by Fernando et al. (2021) confirmed currently known *Epichloë*-derived antimammalian and insecticidal alkaloids and their intermediates (peramine, *n*-formylloline, *n*-acetylloline, lolitrem B, epoxyjanthitretn I, paxiline, terpendole E, terpendole C, ergovaline) are not responsible for the antifungal activity observed by *Epichloë* endophytes [22].
ergovaline) are not responsible for the antifungal activity observed by Epichloë endophytes [22].

Figure 3. Structures of the fungitoxic compounds isolated from sexual Epichloë species that are listed and referenced in Table 3.
6.4. Untargeted Metabolite Annotation for Antifungal Compound Detection

*Epichloë* endophytes and their host plants are complex and sophisticated multicellular organisms, thus it is not unexpected that the metabolic profile of endophytes in planta is vastly different and more complex than endophytes in vitro [36,37]. In endophyte-infected plants, both the host plant and endophyte metabolome trigger secondary metabolite biosynthesis machinery that would not be otherwise observed individually [117]. It is important to study the metabolic profiles of these endophyte strains both in vitro, in planta and upon infection with diseases to understand the complex biological process involved in endophyte mediated disease resistance (Figure 2d).

With recent whole-genome sequencing strategies revealing that the number of genes encoding the biosynthetic enzymes in various fungi and bacteria are undoubtedly greater than the known secondary metabolites of these microorganisms, it is highly likely that most endophytes might actually express only a subset of their biosynthetic genes under standard in vitro laboratory conditions, such that only a minor portion of their actual biosynthetic potential is harnessed [118,119]. Thus, selection of the most appropriate method for metabolic profiling and isolation of metabolites, as well as consideration of the environmental conditions the endophytes are grown in, is important to identify the most agronomically important endophytes in the ecosystem.

Metabolomics is the study of metabolite profiles of a cell, tissue or an organism under given conditions [120]. Investigation of metabolic profiles of endophytes in response to a pathogen infection is the first step in understanding the mode of action and enabling their use efficiently against pathogens. Metabolome analysis may entail either a targeted analysis of a certain class of metabolite, or total (untargeted) metabolite profiling of a given sample. Preliminary screening of the total metabolome provides a view on overall performance, whereas targeted bioassay-guided isolations provide more specific details about metabolites and their potential uses [98]. It is important to realise that during the isolation procedure, bioactivity can disappear due to degradation of the bioactive compounds in the extract. Furthermore, the activity can be diluted due to inadequate chromatographic separation and poor fractionation, thus reducing the effective concentration [98].

The most widely used techniques for microbial extracts are High Performance Liquid Chromatography (HPLC), Gas Chromatography (GC), Mass Spectrometry (MS) and Nuclear Magnetic Resonance (NMR) [120]. The more advanced methods couple high resolution mass spectrometers to other analytical techniques, such as chromatography or collision-induced fragmentation, to obtain precision and structural information of compounds in a short period of time with less effort, for example LC-MS, MS2 and GC-MS [120,121]. A qualitative, as well as quantitative, understanding of microbial metabolites requires knowledge of both extracellular and intracellular metabolites [121–123]. Hence the microbes, as well as the media they are grown on, are analysed for metabolites. To identify these metabolites, it may be necessary to extract the microbial metabolites into a solvent that is suitable for the specific technique. To extract microbial metabolites, many techniques can be used and there is also a range of possible solvents available depending on the polarity of the metabolites of interest [123].

Pini et al. describes metabolic footprint analysis as the global identification and quantification of the metabolites present in the spent culture medium of microbial cells using different analytical techniques. Both extracellular as well as intracellular metabolites are specific to a time point under a certain set of environmental conditions while the microbes are growing. Capturing these timepoints is important to conduct a detailed descriptive study; thus, quenching (stopping further biological activity) becomes an important step in microbial metabolite analysis. Depending on the aims, the study could be dynamic (gathering data on microbial growth at multiple time points) or time resolved (on a single time point) [124–126]. Selection of an appropriate extraction solvent is important because it affects the yield and polarity of metabolites extracted. Thorough metabolic profiling can unravel the potential of a microorganism (biocontrol agent, natural product uses) and complement genomic studies of the organism.
Untargeted metabolomic study approaches have been used to understand the metabolic potential of *Epichloë* strains relating to antifeeding (insect/invertebrate) activity [127–129]. Other studies have investigated the effect of endophytes on the root exudate metabolome [130] and in response to different field nutrient or environmental conditions [128,131]. Untargeted metabolite fingerprinting coupled to spectral data analysis software could be the solution to understanding the complex process of endophyte-host plant-phytopathogen relationships and the metabolic response or production of bioactive metabolites in the presence of a pathogen (Figure 2d). Green et al. (2020) studied the *Lolium perenne* apoplast to identify *Epichloë festucae*-derived novel metabolites [73] and recently Fernando et al. (2021) conducted bioassay guided extraction of antifungal metabolites and metabolite annotation of bioactive extracts from *Epichloë* strains [36].

The analytical tools used to detect metabolites in these studies include ultra-high performance liquid chromatography systems (UHPLC), photodiode array detector, Q Exactive Plus high-resolution mass spectrometer (QE-MS) and high-resolution orthogonal time-of-flight MS (e.g., HRqTOF-MS). There are many data mining software platforms available to obtain a list of metabolites or features including vendor specific software such as MARKERLYNX XS for MASSLYNX v.4.1. (Waters, MA, USA), or vendor neutral software such as Refiner MS and Analyst modules of Genedata Expressionist® (Genedata, Basel, Switzerland). Metabolites can be annotated based on accurate mass information and their identity defined by database searches. Green et al. (2020) used databases (KEGG, BioCyc and in-house databases) (http://www.kegg.jp) (http://biocyc.org) to annotate metabolites and identified a novel amino acid glycoside, a set of *Epichloë* cyclins, peramine, and putatively identified two peptides, a Kojibiose-related metabolite (322.0643 Da), N-(hydroxypentyl) acetamide (145.1101 Da), and a compound with an accurate mass or m/z of 471.1952 Da [73]. Fernando et al. annotated metabolites from two bioactive strains of asexual *Epichloë* species and used them as biomarkers for detection in planta. Bioassay guided fractionation, followed by metabolite analysis, identified 61 “known unknown” prospective antifungal metabolites (out of more than 20,000 in planta) that, either singly or in combination, are responsible for the observed bioactivity. The workflow developed in this study allows testing of endophyte bioactivity while ensuring that the metabolite is expressed in planta and so useful for field deployment (Figure 2).

These studies show how metabolite fingerprinting can be used for bioprospecting antifungal metabolites from *Epichloë* endophytes. While isolation and full chemical characterisation is still necessary to understand the role and mode of action of the bioactive metabolites, these robust novel methods help visualise the potential of novel *Epichloë* strains as biocontrol agents without conducting field or pot trials. Based on this, we propose a methodology of exhaustive testing and analysis to identify biologically active compounds (Figure 2).

### 6.5. Qualitative and Quantitative Confirmation of Antifungal Metabolites in Planta

It is a common practice to conduct routine alkaloid testing for the ‘known known’ *Epichloë*-derived toxic alkaloids before field deployment of novel associations [26,27,34]. Comprehensively characterized and purified compounds can be applied in routine diagnostics for the presence and abundance of *Epichloë*-derived antifungal metabolites in planta (Figure 2e) [36]. Availability of well tested antifungal compound standards, or availability of isolation methods through scientific publication, enables further investigation of abundance of antifungal compounds in response to environmental conditions, such as disease challenge by phytopathogens, and will confirm their role in improving host plant disease resistance.
| Fungitoxic Compound | Chemical Characteristics | Chemical Formula | \( m/z \) or Molecular Weight | Epichloë sp. | Host Grass | Source Material | Tested Pathogens | Reference |
|---------------------|--------------------------|------------------|-----------------------------|--------------|------------|---------------|------------------|-----------|
| Indole-3-acetic acid| IAA derivative           | C\(_{10}\)H\(_9\)NO\(_2\) | 176.0667                    | E. festucae  | Festuca rubra | purified mycelia | Lactisaria fusiformis, Magnaportha poae, Rhizoctonia solani | [112]    |
| Indole-3-ethanol    | IAA derivative           | C\(_{10}\)H\(_{11}\)NO | 162.0874                    | E. festucae  | Festuca rubra | purified mycelia | L. fusiformis, M. poae, R. solani | [112]    |
| Methylindole-3-carboxylate | IAA derivative        | C\(_{10}\)H\(_9\)NO\(_2\) | 175.0594                    | E. festucae  | Festuca rubra | purified mycelia | L. fusiformis, M. poae, R. solani | [112]    |
| Indole-3-carboxaldehyde | IAA derivative         | C\(_{9}\)H\(_7\)NO | 159.0684                    | E. festucae  | Festuca rubra | purified mycelia | L. fusiformis, M. poae, R. solani | [112]    |
| Cyclonerodiol       | sesquiterpinoid          | C\(_{15}\)H\(_{28}\)O\(_2\) | 165.0507                    | E. festucae  | Festuca rubra | purified mycelia | L. fusiformis, M. poae, R. solani | [112]    |
| Chokol A             | sesquiterpinoid          | C\(_{12}\)H\(_{25}\)O\(_2\) | 199.1693 239.2006 239.2006 271.2288 241.1798 185.1536 | E. typhina   | Phleum pratense | Epichloë infected plant material (choke) | Cladosporium herbarum, Cladosporium phlei | [114,132] |
| Chokol B             |                          | C\(_{15}\)H\(_{26}\)O\(_2\) |  | E. sylvatica | Sylvesticum | Unfertilized stromata and unfertilized stromata head space | Stagonospora nodorum, Mycosphaerella graminicola | [102]    |
| Chokol C             |                          | C\(_{15}\)H\(_{26}\)O\(_2\) |  | E. clarkii  | Sylvesticum  | Unfertilized stromata and unfertilized stromata head space | Stagonospora nodorum, Mycosphaerella graminicola | [102]    |
| Chokol D             |                          | C\(_{15}\)H\(_{26}\)O\(_2\) |  |  | Holcus lanatus | Unfertilized stromata and unfertilized stromata head space | Stagonospora nodorum, Mycosphaerella graminicola | [102]    |
| Chokol E             |                          | C\(_{15}\)H\(_{25}\)O\(_3\) |  |  |  | Unfertilized stromata and unfertilized stromata head space | Stagonospora nodorum, Mycosphaerella graminicola | [102]    |
| Chokol F             |                          | C\(_{14}\)H\(_{24}\)O\(_3\) |  |  |  | Unfertilized stromata and unfertilized stromata head space | Stagonospora nodorum, Mycosphaerella graminicola | [102]    |
| Chokol G             |                          | C\(_{11}\)H\(_{20}\)O\(_2\) |  |  |  | Unfertilized stromata and unfertilized stromata head space | Stagonospora nodorum, Mycosphaerella graminicola | [102]    |
| Chokol K             | sesquiterpinoid          | C\(_{15}\)H\(_{26}\)O | 1 222.1984 [M-H\(_2\)O\(^-\)] = 204 | E. sylvatica | Sylvesticum | Holcus lanatus | Unfertilized stromata and unfertilized stromata head space | Stagonospora nodorum, Mycosphaerella graminicola | [102]    |
| N,N-diacetamide      | diactamide               | C\(_{4}\)H\(_7\)NO\(_2\) | 102.0610                     | E. festucae  | Festuca rubra | Epichloë infected plant material (leaves) | L. fusiformis, M. poae, R. solani | [112]    |
| Gamahonolide A       | gamahonolide             | C\(_{12}\)H\(_{23}\) | 213.1502 234.1952 222.0881 | E. typhina   | Phleum pratense | Epichloë infected plant material (choke) | [133]    |
Table 3. Cont.

| Fungitoxic Compound | Chemical Characteristics | Chemical Formula | m/z or Molecular Weight | Epichloë sp. | Host Grass | Source Material | Tested Pathogens | Reference |
|---------------------|--------------------------|------------------|-------------------------|--------------|------------|----------------|----------------|----------|
| 5-hydroxy-4-phenyl-2(5H)-furanone | | C_{12}H_{10}O_{3} | 177.0546 | E. typhina | Phleum pratense | Epichloë infected plant material (choke) | | [133] |
| Trans-p-coumaric acid | Cis-p-coumaric acid | | | | | | | |
| p-hydroxybenzoic acid | p-hydroxyphenylacetic acid | | | | | | | |
| Tyrosol | phenolic acid derivatives | C_{9}H_{8}O_{3} | C_{9}H_{8}O_{3} | C_{7}H_{6}O_{3} | C_{8}H_{10}O_{2} | | | |
| | | 164.1580 164.1580 139.1220 153.0507 168.0980 | E. typhina | Phleum pratense | Epichloë infected plant material (choke) | C. phlei | | [134] |
| Epichlicin | Cyclic peptide | C_{48}H_{74}N_{12}O_{14} | ^{3} [M+Na+17]^+ = 1082 | E. typhina | Phleum pretense | purified mycelia | C. phlei | [135] |
| Fatty acids | C-18 and C-19 fatty acid | C_{18}H_{32}O_{3} | C_{19}H_{34}O_{3} | C_{19}H_{36}O_{3} | C_{19}H_{36}O_{3} | | | |
| | | 297 | 311 | 312 | 312 | E. typhina | Phleum pratense | Epichloë infected plant material (choke) | C. herbarum, C. phlei | [115] |
| Efe-AfpA | protein | 55 amino acids | 6278 Da | E. festucae | Festuca rubra subsp. rubra | Epichloë infected plant material (tiller) | Sclerotinia homoeocarpa | Alternaria alternata, Bipolaris sorokiniana, Fusarium avenaceum Curvularia lunata | [105,136] |
| Cyclosporin T | peptides | C_{61}H_{109}N_{11}O_{12} (11 amino acids) | 1188.6 g/mol (2 MW) | E. bromicola | Elymus tangutorum | purified mycelia | | [136] |
| ε-poly-L-lysines | peptides | 28–34 lysine sub-units | E. festucae | Festuca pulchella | purified mycelia | Drechslera crythropsila Phytophthora capsici | | [116] |

^{1} m/z when ionised to [M-H_{2}O]^-, ^{2} Molecular weight indicated in g/mol, ^{3} m/z when ionised to [M+Na+17]^+. 
7. Future Directions

In the past 25 years, the potential of asexual Epichloë sp. to provide bioprotection from fungal pathogens to the host grass has been noted frequently but not pursued. Wildtype strains have been shown to improve disease control in pot trials and field scenarios, often when outbreaks have occurred rather than by design. There is a strong market globally for novel, animal safe, endophyte strains that provide insect and disease control. Some of these strains exhibit strong antifungal activity against phytopathogens, as observed using in vitro bioassays. With the discovery of novel endophytes with favourable ‘known known’ alkaloid profiles, and significant advances in high-throughput analytical techniques and data analysis, the opportunity now arises to investigate endophyte-mediated disease resistance. This knowledge can then be applied to select superior strains for the pasture and turf industries, improving animal health and host grass performance through enhanced disease resistance. It is anticipated that the outcomes of these studies would expand the current screening methods for Epichloë sp. strains to include bioprotective compounds and their respective genes.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/microorganisms10010064/s1, Supplementary Table S1: List of peer reviewed scientific reports in the last 25 years (1996–2021) with the terms “Epichloë” OR “Neotyphodium” AND “fungitoxic” OR “antifungal” in the title, abstract, or keywords.

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