Structure and Biosynthesis of Fungal Secondary Metabolites

Studies of the Root Rot Pathogen
Heterobasidion annosum s.l. and the Biocontrol Fungus Phlebiopsis gigantea

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Structure and Biosynthesis of Fungal Secondary Metabolites. Studies of the Root Rot Pathogen *Heterobasidion annosum* s.l. and the Biocontrol Fungus *Phlebiopsis gigantea*.

**Abstract**

The root rot pathogen *Heterobasidion annosum* s.l., i.e. *H. abietinum*, *H. parviporum*, *H. annosum* s.s., *H. irregulare* and *H. occidentale*, and the biocontrol fungus *Phlebiopsis gigantea* were investigated regarding their secondary metabolites.

Thirty-three compounds, in total, were identified from *H. annosum* s.l. by HRMS and NMR, including six new fomannosin related sesquiterpenes (illudolone A and B, illudolactone A and B and deoxyfomannosin A and B), one new fomajorin-type compound and seven previously unreported natural products with fomannoxin related structures. The new fomannosin related compounds were proposed to be intermediates in the biosynthesis of the known phytoxin fomannosin.

Fomannoxin is a benzohydrofuran that previously has been suggested to be involved in the pathogenicity of *H. annosum* s.l. The biosynthesis of fomannoxin was investigated through an isotopic enrichment study utilizing [1-13C]glucose as metabolic tracer. The results showed that fomannoxin is produced by a combination of the MVA pathway and the shikimic acid pathway.

The secondary metabolite production of the species within *H. annosum* s.l. was analyzed by LC-HRMS. Subsequent principal component analysis showed that the samples from the five species grouped in accord with the respective species, preferred host and previously established phylogeny.

The biocontrol fungus *P. gigantea* is used to prevent root rot caused by *H. annosum* s.l. Its mode of action was studied by investigating the production of secondary metabolites. Five secondary metabolites were isolated and identified by HRMS and NMR, out of which three were new compounds (phlebiopsin A-C), one was a new natural product (methyl-terfestatin A) and one was a known compound (o-orsellinaldehyde). Only o-orsellinaldehyde showed growth inhibiting activity against *H. occidentale*.

**Keywords:** *Heterobasidion annosum* s.l., *Phlebiopsis gigantea*, secondary metabolites, biosynthesis, root rot, biocontrol, MS, NMR, HPLC.

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Dedication

Till Therese

Det är bättre att fråga en gång för mycket än att vara dum hela livet.
Jan-Ove Hansson
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References

Acknowledgements
List of Publications

This thesis is based on the work contained in the following papers, referred to by Roman numerals in the text:


III Hansson, D., Wubshet, S. G., Olson, Å., Karlsson, M., Staerk, D. and Broberg, A. Secondary metabolite comparison of the species within the Heterobasidion annosum s.l. complex (manuscript).


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Paper not included in the thesis:

The contribution of David Hansson to the papers included in this thesis was as follows:

I Planning the work together with the co-authors. All chemical work (except the HRMS analysis) and writing the majority of the chemical related part. Interpretation of chemical data was made together with Anders Broberg. Overall conclusions were made together with all co-authors.

II Planning the work together with the co-authors. All chemical work (except the HRMS analysis) and writing the majority of the chemical related part. Interpretation of chemical data was made together with Anders Broberg. Overall conclusions were made together with all co-authors.

III Planning the work together with the co-authors. Did the LC-HRMS analysis but the principal component analysis and interpretation of these data was done together with Anders Broberg. Isolation and NMR analysis as well as interpretation of these data were done together with Sileshi G. Wubshet and Anders Broberg. Writing much of the chemical related part together with Anders Broberg and Sileshi G. Wubshet. Overall conclusions were made together with Anders Broberg, Magnus Karlsson and Åke Olson.

IV Planning the work together with the co-authors. Majority of the chemical work (except GC-MS analysis), the interpretation of data and the writing.
Abbreviations

ACP    acyl carrier protein
APCI   atmospheric-pressure chemical ionization
CDCl₃  deuterated chloroform
CoA    coenzyme A
COSY   correlation spectroscopy
DMAPP  dimethylallyl diphosphate
E4P    erythrose-4-phosphate
EI     electron ionization
Enz    enzyme
ESI    electrospray ionization
FAB    fast atom bombardment
FID    free induction delay
FPP    farnesyl diphosphate
GC     gas chromatography
GFPP   geranyl-farnesyl diphosphate
GGPP   geranylgeranyl diphosphate
GPP    geranyl diphosphate
HMBC   heteronuclear multiple bond correlation
HMG    3-hydroxy-3-methylglutaryl
HPLC   high performance liquid chromatography
HR     high resolution
HSQC   heteronuclear single quantum coherence
IPP    isopentenyl diphosphate
KS     ketoacyl synthase
LLE    liquid-liquid extraction
m/z    mass-to-charge ratio
MALDI  matrix-assisted laser desorption
MeCN   acetonitrile
MEP    methylerthyritol phosphate
MS  
MVA  
NMR  
NOE  
NOESY  
NRPS  
P  
PCA  
PEP  
PKS  
PP  
ppm  
Q  
QqQ  
ROESY  
RP  
s.l.  
s.s.  
SPE  
TMS  
TOF  
UHPLC  
t_R  
δ
1 Introduction

1.1 Fungal secondary metabolites

The science of natural product chemistry covers the study of all different compounds produced by living organisms, including polymeric macromolecules such as nucleic acids, proteins, and carbohydrates, as well as, low molecular weight compounds. The latter includes compounds called secondary metabolites and is what the present thesis focuses on. Secondary metabolites are loosely defined as organic compounds that are not directly involved in primary metabolic processes such as cell growth, cell division, cell respiration or photosynthesis. Furthermore, secondary metabolites are derived from a few common biosynthetic pathways which branch off the primary metabolic pathways and are often produced as families of related compounds, often specific for a group of organisms (Dewick, 2009; Hartmann, 2007; Hanson, 2003).

Many organisms produce large numbers of secondary metabolites and the complexity and diversity is sometimes astonishing. Secondary metabolites have had great impact on society for centuries in traditional medicine and in modern times, as pharmaceuticals, fragrances in cosmetics, flavouring in foods and drinks, agrochemicals, etc. But what is the biological reason for producing these compounds? Why do organisms put huge amounts of energy on producing them?

Secondary metabolites perform many different functions for their producer including functions as (1) volatile alarm pheromones, (2) sex attractants in many insects, (3) defense against predators and (4) weapons for establishment in their respective ecological niche, just to name a few. However, many of the biological functions are still to be determined.

Fungi are a rich source of secondary metabolites and have been of interest for humans for thousands of years. However, it was not until when Alexander
Fleming discovered the antibiotic penicillin (penicillin F) 1929 and its development into a medicine by Florey and Chain in the 1940s (penicillin G), that the attention to fungal secondary metabolites became widespread. Since then, huge research efforts have been made and resulted in thousands of characterized compounds with a range of different biological activities. For example: a literature study on 1500 fungal secondary metabolites isolated and characterized between 1993 and 2001 showed that more than half of these compounds had antibacterial, antifungal or antitumor activity (Selim et al., 2012; Keller et al., 2005). Besides these effects, fungal secondary metabolites are also suggested to be crucial in the disease-causing interaction with plants. These fungi, causing disease on plants, are called pathogens and their secondary metabolites often secreted during infection are called mycotoxins, phytotoxins or more generally toxins (Meinwald, 2011; Fox & Howlett, 2008).

![Figure 1. The structure of penicillin F (left) and penicillin G (right).](image)

1.2 Root rot and *Heterobasidion annosum* s.l.

The fungi within the species complex *Heterobasidion annosum* sensu lato (s.l.) are aggressive plant pathogens that cause the disease symptom called root rot, which is a serious problem for forestry. They affect mainly conifers causing growth reductions, decay of the interior parts of the stem which leads to devaluation of timber, increased risk of wind-throw and in the worst case, death of trees. The economic losses to forestry as a result of *H. annosum* s.l. infections are high and have been estimated to be approximately 54 and 790 million € in Sweden and Europe, respectively (Woodward et al., 1998; Hellgren & Stenlid, 1995).

*H. annosum* s.l. has since its discovery in 1821 by Elias Fries, been known by many different names, from *Polyporus annosus* and *Fomes annosum* to names connected to preferred host plant. At the present time, five species have been identified within *H. annosum* s.l. 1) *H. abietinum* Niemelä & Korhonen, 2) *H. parviporum* Niemelä & Korhonen, 3) *H. annosum* sensu strictro (s.s.), (Fr.) Bref., 4) *H. irregulare* Otrosina & Garbelotto, 5) *H. occidentale* Otrosina & Garbelotto (Otrosina & Garbelotto, 2010; Niemelä & Korhonen, 1998).
The species within the species complex have different distributions and host preferences. *H. irregulare* and *H. occidentale* are North American species whereas the other three are appearing in Europe. Generally, *H. parviporum*, *H. abietinum* and *H. occidentale* infect spruce and *H. annosum* s.s. and *H. irregulare* pine. However, overlap does occur and other tree species can be affected as well (Otrosina & Garbelotto, 2010).

*H. annosum* s.l. infections are initially spread by airborne basidiospores to freshly cut stumps or wounds on the roots or stem. The fresh surface is subsequently colonized and the infection spreads rapidly down the stem to the root system. Once the infection has reached the root system it can, by root to root contacts infect nearby healthy trees. The infection grows subsequently up through the trunk and can reach several meters high (Stenlid & Redfern, 1998).

### 1.3 Biocontrol and *Phlebiopsis gigantea*

Major efforts have been made to prevent *H. annosum* s.l. root rot resulting in many different control methods including biological, chemical and silvicultural methods. The biological method includes the use of other microorganisms to suppress disease and is referred to as biocontrol. The fungus *Phlebiopsis gigantea* is such a biocontrol agent used to prevent *H. annosum* s.l. infections.

*P. gigantea* is a highly competitive fungus and is, like *H. annosum* s.l., an early colonizer of freshly cut conifer wood. The ability to outcompete *H. annosum* s.l. with *P. gigantea* in pine stumps and roots was observed already in the fifties of the last century by Rishbeth (1951), but it was not until a decade later that it was used as a biocontrol in practical use. A arthroconidia-based powder formulation of *P. gigantea*, based on a Finnish strain isolated by Korhonen *et al.* (1994), was later developed into a commercially available product, Rotstop®. Rotstop® is now the most widely applied stump treatment against *H.annosum* s.l. infections in Europe, used on more than 200 000 ha annually (Menkis *et al.*, 2012; Holdenrieder & Greig, 1998).

Stump treatment with aqueous solution of urea or disodium octaborate tetrahydrate are two chemical methods that are in use, especially in Great Britain and in North America, but less frequently in Sweden. Urea acts by increasing the pH of the treated surface which can favor other competitive fungi but it is also proposed to be poisonous for *H. annosum* s.l. (Johansson *et al.*, 2002). Boron is poisonous for fungi in general but also for other organism, for example insects. It acts by blocking the growth of cells in *H. annosum* s.l. (Sturesson *et al.*, 2011).

Silvicultural methods that have been developed to reduce *H. annosum* s.l. infections are, among others, stump removal, choice of tree species to less
susceptible species, mixed stands (mix of differently susceptible species) and thinning in the winter when temperatures are below 5 °C and germination of basidiospores is poor (Cleary et al., 2013; Korhonen et al., 1998).

1.4 Aims and objectives

Sweden is one of the world’s largest exporters of sawn timber, pulp and paper and the forest is a cornerstone in Swedish industry. A severe threat against healthy tree stands are root rot caused primarily by the root rooting fungi *H. annosum* s.l. The overall aim of the present thesis was to further our understanding of interactions between *H. annosum* s.l., their host and the biocontrol fungus *P. gigantea*. As part of this work, it is important to gain knowledge about involved secondary metabolites, their structure and biosynthesis. The specific objectives of the individual projects were:

- *H. annosum* s.l. produces several bioactive secondary metabolites including the benzofuran fomannoxin and the sesquiterpene fomannosin. The objective of this study was to further investigate bioactive secondary metabolites produced by *H. occidentale*. (paper I)

- Fomannoxin is a biologically active benzohydrofuran produced by *H. annosum* s.l. and has been proposed to be involved in the fungus pathogenicity. The objective of this study was, by incorporation of $^{13}$C, to investigate the biosynthesis of fomannoxin. (paper II)

- The objective of this study was to investigate if there are differences within the *H. annosum* s.l. species complex regarding the production of secondary metabolites and if this could be related to host preferences. (paper III)

- Little is known about how the biocontrol fungus *P. gigantea* act as a biocontrol against *H. annosum* s.l. The objective of this project was to characterize *P. gigantea* regarding the production of secondary metabolites and investigate whether these compounds could contribute to its biocontrol effect against *H. annosum* s.l. (paper IV)
Secondary metabolites are biosynthesized of building blocks that are put together in various metabolic pathways. The pathways are usually named after enzymes or intermediates involved and are commonly also used to classify secondary metabolites. The diversity and complexity of the structures that these relatively few building blocks can provide is both surprising and fascinating.

This section aims to briefly explain the different pathways with focus on the pathways relevant for the compounds within this thesis. For further reading see comprehensive textbooks on the biosynthesis subject e.g. Dewick (2009) or Mann et al. (1994).

2.1 The MVA and MEP pathway: Terpenoids

Terpenoids are found in essentially all forms of life and they form a large group of secondary metabolites with more than 40 000 structures (Bohlmann & Keeling, 2008). They are built up from five-carbon segments, so called “isoprene units”, which can combine to form different classes of terpenoids: hemi- (C_5), mono- (C_10), sesqui- (C_15), di- (C_20), sester- (C_25), tri- (C_30) and tetraterpenes (C_40). The name “isoprene units” might be misleading since isoprene itself is a hemiterpene. The actual fundamental building block to all terpenes and terpenoids is isopentenyl diphosphate (IPP) and its isomer dimethylallyl diphosphate (DMAPP). Depending on the producing organism, IPP and DMAPP are biosynthesized either via the mevalonic acid (MVA) pathway or the methylerythritol phosphate (MEP) pathway. Fungi and animals have only access to the MVA pathway whereas algae and most bacteria use the MEP pathway. Plants and some bacteria have the possibility to use both pathways (Grawert et al., 2011; Walter et al., 2000; Rohmer, 1999).
The MVA pathway (Figure 2) starts by combining three acetyl-coenzyme A (acetyl-CoA) to form the six-carbon compound MVA. One acetyl-CoA binds to the enzyme and is combined with acetyl-CoA by a Claisen condensation to give acetoacetyl-CoA. A third enzyme-bound acetyl group is subsequently attached by an aldol addition and results in the formation of 3-hydroxy-3-methylglutaryl-CoA (HMG-CoA). A three step reduction, performed by HMG-CoA reductase, is followed and MVA is synthesized. MVA is finally transformed to the five-carbon unit, IPP, by a series of reactions including phosphorylation, decarboxylation and dehydration. IPP can then isomerize to DMAPP by incorporation of a proton at C-4 and stereospecific loss of another proton at C-2.

Pyruvic acid and glyceraldehyde 3-phosphate are the starting material to IPP and DMAPP via the MEP pathway (Figure 3). Deoxyxylulose 5-phosphate is formed after decarboxylation of pyruvic acid and nucleophilic attack onto the aldehyde of glyceraldehyde 3-phosphate, a process which requires the cofactor thiamine diphosphate. A subsequent rearrangement reaction then transforms deoxyxylulose phosphate into MEP. Several reaction steps follow, including two reductions, finally leading to the major end product IPP and the minor
product DMAPP. If needed, DMAPP can also be formed by isomerization of IPP as in the MVA pathway.

Figure 4. Examples of C5-C25 terpenoids and the formation of respective precursors.

IPP has, with its terminal double bond, a nucleophilic character. In contrast, DMAPP is electrophilic when diphosphate is lost. These basic differences in reactivities between IPP and DMAPP are the basis for constructing larger terpenoids. The allylic resonance stabilized carbocation formed from DMAPP
can go on to form different hemiterpenes, exemplified in Figure 4 by isoprene and methybutenol, or it can react with IPP and form, after loss of a proton, geranyl diphosphate (GPP). GPP, in turn, is the precursor of monoterpenes but can also be extended by addition of one further IPP to form farnesyl diphosphate (FPP), the precursor of sesquiterpenes. Precursors of diterpenes and sesterterpenes, geranylgeranyl diphosphate (GGPP) and geranylfarnesyl diphosphate (GFPP), respectively, are formed in analogous additions reactions of IPP to FPP and GGPP, respectively, as illustrated in Figure 4. Electrophiles as DMAPP, GPP, FPP etc. can also act as alkylating agents in the construction of meroterpenoids, i.e. other secondary metabolites that contain terpenoid moieties. Alkylation by the C5 carbon segment from DMAPP is, for example, a very common reaction to extend the diversity of aromatics in bacteria, fungi and plants (Saleh et al., 2009; Yazaki et al., 2009).

Several examples of C5-C25 terpenoids are given in Figure 4 and by looking at the structure of these molecules one can see some of all the different modifications that can be done to respective precursor, contributing to the diversity of terpenoids. From different carbon skeletons; linear, mono-, bi and multicyclic to various functional groups e.g. carboxylic acids, alcohols, aldehydes, ketones, ester etc.

Terpenoids possess many biological properties and they are widely used as flavors, fragrances, pharmaceuticals, food additives etc. For example, the monoterpene limonene has a strong smell of oranges and is apparent in considerable amounts in many citrus fruits. Citronellol, found in rose oil, is like limonene used as fragrance in many perfumes but also as insecticide (Forster-Fromme & Jendrossek, 2010; Dewick, 2009). Another monoterpene, eucalyptol, the main constituent of the Eucalyptus plant, is used as an antibacterial, anti-inflammatory, antihypertensive agent etc. (Liapi et al., 2007). The sesquiterpene illudin S, produced by the mushroom Omphalotus illudens, is an extremely toxic compound. The toxicity is proposed to be due to the presence of the strained cyclopropyl group which makes it reactive against nucleophiles and has been shown to alkylate DNA (Schobert et al., 2011). Another sesquiterpene, humulene, found in hops is an important fragrance of beer, but it is also a key intermediate in the formation of e.g. fomannosin (paper I) and protoilludanes (Abraham, 2001). Forskolin, isolated from roots of Coleus forskohlii, an Indian plant, has been used in traditional medicine and has been shown to lower blood pressure and has cardioprotective properties (Asada et al., 2012). A well-known plant hormone, gibberellic acid, first isolated from the fungus Gibberella fujikuroi, as a phytotoxin, is produced commercially and used, for example, in the production of seedless grapes (Hanson, 2003). The last example, Ophiobolin A, a sesterterpene found in the
plant pathogen *Helminthosporium maydis*, has a broad spectrum of biological activity against bacteria, fungi and nematodes (Dewick, 2009).

All different structures and properties of terpenoids are the results of modifications accomplished via various enzymatic reactions such as changes to the oxidation state of a molecule by oxidation and reduction reactions, alkylations, decarboxylations, glycosylations, rearrangements and cyclization reactions *etc*. Many of them, as for example, rearrangement reactions and cyclization reactions are often carbocation driven. The cyclization of *E,E*-FPP to humulene can serve as an example and is illustrated in Figure 5. The cyclization to the 11-membered ring system of the humulyl cation is the result of a *S_N*1 reaction, involving a carbocation intermediate formed upon loss of diphosphate from FPP. Subsequent loss of a proton leads to humulene. Humulene can then be further transformed to structurally complex bicyclic and tricyclic sesquiterpenes, with the most important pathway generating the protoilludane skeleton (Abraham, 2001) present in for example illudolone A and B, (Paper I).

![Figure 5. Cyclization of humulene.](image)

Triterpenoids and tetraterpenoids are not formed by adding further IPP building blocks to GFPP, instead pairwise combination of FPP and GGPP, respectively, give rise to these two classes of terpenoids (*Figure 6*).

Squalene, first isolated from shark liver oil, is the precursor of triterpenes and is formed, after a sequence of reactions catalyzed by squalene synthase, from two FPP molecules. Cyclization of squalene then leads, via the intermediate protosteryl cation, to lanosterol or cycloartenol. Lanosterol, formed in fungi and animals, is in turn the precursor for many triterpenoids and modified triterpenoids *e.g.* steroids and sterols. Cycloartenol has the corresponding function as lanosterol, but is formed exclusively in plants (Dewick, 2009).
Tetraterpenoids originates from lycopene which in turn is formed by condensation of two GGPP molecules (Figure 6). There is only one group of tetraterpenoids, the carotenoids, but this group includes several hundred structural variants. They are recognizable as the dominant pigments in many fruits and flowers e.g. carrots, tomatoes and red pepper fruits (Dewick, 2009).

Figure 6. The formation of tri- and tetraterpenoids.
2.2 The shikimic acid pathway: Aromatic amino acids and phenylpropanoids

The shikimic acid pathway is an important biosynthetic route in fungi, plants and other microorganisms for the production of aromatic compounds, particularly the aromatic amino acids L-phenylalanine, L-tyrosine and L-tryptophan. The shikimic acid pathway is, however, not present in animals and therefore are these amino acids classified as essential for humans. Phenylalanine and tyrosine are in turn important precursors for many alkaloids but also for phenylpropanoids, recognized by their C₆C₃ carbon skeleton and found in many structurally diverse secondary metabolites.

The initial step in the shikimic acid pathway (Figure 7) is the condensation of phosphoenolpyruvate (PEP) and erythrose 4-phosphate (E4P) to give shikimic acid by two aldol-type reactions, one intermolecular and one intramolecular, followed by elimination of water and a reduction reaction. Shikimic acid is then phosphorylated and another PEP molecule is attached to the 5-hydroxy group of shikimic acid 3-phosphate. Two sequential eliminations of phosphoric acid follow and lead to the end product of the shikimic acid pathway, namely chorismic acid (Dewick, 2009; Hanson, 2003).

Figure 7. Formation of chorismic acid via the shikimic acid pathway.

The initial step in the shikimic acid pathway (Figure 7) is the condensation of phosphoenolpyruvate (PEP) and erythrose 4-phosphate (E4P) to give shikimic acid by two aldol-type reactions, one intermolecular and one intramolecular, followed by elimination of water and a reduction reaction. Shikimic acid is then phosphorylated and another PEP molecule is attached to the 5-hydroxy group of shikimic acid 3-phosphate. Two sequential eliminations of phosphoric acid follow and lead to the end product of the shikimic acid pathway, namely chorismic acid (Dewick, 2009; Hanson, 2003).
Figure 8. Chorismic acid as an important precursor for the formation of e.g. the C₆C₃ carbon skeleton and benzoic acid derivates.

Chorismic acid is a key intermediate and a starting point in the formation of many important compounds (Figure 8). It is transformed into prephenic acid upon a Claisen rearrangement reaction. Decarboxylation, aromatization and loss of the hydroxy group of prephenic acid result in phenylpyruvic acid, which after transamination becomes phenylalanine. Tyrosine is formed from prephenic acid via 4-hydroxyphenylpyruvic acid in a similar reaction sequence, but with an additional oxidation that retains the 4-hydroxy group of prephenic acid. Elimination of ammonia by E2 mechanisms results in cinnamic acid and 4-coumaric acid from phenylalanine and tyrosine, respectively. These two compounds are precursors for phenylpropanoids, which as mentioned above, are found in many structures of secondary metabolites e.g. lignans, phenylpropenes and coumarins. They are, in addition, found in secondary metabolites that are biosynthesized by combination of building blocks, such compounds could be exemplified by styrylpyrones, flavanoids and stilbenes (Dewick, 2009; Hanson, 2003).

Many compounds, containing the C₆C₁ unit such as simple benzoic acid derivates, are originating from 4-hydroxybenzoic acid, which in turn can be formed by side-chain shortening of 4-coumaric acid. As shown in Figure 9, 4-hydroxybenzoic acid can also be formed directly from chorismic acid by
elimination of pyruvate, a reaction catalyzed by the enzyme chorismate pyruvate-lyase. This pathway is used by bacteria, however, fungi (paper II) may also use this pathway as an alternative route to 4-hydroxybenzoic acid and similar aromatic compounds (Stadthagen et al., 2005; Meganathan, 2001). Ubiquinones are example compounds that can be formed from 4-hydroxybenzoic acid. These compounds are present in almost all organisms and many of them have important functions in electron transport systems for respiration or photosynthesis but they have also additional functions such as antioxidants (Nowicka & Kruk, 2010).

The biosynthesis of the indole containing amino acid L-tryptophan is complex and involves several reaction steps. Amination of chorismic acid followed by elimination of pyruvic acid results in the intermediate anthranilic acid. The result of the sequence of reactions that follows is first the formation of the indole skeleton, then the amino acid L-serin attached to finally make the carbon skeleton of tryptophan complete. Tryptophan is an important precursor for many alkaloids containing the indole ring system but there are also examples where anthranilic acid itself functions as alkaloid precursor (Dewick, 2009; Hanson, 2003).

Figure 9. Chorismic acid as precursor in the formation of e.g. L-tryptophan and benzoic acid derivates.
2.3 The acetate pathway: Polyketides

The polyketides is a large family of secondary metabolites found in bacteria, fungi and plants. They are particularly important for fungi as the most abundant fungal secondary metabolites and many of them have important biological activities. The cholesterol lowering compound lovastatin, produced by the fungi *Monascus ruber* and *Aspergillus terreus*, serves as a good example. It works as an inhibitor of the enzyme HMG-CoA reductase which is active in the formation of MVA and was the first statin to be marketed (Dewick, 2009; Keller et al., 2005).

Polyketides are derived from poly-\(\beta\)-keto chains, formed by stepwise condensation of mainly acetyl-CoA (starter unit) and malonyl-CoA (extender unit). The reactions involved in the construction of the poly-\(\beta\)-keto chain are catalyzed by polyketide synthase (PKS), a family of multidomain enzymes or enzyme complexes. There are several types of PKSs found in different organisms but the principal of constructing the poly-\(\beta\)-keto chain are the same in all PKSs (*Figure 10*). The initial step is the loading, as thioesters, of both the starter unit and the extender unit to the respective site on the PKS. Acetyl-CoA is bound to a ketoacyl-CoA synthase (KS) domain whereas the extender unit is bound to an acyl carrier (ACP) domain. Condensation then occurs between malonyl-ACP and acetyl-KS by a Claisen-type reaction and simultaneous decarboxylation of the ACP-bound extender unit. The resulting \(\beta\)-ketothioester bound to the ACP domain can then be transferred to a KS domain and extended by another malonyl-ACP. This cycle repeats until the growing \(\beta\)-keto chain has reached the desired length. Once the chain is completed it can be folded and activated at desired positions to allow for intramolecular reactions. This is exemplified in *Figure 10* by the biosynthesis of orsellinic acid and phloracetophenone. Other more complicated aromatic compounds like the anthraquinones can be synthesized in the same manner but with a longer \(\beta\)-keto chain as precursor (Dewick, 2009; Hanson, 2003).
The carbon skeleton of macrolides, another type of polyketides characterized by large lactone rings, is formed by the same general process but with some modifications. The modifications include reductions, dehydrations, etc., of the β-carbonyl group and may take place after each condensation step and before the next chain extension (Figure 11). The carbon chain of the polyketide leading to macrolides is thus partially reduced. The biosynthesis of macrolides is exemplified in Figure 11 by zearalenone, a mycotoxin produced by e.g. several *Fusarium* species (Dewick, 2009; Elsharkawy & Abulhajj, 1987).

In addition to the variability that can be achieved through the use of the above mentioned capabilities, (number of extender units, different folding, reduction reactions etc.), the use of other starter and/or extender units than acetyl-CoA and malonyl-CoA further extend the structural diversity of polyketides. Flavanoids and stilbenes are, for example, biosynthesized from a cinnamoyl-CoA starter unit (derivied from the shikimic acid pathway) with chain extension using three malonyl-CoA units.

*Figure 10. Polyketide synthesis.*
2.4 Alkaloids, carbohydrates, fatty acids, peptides and proteins

2.4.1 Alkaloids

Alkaloids are cyclic organic compounds that contain one or more nitrogen atoms and are often basic. They have usually pronounced effects on the nervous system of humans and other animals. A few well-known examples of alkaloids are presented in Figure 12.

Figure 12. Structure of various familiar alkaloids.
Alkaloids are commonly produced from amino acids such as ornithine, lysine, tryptophan and tyrosine but other building blocks, e.g. terpenes or acetate pathway derived moieties, are also often incorporated into structures of alkaloids.

Ergolines are a group of alkaloids containing the indole ring system and have been found in several fungal genera e.g. Claviceps. These fungi are responsible for a fungal disease called ergot that affect cultivated grass such as wheat and rye and can be poisonous for humans and animals upon feeding. D-(+)-lysergic acid (Figure 13) is one of more than 50 alkaloids characterized from these fungi and are formed from tryptophan and DMAPP, originating from the shikimic acid pathway and the MVA/MEP pathway, respectively (Dewick, 2009; Schiff, 2006).

![Figure 13. Formation of lysergic acid from tryptophan.](image)

2.4.2 Carbohydrates

Carbohydrate biosynthesis and degradation are essential for all organisms and are typical components of the primary metabolism. However, some of these carbohydrates e.g. glucose, are attached to secondary metabolites in glycosides. The non-carbohydrate moiety of such a compound is known as the aglycone and may originate from one or more biosynthetic pathway e.g. the shikimic acid pathway and the acetate pathway. Fusicoccin A (Figure 14), a fungal secondary metabolite from Fusicoccum amygdale, is an example of a glucopyranoside of a diterpene that has, in a recent study, been shown to have growth inhibiting effects on the deadly brain tumor, glioblastoma multiforme (Bury et al., 2013).
2.4.3 Fatty acids

Common fatty acids are mostly regarded as primary metabolites as they are important components of some vital functions in organisms. They are usually combined as esters with glycerol, forming glycerides. Triglycerides as an example, form fats and oils, depending of the individual fatty acids, and are an important source of energy upon β-oxidation.

Fatty acids are biosynthesized via the acetate pathway in a very similar way as the polyketides, although different enzymes are working. Fatty acids may also, as carbohydrates, be attached to other metabolites. One example is the 16-palmityl ester of cucurbitacin B (Figure 15), found in the fungus *Leucopaxillus gentianeus* (Clericuzio et al., 2004).

2.4.4 Peptides and proteins

Peptides and proteins consist of amino acids that are linked to form chains with different lengths, the length determine whether the product will be referred to as peptide (up to 40-50 amino acids) or protein (generally more than 40-50 amino acids). Peptides and proteins, as a group, are difficult to classify as primary or secondary metabolites since many of them are large in size and found in materials that are widely distributed and occur in many different organisms while other are small and restricted in occurrence.
Peptides and proteins are produced by either ribosomal (produced on the ribosome) or non-ribosomal biosynthesis. Non-ribosomal peptides are synthesized by multidomain, multimodular enzymes called non-ribosomal peptide synthases (NRPSs) and are, in contrast to ribosomal synthesis, not dependent of mRNA. The sequential extension in non-ribosomal peptide synthesis can be compared to and is similar to polyketide synthesis. Each module in an NRPS contains several domains and each domain has specific function e.g. recognition of amino acid, bond formation between amino acids, or release of synthesized peptide (Dewick, 2009; Keller et al., 2005).

δ-(L-α-aminoacidipyl)-L-cysteinyl-D-valine synthetase, the first identified fungal NRPS, catalyses the assembly of the L-aminoacids and the epimerization of L-valine to D-valine in the formation of the tripeptide that, after cyclization by isopenicillin N synthase, become isopenicillin N (Figure 16). Isopenicillin N, in turn, is the precursor of β-lactam antibiotics e.g. penicillin G (Keller et al., 2005; Smith et al., 1990).

![Figure 16. Biosynthesis of isopenicillin N and subsequent modification to penicillin G.](image-url)
3 Experimental

This chapter aims to briefly explain the principal of techniques and methods used during this thesis work. Examining the structure and biosynthesis of secondary metabolites involves (1) sample preparation, (2) separation and isolation, (3) structure determination and characterization.

There is also, at the end, a brief description of the isotopic labeling method used for studying the biosynthesis of fomannoxin.

3.1 Sample preparation

Sample preparation isolates analytes of interest and simultaneous reduces impurities, from a matrix that is sometimes both large and complex. The aim of sample preparation is a purified and concentrated sample ready for further analysis.

3.1.1 Solvent extraction

Solvent extraction is one of the oldest and most common extraction methods. It was used for extracting semi-solid samples in this work but is generally applicable for both solid and liquid samples. If a liquid is to be extracted, the extraction method is commonly referred to as liquid-liquid extraction (LLE). Before extraction of solids, the samples are often homogenized by chopping or pulverizing. The homogenized solid or liquid sample is then repeatedly extracted with a suitable organic solvent. Following extraction, the extract is commonly filtrated or centrifuged and concentrated by rotary evaporation (Zhang et al., 2012; Novakova & Vlckova, 2009).

Drawbacks of solvent extraction is that this method is relative time-consuming and large amounts of environmentally harmful organic solvents are used. However, alternative extraction methods for liquids or dissolved solids have been developed e.g. solid phase extraction.
3.1.2 Solid phase extraction

Solid phase extraction (SPE) is nowadays the most popular sample preparation method and has been used extensively in this thesis for e.g. sample concentration and clean up. The general SPE procedure is simple: a sample commonly containing lipophilic analytes in an aqueous solution, are subjected to a short column which contains the solid phase. The column has previously been activated and equilibrated by e.g. methanol and water, respectively. Activation solvates the dry solid phase and makes it available for the analytes whereas the equilibration aims to prevent losses upon sample loading. The lipophilic analytes are adsorbed to the solid phase whereas the impurities and solvent are passed through. The column is then washed, with water, in order to remove impurities and the analytes are then finally desorbed and eluted by adding a small amount of an organic solvent e.g. methanol or acetonitrile (Zhang et al., 2012).

There are several different solid phases, also referred to as packing material or sorbents, available and the choice strongly depends on the analytes and their properties. Common SPE sorbents include alkyl-bonded silica particles such as C_{18} (Figure 17) and C_{8}, ion-exchange materials, polymeric materials and graphitized carbon. Columns packed with C_{18} or some polymeric materials such as Oasis HLB are good choices for compounds with a wide lipophilic range existing in aqueous solution. Furthermore, these packing materials are available in columns with several different formats, from bigger 10 g (C_{18}) columns to small cartridges containing only a few mg (Buszewski & Szultka, 2012; Novakova & Vlckova, 2009).

![Figure 17. Schematic illustration of a C_{18} silica particle.](image)

Small cartridges are well suited for preparation of samples to be analyzed by NMR directly after elution, since only small volume of solvent, typically 500-600 µl, are necessary for elution. There are, however, some modifications that should be considered in the preparation of NMR samples. Firstly, the washing step, usually performed with water, is preferably replaced with deuterated water and secondly the elution must be done with deuterated solvent. In this way, SPE is helpful when dealing with volatile compounds that otherwise might be lost upon concentration by evaporation. It may also be advantageous since some compounds are difficult to redissolve after drying.
3.2 Separation and isolation

A fungus may produce tens or perhaps hundreds of secondary metabolites in different amounts and with various physical and chemical properties. To facilitate structure elucidation and characterization of individual secondary metabolites, the compounds need to be isolated. Chromatography has become an excellent tool for such a task and several chromatographic methods have been developed since it first was described about hundred years ago by the Russian botanist Michail Tswett (Ettre, 2003). High performance liquid chromatography (HPLC) and gas chromatography (GC) are the two most important chromatographic techniques for low molecular weight compounds. GC uses an inert gas e.g. helium as mobile phase whereas HPLC uses a liquid. Chromatography can be performed either for analytical purposes or for preparative purposes. Analytic chromatography aims to separate, identify and/or quantify components of a sample and can be performed by both GC and HPLC. Preparative chromatography, on the other hand, aims to not only separate but also physically isolate components by collecting them as they are eluted from the column and cannot be performed by GC. In addition, samples to be analyzed by GC must be volatile, a characteristic lacking in many components of a fungal extract. Therefore, HPLC has been the major chromatographic technique used in this thesis.

3.2.1 High performance liquid chromatography

Among the various HPLC techniques; normal phase HPLC, reversed phase HPLC (RP-HPLC), ion exchange and size exclusion HPLC, RP-HPLC is by far the most common. RP-HPLC is also the techniques that has been used during this thesis work and is from now on referred to as simply HPLC.

A typical HPLC system consist of a high-pressure pump system, an automated sample injector, a separation column, a detector and a computer for displaying the results and controlling the system. Additional common components in a modern HPLC system include solvent degasser, pre-column and column oven. An injected sample is transported by the mobile phase to the column where the actual separation begins. The column is packed with a non-polar stationary phase through which a moderately polar mobile phase is pumped. The compounds are separated because they interact with the stationary phase and are retained differently. Compounds, poorly retained, travel faster through the column and are said to have shorter retention time ($t_R$) than compounds that are strongly retained by the stationary phase.

There are several factors that influence the retention and thus the separation e.g. selected stationary phase, mobile phase modifier, mobile phase
composition, temperature, and flow rate. The most commonly used stationary phases are based on microporous silica particles (typically 3-5 μm in diameter) to which alkyl chains e.g. C₁₈, C₈ or substituted alkyl chains e.g. (CH₂)$_₃$NH₂, (CH₂)CN are attached. Water miscible organic solvents such as methanol and acetonitrile are typical mobile phase modifiers that are used in different composition with water. The composition of the mobile phase can either be kept constant as in isocratic elution or it can be continuous changed during the chromatography to increase the eluent strength. The latter method, called gradient elution, is often a good starting point for complex samples and a typical gradient profile starts at 5% organic modifier and progress to 95% within 10-40 minutes. An optimized isocratic elution method, on the other hand, is sometimes a good alternative for compounds with similar $t_R$ and difficult to separate with a gradient method. For further reading of HPLC, see for e.g. Miller (2009).

Fast analysis and high efficiency are desired in HPLC and have driven the development of, mainly, columns and pump systems. Smaller columns with particle size less than 2 μm are nowadays available that permit separation of rather complex mixtures in just a few minutes. However, these columns need pump systems able to perform at higher pressure and these new systems, called ultra high performance LC (UHPLC), are rather expensive compared to regular HPLC systems. Superficially porous silica particle columns also referred to as fused core columns are a new type of columns with a solid inner core (typically 1.6-1.9 μm) covered by a porous silica shell (typically 0.5 μm). Columns packed with fused core material create a much lower back-pressure and allow high flow rates to be maintained resulting in shorter analysis time without losing chromatographic quality. Furthermore, these columns can be used by regular HPLC systems and could be an alternative to UHPLC (Ali et al., 2012; Novakova & Vlckova, 2009).

### 3.3 Structure determination and characterization

#### 3.3.1 Mass spectrometry

Mass spectrometry (MS) can be divided into three basic steps: (1) ionization, (2) mass analysis and (3) detection.

Ionization takes place in the ion source and aims to convert analytes into gas-phas ions. There are a number of different methods and the choice depends on the sample phase e.g. liquid or solid and the nature of the analytes. Common ionization methods includes electrospray ionization (ESI), electron ionization (EI), fast atom bombardment (FAB), matrix-assisted laser desorption ionization (MALDI), atmospheric-pressure chemical ionization (APCI) among
others. One of the most common methods, particularly in combination with HPLC is the ESI method. In the following, this method is described in further details.

The ESI process (Figure 18) starts at the end of a capillary, which is fixed to the ion source chamber. The capillary in turn can be connected to e.g. the HPLC outflow. The tip of the steel capillary is held at a high electric potential with respect to a counter electrode sitting at the inlet of a mass analyzer. The potential difference leads to separation of ions in the solution. Cations, are in positive mode, accumulated at the tip of the capillary and are attracted by the negatively charged counter electrode. If the electric force is strong enough, a cone is formed that eventually breaks and forms highly charged droplets. Nitrogen gas that is continuously pumped into the heated ion source allows these droplets to concentrate resulting in smaller droplets with higher and higher charge density. When the surface tension is overcome droplet fission occurs and concentration of these newly formed droplets starts. This sequence repeats until gas phase ions are formed that subsequently enter the mass analyzer.

Figure 18. Schematic illustration of the ESI process.

The ions entering the mass analyzer are separated and mass analyzed on the basis of their mass-to-charge ratio (m/z). Magnetic and/or electric fields are used to control the motion of the ions and different instruments have their own method and design to accomplish this. Common mass analyzers among others are quadrupole (Q), time-of-flight (TOF) and ion-trap. Different combinations of analyzers, tandem mass spectrometers e.g. triple quadrupole (QqQ) and hybrid instrument e.g. Q-TOF, have in recent time become popular. A
schematic illustration of one of the most frequently used instruments in this work, a Q-TOF, is shown in Figure 19.

Figure 19. Schematic illustration of a Q-TOF mass spectrometer.

Ions are transferred from the ion source via an ionfunnel into an analytical quadrupol consisting of two parts: a quadrupole and a collision cell. The quadrupole uses an oscillating electric field to separate ions according to their \textit{m/z}. Only ions with stable trajectory pass the quadrupole and progress to the collision cell. By letting ions collide with an inert gas \textit{e.g.} nitrogen, fragments can be formed that can be analyzed by the following TOF analyzer. The TOF analyzer consists of a long field-free flight tube with a detector at the end. Ions are pushed into this tube by a short electric pulse given by an accelerator. Ions with different \textit{m/z} travel at different velocities, assuming that the same kinetic energy was given by the pulse, and are thus separated. The time between the pulse and the detection gives a measure of the weight of the ion. However, some ions with the same \textit{m/z} that enter the flight tube will have slightly different kinetic energy. Many modern TOF analyzer, including the Bruker maXis impact (Bruker, 2011) used in this work, are also equipped with a reflectron. The reflectron device is used to compensate for this since these ions will penetrate the reflectron to different depths. The beam of ions in the flight tube will finally reach the detector, an electron multiplier, which converts the ion beam into an electrical signal that can be amplified and recorded by a computer (Bruker, 2011; de Hoffmann & Stroobant, 2007; Harris, 2002).

Depending of the purpose of the experiment, the Q-TOF mass analyzer can be set in different modes. The Q part can be used as an ion funnel letting the TOF work as a standalone MS (MS experiments) or it can be set to select
specific ions for fragmentation and let the TOF analyze the fragments in MS/MS experiment mode. Since the TOF has high resolution and mass accuracy, determination of elemental composition of both fragment ions and precursor ions are feasible. This information is highly valuable for structure elucidation of unknown compounds. It is, however, important to calibrate the system when accurate mass determination is the aim of the analysis. The Q-TOF system used was calibrated by injecting a sodium formate solution. Sodium formate forms a series of cluster over a rather large mass range and is suitable for analysis of low molecular weight compounds with ESI.

The result of a MS analysis is presented in a mass spectrum and when the MS is connected to a HPLC, also a chromatogram. The chromatogram can be based on different types of data e.g. UV data, total ion current etc. As an example of a typical HPLC-HRMS (high resolution MS) analysis, the chromatogram and the mass spectra of phlebiopsin A and B from a 20% acetonitrile SPE eluate of *P. gigantea* are presented in Figure 20.
3.3.2 Nuclear magnetic resonance

Nuclear magnetic resonance (NMR) is a powerful method for determination of molecular structures. It is a technique that takes advantage of the fact that nuclei of different atoms in molecular structures absorb and re-emit electromagnetic radiation differently. Provided, however, that the molecule under study is placed in a strong magnetic field. The NMR data recorded can...
then be used to elucidate the surrounding of each atom and ultimately the molecular structure.

Briefly, many atomic nuclei have what is called nuclear spin, i.e. the nuclei spin around their own axis and generate a small magnetic field and a magnetic moment that is oriented along the axis of spin. The magnetic moment is a vector and has both a direction and a magnitude. The direction of the magnetic moment in the atoms studied during this thesis work, $^1\text{H}$ and $^{13}\text{C}$, can have two directions in the presence of a strong magnetic field. That is, either with or against the external magnetic field. Nuclei whose magnetic moment are aligned with the external field have slightly lower energy than the nuclei aligned against, resulting in a net magnetization that is aligned with the external magnetic field (+z). The system is in this situation in equilibrium. Furthermore, the force from the external magnetic field causes the magnetic moments to precess or circulate around the z-axis with a specific frequency, the Larmor frequency. If a pulse of electromagnetic radiation with a frequency that match the Larmor frequency now is applied along the x-axis, the net magnetization will be rotated around the x-axis in the yz-plane. The extent of rotation depends upon the given pulse. A 90 degree pulse, for example, will rotate the net magnetization so that it is along the y-axis. Once the pulse has been terminated, the nuclei start to relax and return to equilibrium. It is during this process that a signal is recorded. The magnetic moment starts, immediately after the pulse, to precess around the z-axis with the Larmor frequency but now with magnetic moment in the xy-plane. This motion creates a signal that is detected by a receiver coil commonly located along the x-axis. As the relaxation proceeds the magnetization in the xy-plane steadily decrease resulting in a signal that look like a decaying sinus wave. The recorded signal is called free induction decay (FID) and is transformed by Fourier transformation to a NMR spectrum (Lambert & Mazzola, 2003; Hore, 1995; Breitmaier, 1993).

The reason NMR is useful for structure elucidation is that many nuclei in a molecule have slightly different magnetic environments, depending of the electron density around them. This will result in signals at different frequencies which are, in the NMR spectrum, expressed as chemical shifts (δ) in parts per million (ppm) compared to the signal of tetramethylsilane (TMS). Nuclei do also affect each other and are said to couple to different extent. The major factor on the size of a coupling is the number of bonds between the interacting nuclei and the dihedral angle. The coupling results in a splitting of an NMR signal. For example, a coupling between two aromatic ortho protons will result in two doublets, with a mutual coupling constant in the order of 6-9 Hz.
3.3.3 The process of structure elucidation

This section aims to generally describe the process of how data from MS and various NMR experiments can be used to elucidate the chemical structure of an unknown compound. Structure elucidation normally involves (1) calculation of molecular formula, (2) determination of connectivity between atoms. This step can be divided into (i) determination of small fragments (ii) combining small fragments, (3) determination of relative configuration, (4) determination of absolute configuration.

As mentioned earlier, the data from high resolution MS can be used to calculate the molecular formula. This is commonly the first step in a structure elucidation analysis and can give a hint of the compound under investigation. For example, a molecular formula of C\textsubscript{15}H\textsubscript{22}O\textsubscript{3}, may suggest that it is a sesquiterpene, particularly if the organism is known to produce this type of compounds. More importantly, the molecular formula can be used to calculate the unsaturation index (five in C\textsubscript{15}H\textsubscript{22}O\textsubscript{3}) which is the sum of the number of rings and multiple bonds in the molecule.

There are a large amount of NMR experiments available that give different information. However, for routine structural elucidations a basic set of relatively simple, sensitive and fast experiments is most often sufficient. A basic set of experiments includes, for example, one dimensional \textsuperscript{1}H and \textsuperscript{13}C, \textsuperscript{1}H-\textsuperscript{1}H correlation spectroscopy (COSY), \textsuperscript{1}H-\textsuperscript{13}C heteronuclear single quantum coherence (HSQC), \textsuperscript{1}H-\textsuperscript{13}C heteronuclear multiple bond correlation (HMBC), \textsuperscript{1}H-\textsuperscript{1}H nuclear Overhauser effect spectroscopy (NOESY) or rotating frame nuclear Overhauser effect spectroscopy (ROESY). A good quality 1D \textsuperscript{13}C spectrum is, however, not always feasible to obtain as it requires more sample material compared to e.g. an HMBC experiment.

Except considering the molecular formula, a sample elucidation starts by determining the connectivity between the atoms in the molecule and the first analysis is often of the 1D \textsuperscript{1}H spectrum. The \textsuperscript{1}H spectrum might give an idea of the types of protons present in the molecule e.g. aldehyde, aromatic or methyl. In addition, each signal can be integrated to give the number of protons related to a certain signal. The number of protons should of course also be compared to the molecular formula. Missing signals could be a valuable source of information as exchangeable protons are deuterated in a protic deuterated solvent such as methanol-d\textsubscript{4}.

The next step is often to determine small fragments in the molecule by using a combination of data from COSY, HSQC and if recorded also 1D \textsuperscript{13}C. HSQC is a phase-sensitive two dimensional experiment which is used to identify the carbons and their chemical shifts to which certain protons are attached. This is possible since each peak in the HSQC spectrum represent the
chemical shift of every proton and its directly attached carbon i.e. one bond $^1$H-$^{13}$C. Another advantage of the HSQC experiment is that the phase of each peak shows whether its corresponding carbon atom is bonded to an even or odd number of protons. The remaining carbons not bonded to any proton could now be identified from the 1D $^{13}$C data. Obtained $^1$H and $^{13}$C chemical shifts can be compared to tabulated chemical shifts available in for example many books (Lambert & Mazzola, 2003; Pavia et al., 2000). These can give a good picture of functional groups present in the molecule. COSY is a proton-proton correlated experiments where each peak represent a through bond correlation between certain protons. These correlations could be two (geminal), three (vicinal) or four bonds long (long-range) and sometimes even longer. Geminal couplings can be excluded from the information obtained from HSQC data and long-range couplings are often weaker and could therefore also be temporary excluded. Vicinal couplings are therefore left and small fragments can be determined.

HMBC data shows the connection between a certain proton and the carbons two and three bonds away, sometimes longer depending on the molecular structure. This is a very useful experiment to combine small fragments and finally determine the overall connectivity of the molecule. In addition, since 1D $^{13}$C data sometimes can be difficult to obtain, HMBC data can be sufficient to extract all the carbon chemical shifts in the molecule. The number of rings and multiple bonds as calculated from the molecular formula is, at this point, an excellent source of information to support in the puzzle of putting pieces together.

Once the connectivity is determined, the next step is to determine the relative configuration of the stereogenic carbons. NOESY and ROESY experiments are often used for this purpose. These two dimensional experiments show cross-peaks for protons close to each other in space.

Finally, when both the connectivity and the relation between groups on stereogenic carbons, i.e. relative configuration, have been established only absolute configuration remains to be determined. This is often one of the most difficult parts in the elucidation process but is needed for full structure determination.

Several methods exists e.g. X-ray crystallography, circular dichroism and different NMR based methods. The method used in this work was based on biosynthetic considerations. For example, a precursor involved in the biosynthesis of a certain secondary metabolite is known to be produced with a certain configuration. Logical reasoning could then be used to propose the absolute configuration of at least one stereogenic carbon of the investigated
compound and then, by considering the relative configuration, the absolute configuration of all stereogenic carbons.

3.4 Stable isotope labeling

Feeding living systems with isotopically labeled substrates is a common method to elucidate which metabolic pathways that have been used in the construction of a certain secondary metabolites. The method is based on that an organism is grown in the presence of a labeled substrate which is incorporated into the molecular structure of the metabolite under study. The metabolite is then analyzed to detect the presence of the labeling. It is thus essential that the organism produce the metabolite during the feeding period.

Radioactive isotopes such as $^{14}$C and $^{3}$H were frequently used in the middle of the 20th century but biosynthetic elucidations using these isotopes were dependent on careful laborious chemical degradation of the product to locate the labeling. $^{14}$C and $^{3}$H were later displaced with substrates labeled with stable isotopes e.g. $^{13}$C. $^{13}$C is an excellent tracer since it can be studied by NMR. $^{13}$C-labeled positions are identified by their enhanced signals compared to unlabeled positions. Assignment of NMR signals will, in addition, give information of which carbons that have been labeled without any chemical degradation (Bacher et al., 1998).

All biosynthetic pathways leading to secondary metabolites use starting materials from the central metabolic pathways where glucose is a key compound. $^{13}$C-labeled glucose e.g. $[1^{13}$C]glucose, used in the biosynthetic study of fomannoxin (paper II), is therefore a qualified and common tracer to elucidate biosynthetic pathways. Depending on which metabolic pathways organisms use to incorporate glucose derived carbons, the $^{13}$C-labeling of glucose end up at characteristic positions in the metabolites. The observed labeling pattern of a metabolite under study can thus reveal which biosynthetic pathways that have been used for the formation.
4 Results & Discussion

4.1 Secondary metabolites of *H. annosum* s.l. (Papers I-III)

One of the objectives of this thesis work was to further investigate the secondary metabolites produced by *H. annosum* s.l. This subsection aims to conclude all known secondary metabolites with focus on the compounds that have been identified during this work and are described in papers I-III. In addition, the differences and/or similarities, regarding the production of secondary metabolites, between the species within the *H. annosum* s.l. that were investigated in paper III are concluded and discussed.

4.1.1 Sesquiterpenes – Fomannosin-type

Fomannosin is a sesquiterpene secondary metabolite that has previously been isolated from *H. annosum* s.l. and was first described 1967 by Kepler *et al.* (1967). The phytotoxic properties of fomannosin presented in paper I, was in accord with previously published data where fomannosin was shown to have activity against the pine *Pinus taeda* (Bassett *et al.*, 1967). Because of its phytotoxicity, fomannosin has been claimed to be involved in the plant pathogenic activity of *H. annosum* s.l.

With the interest to further investigate *H. annosum* s.l. regarding bioactive secondary metabolites, a small scale experiment were set up and all five species within the *H. annosum* s.l. species complex were cultivated in different medium (Hagem or Melin Norkrans medium) and with or without stirring. The liquid cultures were subsequently analyzed with LC-MS/UV to determine which species, strain and cultivating parameters that produce most secondary metabolites. *H. occidentale* strain TC122-12, cultivated in Hagem medium at stationary conditions was shown to be a promising candidate.

Hence, *H. occidentale* strain TC122-12 were cultivated in liquid Hagem media, filtered and subjected to SPE. The 95% MeCN eluate was subjected to
gradient semi-preparative HPLC (Figure 21) and fractions were collected in deep-well plates. All HPLC fractions were then tested for growth inhibiting activity against four fungi: *Fusarium oxysporum*, *P. gigantea*, *Phanerochaete chrysosporium* and *H. occidentale*; against one bacterium: *Variovorax paradoxus* and against one moss: *Physcomitrella patens*. These bioassays were used as a guide to select which collected fractions to be subjected to further chemical characterization. In addition to active fractions, some other major UV-peaks were collected, pooled and subjected to further analysis in the same manner as the active fractions.

Small aliquots, of pooled fractions, were analyzed by LC-MS to obtain information of molecular mass. Notable, many of the compounds corresponding to active fractions behaved similar in the MS and formed the following adduct ions: [M-H$_2$O+H]$^+$, [M+H]$^+$, [M+Na]$^+$, [2M+H]$^+$ and [2M+Na]$^+$. [M+H]$^+$ was usually observed as the minor ion. Pooled fractions, were then concentrated by evaporation, lyophilized and subsequently redissolved in CDCl$_3$. The samples were analyzed by NMR and their structures

![Figure 21. Preparative chromatogram from isolation of fomannosin-type sesquiterpenes and fomannoxin from *H. occidentale* (active fractions marked with black boxes).](image)
were elucidated, resulting in the identification of fomannosin and the benzohydrofuran fomannoxin as well as the novel sesquiterpenes illudolone A and B, illudolactone A and B and deoxyfomannosin A and B (Figure 22). The molecular formulas of respective compound were later determined by HRFABMS or HREIMS and the compounds were further characterized by polarimetry and UV.

Figure 22. Structure of fomannosin and fomannosin type compounds (paper I).

4.1.2 Sesquiterpenes – Fomajorin-type

In order to investigate how the secondary metabolite production of *H. occidentale* are affected by the biocontrol fungus *P. gigantea*, *H. occidentale* and *P. gigantea* were grown in liquid dual cultures and subsequently extracted by SPE. Chromatographic separation of the SPE extract resulted in the isolation of, among others, three fomajorin-type sesquiterpenes (unpublished results), namely, fomajorin D, 9-hydroxy-7-(hydroxymethyl)-5,7-dimethyl-7,8-dihydrocyclopenta[g]isochromen-1(6H)-one (1) and 9-hydroxy-5-(hydroxymethyl)-7,7-dimethyl-7,8-dihydrocyclopenta[g]isochromen-1(6H)-one (2, Figure 23). Structure elucidations of the compounds were done by one- and two-dimensional NMR spectroscopy and HRMS.

Figure 23. Structure of the fomajorins.

Compound 2 has never been described before whereas fomajorin D and compound 1 have previously been reported from *H. annosum* s.l. together with
fomajorin S (Sonnenbichler et al., 1989; Donnelly et al., 1982). Furthermore, fomajorin D was later also identified in paper III.

The $^1$H NMR spectrum of compound 2 (Figure 24) contained signals at almost identical chemical shifts as fomajorin D, except that the signal from the aromatic methyl group of fomajorin D at $\delta_H$ 2.24 ppm (CDCl$_3$, 30 °C) was replaced with an signals at $\delta_H$ 4.73 that integrated for two protons. HMBC experiments further demonstrated that this methylene group was attached to the aromatic ring, which further strengthen the proposed structure of compound 2.

![Figure 24. $^1$H NMR spectrum of compound 2.](image)

4.1.3 Sesquiterpenes – Drimane-type

A third type of sesquiterpenes, the drimane-type, was also identified to be produced by *H. annosum* s.l. (paper III). The identification of 7α,8β,11-trihydroxydrimane (Figure 25) was done by NMR and HRMS. 7α,8β,11-Trihydroxydrimane has previously been described in *H. annosum* s.l. and the recorded NMR data were in agreement with literature data (Liu et al., 2008; Donnelly et al., 1980). Epoxydrimenol (Figure 25) was proposed to be a true secondary metabolite of *H. annosoum* s.l. although no NMR data was recorded for this compound. It was indirectly identified as the epoxide of 7α,8β,11-trihydroxydrimane (paper III). The molecular formula of epoxydrimenol,
C_{15}H_{26}O_2 (obtained by LC-HRMS), was not in agreement with the recorded NMR data of the same sample. The NMR data was, however, shown to be in accord with the NMR data recorded for the drimanetriol. Re-analysis by LC-HRMS of the epoxide confirmed that the epoxide had been transformed to the drimanetriol (C_{15}H_{28}O_3). Both the retention time and the molecular formula were in agreement with the drimanetriol. Epoxydrimenol was thus proposed to open during work-up in acidic conditions and form the 7α,8β,11-trihydroxydrimane.

![Figure 25. Structure of drimane-type sesquiterpenes.](image)

4.1.4 Fomannoxin and fomannoxin related compounds

Fomannoxin (Figure 26) is a benzohydrofuran first described by Hirotani et al. (1977) when it was isolated from *H. annosum* s.l. Fomannoxin has been identified in infected plant tissue and has been shown to be much more phytotoxic than *e.g.* fomannosin (Heslin *et al.*, 1983; Hirotani *et al.*, 1977). Fomannoxin and eighteen additional fomannoxin related compounds were identified during this thesis work (Figure 26). Paper II describes four of these (3-4 and 12-13), in addition to fomannoxin. Compound 13 was a new natural product and the structure elucidation, done by NMR and HRMS, is described in paper II. Compounds 5-11 and 14-20 were, in addition to 3-4 and 12, identified in paper III by NMR and/or LC-HRMS. Compounds 8-9 and 15 were found to be new compounds and their structure elucidations are described in paper III for the first time, whereas compounds 14, 17-18 had been described earlier but not reported as natural products.
4.1.5 Indole-type

Tryptophan and 2,3,4,9-tetrahydro-1\textit{H}-β-carboline-3-carboxylic acid (21) were identified as two compounds that differ in either relative concentration or occurrence between the species within the \textit{H. annosum} s.l. complex (paper III). Both compounds were identified by HRMS and NMR. The structure of tryptophan was confirmed by comparing \textit{1H} NMR data with reference \textit{l-tryptophan} whereas NMR data of 2,3,4,9-tetrahydro-1\textit{H}-β-carboline-3-carboxylic acid was compared to literature data (Kotanen \textit{et al.}, 2003).

![Figure 27. Structures of identified indole-type compounds.](image)

4.2 Secondary metabolites comparison between the \textit{H. annosum} s.l. species (Paper III)

With the aim to investigate if the species within the \textit{H. annosum} s.l. complex could be distinguished on their secondary metabolite profiles, all five species
were grown on separate agar plates, extracted with methanol and analyzed by LC-HRMS. The data set were subsequently evaluated by principal component analysis (PCA). As can be seen from the score plot (Figure 28), the samples were separated into five groups in PC1 and PC2 corresponding to the five species.

![Figure 28. PCA score plot (PC1 and PC2) on LC-HRMS data of five different species within the *H. annosum* s.l. species complex.](image)

PC1 describes most of the differences and showed that the pine infecting species, *H. annosum* s.s. and *H. irregulare*, grouped close to each other and well separated from the non-pine infecting species. *H. parviporum* grouped close to the other non-pine infecting species, *H. occidentale* and *H. abietinum* in PC1 but was separated from these in PC2. Furthermore, *H. annosum* s.s./*H. irregulare* and *H. occidentale/H. abietinum* were separated in PC3 and PC4, respectively (paper III).

Compounds important for the separation were targeted for isolation and identification. Twenty-three compounds, in total, were identified by NMR and HRMS or by comparison with references and eleven compounds were assigned molecular formula. In addition, three compounds were identified based on major UV-peaks in preparative HPLC chromatograms (paper III).

Several known compounds including fomannoxin, fomannoxin acid, fomannosin, deoxyfomannosin A and B, epoxydrimenol, tryptophan etc. were identified as compounds important for the separation of the five species within the *H. annosum* s.l. species complex (paper III). The production of fomannoxin was particularly important as the main difference between the pine infecting species and the non-pine infecting species. Earlier studies of for example *H. occidentale* (paper I-II) have, however, shown that other species than the pine-infecting are able to produce fomannoxin which indicate that the difference observed in this study is not on the genome level.
Nevertheless, the study revealed that it is feasible to differentiate the species by their respective metabolite pattern, but whether these differences remain when the fungi are grown on other medium as e.g. their respective host is uncertain. Furthermore, the samples were separated in accord with the previously identified phylogeny (Dalman et al., 2010) and host preferences.

4.3 Biosynthesis of secondary metabolites produced by *H. annosum* s.l. (Papers I-III)

4.3.1 Biosynthesis of sesquiterpene compounds

The biosynthesis of fomannosin and fomajorin D has previously been investigated and has been proposed to proceed from *trans, trans*-FPP via humulene and the protoilludyl cation. The three isoprene building blocks needed for the assembly of FPP were shown, by labeling experiments of [1,2-\textsuperscript{13}C]acetate, [1-\textsuperscript{13}C]acetate and [2-\textsuperscript{13}C], to originate from the MVA pathway (Donnelly et al., 1987; Donnelly et al., 1983; Cane & Nachbar, 1978; Cane & Nachbar, 1976). By breaking appropriate C-C bonds of the protoilludyl cation (*Figure 29*) the carbon back-bone of both fomannosin and the fomajorins could be formed.

*Figure 29.* Proposed biosynthetic pathway to fomannosin and fomajorins.

Illudolone A and B, illudolactone A and B, and deoxyfomannosin A and B (paper I), were all suggested to be intermediates in the biosynthesis of fomannosin (*Figure 30*). Loss of a proton of protoilludyl cation forms (−)-\Delta^6- protoilludene which, upon monooxygenase and dehydrogenase oxidation may result in illudolone A. Illudolone B is then formed upon elimination of water. A proposed Baeyer-Villiger oxidation of illudolone B would break the C4-C4a carbon bond, necessary for formation of fomannosin and form the seven-membered lactone ring present in illudolactone A and B. Dehydrogenase oxidation of the cyclopentyl ring of a proposed lactone intermediate would introduce the double bond present in illudolactone A and B. Hydrolysis of the lactone rings followed by keto-enol tautomerization would result in deoxyfomannosin A and B. Finally, hydroxylation and lactonization would lead to fomannosin.
The absolute configuration assigned for illudolone A and B, illudolactone A and B, and deoxyfomannosin A and B were based on the above described biosynthetic pathway and that they all share the protoilludyl cation as a common precursor. Furthermore, it was assumed that the configuration of the stereogenic carbon in the lactone ring of fomannosin is set during the cyclization of humulene.

Figure 30. Proposed biosynthetic pathway for fomannosin (paper I) and the fomajorins from the common precursor, \((-\Delta^6\)-protoilludene).
The biosynthesis of the fomajorins has, as mentioned above, been suggested to proceed from the protoilludyl cation, however, no intermediates have been reported. Hence, a proposed biosynthetic route to the fomajorins is presented (Figure 30). \((-\Delta^6\)-protoilludene is formed upon loss of a proton of the protoilludyl cation as in the biosynthesis of fomannosin and the fomannosin intermediates. A hydroxylation, followed by elimination of water promote for later aromatization. These steps could alternatively be performed by dehydrogenase oxidation to introduce the double bond directly. Likewise the proposed biosynthesis of fomannosin, a Baeyer-Villiger oxidation might be involved to break the C-C bond necessary for formation of the fomajorin carbon skeleton. A sequence of reactions including reduction, aromatization, lactonization and oxidations would finally lead to the formation of fomajorin D. The other fomajorin type sesquiterpenes; fomajorin S, 9-hydroxy-7-(hydroxymethyl)-5,7-dimethyl-7,8-dihydrocyclopenta[g]isochromen-1(6H)-one (1) and 9-hydroxy-5-(hydroxymethyl)-7,7-dimethyl-7,8-dihydrocyclopenta[g]isochromen-1(6H)-one (2) are most likely oxidation products of fomajorin D.

The suggested biosynthetic pathways for fomannosin, fomajorin and related intermediates as illudolactone A and B, include several enzyme-catalyzed reactions. In the closely related species, \(H. \text{irregulare}\), several genes encoding for such enzymes have been identified. For example: several monooxygenases, three terpene cyclases and three Baeyer-Villiger monooxygenases have been identified. These findings further strengthened the proposed biosynthetic pathways for these compounds.

\[\text{trans, trans-FPP} \rightarrow \text{OPP} \rightarrow \text{OH} \rightarrow \text{epoxydrimenol} \rightarrow \text{7α,8β,11-trihydroxydrimane}\]

*Figure 31. Proposed biosynthesis to epoxydrimenol and 7α,8β,11-trihydroxydrimane.*

The biosynthesis of 7α,8β,11-trihydroxydrimane, previously found in \(H. \text{annosum}\) s.l., has not been investigated. However, since the other sesquiterpenes, fomannosin and the fomajorins, are biosynthesized from MVA via \(\text{trans, trans-FPP}\), it seems likely that also epoxydrimenol and 7α,8β,11-trihydroxydrimane are biosynthesized via similar pathways. A proposed biosynthetic route to these sesquiterpene compounds is presented in Figure 31. Cyclization of \(\text{trans, trans-FPP}\), driven by the addition of a proton at C-10 of FPP, would result in the formation of the two six-membered rings present in
the carbon backbone of drimanes. Loss of a proton to neutralize the positive charge forms a double bond, which upon oxidation and hydrolysis of diphosphate leads to epoxydrimenol. 7α,8β,11-Trihydroxydrimane could finally be formed after opening of the epoxide.

4.3.2 Biosynthesis of fomannoxin and fomannoxin related compounds

Fomannoxin is a known phytotoxin of *H. annosum* s.l. and is one of the major secondary metabolites produced by several of the species within the species complex. One of the objectives of this thesis work was to investigate the biosynthesis of this compound (for details see paper II).

Liquid cultures of *H. occidentale* were grown in Hagem media and small aliquots were regularly taken and analyzed by LC-MS. When the presence of fomannoxin was observed (after ca. three weeks of growth), by the detection of an ion at \( m/z \) 189 corresponding to the \([M+H]^+\) ion of fomannoxin, [1-\(^{13}\)C]glucose was added. The labeled cultures were then grown for another two weeks before they were harvested by filtration and extracted by SPE. Unlabeled cultures were cultivated in parallel. The SPE extracts were subsequently fractionated by preparative HPLC and fractions corresponding to fomannoxin and other major UV-peaks were pooled and prepared for NMR analysis. \(^{13}\)C NMR spectra of fomannoxin, cultivated in both labeled and unlabeled media, are presented in Figure 32.
The $^{13}$C NMR spectra revealed that five positions had been enriched; C-4 and C-6, belonging to the aromatic ring, and C-2, C-2’ and C-3’ of the five-carbon side-chain. Suggested by the structure of fomannoxin, this side-chain may be a result of alkylation of an aromatic ring-containing substrate such as 4-hydroxybenzaldehyde with an isoprene unit. As discussed in chapter 2, isoprene units (IPP or DMAPP) might be formed by either the MVA pathway or by the MEP pathway. When cultivation is performed in the presence of a selectively $^{13}$C-labeled carbon source such as [$1^{13}$C]glucose, the $^{13}$C-labeling patterns will reflect the metabolic pathway used for the construction. If the isoprene unit is biosynthesized via the MVA pathway, three carbons would be labeled (corresponding labeled carbons in fomannoxin; C-2, C-2’ and C-3’, Figure 33), whereas the MEP pathway would yield two labeled positions (corresponding labeled carbons in fomannoxin; C-3 and C-2’). Thus, the result clearly suggested that the side-chain is derived via the MVA pathway.

Aromatic rings are commonly biosynthesized either via the shikimic acid pathway or as a polyketide via the acetate pathway (see section 2.2 and 2.3). Polyketide originating aromatics would have three ring carbons enriched, irrespective of folding of the poly-β-keto chain. The observed labeling pattern of fomannoxin, showed that only two carbons had been enriched and the
aromatic ring could, thus, not been formed via the acetate pathway. A shikimic acid derived aromatic compound would, on the other hand, have only two carbons enriched in the aromatic ring, as observed in the present experiment. Depending on if 4-hydroxybenzaldehyde is formed via a C₆C₃ unit followed by chain shortening or if it is directly formed from chorismic acid by elimination of pyruvate, the aldehyde carbon would be labeled or not. The observed labeling pattern clearly suggested the latter pathway. Thus, fomannoxin was suggested to be formed by a combination of biosynthetic pathways. The aromatic moiety from the shikimic acid pathway via elimination of pyruvate from chorismic acid and the isoprene side-chain via the MVA pathway.

A key step in the biosynthesis of fomannoxin is the alkylation of the five-carbon side-chain, a reaction type commonly performed by the action of prenyltransferases. One gene encoding a putative aromatic prenyltransferase was found in the genome of *H. irregulare*, one of the other fungi within the *H. annosum* species complex. Furthermore, a new natural compound (13) was identified and, based on the above suggested biosynthetic pathway, suggested to be a key intermediate in the formation of fomannoxin and related compounds. In addition, the labeling pattern of compound 13 was similar to the pattern of fomannoxin. Other fomannoxin related compounds is suggested to be intermediates in the biosynthesis of fomannoxin or formed by simple modifications as oxidations and/or reductions, of fomannoxin intermediates or fomannoxin itself (compounds 3-12, Figure 26). Compounds 17-20 (Figure 26) have an extra carbon substituent ortho to the hydrox group of 4-hydroxybenzaldehyde/4-hydroxybenzoic acid and are suggested to be formed either by methylation or by degradation of the fomannoxin carbon skeleton. Compounds 14-15, containing the benzohydrofuran moiety, might also be formed by degradation of appropriate fomannoxin analogs. Oxidative cleavage of compound 13 followed by ring cyclization would for example, be a plausible route to these compounds.
Figure 33. Biosynthesis of fomannoxin. Dots indicates $^{13}$C enriched positions arising from [1-$^{13}$C]glucose.

4.3.3 Biosynthesis of indole-type compounds

The biosynthesis of tryptophan was described in section 2.2. The other indole-type compound identified in paper III, compound 21 (Figure 27), belongs to a group of compounds called β-carbolines. Herraiz (2000), suggested compound 21 to be biosynthesized by condensation between tryptophan and formaldehyde. The biosynthesis is depicted in Figure 34. The extra carbon needed for the cyclization to the six-membered ring is attached to the amine function by a Schiff-base formation reaction. C-2 of the indole system is then attacking the Schiff-base and forms, after loss of a proton (H-2) of the indole system and restoration of the aromaticity, compound 21.

Figure 34. Biosynthesis of compound 21 from tryptophan.
4.4 Secondary metabolites of *P. gigantea* (paper IV)

*P. gigantea* is used as biocontrol fungus against *H. annosum* s.l. but the mechanism of the biocontrol action is not fully understood. The objective of this study was to investigate *P. gigantea* with regard to its production of secondary metabolites and to test whether these compounds have an effect on *H. occidentale* and thus, could be involved in the *in vivo* biocontrol action between *P. gigantea* and the *H. annosum* s.l.

*P. gigantea* was cultivated in Melin Norkrans media, harvested by filtration and extracted by SPE. The SPE extracts were then fractionated by preparative HPLC and major UV-peaks were collected and analyzed by one- and two-dimensional NMR spectroscopy and HRMS. Five compounds were identified, phlebiopsin A-C, methyl-terfestatin A and *o*-orsellinaldehyde (*Figure 34*).

*Figure 35. Structure of secondary metabolites identified in *P. gigantea*.*

Phlebiopsin A-C are structurally similar to a group of compounds that are rarely found in nature called diarylcyclopentenones. These compounds have been described as responsible for the bluing reactions observed when the fruiting bodies of certain mushrooms of the order Boletales are bruised. Their activities against three fungi including *H. occidentale* were tested. However, no activities against either of the fungi were observed at the tested concentrations.

The aglycone of the identified compound methyl-terfestatin A, is very similar to 2’,3’,5’-trimethoxy-α-terphenyl, a compound that was not found in this study but has been reported from *P. gigantea* previously (Briggs et al., 1975). Methyl-terfestatin A is also very similar to terfestatin A, the only difference being the methoxy group at C-3’. Terfestatin A has previously been isolated from *Streptomyces* bacteria and has shown to inhibit auxin signaling in
plants (Yamazoe et al., 2005; Yamazoe et al., 2004). Methyl-terfestatin A did, however, not show any activity against the fungi tested in this study.

In addition to the compounds above, which all were new natural products, the known compound o-orsellinaldehyde was identified. o-Orsellinaldehyde was shown to inhibit the conidial germination of the tested fungi, but at rather high concentration. The conclusion was, thus, that neither of the identified compounds seems to be important for the biocontrol action of P. gigantea against H. annosum s.l. and that the previously suggested mode of action in biocontrol of H. annosum s.l., i.e. by resource competition, is supported.

4.5 Biosynthesis of secondary metabolites produced by P.gigantea

The biosynthesis of orsellinic acid is known to proceed via the acetate pathway by folding of the appropriate poly-β-ketoester (section 2.3). o-Orsellinaldehyde is then probably formed by reduction of the carboxylic acid to an aldehyde. The aglycone of methyl-terfestatin A, belongs to a structural class called para-terphenyls, identified by their chain of three benzene rings. p-Terphenyls have been proposed to be biosynthesized (Figure 36) by condensation of two phenylpyruvic acid molecules (Zhou & Liu, 2010) via a ketone intermediate and polyporic acid. Subsequent reduction and aromatization would form 2´,3´,5´-trihydroxy-p-terphenyl, which upon glycosylation and methylation of the hydroxyl group at C-2´and C-3´, respectively, would form methyl-terfestatin A. 2´,3´,5´-Trimethoxy-p-terphenyl, previously identified from P. gigantea (Briggs et al., 1975) is probable formed by methylation of the hydroxyl group of 2´,3´,5´-trihydroxy-p-terphenyl. Phenylpyruvic acid has also been proposed to be the precursor of pulvinic acid (Yun et al., 2001). Pulvinic acid in turn, has been suggested to be the precursor of several diarylpentenones e.g. chamomixin, involutin, gyroporin and gyrocyanin (Velisek & Cejpek, 2011). These compounds have similar structures as phlebiopsin A-C.
Figure 36. Biosynthesis of methyl-terfesstatin A.
5 Conclusions and suggestions for further research

The objectives of the present thesis work included investigating *H. annosum* s.l. and *P. gigantea* regarding their production of secondary metabolites, how they are biosynthesized and their biological activity.

In total, 33 compounds were identified during the investigation of *H. annosum* s.l. Six new sesquiterpenes (illudolone A and B, illudolactone A and B and deoxyfomannosin A and B) belonging to the fomannosin class of compounds were identified and described. Furthermore, three sesquiterpenes belonging to the fomajorins and two drimanes were identified out of which one was a new compound (2). The benzohydrofuran fomannoxin and several (19) fomannoxin related compounds were identified. Three of these were shown to be previously unknown compounds and four had not been described from any natural sources. In addition, tryptophan and another indole-containing compound, a β-carboline, were identified.

Fomannoxin has been suggested to be important for the pathogenicity of *H. annosum* s.l. Its biosynthesis was investigated by an isotopic enrichment study utilizing [1-13C]glucose as metabolic tracer. The labeling pattern, analyzed by 13C-NMR analysis, showed that fomannoxin is biosynthesized by a combination of the MVA pathway and the shikimic acid pathway.

The species within *H. annosum* s.l. were compared regarding their production of secondary metabolites. The results showed that the five species could be separated by their respective metabolite profile and that the separation was in accord with both host preference and previously established phylogeny.

The biocontrol fungus *P. gigantea*, was investigated regarding its production of secondary metabolites. The study resulted in five identified compounds out of which three were new compounds (phlebiopsin A-C), one was a new natural product (methyl-terfestatin A) and one (*o*-orsellinaldehyde) was a known compound. *o*-Orsellinaldehyde showed weak activity against *H.*
occidentale whereas none of the new natural products showed any activity at the tested concentrations.

*H. annosum* s.l. and *P. gigantea* produce, for sure, a lot more secondary metabolites than were identified during this work. For example, recent studies of *H. annosum* s.l. have shown that there are several NRPSs, PKSs and halogenases present in the genome (Olson *et al.*, 2012). However, neither NRPS, PKS derived nor halogenated secondary metabolites were identified to be produced in *H. annosum* s.l. during this work. Possibly, other “secondary metabolite genes” are expressed in the presence of natural competitors or during infection of host *in vivo*. An emerging technology within the natural product chemistry field is epigenetics. Epigenetics manipulate which genes that are expressed. The result could be that earlier “silent genes” became active and that other secondary metabolites are formed. Regardless of how the previously silent genes are made active, it would be interesting to investigate the resulting metabolic profiles.

A natural continuation on the study of *P. gigantea* would be to investigate the biosynthesis of the diarylcyclopentenones that were identified. Previously suggested pathways have not been established by labeled substrates, but only by discussions based on structure similarities.
References


*chrysogenum* is a 421 073 Dalton protein similar to *Bacillus brevis* peptide antibiotic synthetases. *Embo Journal* 9(9), 2743-2750.


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