Antarctic Microfungi as a Potential Bioresource

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Declaration

The research presented in this thesis is original work performed between March 1997 and March 2003 by the author. This research has not been submitted to any other university or institution as part of the requirements for any higher degree or course.

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Abstract

The Antarctic occupies that region of the planet that falls below the 60th parallel of South latitude. Although it has been frequented by adventurers, journeyman scientists and tourists for the past 100 years, the Continent has remained virtually unoccupied. The intense cold, the absence of human occupation and the limited range of local higher animal species have combined to create the impression that the Continent is virtually devoid of life.

Although the microbiota of the Antarctic has attracted some small level of attention in the past, the examination of filamentous microfungi has been largely overlooked and fallen to a small group of dedicated investigators. In this study it will be shown that far from being an insignificant component of the Antarctic network, microfungi represent a potentially large and so far untapped bioresource.

From just 11 bryophyte samples collected at four sites in the Ross Sea/Dry Valleys region of Southern Antarctica, some 30 microfungal isolates were recovered. Using molecular techniques, the internal transcribed spacer (ITS) region of the nuclear ribosomal DNA (nrDNA) was sequenced to reveal no less than nine unique microfungal species. For only two of these species did the ITS sequence data produce a 100% match with records held on the public databases. This investigation also highlighted the problems inherent in the traditional morphological identification system which are now being perpetuated in the molecular database records.

A set of seven notionally identified isolates obtained from ornithogenic soil samples gathered in the Windmill Islands in Eastern Antarctica (offshore from the Australian
Antarctic Division’s Casey Station) were also subjected to molecular identification based on ITS sequence data. Each of the seven isolates was identified as a unique species; six were cosmopolitan in nature and the one remaining bore very little resemblance at the molecular level to any of the recorded species although it was provided with an epithet commonly used in the identification of Antarctic microfungal species.

To evaluate their potential as a bioresource, samples of Antarctic microfungi were examined to determine if the same physiological factors common to mesophilic species also applied to their Antarctic analogues. It is known that when placed under stress, trehalose can act as a protectant against cold (cryoprotection) and dehydration in mesophilic yeasts and fungi. The level of trehalose produced by the Antarctic isolates and their mesophilic analogues when subjected to stress was compared. A similar comparison was made for the production of glycerol which is well established as a compatible solute providing protection to mesophilic species against osmotic stress. Only in the case of trehalose production by an Antarctic *Embellisia* was there any indication that either of these two compounds could play a significant role in providing protection to the Antarctic fungi against the rigours of their environment, which leaves open to question what in fact does.

In the course of investigating the means by which Antarctic microfungi guard against the damage which can ensue when subjected to oxidative stress, flow cytometry was introduced as an investigatory tool. It was established that there is a window of opportunity during which flow cytometry can be used to undertake a detailed analysis of the early stages of fungal growth from germination through hyphal development.
Of major significance in determining the potential of Antarctic microfungi as a resource is their ability to produce new and novel enzymes and proteins. The microfungal isolates were screened for hydrolytic activity on solid media containing indicative substrates and proved to be a fruitful source of enzymes active over a range of temperatures. A detailed characterisation of two hemicellulases, β-mannanase and xylanase, secreted into a liquid medium by a subset of the Antarctic fungi and a high producing mesophilic reference strain permitted direct comparisons to be made. It was shown that the maximum hemicellulase activity of the Antarctic strains occurred at least 10ºC and as much as 30ºC lower than that of the reference strain and that mannanase activity for two of the Antarctic isolates exceeded 40% of their maximum at 0ºC. These assay results highlight the potential of Antarctic microfungi to yield novel cold-active enzymes.

As a final measure of the capacity of the Antarctic to yield novel enzymes from its microfungal stock, a lipase gene was selected as a target for isolation and expression in a heterologous fungal host. Using PCR techniques, the gene of interest was isolated from an Antarctic isolate of *Penicillium allii*, transformed into the mesophilic production host *Trichoderma reesei* and the active protein successfully produced in the growth medium. The recombinant lipase was assayed and found to exhibit novel characteristics consistent with a cold-adapted enzyme.
Preface

There are many people that should be acknowledged for the support they have given me over a long period of time. Firstly, I give my heartfelt thanks to my supervisor, Associate Professor Helena Nevalainen, for her friendship, guidance, encouragement over many years during the course of this study and throughout my undergraduate years. Her help and constructive criticism, particularly in the preparation of this document and other papers, was warmly appreciated. Special thanks also to my Associate Supervisor, Associate Professor Michael Gillings for his valued guidance in matters molecular, sequencing and in the preparation of this document and other papers.

I also extend my warmest thanks to all of my colleagues in the EDGE Laboratory at Macquarie University for providing such a pleasant working environment and in particular to Professor Peter Bergquist, Drs. Morland Gibbs, Junior Te’o, Roberto Anatori, Anwar Sunna and Noosha Ehya and Ms Roz Reeves and Ms Natalie Curach each of whom has helped me in some way during the course of this study, with either advice or constructive comment. I would also like to thank Professor Duncan Veal and Dr. Paul Attfield for their guidance and help with the flow cytometer, Dr Robert Willows for advice on biochemistry and my co-authors Dr. Philip Bell and in particular Dr. Rani Sidhu who provided invaluable assistance in the tedious task of assaying for hemicellulase activity.

Finally, I express my warmest thanks to my wife Dawn for her love, patience and support throughout this work and to my son Alexander who has helped me greatly with some of the more mystifying points of computer software.
List of original publications

This work is based on the following articles, referred to in the text by the Roman numerals given below. Additional unpublished data is also presented.


The author of this thesis had the main responsibility for the work contained in each of these publications and also for planning the experiments and writing the articles. The role of Assoc. Prof. Helena Nevalainen (publications I – V) and Assoc. Prof. Michael Gillings (I & II) was to act in their capacity as my supervisors, providing me with overall support and guidance and participated in the planning and evaluation of the experiments. Dr Rani Sidhu provided technical assistance in assaying for hemicellulase activity (II) and together with Miss Beta Yee, assisted in the isolation of the fungal material associated with the bryophytes collected in Antarctica by Dr. Patricia Selkirk and Dr. Mary Skotnicki (III). Dr. Philip Bell designed the suite of PCR primers used to identify the lipase gene in the Antarctic microfungus and Dr. Junior Te’o was responsible for engineering the plasmid utilised in the transformation of the heterologous fungal host and provided support with the biolistic transformation system (V). Sequencing with the ABI Prism automated fluorescent DNA sequencer was undertaken by the the staff of the facility at Macquarie University.
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Abbreviations

aa  amino acid
Ala  alanine
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Arg  arginine
Asn  asparagine
BAC  bacterial artificial chromosome
bp  base pairs
DHE  dihydroethidium (a fluorescent stain)
FC  flow cytometry/cytometer
FL  fluorescence
FSC  forward scatter
gDNA  genomic DNA
Glu  glutamic acid
HI  hexidium iodide
IPTG  isopropyl-β-D-thiogalactosidase
ITS  internal transcribed spacer region
Kbp  kilo base pairs
kcat  dissociation rate (s⁻¹) - Michaelis-Menten kinetics
kDa  kilo Dalton
Kₘ  Michaelis constant (mol L⁻¹) - Michaelis-Menten kinetics
lat  latitude
LB  Luria-Bertani (medium)
L-broth  Luria-Bertani broth
Lys  lysine
nrDNA  ribosomal DNA (nuclear)
nt  nucleotide
PCR  polymerase chain reaction
PD  potato dextrose
PDA  potato dextrose agar
PMT  photomultiplier tube
RAPD  randomly amplified polymorphic DNA
RFLP  restriction fragment length polymorphism
ROS  reactive oxygen species
R/T  room temperature
S.E.  Standard Error
sq km  square kilometre
SSC  side scatter
SSU  small sub-unit
uv  ultra violet light
UP-PCR  universally primed PCR
Val  valine
v/v  volume/volume
w/v  weight/volume
X-gal  5-bromo-4-chloro-3-indolyl-β-D-galactosidase