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**PHYSIOLOGY AND PRODUCTION OF
INULINASE AND GLUCOAMYLASE
ENZYMES OF *RHIZOPUS OLIGOSPORUS***

Thesis

Submitted in Fulfillment of The Requirements for The
Degree of Doctor of Philosophy

**In
Pharmaceutical Sciences
(Microbiology)**

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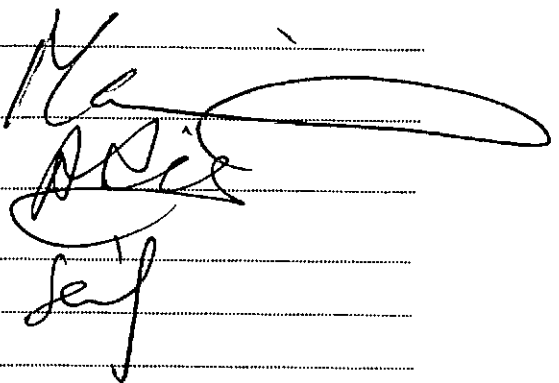
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بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

«قَالُوا سُبْحَانَكَ لَا عِلْمَ لَنَا
إِلَّا مَا عَلَّمْتَنَا إِنَّكَ أَنْتَ
الْعَلِيمُ الْحَكِيمُ»

صَدَقَ اللَّهُ الْعَظِيمُ

سورة البقرة - آية ٢٢

This Thesis is dedicated
to
My children and my
husband

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