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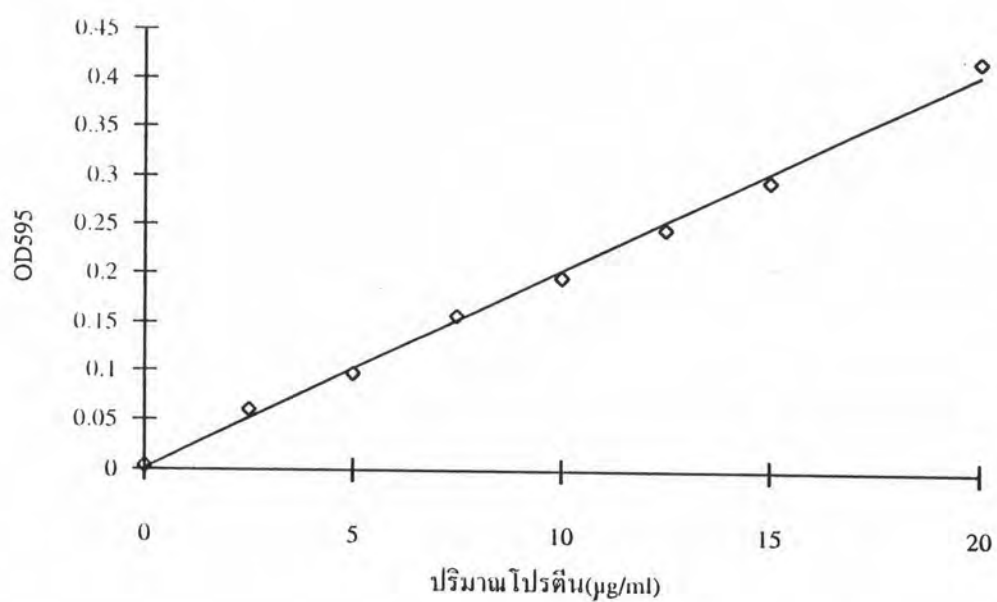
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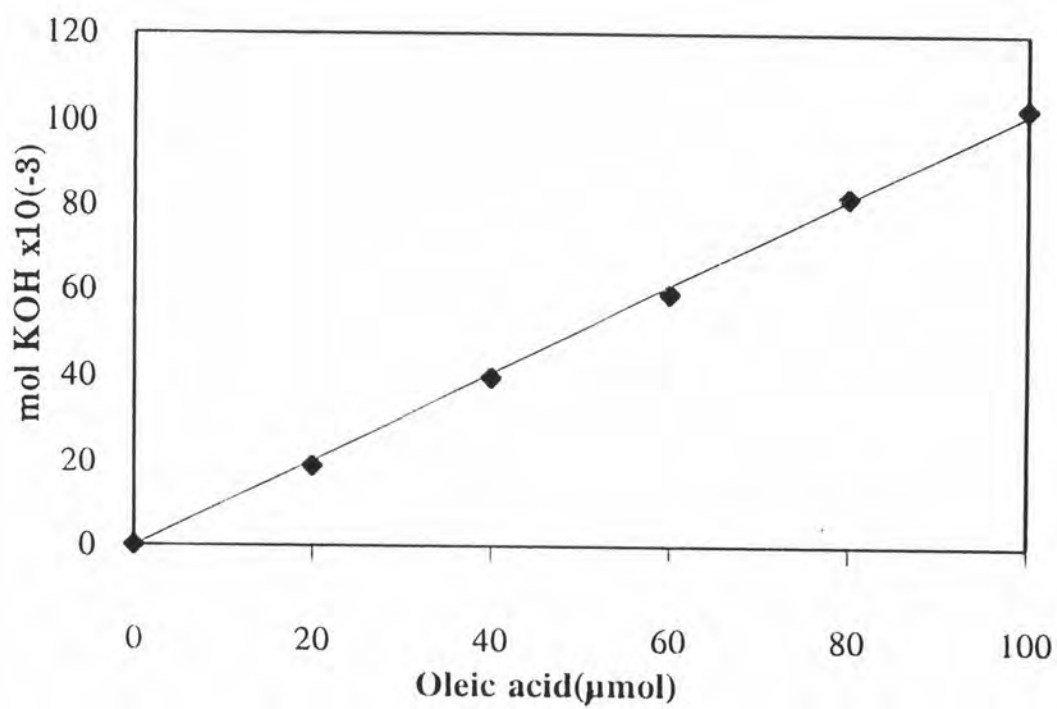
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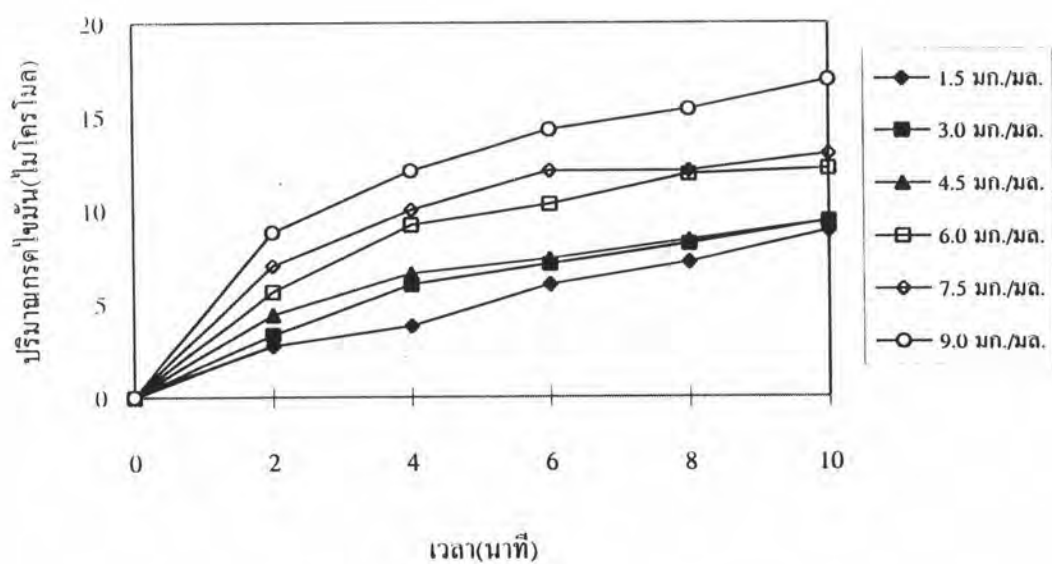
ภาคผนวก



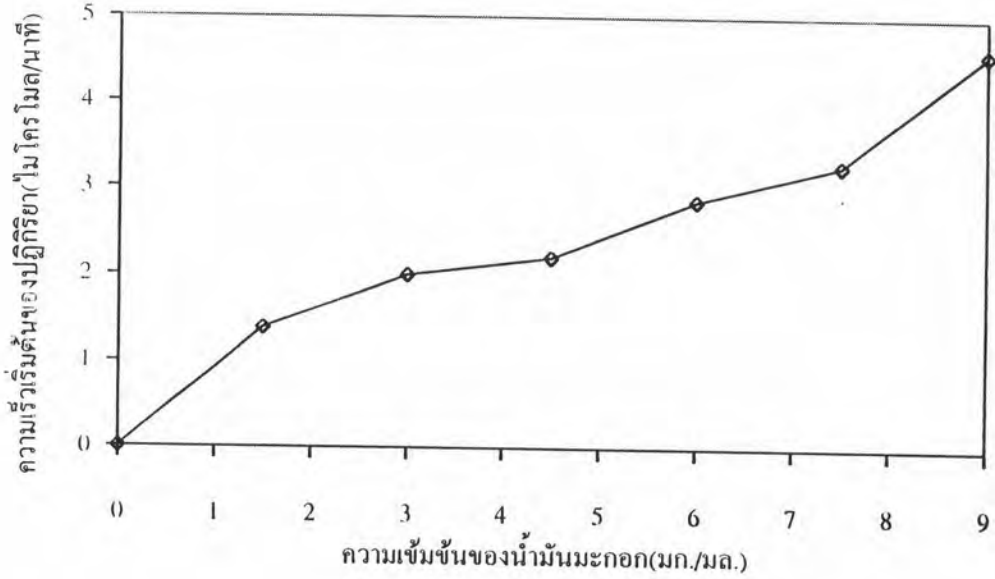
ภาคผนวกที่ 1 กราฟมาตรฐานสำหรับวิเคราะห์โปรตีนด้วยวิธีเบรดฟอร์ด
แปรความเข้มข้นของโปรตีนมาตรฐานคือ BSA ในช่วง
0-20 ไมโครกรัม(รายละเอียดวิธีทดลองตามข้อ 3.6)



ภาคผนวกที่ 2 กราฟมาตรฐานของกรดโอเลอิก แปรผันปริมาณความเข้มข้นตั้งแต่ 0-100 ไมโครโมล ซึ่งเมื่อหาความชันของกราฟได้เท่ากับ 1.02×10^{-3}



ภาพที่ 3 แสดงผลของความเข้มข้นของสับสเตรตต่ออัตราการเกิดปฏิกิริยา โดยใช้ไขมัน
มะกอกความเข้มข้นต่างๆกันเป็นสับสเตรต ดังวิธีทดลองข้อ 3.9.8



ภาพผนวกที่ 4 แสดงผลของความเข้มข้นของสับสเตรทต่อความเร็วเริ่มต้นของปฏิกิริยา

ประวัติผู้เขียน

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ศึกษาต่อระดับปริญญาโทที่ภาควิชา ชีวเคมี คณะวิทยาศาสตร์ จุฬาลงกรณ์มหาวิทยาลัย
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