

**CURETALL™**

**PROTOCOL NUMBER PH123012**

**A MULTICENTER DOUBLE-BLIND RANDOMIZED  
STUDY TO EVALUATE THE SAFETY AND EFFICACY  
OF CURETALL IN PATIENTS WITH ALLERGIES  
SUFFERING FROM CHRONIC ASTHMA**

**PROCEDURES IN CASE OF EMERGENCY****Table 1: Emergency Contact Information**

<b>Role in Study</b>	<b>Name</b>	<b>Address and Telephone number</b>
Clinical Study Leader	Jocelyn M. Taget, RN	2976 S. Market Street West Chester, PA 88765 215-787-5668 215-455-2323 (Cell)
Responsible Physician	Franklin L. Researcher, M.D.	2976 S. Market Street West Chester, PA 88765 215-787-5467
Drug Safety Physician	Rebecca L. Tanaka, M.D.	2743 N. Covington St. Allentown, PA 18897 610-446-5566
24-Hour emergency contact	Jocelyn M. Taget, RN	2976 S. Market Street West Chester, PA 88765 215-787-5668 215-455-2323 (Cell)

## 2. SYNOPSIS

<b>Name of Sponsor/Company:</b> <b>Pharma Corporation</b>	
<b>Name of Investigational Product:</b> <b>CuretALL</b>	
<b>Name of Active Ingredient:</b> <b>Sugar</b>	
<b>Title of Study:</b> A Multicenter Double-Blind Randomized Study to Evaluate the Safety and Efficacy of CuretALL in Patients with allergies suffering from chronic asthma	
<b>Study center(s):</b> <b>American Medical Research, Inc.</b> <b>1234 S. Lakeshore Blvd.</b> <b>Chicago, IL 99876</b>	
<b>Principal Investigator: Harry L. Tanaka, M.D.</b> <b>Investigators: Janet Jones, M.D., Lionel N. Winter, M.D., John Jones, R.N.</b>	
<b>Studied period (years):</b> Estimated date first patient enrolled: 01-September-2007 Estimated date last patient completed: 30-April-2008	<b>Phase of development:</b> III
<b>Objectives:</b> <b>Primary:</b> <ul style="list-style-type: none"> <li>To evaluate the efficacy of oral CuretALL across a range of doses for the treatment of allergies in patients suffering from chronic asthma</li> <li>To evaluate the safety of the long-term use of oral 5.0 mg CuretALL for the treatment of allergies in patients suffering from chronic asthma</li> </ul> <b>Secondary:</b> <ul style="list-style-type: none"> <li>Evaluation of tolerability and evaluation of long-term</li> </ul>	
<b>Methodology:</b> The detailed methodology is attached as Appendix 1.1	
<b>Number of patients (planned):</b> It is anticipated that a total of 800 patients will be randomized from approximately 40 centers in the United States.	

<p><b>Diagnosis and main criteria for inclusion:</b></p> <p>Patients suffering from allergies that require on-going medical monitoring with a minimum of 4 asthma attacks per year and a maximum of 18 asthma attacks per year.</p>
<p><b>Investigational product, dosage and mode of administration:</b></p> <p>CuretALL 5mg tablet daily compared to placebo</p>
<p><b>Duration of treatment:</b></p> <p>12 months</p>
<p><b>Reference therapy, dosage and mode of administration:</b></p> <p>Placebo tablet</p>
<p>Criteria for evaluation:</p> <p><b>Efficacy:</b></p> <p>Primary variable: 2-point scale allergy response at 1 hour</p> <p>Secondary variables: Secondary variables included measurement of 2-point scale allergy response at 30 minutes. 4-point scale allergy relief; change from baseline in allergy intensity symptoms at 30 minutes, 1 hour and 2 hours.</p> <p><b>Safety:</b></p> <p>Data will be collected and recorded for all adverse events, specified clinical laboratory tests, electrocardiogram (ECG) and vital signs, and medical history and physical examination findings.</p>
<p><b>Statistical methods:</b></p> <p>Binary response data will be analyzed using logistic regression for the primary and selected secondary efficacy variables; including terms for treatment, region, and baseline (either baseline severity 2-point scale or continuous baseline as appropriate) will be fitted irrespective of significance because intensity is known to influence efficacy. Analyses will be done primarily on the intent-to-treat (ITT) population with some analyses being done on the per-protocol (PP) population. All formal statistical tests for treatment difference will be performed using a 2-sided hypothesis test with a significance level of 0.030.</p>

### **3. TABLE OF CONTENTS, LIST OF TABLES, AND LIST OF FIGURES**

#### **TABLE OF CONTENTS**

1.	TITLE PAGE.....	1
2.	SYNOPSIS .....	3
3.	TABLE OF CONTENTS, LIST OF TABLES, AND LIST OF FIGURES .....	5
4.	LIST OF ABBREVIATIONS AND DEFINITIONS OF TERMS.....	9
5.	INTRODUCTION .....	10
6.	TRIAL OBJECTIVES AND PURPOSE.....	12
6.1.	Primary objective.....	12
6.2.	Secondary objectives .....	12
7.	INVESTIGATIONAL PLAN.....	13
7.1.	Overall Study Design and Plan: Description .....	13
8.	SELECTION AND WITHDRAWAL OF SUBJECTS.....	16
8.1.	Subject Inclusion Criteria .....	16
8.2.	Subject Exclusion Criteria .....	16
8.3.	Subject Withdrawal Criteria .....	16
9.	TREATMENT OF SUBJECTS.....	17
9.1.	Description of Study Drug.....	17
9.2.	Concomitant Medications .....	17
9.3.	Treatment Compliance.....	17
9.4.	Randomization and Blinding .....	17
10.	STUDY DRUG MATERIALS AND MANAGEMENT .....	18
10.1.	Study Drug.....	18
10.2.	Study Drug Packaging and Labeling .....	18
10.3.	Study Drug Storage.....	18
10.4.	Study Drug Preparation .....	18
10.5.	Administration .....	18
10.6.	Study Drug Accountability .....	18
10.7.	Study Drug Handling and Disposal .....	18
11.	ASSESSMENT OF EFFICACY .....	19
12.	ASSESSMENT OF SAFETY.....	20

12.1.	Safety Parameters .....	20
12.2.	Relationship to Study Drug .....	20
12.3.	Recording Adverse Events .....	20
12.4.	Reporting Adverse Events .....	20
13.	STATISTICS .....	21
14.	DIRECT ACCESS TO SOURCE DATA/DOCUMENTS.....	22
14.1.	Study Monitoring.....	22
14.2.	Audits and Inspections.....	22
14.3.	Institutional Review Board (IRB).....	22
15.	QUALITY CONTROL AND QUALITY ASSURANCE .....	23
16.	ETHICS .....	24
16.1.	Ethics Review .....	24
16.2.	Ethical Conduct of the Study .....	24
16.3.	Written Informed Consent .....	24
17.	DATA HANDLING AND RECORDKEEPING .....	25
17.1.	Inspection of Records .....	25
17.2.	Retention of Records .....	25
18.	REFERENCES .....	26
19.	APPENDICES .....	27

**LIST OF TABLES**

Table 1:	Emergency Contact Information.....	2
Table 2:	Abbreviations and specialist terms .....	9
Table 3:	Study design and schedule of assessments .....	15

## **LIST OF FIGURES**

Figure 1: Study design and schedule of assessments .....	14
----------------------------------------------------------	----



#### 4. LIST OF ABBREVIATIONS AND DEFINITIONS OF TERMS

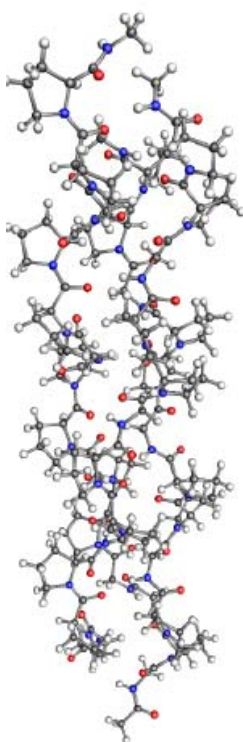
**Table 2: Abbreviations and specialist terms**

Abbreviation or specialist term	Explanation
AE	Adverse event
GCP	Good Clinical Practice
ICH	International Conference on Harmonization
IEC	Independent Ethics Committee
IRB	Institutional Review Board
OAE	Other significant adverse event
PI	Principal Investigator The investigator who leads the study conduct at an individual study center. Every study center has a principal investigator.
SAE	Serious adverse event

## 5. INTRODUCTION

### Background Information

The investigational product is CuretALL which is a compound created by Pharma Corporation for the purpose of relieving severe allergic symptoms and can be used by patients suffering from chronic asthma. The chemical structure for the compound is as follows:



### **Chemical Structure of CuretAll**

#### Nonclinical Summary

The findings from 12 nonclinical studies (Appendices 2.1 through 2.12) indicate acceptable results for development of CuretALL for the indications stated.

#### Potential Risks and Benefits

There are no known potential risks to humans based on studies to date. There is a potential benefit to patients suffering from chronic asthma who are also suffering from allergies.

#### Route of Administration, Dosage, Dosage Regimen and Treatment Periods

We believe the study will support the premise that a 5mg tablet taken orally upon rising will deliver the most efficacious dosage to study subjects.

GCP Requirements

The study will be conducted in compliance with the details of this Protocol and in accordance with current GCP requirements.

Population Studied

It is anticipated that a total of 800 patients will be randomized from approximately 40 centers in the United States providing a population of 750 patients who will actually complete the study.

Literature

A listing of relevant literature is attached as Appendix 3.1.

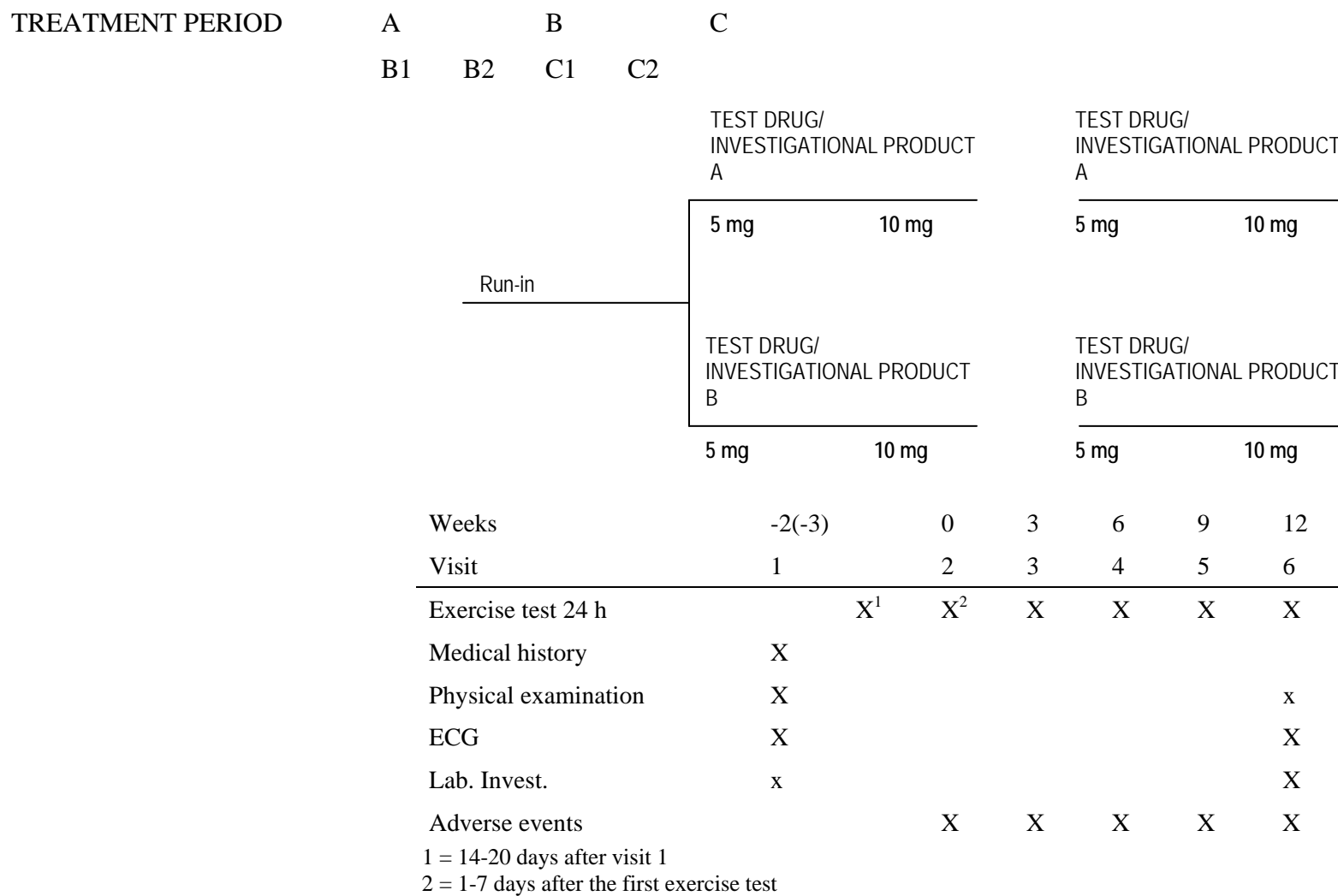
## **6. TRIAL OBJECTIVES AND PURPOSE**

- 6.1. Primary objective: To evaluate the efficacy of oral CuretALL across a range of doses for the treatment of allergies in patients suffering from chronic asthma.
- 6.2. To evaluate the safety of the long-term use of oral 5.0 mg CuretALL for the treatment of allergies in patients suffering from chronic asthma.
- 6.3. Secondary objectives: Evaluation of tolerability and evaluation of long-term

## **7. INVESTIGATIONAL PLAN**

### **7.1. Overall Study Design and Plan: Description**

The following is a full description of the study design and plan. This document would include specific details if this were a real Protocol.

**Figure 1: Study design and schedule of assessments**

**Table 3: Study design and schedule of assessments**

Assessment	Screening	Run-in	Baseline	Treatment					Follow-up	
Study Week	-2	-1	0	1	2	3	4	5	6	8
Informed Consent	X									
Inclusion/ Exclusion	X									
Medical History	X									
Physical Exam.	X									X
<u>Effectiveness</u>										
Primary variable	X	X	X	X	X	X	X	X	X	X
Secondary variable	X	X	X	X		X			X	X
<u>Safety</u>										
Adverse events	X	X	X	X	X	X	X	X	X	X
Lab. Tests	X		X	X			X		X	X
Body weight	X		X						X	X

## **8. SELECTION AND WITHDRAWAL OF SUBJECTS**

### **8.1. Subject Inclusion Criteria**

**Patients suffering from allergies that require on-going medical monitoring with a minimum of 4 asthma attacks per year and a maximum of 18 asthma attacks per year.**

### **8.2. Subject Exclusion Criteria**

**Patients that do not require on-going medical monitoring or patients that have <4 or >18 asthma attacks per year.**

### **8.3. Subject Withdrawal Criteria**

**Patients that do not report for study procedures, or voluntarily withdraw from the study. Subjects that withdraw from the study will not be followed.**



**9. TREATMENT OF SUBJECTS**

**9.1. Description of Study Drug**

**9.2. Concomitant Medications**

**9.3. Treatment Compliance**

**9.4. Randomization and Blinding**

## **10. STUDY DRUG MATERIALS AND MANAGEMENT**

### **10.1. Study Drug**

### **10.2. Study Drug Packaging and Labeling**

### **10.3. Study Drug Storage**

### **10.4. Study Drug Preparation**

### **10.5. Administration**

### **10.6. Study Drug Accountability**

### **10.7. Study Drug Handling and Disposal**

## **11. ASSESSMENT OF EFFICACY**

## **12. ASSESSMENT OF SAFETY**

### **12.1. Safety Parameters**

### **12.2. Relationship to Study Drug**

### **12.3. Recording Adverse Events**

### **12.4. Reporting Adverse Events**

## **13. STATISTICS**

## **14. DIRECT ACCESS TO SOURCE DATA/DOCUMENTS**

### **14.1. Study Monitoring**

### **14.2. Audits and Inspections**

### **14.3. Institutional Review Board (IRB)**

## **15. QUALITY CONTROL AND QUALITY ASSURANCE**

## **16. ETHICS**

### **16.1. Ethics Review**

### **16.2. Ethical Conduct of the Study**

### **16.3. Written Informed Consent**

Each subject will be required to agree in writing to an Informed Consent which is attached as Appendix 4.1.



## **17. DATA HANDLING AND RECORDKEEPING**

### **17.1. Inspection of Records**

### **17.2. Retention of Records**

## **18. REFERENCES**

## **19. APPENDICES**

### **1. Methodology**

#### **1.1 Detailed Methodology**

### **2. Non-Clinical Studies**

- 2.1. Name, date and title of Non-Clinical Study #1
- 2.2. Name, date and title of Non-Clinical Study #2
- 2.3. Name, date and title of Non-Clinical Study #3
- 2.4. Name, date and title of Non-Clinical Study #4
- 2.5. Name, date and title of Non-Clinical Study #5
- 2.6. Name, date and title of Non-Clinical Study #6
- 2.7. Name, date and title of Non-Clinical Study #7
- 2.8. Name, date and title of Non-Clinical Study #8
- 2.9. Name, date and title of Non-Clinical Study #9
- 2.10. Name, date and title of Non-Clinical Study #10
- 2.11. Name, date and title of Non-Clinical Study #11
- 2.12. Name, date and title of Non-Clinical Study #12

### **3. Relevant Literature**

#### **3.1 List of Relevant Literature**

### **4. Study Documents**

- 4.1 Informed Consent
- 4.2 Sample Case Report Forms

## Appendix 1.1 – Detailed Methodology

This document will provide details concerning the Methodology that will be used to conduct the following study:

CURETALL™

PROTOCOL NUMBER PH123012

A MULTICENTER DOUBLE-BLIND RANDOMIZED STUDY TO EVALUATE THE SAFETY AND EFFICACY OF CURETALL IN PATIENTS WITH ALLERGIES SUFFERING FROM CHRONIC ASTHMA

The number of subjects to be included in the study is: 800

Criteria for Inclusion: Patients suffering from allergies that require on-going medical monitoring with a minimum of 4 asthma attacks per year and a maximum of 18 asthma attacks per year.

The Investigational product is CuretALL 5mg tablet daily compared to placebo

The duration of treatment for all subjects is 12 months

The criteria for evaluation will be:

Efficacy:

Primary variable: 2-point scale allergy response at 1 hour

Secondary variables: Secondary variables included measurement of 2-point scale allergy response at 30 minutes. 4-point scale allergy relief; change from baseline in allergy intensity symptoms at 30 minutes, 1 hour and 2 hours.

Safety:

Data will be collected and recorded for all adverse events, specified clinical laboratory tests, electrocardiogram (ECG) and vital signs, and medical history and physical examination findings. Statistical methods: Binary response data will be analyzed using logistic regression for the primary and selected secondary efficacy variables; including terms for treatment, region, and baseline (either baseline severity 2-point scale or continuous baseline as appropriate) will be fitted irrespective of significance because intensity is known to influence efficacy. Analyses will be done primarily on the intent-to-treat (ITT) population with some analyses being done on the per-protocol (PP) population. All formal statistical tests for treatment difference will be performed using a 2-sided hypothesis test with a significance level of 0.030

## Appendix 2.1 – Non-Clinical Study #1

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

## Appendix 2.10 – Non-Clinical Study #10

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

## Appendix 2.2 – Non-Clinical Study #2

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

### Appendix 2.3 – Non-Clinical Study #3

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.



#### Appendix 2.4 – Non-Clinical Study #4

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

## Appendix 2.5 – Non-Clinical Study #5

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

## Appendix 2.6 – Non-Clinical Study #6

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

## Appendix 2.7 – Non-Clinical Study #7

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

## Appendix 2.8 – Non-Clinical Study #8

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

## Appendix 2.9 – Non-Clinical Study #9

This document would list all of the relevant information about the Non-Clinical Study, including title, synopsis, methodology, findings, results, summary and appendices.

### Appendix 3.1 – List of Relevant Literature

- 3.1.1 Article 1 (include title, author, date, synopsis)
- 3.1.2 Article 2 (include title, author, date, synopsis)
- 3.1.3 Article 3 (include title, author, date, synopsis)
- 3.1.4 Article 4 (include title, author, date, synopsis)
- 3.1.5 Article 5 (include title, author, date, synopsis)
- 3.1.6 Article 6 (include title, author, date, synopsis)
- 3.1.7 Article 7 (include title, author, date, synopsis)
- 3.1.8 Article 8(include title, author, date, synopsis)
- 3.1.9 Article 9 (include title, author, date, synopsis)
- 3.1.10 Article 10 (include title, author, date, synopsis)



**Investigational Study for Curetall – Protocol # \_\_\_\_ (Harry use the same Protocol Number that we are using for other documents)**

**Subject ID \_\_\_\_\_**

### **INFORMED CONSENT**

To be completed and signed by the subject  
and signed by the Investigator

Read each item below and sign in the space provided if you understand each item and agree to follow the instructions as outlined in the Protocol and presented to you by the study investigator or designate.

**Do not sign this Informed Consent and do not participate in this study if there is anything that you do not understand or about the information you have received about participating in this study.**

I understand that the therapies distributed in this study are medicines used to treat severe asthma and allergies.

The nature and purpose of this Clinical Study including its benefits and risks, have been fully explained to me by the Study Investigator. I have been given the opportunity to ask any questions regarding my participation in this study. All of my questions have been answered to my satisfaction.

I have been fully informed about the treatments provided in the study.

I understand that there are side effects that may happen while I am participating in the study. These have been explained to me.

I understand that I may receive the therapy being investigated, a comparable therapy or a placebo while participating in this study. Neither my physician nor the investigator conducting the study will have knowledge of the therapy that I am receiving.



All costs associated with physical examinations, treatment therapies, diagnostic tests, office visits, laboratory tests and any other medical treatment required by virtue of my participation in this study will be covered by Health Corporation.

I agree to comply with all restrictions as outlined in the Protocol and agree not to take any medications not expressly permitted under the Protocol as explained to me by the investigator.

I agree to be available for all scheduled visits and to strictly comply with the dosing routine as outlined in the Protocol.

I acknowledge that I have received all relevant information and copies of documents (including a copy of this Informed Consent), as outlined in the GCP's.

Subject signature \_\_\_\_\_ Date \_\_\_\_\_

Witness signature: \_\_\_\_\_ Date \_\_\_\_\_