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Controlled Release of Insulin and Modified Insulin from a Novel Injectable Biodegradable Gel

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CONTROLLED RELEASE OF INSULIN AND MODIFIED INSULIN FROM A NOVEL INJECTABLE BIODEGRADABLE GEL

A Dissertation
Presented for
The Graduate Studies Council
The University of Tennessee
Health Science Center

In Partial Fulfillment
Of the Requirements for the Degree
Doctor of Philosophy
From The University of Tennessee

By
Om Anand
December 2008
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DEDICATION

This dissertation is dedicated to my parents, Mr. Naresh Chandra and Mrs. Roshni Devi and my wife Mrs. Geetanjali Dabas and my son Ishan, for their endless love and support.
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ABSTRACT

The objective of the study was to develop a controlled release dosage form of insulin, which can provide basal concentrations of insulin in diabetic rats for 1 to 2 weeks after a single subcutaneous injection.

A biodegradable injectable drug delivery gel was prepared by dissolving a biodegradable polymer, polylactic-co-glycolic acid (PLGA), in biocompatible plasticizer(s), triethyl citrate (TEC) and/or acetyl triethyl citrate (ATEC). Insulin was then loaded into the blank gel to form an insulin suspension in the polymer solution. After the insulin-loaded gel was injected subcutaneously, the plasticizer(s) dissolved in the aqueous media and were gradually taken away from the gel. The polymer precipitated after the plasticizer(s) were extracted by the aqueous medium and a solid depot of insulin was formed. The insulin was released slowly from the depot by a combination of drug diffusion and erosion of the polymer.

In the first part of this study, the effect of different water-soluble and water-insoluble zinc salts on blood glucose lowering effect of insulin in type-2 diabetic ZDF rats was investigated. Insulin formulations containing varying concentrations of different water-soluble and water-insoluble zinc salts were prepared and injected subcutaneously in type-2 ZDF rats and blood glucose concentration lowering effect was studied. Insulin in presence of water-soluble salts of zinc could suppress blood glucose concentrations in ZDF rats for up to 16 hours.

Insulin was loaded into different gel formulations (5% PLGA (i.v. 0.09, acid end group), ATEC:TEC (3:1) and 4% insulin) and tested in vivo. However, these insulin-loaded gel formulations only suppressed the blood glucose concentrations in the ZDF rats for 1 day after a single subcutaneous injection. In order to achieve longer control over the release of insulin from the gels, a water-soluble salt, zinc sulfate was incorporated in these insulin containing gels at different concentrations. A biodegradable injectable gel formulation prepared with zinc sulfate was able to maintain low blood glucose concentrations for up to 8 to 10 days following a single subcutaneous injection.

In order to achieve better glucose control after the release of insulin from the gels, insulin glargine particles were purified from commercially available Lantus® formulation. The freeze dried insulin glargine particles were then loaded into the blank gels and tested in vivo. The formulation prepared with 5% PLGA (i.v. 0.09, acid end group), ATEC:TEC (3:1) and 4% insulin glargine was able to suppress the blood glucose concentrations of the ZDF rats significantly for 10 days after a single subcutaneous injection. The concentration of insulin glargine was maintained between $260 \pm 134.9$ mIU/L and $188 \pm 55.9$ mIU/L until day 10 after single subcutaneous injection. The addition of zinc sulfate to the formulations prepared with purified insulin glargine particles further slowed down the drop in blood glucose concentrations.
# TABLE OF CONTENTS

## CHAPTER 1. INTRODUCTION .......................................................................................... 1

1.1 Diabetes mellitus ........................................................................................................ 1
   1.1.1 Type-1 diabetes .................................................................................................. 1
   1.1.2 Type-2 diabetes ................................................................................................ 2

1.2 Insulin .......................................................................................................................... 3
   1.2.1 The structure of human insulin ......................................................................... 4
   1.2.2 Insulin and its aggregates ................................................................................ 6
   1.2.3 Insulin zinc and different state ........................................................................ 7
   1.2.4 Insulin zinc interaction and pH ........................................................................ 9
   1.2.5 Insulin mechanism of action and degradation ................................................ 9

1.3 Treatment of diabetes .................................................................................................. 10

1.4 Non-invasive insulin delivery systems ........................................................................ 14
   1.4.1 Oral insulin delivery of insulin ......................................................................... 14
   1.4.2 Nasal delivery of insulin .................................................................................. 17
   1.4.3 Pulmonary-lung delivery of insulin .................................................................. 18
   1.4.4 Ocular delivery of insulin ................................................................................. 18
   1.4.5 Rectal delivery of insulin ................................................................................ 19

1.5 Controlled release of insulin ....................................................................................... 19
   1.5.1 Artificial pancreas ............................................................................................ 19
   1.5.2 Glucose sensitive hydrogels .............................................................................. 20
   1.5.3 Extended release of insulin .............................................................................. 20
   1.5.4 Insulin pumps .................................................................................................. 21
   1.5.5 Insulin gels ........................................................................................................ 21

## CHAPTER 2. PROLONGED EFFECT OF INSULIN IN PRESENCE OF WATER SOLUBLE AND WATER INSOLUBLE SOLUBLE ZINC SALTS ........................................................................ 23

2.1 Introduction .................................................................................................................. 23

2.2 Materials and methods ............................................................................................... 24
   2.2.1 Materials ........................................................................................................... 24
   2.2.2 Methods ............................................................................................................ 24

2.3 Results and discussion ............................................................................................... 31
   2.3.1 Pharmacodynamic studies of insulin in ZDF rats .............................................. 31
   2.3.2 In vitro release of insulin from insulin formulations with varying concentrations of zinc sulfate ................................................................. 38

2.4 Conclusions .................................................................................................................. 46

## CHAPTER 3. CONTROLLED RELEASE OF INSULIN FROM NOVEL BIODEGRADABLE INJECTABLE GELS .............................................................................. 49

3.1 Introduction .................................................................................................................. 49
3.2   Materials and methods ................................................................. 50
   3.2.1  Materials ................................................................. 50
   3.2.2  Methods ................................................................. 50
3.3   Results and discussion .......................................................... 56
3.4   Conclusions .............................................................................. 67

CHAPTER 4. CONTROLLED RELEASE OF MODIFIED INSULIN
GLARGINE FROM NOVEL BIODEGRADABLE INJECTABLE GELS ..........70
4.1   Introduction .................................................................................. 70
4.2   Materials and methods ................................................................. 71
   4.2.1  Materials ................................................................. 71
   4.2.2  Methods ................................................................. 71
4.3   Results and discussion .............................................................. 76
   4.3.1  Purity of insulin glargine particles .............................................. 76
   4.3.2  Pharmacodynamic studies of insulin formulations in ZDF rats ....... 79
   4.3.3  In vivo studies of gel formulations loaded with insulin glargine
          particles .............................................................................. 79
4.4   Conclusions .............................................................................. 95

LIST OF REFERENCES ................................................................................. 97
VITA ..................................................................................................................119
LIST OF TABLES

Table 1-1  Insulin and modified insulin preparations.......................................................11
Table 2-1  Composition of insulin formulations containing zinc sulfate.........................25
Table 2-2  Composition of insulin formulations containing zinc chloride........................26
Table 2-3  Composition of insulin formulations containing zinc carbonate.......................27
Table 2-4  Composition of insulin formulations containing zinc oxide...........................28
Table 3-1  Formulations of insulin suspensions...............................................................51
Table 3-2  Formulation composition of PLGA gel containing insulin and zinc ..............52
Table 3-3  Formulation composition of PLGA gel containing insulin and zinc sulfate
physical mixtures............................................................................................54
Table 3-4  Zinc contents in insulin-zinc co-precipitates ..................................................55
Table 3-5  Formulation composition of PLGA gel of insulin..........................................57
Table 4-1  Composition of gel formulations used for in vivo studies .............................74
LIST OF FIGURES

Figure 1-1  Structure of human insulin ............................................................... 5

Figure 2-1  Blood glucose concentrations of ZDF rats after a single subcutaneous injection of insulin at a dose of 10 IU/kg (Mean ± SEM, n = 6) .................32

Figure 2-2  Comparison of blood glucose concentrations in ZDF rats after a single subcutaneous injection of insulin with varying concentrations of zinc sulfate (Insulin dose-10 U/kg) (Mean ± SEM, n = 4) ........................................33

Figure 2-3  Comparison of blood glucose concentrations in ZDF rats after a single subcutaneous injection of insulin with varying concentrations of zinc chloride (Insulin dose-10 U/kg) (Mean ± SEM, n = 4) ..................................................34

Figure 2-4  Comparison of blood glucose concentrations in ZDF rats after subcutaneous injection of insulin, ultralente and insulin with varying concentrations of zinc carbonate (Insulin dose-10 U/kg) (Mean ± SEM, n = 4) ..................................................................................36

Figure 2-5  Comparison of blood glucose concentrations in ZDF rats after a single subcutaneous injection of insulin with varying concentrations of zinc oxide (Insulin dose-10 U/kg) (Mean ± SEM, n = 4) ..................................................37

Figure 2-6  Comparison of blood glucose concentrations in ZDF rats after a single subcutaneous injection of insulin with water-soluble and water-insoluble zinc salts (Insulin dose-10 U/kg) (Mean ± SEM, n = 4) ..................................................39

Figure 2-7  Comparison of in vitro insulin release from insulin suspension formulations with and without zinc sulfate .........................................................40

Figure 2-8  Comparison of in vitro insulin release from insulin suspension formulations with zinc sulfate .................................................................41

Figure 2-9  Percentage of insulin remaining in the dialysis bags after completion of in vitro release study of insulin formulations with varying concentrations of zinc ..............................................................................42

Figure 2-10 Concentrations of insulin in the supernatant obtained from formulations of insulin alone or insulin with zinc sulfate (0.24mg/mL) at different pH values .................................................................44

Figure 2-11 pH and concentration of insulin in the supernatant obtained from formulations of insulin (RHI) with varying concentrations zinc sulfate .......45
Figure 2-12  Concentration of insulin (RHI) in the supernatant obtained from formulations of insulin alone at different pH, or formulations of insulin with varying concentrations zinc sulfate.................................47

Figure 2-13  Possible mechanism of extended \textit{in vivo} effect of formulations of insulin with varying concentrations zinc sulfate.................................................................48

Figure 3-1  Changes in blood glucose concentrations of ZDF rats after a subcutaneous injection of PLGA (5% and 10%) gel formulations loaded with 4% insulin (Formulations OA-3-1 and OA-3-2) (Mean ± SEM, n = 5) .....................................................................................................................58

Figure 3-2  Changes in body weights of ZDF rats after subcutaneous injections of PLGA (5 and 10%) gel formulations loaded with 4% insulin (Formulations OA-3-1 and OA-3-2) (Mean ± SEM, n = 5) ......................60

Figure 3-3  Changes in blood glucose concentrations of ZDF rats after subcutaneous injections of PLGA (5%) gel formulations loaded with 4% insulin alone (Formulation OA-3-1) or insulin-zinc co-precipitates (Formulations OA-3-7 and OA-3-8) (Mean ± SEM, n = 5) .................................................................61

Figure 3-4  Changes in blood glucose concentrations of ZDF rats after subcutaneous injections of PLGA (5%) gel formulations loaded with 4% insulin alone (Formulation OA-3-1) or insulin-zinc co-precipitates (Formulations OA-3-9 and OA-3-10) (Mean ± SEM, n = 5) ................................................................................................................63

Figure 3-5  Changes in body weights of ZDF rats after subcutaneous injections of PLGA (5%) gel formulations loaded with 4% insulin-zinc co-precipitates (Formulations OA-3-7, OA-3-8, OA-3-9 and OA-3-10) (Mean ± SEM, n = 5) ................................................................................................................65

Figure 3-6  Changes in blood glucose concentrations of ZDF rats after subcutaneous injections of insulin alone (Formulations OA-3-1) or physical mixtures of insulin (4%) and zinc sulfate (1 or 2%) loaded in PLGA (5%) gel formulations (Formulations OA-3-11 and OA-3-12) (Mean ± SEM, n = 5) .................................................................................................66

Figure 3-7  Changes in blood glucose concentrations of ZDF rats after subcutaneous injections of insulin loaded in PLGA (5%) gel formulations with different concentrations of zinc sulfate (2.8% and 5.4%) (Formulations OA-3-13 and OA-3-14) (Mean ± SEM, n = 5) ..............................................................................................................68

Figure 3-8  Changes in body weights of ZDF rats after subcutaneous injections of PLGA (5%) gel formulations loaded with 4% insulin and zinc sulfate mixtures (Formulations OA-3-11, OA-3-12, OA-3-13 and OA-3-14) (Mean ± SEM, n = 5) ..............................................................................................................69
Figure 4-1  HPLC chromatogram of placebo of Lantus® formulation ........................................77

Figure 4-2  HPLC chromatogram of Lantus® formulation .........................................................78

Figure 4-3  HPLC chromatogram of insulin glargine particles dissolved in HCl .................80

Figure 4-4  Blood glucose profiles in ZDF rats after subcutaneous injections of insulin (RHI), Lantus® and insulin glargine particles suspended in saline (dose-10 U/Kg) (Mean ± SEM, n = 5) .................................................................81

Figure 4-5  Changes in blood glucose concentrations and body weights of ZDF rats after subcutaneous injections of 4% suspension of insulin glargine particles in PBS (Mean ± SEM, n = 5) .................................................................82

Figure 4-6  Changes in blood glucose concentrations and body weights of ZDF rats after subcutaneous injections of 4% suspension of insulin glargine particles in blends of ATEC:TEC (3:1) (Mean ± SEM, n = 5) .........................................................84

Figure 4-7  Changes in blood glucose concentrations and body weights of ZDF rats after a subcutaneous injection of gel (5% PLGA) formulation loaded with 2% insulin glargine particles (Mean ± SEM, n = 5) .........................................................85

Figure 4-8  Changes in blood glucose concentrations and body weights of ZDF rats after a subcutaneous injection of gel (5% PLGA) formulation loaded with 4% insulin glargine particles (Mean ± SEM, n = 5) .........................................................86

Figure 4-9  Changes in blood glucose concentrations and body weights of ZDF rats after subcutaneous injections of gel (10%) formulations formulation loaded with 4% insulin glargine particles (Mean ± SEM, n = 5) ........................................88

Figure 4-10 Comparison of blood glucose concentrations in ZDF rats after subcutaneous injections of insulin glargine (4% loading) in different formulations (Mean ± SEM, n = 5) ........................................................................89

Figure 4-11  Blood glucose concentrations and concentrations of insulin glargine in serum of ZDF rats after a single subcutaneous injection of Formulation OA-G-1 (Mean ± SEM, n = 7) .................................................................................90

Figure 4-12  Blood glucose concentrations and concentrations of insulin glargine in serum of ZDF rats after a single subcutaneous injection of Formulation OA-G-4 (Mean ± SEM, n = 6) .................................................................................92
Figure 4-13  Comparison of serum glargine concentrations in ZDF rats after subcutaneous injections of 4% insulin glargine loaded in PLGA (5%) gel formulation (Formulation OA-G-4) and PBS .................................................93

Figure 4-14  Comparison of blood glucose concentrations in ZDF rats after injections of insulin glargine (4% loading) in different formulations with various zinc sulfate concentrations (Mean ± SEM, n = 5) .........................................................94
CHAPTER 1. INTRODUCTION

1.1 Diabetes mellitus

Diabetes mellitus is a chronic disorder. It may be defined as “a metabolic disorder of multiple etiologies characterized by chronic hyperglycemia with disturbances of carbohydrate, fat and protein metabolism resulting from defects in insulin secretion, insulin action, or both” (1).

Even though diabetes has been known to humans from early historical periods, it is only recently that the WHO (World Health Organization) has recognized it as first non-infectious, worldwide epidemic (2). The cause of diabetes was unknown for centuries. Then in 1889, Joseph Von Mering and Oskar Minkowski demonstrated in dogs that the disease was associated with pancreas (3). In 1908, a German scientist, Georg Zuelzer prepared a crude extract from animal pancreas and unsuccessfully tried to treat human diabetic subjects (4). In 1921, one of the most phenomenal discoveries of medical history was done by Frederick Banting and Charles Best. They isolated an antidiabetic substance, which was later named as “isletin”, from dog pancreas and injected in two pancreatectomized, hyperglycemic dogs. The blood glucose concentrations of these ‘isletin’ injected dogs reduced significantly. John Macleod and James Collip further purified the extract from pancreas and named ‘isletin’ as insulin. They successfully used insulin to treat a 14 year old diabetic boy (4). This historical treatment marked the beginning of the use of insulin in the treatment of diabetes. In 1923, Frederick Banting and John Macleod were awarded the Nobel Prize in physiology for their discovery of insulin (5). Later, Banting shared his Nobel Prize with Best, and soon after Macleod shared his Noble Prize with Collip (1).

1.1.1 Type-1 diabetes

It is a well established fact now that diabetes mellitus is not a single disease but a combination of different types of diseases. It is broadly categorized as diabetes type-1, diabetes type-2 and gestational diabetes (6).

The diabetes type-1 is caused by deficiency of insulin, and has more common juvenile onset which is active in nature and requires insulin delivery for survival (6). The deficiency of insulin is caused by autoimmune destruction of beta cells in the pancreas.

The type-1 diabetes may further be divided into two classes: type-1 A and type-1 B. Type 1 A disease is due to autoimmune destruction of the beta cells in pancreas, thus reducing the biosynthesis of insulin. Antibodies against the beta cells antigen have been found in those subjects. Type-1 B disease is idiopathic, i.e. no beta cell antibodies are found in the subjects (6).

1.1.1.1 Role of genetics toward susceptibility to type-1 diabetes. It is a well recognized fact that genetics plays an important role in type-1 diabetes. For example, the
concordance rate in monozygotic twins is much higher than the dizygotic twins (6-8). Further, some ethnic groups are more susceptible to type-1 diabetes than others. The first degree relative of type-1 subjects are 15 fold more prone to suffer from type-1 diabetes (6, 9). Davis *et al.* reported that the major genetic determinant of susceptibility to type-1 diabetes is HLA region of chromosome 6 (10). The insulin gene region on chromosome 11 and other confounded genes have also been implicated.

1.1.1.2 Environmental factors. The concordance rate for the development of type-1 diabetes in monozygotic twins is not 100%. Further, there are reports suggesting seasonal peak in the case of type-1 diabetes detection (9). This indicates of some environmental factors or the possible interaction of gene and environmental factors are playing a role in the development of type-1 diabetes (9). One of the Scandinavian studies indicated a correlation between an increase in diabetes incidences and the decline in breast feeding. The study suggested that tolerance to the common antigen might not be developed if children were exposed to cow milk protein in their early ages (11). Certain virus infection, for example Poliovirus, entero virus, and echovirus, have been suggested as the possible reasons for the autoimmune responses against the beta cells of pancreas (12). Exposer to virus infection during early childhood or pregnancy have been associated with type-1 diabetes as well (13-16).

1.1.1.3 Prevention of type-1 diabetes. There are various prevention trial studies going on around the world. The trials may be primary, secondary or tertiary. The aim of these trials is slowing the progression of type-1 diabetes (6). The primary prevention trials are before development of autoimmunity and are conducted in patients who are more prone to diabetes due to their genetic make up. Secondary preventive trials are those which are conducted in subjects showing signs of autoimmunity, but are not diabetic yet. Cyclosporine, an immunosuppressant, insulin and nicotinamide, a vitamin B₃ derivative, alone or in combinations have been injected in these subjects, and development of diabetes has been studied. Some of these trials have been effective, but only for an year (17).

1.1.2 Type-2 diabetes

Type-2 diabetes or non insulin dependent diabetes mellitus is also known as maturity onset diabetes mellitus (18). Type-2 diabetes is a complex disorder that might be due to one or more than one gene, environmental factors or interaction of both (18, 19). In type-2 diabetes, there may be no loss or some loss of beta cells, and insulin concentrations in the blood may be high, normal or low. Type-2 diabetes subjects have a high degree of genetic predisposition, and the onset is generally late (18, 20). Type-2 diabetes, in most cases has a slow progression. The interaction between genetic and environmental factors affects the pathogenesis of type-2 diabetes (18). Genetically predisposed subjects follow a particular trend in the progression of diabetes. The normal subjects develop insulin resistance, hyperinsuleneimia, glucose desensitization, and insulin secretion defects leading to impaired glucose tolerance followed by hyperglycemia leading to type-2 diabetes (18, 21). The type-2 diabetes may also be due to various reasons such as, malfunction of gluco-receptor on the surface of cells, so that
they start responding to only high glucose concentration in the blood. There may be down
regulation of insulin receptors leading to reduction in frequency of these receptors. The
presence of hyperglycemic hormones like glucagon and physical state like obesity may
also result in insulin deficiency leading to type-2 diabetes.

Glucose concentrations in the blood are controlled by three mechanisms; hepatic
 glucose production, peripheral tissue glucose uptake (glucose uptake by muscles
 adipocytes and hepatic cells) and insulin secretion by beta cells (18, 22). There exists a
critical balance between these three mechanisms. Factors affecting any of these
biochemical processes may lead to type-2 diabetes. An increase in hepatic glucose
production by increasing glycogenolysis or gluconeogenesis increases glucose
concentrations (23). This may be due to a decrease in the effectiveness of insulin or low
insulin concentrations, or an increase in the glucagon concentrations. A combination of
the above reasons may also contribute to the increased blood glucose dumping by the
liver, thus leading to diabetes type-2 (24-27).

Genetic components have a strong influence in the development of type-2
diabetes (18). It has been well recognized that type-2 diabetes is multi-genetic in nature.
Different candidate genes that may effect the development and progression of type-2
diabetes have been studied. These genes have been further divided into two categories.
The first category contains candidate genes for insulin secretary defects, while the second
category has candidate gene for insulin resistance (18). Although various candidate genes
affecting the insulin secretion and resistance have been studied, no single gene mutation
that is responsible for the cause and progression of type-2 diabetes has been identified
(18).

1.2 Insulin

Insulin is a metabolic hormone synthesized, stored, and secreted from beta cells of
pancreas. In an important landmark, Abel et al. successfully crystallized insulin in 1927
(28). Scott and Fisher in 1935 studied the role of zinc in insulin crystallization (29-31).
The crystallization techniques helped in improving the yields and purity of insulin. In
1936, Hagedon came out with insulin protamine suspensions, which increased the
duration of antidiabetic effect of insulin and lowered the hypoglycemic reactions (32).
However, the protamine insulin was not stable and had a tendency to stick to the walls of
the vials, thus reducing its potency (33, 34). This problem was solved by Scott and Fisher
by adding zinc salts to the insulin protamine formulation and forming protamine zinc
insulin. This not only stabilized the insulin protamine formulation for more than six
months, but also prolonged the absorption time of insulin (33, 34). In 1951, Hallas-
Moller et al. came out with an idea of Lente insulins. Semilente insulin was a suspension
of amorphous insulin and zinc particles, and had a longer action than normal insulin.
Ultralente was a crystalline modification (crystallized in acetate buffer) with a longer
action similar to protamine zinc insulin. Lente insulin formulation contained a mixture of
semilente and ultralente for rapid onset and prolonged action (35-40). In 1956, Frederick
Sanger and co workers elucidated the amino acid sequence of insulin, which was found to
be a two chain polypeptide having 51 amino acids (41, 42). The A-chain has 21 amino acids and the B chain has 30 amino acids. In 1958, Sanger was awarded the Nobel Prize in Chemistry for this work (43). Until then, the main source of insulin was pork or beef. Pork and beef insulin are found to be slightly different from the human insulin. Pork insulin has alanine in place of threonine at the B-30 position, whereas, the beef insulin differs in three amino acids. Thus pork insulin is more homologous to human insulin. However, due to differences in the amino acid sequences and presence of other impurities, these insulin formulations were responsible for a lot of hypersensitivity reactions (41, 42).

The knowledge of amino acid sequence made it possible to chemically synthesize human insulin. In 1963, Meienhofer et al. and Katsoyannis et al. successfully synthesized human insulin (44-49). Kung et al. also synthesized human insulin in 1966 (50-53). However, the amounts were never sufficient for a clinical trial and therefore, pork and bovine insulin remained in continuous use. The development in analytical and purification techniques in the 1960s led to the production of a much purer form of these animal insulins, thus reducing the cases of hypersensitivity and antigenicity in the human beings. In 1967, Steiner and Oyer published the biosynthesis of insulin and described that insulin is synthesized as proinsulin in the beta cells of pancreas, and the enzymatic conversion of proinsulin in the storage vessels produces insulin (54, 55). Dorothy Crowfoot Hodgkin, who had been awarded a Nobel Prize in Chemistry in 1964 for the development of crystallography, later elucidated the spatial conformation of the insulin molecule (43). X-ray crystallography was further used by Adams et al. to determine the structure of insulin crystals (56).

Radioimmunoassay, for determination of very low concentrations of insulin in diabetic subjects, was developed by Rosalyn Sussman Yalow and Solomon Berson. In 1977, Rosalyn Sussman Yalow was awarded the Nobel Prize in Medicine for her work on insulin (5). In 1979, Inouye et al. completed enzymatic semi-synthesis of human insulin by using porcine insulin as a starting material. This technique immobilized trypsin, which was helpful for continuous production of insulin (57-59). Hence, industrial production of human insulin was finally feasible.

The 1970s was also an era of rapid advances in recombinant DNA technology which helped in the development of new methods for biosynthesis of human insulin. Genentech Inc. in 1978 used genetically engineered Escherichia coli to produce two chains of insulin separately and then joined the two chains chemically to produce human insulin (60). Novo Nordisk in 1982 commercially launched the first human insulin which was produced by the semi synthetic method (61). However, soon after, in the same year, Eli Lilly used Genentech’s method and got approval from the US FDA to market the human insulin - the first human protein produced by the recombinant DNA technology.

1.2.1 The structure of human insulin

The primary structure of human insulin consists of two polypeptides (Figure 1-1). The shorter chain A, which consists of 21 amino acids (A1-A21) and the longer chain B
Figure 1-1 Structure of human insulin
consists of 30 amino acids (B1-B30). The chains are connected together by two interchain disulfide bonds between A7 and B7, and A20 and B19 cystines. In the A chain, an intra-chain disulfide bond is also present between A6 and A11 cystines. The amino acids of the two chains also interact with one another by non covalent bonds.

In three dimensional conformation of human insulin, the A chain contains two alpha helix segments—one from A2 to A8, and the another from A13 to A20. These two alpha helices are antiparallel. The N-terminal of B chain contains an alpha helix from B9 to B19, whereas, B21 to B30 forms a beta strand (62, 63). Wollmer et al. and Derewenda et al. explained that the A chain and the B9 to B19 structural units of B chain are stable, whereas, the conformations of B1 to B8 and B25 to B30 change in the presence of certain chemicals like phenols. In presence of phenol, the B1 to B8 segment extends to form the alpha helix (64-66).

Human insulin exists in different forms, depending on the insulin concentration, pH and presence of zinc ions. Monomers, i.e. single unit of insulin are only present at very low concentration (<0.1 µM or 0.6 mg/L) (67).

1.2.2 Insulin and its aggregates

The insulin solution exits in a complex and dynamic state of equilibrium. Based on the insulin concentration, pH and presence of other ions, insulin may be present as monomers, dimers, tetramers or hexamers, which may further form insulin hexamer associates (67-71).

Insulin in solution at very low concentrations (< 0.1 µmol ~ 0.6 µg/ml) exists as monomers (single units) (63, 67, 71). Insulin in blood is also found in predominantly monomeric form, which is also considered as the physiologically relevant form. Insulin monomer, due to its non-polar surface forces and hydrogen bond formation potentials, has a strong tendency to associate and form compact dimers (71). The dimers exist in a state of equilibrium with monomers and hexamers (71). The major interaction in monomers to form dimers occurs at B8, B9, B12, B13, B16, and C terminus segment B23 to B28. The two COOH-terminal strands of B chain are arranged in the form of antiparallel beta sheets.

The major interaction in monomers to form dimers occurs at B8, B9, B12, B13, B16, and C terminus segment B23 to B28. The two COOH-terminal strands of B chain are arranged in the form of antiparallel beta sheets. This allows close association of non-polar forces and formation of four hydrogen bonds between B24 and B26 amino acids (71). In acidic solutions (below pH 4), the insulin molecules are predominantly in dimeric state. The dimers are flexible in nature and are prone to degradation, deamidation and fibrillation in an acidic environment. This may be due to the interaction of exposed non-polar surfaces which may interact and form heterogeneous aggregates and fibrils (67, 71).

The insulin hexamer, formed by the association of three dimers, is a nearly spherical structure with a height of approximately 35Å and a diameter of 50 Å (71).
center of this spherical hexamer form is a polar channel. Two zinc ions form a coordination bond with the imidazole group of B10 histidine, which is present in the centrally located polar channel of the hexamer. The addition of zinc ions or other divalent transition metals promote the formation of hexamers (72, 73).

In solution, the insulin hexamers bind with metal ions and exist in three conformational states, namely, T6, T3R3 and R6. In solution, the three states of hexamers exist in a dynamic equilibrium. Addition of inorganic anions or adequate concentration of phenol like molecule shifts this dynamic equilibrium from T6 to T3R3 (72). Higher concentration of phenols further strengthens the hexamers by shifting the equilibrium completely toward the conformational state R6 (74). In all of these three conformations, the A chains folds into a helix-loop-helix motif, A2 to A8 and A12 to A19 forms two helical structures (74). The B chain can take two different conformations. The B chain in T state contains “an extended N-terminal arm”, a central α-helix (B9 to B19), a type I turn (B20 to B23) and a C terminal B strand “In the R state, residue B1 to B9 takes up a helical conformation to form a region of alpha-helix contiguous from B1 to B9” (74).

1.2.3 Insulin zinc and different state

In T6 state, the six monomeric units exists in the T fold, whereas, in T3R3 state, three monomeric units exist in T fold and three exists in R fold. In the R state, the all six monomeric units exist in the R fold. Brange et al. reported that A13, A14, A17 residue of A chain are involved in hexamer formation (71). In B chain, B1, B2, B4, B10, B13, B14, B17, B18, B19, B20 residues play a major role in hexamer formation. Zinc ions are key to formation of hexamers and further stabilizing them. Kadima et al. reported that T6 or two zinc “insulin hexamer is an assembly of three equivalent dimers related by 3 fold axis of symmetry” and an identical symmetry is expected in solution (75). In crystalline rhombohedral state of T6, there are two equivalent zinc binding sites. These two sites are present along the three fold axis of symmetry. In each of these two sites, there are three equivalent histidine B10 residues. Zinc forms coordinate bonds with nitrogen of imidazole side chain of these histidine. The other coordinate positions of the zinc are occupied by three water molecules or other ligands (75, 76). Hence, in T6 conformation state, each of the two zinc ions are coordinated with nitrogens from three His-B10 imidazole groups and three water molecules. However, Emdin et al. reported that T6 states are capable of binding more than two zinc ions. The middle of the hexamer also contains a calcium binding site. The calcium binding site is formed by the six carboxylate groups of B13 glutamate residue, and can bind three calcium ions (75).

The dimer in T6 hexamer also interacts through non-polar surfaces. The alanine B14 and leucine B17 residues of the dimer in T6 hexamer create non-polar surfaces, which further help in contact of dimers. The B1 to B8 residues are in an extended or non-helical conformation. It has also been reported that phenylalanine B1 from one dimer interacts with leucine A13 and tyrosine A14 from the other dimer (74). Phenols and similar aromatic alcohols like cresols, and chloride ions in concentration above 6% dramatically change the structure of insulin hexamer (67, 76).
Insulin crystals can be grown in the presence of chloride ions, and these crystals are similar to crystals of two zinc insulin. However, chloride ions induce a helical structure in extended residues, B1-B8 of three monomers in a hexamer, thus producing T3R3 conformation (67, 76). Some other lycotrophic anions like SCN-, I-, Br- (INS-28, 11) have also been reported to induce this T to R conformation in three of the six monomers, thus giving a T3R3 conformation. This T3R3 conformation gives a new conformation to histidine at B5, which creates an additional zinc binding site (67). Two more zinc ions can coordinate with nitrogen of insulin hexamer in T3R3 conformation and hence T3R3 is also known as four-Zn insulin. However, Coffman et al. reported that in crystals, the four-Zn insulin hexamer or T3R3 state does not bind four zinc ions. They further reported an occupancy of 2.67 (77).

Monoclinic crystals are obtained on crystallization of insulin in the presence of 0.5% phenol (67). Phenols induce T to R conformation change in all six monomers of the hexamer, where the hexamer in these monoclinic crystals are in the R6 state. In helical ‘R’ state, all the insulin monomer subunits have the B1 to B19 helical structure (78). Since all the subunits are identical in R6 conformation, the two fold symmetry axis are restored (67). R6 conformation has two zinc ion binding sites, which lies on the symmetry axis as in T6. These two sites are again formed by His B10 imidazole, but interact with only one water molecule (78).

Phenol induced R6 hexamer also contains six identical hydrophobic pockets for binding of six phenol molecules (78). One side of the pocket is bound by B chain helix and A-chain bound the other three sides (78). Phenol molecules specifically bind to each of the six subunits at these pockets. The carbonyl oxygen of cystine A6 and amide nitrogen of A11 cystine forms hydrogen bond with hydroxyl group of phenol (67). Each of the His B5 of R6 interacts with the aromatic ring of phenols through Van der Waals forces. This blocks the axial zinc binding sites of His B5 in R6 conformation. Therefore, only two zinc binding sites are available (67).

It has been reported by several groups that T (extended) and R (helical) conformation interconvert in solution (78). Furthermore, these different conformations (T6, T3, R3, R6) exhibit different spectroscopic properties (78). Based on various studies, Kaarsholm et al. also believes that the structure of T6 and R6 are identical in solution and in the crystalline state (78).

In 1961, Schlichtkull reported that a number of metal ions such as Cd$^{2+}$, Co$^{2+}$, Cu$^{2+}$, Ni$^{2+}$, Fe$^{2+}$, and Mn$^{2+}$ can be accommodated by the zinc binding site of T6 (2-Zn insulin hexamer) (79). Roy et al. in 1990, investigated the existence of a zinc-free R6 hexamer in solution (80). Kadima et al. reported the formation of Cd$^{2+}$, Ni$^{2+}$ and Fe$^{2+}$ substituted R state hexamers (75). In 1987, Wollmer et al. showed that placing nickel (II) in zinc sites impeded the transition of T to R conformation (81). In 1956, Schlichtkrull was studying the growth of very small insulin crystals, which could be injected for the treatment of diabetes, when he discovered – insulin hexamer (82-85). The monomer of insulin has 23 polar residues. All of these polar residues are present on the surface of the
the hexamer, the two zinc ions are located on the 3-fold axis of symmetry. One is located 8Å above and the other 8Å below the 2 fold axis (71, 86).

1.2.4 Insulin zinc interaction and pH

It is well established that insulin in the solution in neutral and alkaline pH range, and in the presence of zinc, predominantly exists as hexamers of molecular weight of approximately 36,000 (86). The proportion of hexamers also increases in this pH range as the zinc concentration increases up to two zinc ions per hexamer (86). Hallas–Mollar et al. and Cunningham et al. reported that as the pH increases from pH 4.5 to 8.0, additional seven ions of zinc can associate with a hexamer (86, 87). Blundell et al. reported that at neutral pH, as the zinc concentration increases above two zinc atoms of zinc per hexamer, a 72,000 dalton unit starts appearing (86). They further reported that as the zinc amount increased above six atoms per hexamer, the distribution of insulin units became polydisperse and units with 200,000-3000,000 daltons occurred predominantly. As the pH was increased above pH 8.0, even in high concentration of zinc, these polymers dissociated in 36,000 dalton hexamers in the pH range of 9-10 (86).

1.2.5 Insulin mechanism of action and degradation

Insulin binds to insulin receptors, which are located on cell membranes. The insulin receptors are present on all the cells; however, their density is highest on liver and the adipose cells. The insulin receptors are heterotetrametric glycoprotein consisting of two extracellular α and two transmembrane β subunits. These subunits are linked together by disulfide bonds. The insulin binding sites are present on α subunits, while the have tyrosine protein kinase activity. The insulin binds to α subunits and induces aggregation and internalization of receptor and the bound insulin molecules. This activates a cascade of reactions resulting in the metabolic activities of insulin (88-92).

The internalized receptor insulin complex is either degraded intracellularly by insulin-degrading enzyme (IDE) or this complex is translocated back to the surface of the cell membrane, from where it may be released in systemic circulation. Insulin has plasma half life of approximately 4 to 5 minutes (93, 94). The liver and the kidneys are two major sites for insulin clearance from the systemic circulation (95-99). The receptor mediated clearance of insulin by hepatic and renal cells follow similar cellular mechanisms (100, 101). The insulin-degrading enzyme (IDE) is the major enzyme responsible for degradation of insulin. The IDE mediated degradation of insulin generates various products which have been well characterized (93, 102-108). It has been established that IDE recognizes the three-dimensional configuration of the insulin rather than specific peptide bonds (93, 106, 108). Stentz et al. further reported that IDE recognizes α-helical region around leucine-tyrosine bonds, and the major peptide bond cleavage sites of insulin are A 13-14, A 14-16, B 9-10, B 13-14 and B 16-17 (109). These insulin fragments may not only contribute to immunoreactivity, but may also bind to insulin receptors to produce biological activity (110).
1.3 Treatment of diabetes

Diabetes mellitus, depending on the type and stage of the disease, is treated with physical exercise, dietary control, oral hypoglycemics agents alone, or a combination therapy of insulin and oral hypoglycemics agents, or insulin therapy alone (111-114). Diabetics who suffer from the type-1 disease require insulin therapy for their survival. A new class of drug called amylin analogues, e.g. pramlantide acetate has also been recommended as an adjunct to insulin injections for the management of blood glucose concentrations in type-1 diabetic patients (115-117).

Depending on the stage at which the disease is diagnosed, diabetics who suffer from type-2 disease can be treated with an appropriate diet and proper exercise, or a monotherapy or a combination therapy of oral hypoglycemic agents (114). However, most of the type-2 diabetes patients may need insulin at some stage of their treatment for achieving the targeted glycosylated hemoglobin (HbA1c) concentrations of less than 7% (i.e. mean plasma glucose concentration equal to 170 mg/dL or 9.5 mmol/L) (118, 119).

The target of insulin therapy (either alone or in combination with other antidiabetic drugs) is to mimic the physiological release and concentrations of insulin. However, the tendency of regular human insulin to self associate after a subcutaneous injection, cannot allow it to mimic physiological required insulin secretion following food intake (113, 114). This is because of formation of hexamer from regular human insulin at the site of injection. These hexamers cannot cross the capillary barriers due to their size. The hexamers have to dissociate first into dimers and then into monomers to cross the capillary barrier and get absorbed into the blood stream prior to their action on insulin receptors. This process of dissociation of hexamers into monomers leads to a delay in the onset of action (114).

The onset of action with a formulation of regular human insulin is reported to be from 30 min to 1 hour and the duration of action of insulin in the body is reported to be 2 to 6 hours (114). Since the duration of action of insulin is not short, it cannot mimic the rapid decline of the physiological insulin concentrations after the meals. Table 1-1 details various regular human insulin preparations. Intermediate acting and long acting insulin formulations (NPH+ Lente) have been dispensed for basal insulin requirements. However, these formulations are associated with slow onset of action because of further stabilization of slow dissociating hexamers, and these formulations also show a pronounced peak leading to hypoglycemia in diabetic subjects. Furthermore, since the formulation like Lente and Ultralente are suspensions, the duration of action of these formulations varies significantly within and between individuals (114).

The goal of treating diabetes is to maintain a tight control on the blood glucose concentrations. It is difficult to achieve the tight glycemic control in diabetic subjects who are on regular human insulin therapy. This may lead to poor management and further progression of the disease (114). There is a strong need to develop an insulin formulation which can mimic physiological insulin action. The interest in producing the time action profile of injected insulin as per requirement of the diabetic patient is the
<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
<th>Examples</th>
<th>Action (hours)</th>
<th>Onset</th>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapid-acting</td>
<td>Modified insulin</td>
<td>Lispro (Humalog)</td>
<td>&gt;15 min</td>
<td>3-5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Aspart (Novolog)</td>
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<td>Glulisine (Apidra)</td>
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<td>Humulin R</td>
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<tr>
<td>Short-acting</td>
<td>Regular insulin</td>
<td>Novolin R</td>
<td>0.5-1</td>
<td>5-8</td>
<td></td>
</tr>
<tr>
<td>Intermediate-acting</td>
<td>NPH or Isophane Lente</td>
<td>Humulin N</td>
<td>1-3</td>
<td>14+</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>Novolin N</td>
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<tr>
<td></td>
<td></td>
<td>Humulin L</td>
<td>1-2.5</td>
<td>14+</td>
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<td></td>
<td></td>
<td>Glargine</td>
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<tr>
<td>Long-acting</td>
<td>Modified Insulins</td>
<td>Detemir</td>
<td>1-2</td>
<td>20-24</td>
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driving force for the development of modified insulins or insulin analogues (114).

NPH: Neutral Protamine Hagedorn insulin. The US FDA has approved five modified human insulin or insulin analogues. Table 1-1 lists these insulin analog preparations. All the formulations are to be administered via a subcutaneous injection. These insulin analogues can further be divided into two categories: Rapid acting insulin analogues, which attempt to reproduce the rapid secretion of insulin after meals, and the long acting insulin analogues (basal Insulin), which attempt to reproduce the basal concentrations of insulin that is secreted between meals during fasting and nocturnal phases (120, 121).

There are three rapid acting insulin analogues approved by the US FDA. Insulin Lispro (Humulog®, Elli Lilly), Insulin aspart (Novolg®, Novo Nordisk) and Insulin glulisine (Apidra®, Sanofi-Aventis). These insulin analogues bind to the receptor of the native human insulin and produce physiological effects similar to the endogenous insulin (122). These insulin analogues have been generated and produced by using recombinant DNA technology (122). The structure of insulin in the above mentioned analogues has been modified to achieve the desired pharmacokinetic and pharmacodynamic effects (122).

Lispro was the first rapid-acting human insulin analogue that was approved by the US FDA for treating type-1 and type-2 diabetes in humans (113, 123). It is a rapid acting insulin analogue which is effective for a short duration of time. The glucose lowering effect onset is from 5 to 15 minutes, and peaks in 30 to 90 minutes, and lasts for 4 to 6 hours (32, 113). The structure of Lispro is similar to endogenous human insulin except at two amino acid of the B chain. Lispro has been generated by inversion of amino acid lysine (B29) with proline, and proline B28 with lysine (123-125).

These modifications have affected the binding of Lispro with insulin receptor, but only slightly. However, the effects of insulin Lispro on biochemical signaling, glucose and amino acid uptake are similar to regular human insulin (113, 123). The minor change in the structure sequence destabilize the dimers by disrupting hydrophobic interaction and hydrogen bonding between the monomers (123). Hence Lispro monomers have a low propensity to self-associate and form dimers. It has been shown that in the presence of zinc and phenol, which are required to stabilize the Lispro formulation at neutral pH, Lispro forms hexamers (125). However, unlike human insulin where up to six phenol molecules bind to the hexamer, only 3 phenol molecules bind to hexamers of Lispro (125). The association of monomers in Lispro hexamer has been described as relatively weaker than that of regular human insulin hexamers. Therefore, after injection, the phenols diffuse away leading to conversion of loosely associated hexamer directly into monomer (123, 125). The dissociation of hexamer into monomer has been described as the rate limiting step in the absorption of insulin from the site of injection (123). The elimination half-life of Lispro after a subcutaneous injection has been reported as 60 minutes. This is significantly shorter than the half-life of regular human insulin (90 minutes) after a subcutaneous injection (123).
Lispro is recommended for type-1 and type-2 diabetic subjects, and should be injected within 15 minutes before meals and if necessary immediately after meals. Intravenous Lispro injection can be given in case of an emergency to deal with ketoacidosis or surgery (123). In different trials, it has been shown that Lispro is safe for fetus and new born, if it is injected in gestational diabetic subjects; however, its use is not yet recommended. Lispro can be mixed with NPH to form premixed Lispro-NPH. The combination of Lispro-NPH addresses the need of both bolus and basal therapy of diabetic subjects (123).

Insulin Aspart is the second rapid acting insulin analogue approved by the US FDA. Similar to Lispro, it has faster onset and shorter duration of action (125, 126). Insulin Aspart is created by substitution of the proline at B28 with aspartic acid (113, 124, 126). At the physiological pH, a negative charge is generated on the insulin molecule. This negative charge helps in repulsion of the monomers and reduces the tendency of the monomer to self associate into hexamers. Replacement of proline with aspartic acid also leads to a loss of Van der Waals forces, necessary for self association. This leads to rapid dissociation of hexamer to monomer resulting in rapid absorption. Like Lispro, Insulin Aspart is recommended for the treatment of both type-1 and type-2 diabetic subjects. It is recommend to be injected immediately before meals and its duration of action lasts for approximately 2 to 4 hours (126).

Insulin glulisine (Apidra®, Sanofi-Aventis) is another rapid-acting insulin analogue. In insulin glulisine, the asparagine at position B3 is replaced by lysine, and the lysine at position B29 has been replaced by glutamic acid (127). It has also a fast onset and short duration of action. One more modified insulin; is insulin glargine (Lantus®). However, it recommended for maintaining basal insulin concentrations (121, 128). Insulin glargine has been altered at two different positions. In A-chain, at position 21, the asparagine has been substituted with glycine, thus imparting more resistance to deamidation in an acidic environment, and providing more stability (128, 129). The C-terminus of the B-chain has elongated by the addition of two arginine molecules. This addition of two positive charges by the addition of arginines shifted the isoelectric point of the modified insulin glargine from pH 5.4 to 6.7 (129). These modifications have made it possible to formulate insulin glargine into a slightly acidic stable solution, which is easy to inject and improve dose reproducibility. This insulin glargine solution forms stable hexamers, which precipitate at neutral pH at the subcutaneous site of injection, thus forming a depot of microprecipititates of the modified insulin glargine which then dissolves at a steady rate for prolonged period of time (121, 129).

Insulin detemir (Levemir®, Novo Nordisk) is another modified recombinant human insulin whose structure has been altered such that upon subcutaneous injection, it gives a slow and an extended effect. In insulin detemir; the B-30 threonine of human insulin has been deleted (130, 131). A C-14 carbon fatty acid, i.e. myristic acid has been attached to B-29 lysine residue (130, 131). At neutral pH, insulin detemir is soluble and does not precipitate during storage, administration or absorption (132-135). It has been reported that insulin detemir, due to its hydrophobic nature, has a tendency to bind with proteins like albumin, which are present at injection sites and in blood. This reduces its
clearance and increases the half-life; thus leading to the extension in duration of action. Hence, insulin detemir is recommended for basal insulin therapy. It should be injected once or twice daily (132-135). Both insulin detemir and insulin glargine are recommended for the treatment of both type-1 and type-2 diabetic subjects (130).

1.4 Non-invasive insulin delivery systems

Subcutaneous injection of insulin has been a satisfactory mode of insulin therapy for a vast majority of diabetes subjects. However, the use of these subcutaneous injections may cause hyperinsulinemia, which may lead to hypoglycemia and diabetic micro and macro angiopathy (136, 137). Further, these subcutaneous injections of insulin are also associated with pain, inconvenience, physiological stress, high cost, difficulty in handling insulin, localized insulin deposition leading to local hypertrophy, and fat deposition at the site of injection (136, 138). Hence, during recent years, a great deal of interest has been shown for the development of novel ways to deliver insulin noninvasively.

Scientists in academic institutions as well as in the pharmaceutical industry are working on different drug delivery technologies including oral, nasal, buccal, pulmonary, transdermal, rectal and ocular to get a viable and feasible noninvasive insulin dosage form (136, 139, 140).

1.4.1 Oral insulin delivery of insulin

The conventional oral route is the method of choice for delivery of most drugs because of its acceptability by patients and ease of manufacturing. Further, in the case of successful oral insulin delivery, it may mimic physiological pattern of insulin leading to better glucose homeostasis (136, 141). However, efficient oral insulin delivery is challenging because of enzymatic and acidic degradation, thus leading to poor and unpredictable absorption and bioavailability. Therefore, for successful oral delivery, insulin should be protected against acidic and enzymatic degradation. Moreover, its epithelial permeation should be enhanced and bioactivity should be conserved during the formulation process (136). Various groups have tried different pharmaceutical approaches to maximize insulin stability and bioavailability for effective oral insulin delivery. Some of these approaches include enzyme inhibitors, absorption enhancers, mucoadhesive polymer systems, particulate carrier delivery systems and targeted delivery systems.

Enzyme inhibitors: Insulin is extensively degraded by gastrointestinal enzymes such as trypsin, α chemotrypsin and elastase (136, 142). The effects of intestinal degradation of insulin in rats have been evaluated in the presence of different enzyme inhibitors like sodium glycocholate, camostate mesilate, bacitracin, soybean trypsin inhibitor, and aprotinin. None of these enzyme inhibitors have been reported to be effective in the small intestine, and only some of these like sodium glycocholate, camostate mesilate, and bacitracin were found to be effective in showing some
improvement in absorption and bioavailability enhancement from the large intestine of rats. However, in general, these enzyme inhibitors are considered highly toxic and they may also affect the normal degradation and digestion of food proteins.

Absorption enhancers: Absorption enhancers improve the paracellular and transcellular transport of drugs in the gastrointestinal tract (GIT) by changing the membrane fluidity, decreasing mucus viscosity, opening tight junctions, and leakage of proteins from the membranes (136, 143). For improving the absorption of insulin, various non-specific permeation enhancers like bile salts, fatty acids, surfactants, salicylates, chelators, and zonula occluden toxin have been tried (144-147). However, it has been reported by Morishita et al. that the absorption enhancers increase insulin efficacy more effectively in the colon than in small intestine (147). Furthermore, since absorption enhancers increase permeation of cell membrane and open the tight junctions not only for insulin and other peptides, but also for undesirable molecules which are present in the GIT, their use may not be safe for chronic use (136).

Emisphere Technology, Inc, has developed oral capsules, called Eligen™, which use non-acylated amino acids as the carriers for transport of insulin. This oral insulin delivery system is currently undergoing Phase II clinical trials (148, 149).

Mucoadhesive systems for oral insulin delivery have also been developed. These systems increase the residence time of insulin at the site of absorption, strengthen the contact with the mucus, and localize the drug delivery system thus increasing the drug concentration gradient (136). Morishita et al. studied poly (methacrylic acid-g-ethylene glycol) [P(MAA-g-EG) polymers for oral insulin delivery and found that these mucoadhesive systems with smaller sized microparticular insulin loaded polymer system showed approximately 10% bioavailability of insulin after oral administration in diabetic rats (150). In a similar study, lecithin conjugated alginate microparticles, mucoadhesive chitosan nanoparticles of insulin and insulin tablets formulated with mucoadhesive thiolated (thiomers) seemed to increase the absorption of insulin via rat intestinal mucosa (151-154). These mucoadhesive delivery systems are considered to be effective in increasing the absorption of drug molecules susceptible to proteolytic degradation, however, their long term safety data in humans is warranted (155).

Colloid carrier delivery systems like microemulsions (156) liposomes (157) polymeric micelles (158) and polymeric nanoparticles and polymeric microparticles have been studied to improve the insulin delivery via oral route using specific sites like payer’s patches. Other systems for oral administration of insulin that have been studied are dry insulin emulsions, solid-in-oil-in-water emulsion of insulin, and microemulsion of insulin (159-162). These systems have demonstrated an improved efficacy of the orally administered insulin.

Cortecs International, Inc. in collaboration with Provalis PLC has developed oral microemulsions of insulin called Macrulin™, This system is in Phase II clinical trials (162-165).
Different types of liposomes have also been studied for their potential as carrier for oral administration of insulin. For example, Katayama et al. reported double liposomes of insulin with aprotinin, whereas Degim et al. reported similar liposomes of insulin with sodium taurocholate and found that blood glucose concentrations decreased significantly after oral administration in rats (166, 167). Goto et al. prepared fusogenic liposomes of insulin with envelope glycoprotein of Sendri virus (168). These liposomes were directly administered into the lower intestine of rats and were found to improve insulin absorption. Significant decrease in blood glucose concentrations were observed in these rats (162, 168).

Ye et al. reported, layer-by-layer, a new method of insulin loading in alginate – chitosan microcapsules. Nanocapsules of insulin made with polyethy-2-Cyano acrylate for oral delivery have also been reported. Chitosan microspheres with microadhesive properties and permeation enhancing effect have also been studied (169).

Enteric coated microspheres, using Eudragit S100 and nanoparticles using polymethacrylic acid-chitosan- polyethylene glycol have been investigated by Sajeesh et al and reported to be potential carriers for oral insulin delivery (170). Though some success has been achieved with these systems, there are still limitations such as poor drug incorporating efficacy, erratic drug release, particle aggregation, and low bioavailability, which should be addressed prior to further development (170).

Due to benefits of drug delivery to the colon, oral insulin delivery targeted in the colon has been extensively studied. Insulin delivery system with absorption enhancers like sodium glycocholate alone (171) or in combination with polyethylene oxide has shown prolonged absorption of insulin from the colon. Novel azopolymers coated pellets have been developed by Tozaki et al. for colon delivery of insulin (172).

Peyer’s patches, which are present in small intestine and play a major role in absorption, are also potential targets for oral delivery of insulin. Joseph et al. prepared microspheres of PLGA for oral delivery of insulin in mice (173). The microspheres (approximately 1 micron size) were supposed to be absorbed through Peyer’s patches and reported to have very significant effect on lowering of blood glucose concentrations (173).

Receptor mediated transcytosis is one more potential approach for the oral delivery of insulin. This approach does not affect the structure of plasma membranes or the paracellular junctions, and thus may be safer for chronic use. Transferrin receptors are considered good candidates for oral delivery of insulin because of their high density in the human GIT epithelium since transferrin is a natural transport protein. Xia et al. prepared insulin transferrin conjugates and reported prolonged hypoglycemia effect after oral administration in diabetic rats (174).

The buccal route of delivery for peptides and proteins like insulin is also a promising and widely investigated route of administration because of its accessibility to systemic circulation through the jugular vein. Therefore, the hepatic first pass is easily circumvented and higher bioavailability can be achieved through the use of this route.
Moreover, using this route, insulin administration is painless, enzyme activity is low and the system can be easily removed in case of adverse effects. The peptide absorption across the buccal membrane occurs via passive diffusion (175-177). Therefore, absorption enhancer, which are non irritant and not bitter, can be used to improve the permeability and hence enhance the bioavailability of insulin. Morishita et al. reported the preparation of Pluronic F-127 gels consisting of insulin and unsaturated fatty acids for buccal delivery (177). They showed that the gels prepared with Pluronic F-127 containing unsaturated oleic acid improved the bioavailability of insulin significantly in rats (9 ± 7.9%) (177).

Generex Biotechnology, Inc. has developed an oral insulin system called Oral-lyn™. The system is a micellar solution of insulin, which is to used as a buccal spray (148, 178, 179). This system has been marketed in India and Ecuador.

Transferosomes, which are ultra flexible and highly deformable lipid vesicles like liposomes, consisting of cholesterol, soybean phosphotidyl choline and sodium deoxycholate have been developed by Yang et al. for buccal delivery (180). They have shown that the relative bioavailability of insulin from the buccal transferosomes was 19.78% in rabbits. Nagai et al. have shown promising results using mucoadhesive nanoparticles of insulin in beagle dogs (181). Mucoadhesive buccal tablets made of Carbopol-934 with absorption enhancers have been reported by Venugopalan et al. (182). Xu et al. further reported the development of buccal spray of insulin with improved hypoglycemia effect lasting for up to 5 hours (183).

1.4.2 Nasal delivery of insulin

Nasal cavity provides a large surface area, and delivery via nasal route avoids hepatic first pass metabolism. Therefore, nasal insulin delivery systems have been widely studied. Furthermore, it has been reported that nasal delivery of insulin resembles the physiological pulsatile pattern of insulin release (184). Various absorption enhancers like sodium deoxycholate in combination with cyclodextrins, soybean derived sterol mixtures have been used with insulin to increase its absorption via nasal membrane (185, 186) in rats and rabbits. This approach has shown significant absorption promoting effect. However, absorption enhancers may cause irreversible damage to the nasal mucosa upon long term use. Therefore, long term safety of these delivery systems should be studied (187). Nasal mucoadhesive systems may increase the retention and contact time in the nasal cavity, thus improving the total absorption (188, 189). Najafabadi et al. prepared mucoadhesive Carbopol based gel spray and reported that upon nasal administration of the gel, the relative bioavailability of insulin in rabbits was 20.6%. Some other groups have also developed chitosan bioadhesive gels (190), chitosan nanoparticles (191) and chitosan microspheres (192) for nasal delivery of insulin in animals. They have all found that the absolute bioavailability of insulin has been increased significantly after nasal delivery.
Bently Pharmaceuticals, Inc. has developed Nasulin™, a nasal spray of insulin, which utilizes absorption enhancers for increasing the bioavailability of insulin (148, 178, 179). This system is in Phase II clinical trials.

**1.4.3 Pulmonary-lung delivery of insulin**

Pulmonary administration of proteins and peptide is considered very promising because of large surface area (140 m²) offered by the lungs, a thin alveolar epithelium, a lower metabolic activity of enzymes relative to GIT, and by pass of the first pass hepatic effect (136). Spray-dried insulin alone or in combination with other excipients like absorption enhancers and other particulate formulations like liposomes, nanoparticles, and nanospheres have been tried to deliver insulin successfully in animals via pulmonary route (136). Kwo et al. attempted pulmonary delivery of micro crystals of insulin and reported hypoglycemic effect lasting for seven hours in rats (193). Surendrakumar et al. prepared co-spray dried insulin powder and hyalouronic acid for inhalation and studied the insulin concentration in beagle dogs (194). They reported that the insulin-hyalouronic acid-Zn particles, after pulmonary inhalation, lasted for 4 to 5 hours. Mitra et al. studied co-administration of insulin with absorption enhancers like 1-2 dipalmitoyl phosphatidylcholine and showed a significant reduction in blood glucose concentrations after pulmonary delivery (195). Some other studies reported the use of Span 85, bacitracin, and citric acid and cyclodextrin derivatives as absorption enhancers of insulin via pulmonary route (195-197).

PLGA particles of insulin, like PLGA nanospheres, PLGA microparticles with cyclodextrins have also been prepared and reported to be effective in inducing hypoglycemia effect for 4 to 5 hours (198-200).

Exubera®, which was launched by Pfizer in January 2006 as the first non invasive insulin delivery system was a milestone in the development of noninvasive delivery of insulin. However, it was withdrawn from the market in October 2007. Though the major reason for withdrawal of the product from the market was economic loss, other issues such as occurrence of more cases of hypoglycemia (201, 202), dose optimization in smokers and asthmatic patients, issues such as development of antibodies against insulin, and effect of insulin on IGF-1 receptors and proliferation effects should have been more carefully studied and reported before marketing a pulmonary delivery system of insulin like Exubera® (201-205).

**1.4.4 Ocular delivery of insulin**

In 1931, Christie and Hanzal investigated the use of eye as potential delivery route for insulin for the first time (136). Ocular route provides the benefits of fast action, low immunological reactions, and by pass the first pass hepatic metabolism. However, pure insulin eye drops have low bioavailability (136). Hence, various research groups have reported the use of different absorption enhancers for ocular delivery of insulin. Yamamoto et al. studied polyoxyethylene-9-lauryl ether, sodium glycocholate, sodium taurocholate, and sodium deoxycholate as absorption enhancers in eye drops of insulin,
and studied their hypoglycemic effect and safety profiles in rabbits for three months (206). Uses of fusidic acid, dodecylmaltoside, or tetradecyl maltoside as absorption enhancers for ocular delivery of insulin have also been reported (207). One of the issues with ocular delivery of insulin is low retention time of the formulation in the ocular cavity. Positively charged liposomes containing insulin have been formulated and have been reported to reduce the blood glucose concentration in rabbits to 65-70% of initial glucose concentrations, and the effect lasted for five hours (208, 209). Ocular inserts like Gelfoam® have been developed for prolonged release of insulin in the eye and the system has shown lowering of blood glucose concentrations (60% of initial concentrations) for up to 8 hours.

### 1.4.5 Rectal delivery of insulin

The hepatic first pass metabolism can be avoided using rectum as delivery route for insulin and this route is regarded as more physiological route for administration of insulin (136). Insulin suppositories have been developed and tested in rabbits and shown to have promising results (210). Further, absorption enhancers like sodium cholate sodium glycocholate, sodium taurodeoxycholate, sodium taurocholate, sodium deoxycholate and sodium salicylate have been tried in insulin suppositories prepared with Witepsol W 35 as base. Some of these formulations showed up to 50% relative hypoglycemic effect. Snail mucin has also been used as an absorption enhancer in glycerol gelatin suppositories of insulin by Adikwi et al. (211). They showed that blood glucose concentration reduced to 44% within two hours of application of these suppositories (211). Thermoreversible suppositories containing poloxomers, which convert to gels upon rectal application, have also been developed as rectal delivery system for insulin in animals (212).

### 1.5 Controlled release of insulin

Insulin, has been reported to be released in a pulsatile manner from isolated islets, perifused pancreas and in vivo (213). Therefore, for diabetic subjects, the ideal insulin delivery system should be capable of mimicking this natural pattern, i.e. the delivery system should have a sensing and auto shut off capability (113). Various research groups are investigating two kinds of glucose sensing insulin delivery systems. These are glucose sensitive hydrogels and artificial pancreas, i.e. closed loop delivery systems.

#### 1.5.1 Artificial pancreas

These systems are also called closed loop delivery systems (113). These types of systems consist of a sensor, which remains in contact with blood or interstitial fluid, and measure the glucose concentrations in blood, and then communicate with an attached microcomputer. This microcomputer then communicates with the attached pump and modulates the release of insulin from the pump. Hence, the blood glucose concentration determines the release of insulin from the pump. However, there are limitations with
glucose sensing and regarding the glucose turnover and insulin stability in the pumps. These kinds of systems are still in the development phase (113, 214).

### 1.5.2 Glucose sensitive hydrogels

These are also called “smart gels” and have the capability of changing some of their properties in response to the glucose concentration in the surroundings. pH sensitive polymers immobilized with the glucose oxidase have been used in many investigations (113, 215, 216). When these systems encounter glucose, the glucose oxidase present in these systems oxidizes glucose to gluconic acid, thus lowering the pH. The change in pH may lead the polymeric gel to swell and expand or collapse and squeeze. This change in dimension of polymeric gel expansion or the squeezing are proportional to the glucose concentration present in the ambience and this modulates the release of insulin from the gels made up of these polymers. Catalase has been included in these kinds of systems to enhance the oxidation rate of glucose (113, 215, 216).

### 1.5.3 Extended release of insulin

The self regulating glucose sensitive insulin delivery systems are ideal form of insulin dosage form but they are difficult to develop and have certain limitations (113). Therefore, development of an extended release formulation of insulin, which can provide constant release of insulin for a prolonged period of time, is desirable. Long acting insulin like insulin glargine and insulin detemir, and intermediate acting insulin like NPH are available as substitutes of basal insulin. However, a daily injection of these formulations is required. This daily injection may not be tolerated by the diabetic subjects, who would need these injections for rest of their life. This may lead to poor patient compliance, which may further lead to poor management of diabetes. Hence, there is strong need to develop novel insulin formulation having prolonged action, requiring less frequent injections (113).

Non-invasive insulin delivery systems for normal delivery of insulin are still under development. Various parenteral extended release delivery systems are also being considered for the prolong delivery of insulin (113, 217).

Microparticles, including microspheres and microcapsules of nonbiodegradable polymer such as ethylcellulose, or biodegradable polymers such as PLGA have been prepared for prolonged delivery of insulin (113, 218-223). However, more focus is on the development of biodegradable microspheres. Despite being biodegradable, these biodegradable microspheres have some limitations like initial burst leading to hypoglycemia or a lag time in release (113, 221, 222). Yamaguchi et al. reported that addition of hydrophilic excipients like glycerol or water during the preparation of these microspheres can reduce the initial burst (113, 222).

Further, in the microparticles prepared with PLGA, the initial burst is followed by a slow and incomplete release of insulin (218). This may be due to denaturation of insulin in the acidic environment created by the degradation products of PLGA (218, 219). This
denaturation of insulin can be reduced by incorporation of some basic excipients like sodium bicarbonate (113, 218). Incomplete release of insulin from such systems is a major concern and may be a potential hazard and safety issue (218, 219).

Implants are solid systems in form of tablets, rods, wafers, discs or films (113). These implants can be administered using a minor surgery or a trocar device. These implant systems may be biodegradable or non-biodegradable. In the case of non-biodegradable systems, the removal of the system is necessary after depletion of the drug or after completion of the therapy. However, the biodegradable systems will degrade with time and fragment into water soluble products, which dissolve in interstitial fluid and goes away from the implant site (113).

Different kind of implantable systems have been investigated by various groups to achieve an extended release of insulin (224). Ethylene–vinyl acetate copolymers (EVAc) have been used to make discs of insulin implants. These systems could reduce the blood glucose concentrations in diabetic rats from 30 to 105 days, depending on the formulations (224-228). However, in some of these systems, only a part of the loaded insulin could be released. Furthermore, the removal of the system after completion of the release was mandatory. Implants of insulin made up of cholesterol, palmitic acid, chitosan and gellan gum have also been studied in animals (226, 228-230). PLGA and PLA biodegradable implants have also been investigated by various groups (231, 232).

1.5.4 **Insulin pumps**

Insulin pumps are also known as continuous subcutaneous insulin infusion (CSII). Insulin pumps were first devised by Dr. Arnold Kadish in 1960 (233-235). Relative to the initial pumps, the modern pumps are smaller in size and called as micropumps. These micropumps consist of insulin reservoir, which is a replaceable cartridge, a catheter with tubing, and a micropump. To perform, the catheter of the pump is inserted subcutaneously and connected to reservoir through the tubing. The micropump controls the release of insulin from the reservoir. The efficient use of insulin pumps requires frequent blood glucose monitoring, and accordingly, adjustment of the rate of insulin release from the pumps. Since it can cover both basal and bolus needs of insulin, better glycemic control have been reported with these kind of pumps effectively. However, stability of insulin in the reservoir, patient compliance and cost of these pumps are certain issues that are affecting the acceptability of these pumps (113).

1.5.5 **Insulin gels**

Gels are colloid systems consisting of a dispersed phase, which is dispersed in a medium resulting in a semisolid form. The gel used as drug delivery systems consists of a drug, a polymer and a solvent. The drug can be suspended or dissolved in the solvent, depending on the solubility of the drug.

Gels have been widely used for delivery of drugs via topical route. However, a few parenteral gel formulations like Eligard® and Atridox® have been marketed in the
USA. Both of these gels use the Atirgel® technology. The Atirgel® technology combines a biodegradable polymer, PLGA with a biocompatible solvent, NMP. The drug can be dispersed in polymer solution as a suspension or dissolved as a solution. The formulation, upon subcutaneous injection, forms a depot at the site of injection, and releases the drug for a prolonged period of time. A similar formulation, Alzamer®, has been developed by Alza Corporation using either ethyl benzoate or benzyl benzoate as the solvent. However, these systems suffer from initial burst, which must be controlled for successful delivery of insulin (113).

Barichello et al. reported the use of Pluronic (PF 120) gel for delivery of insulin. However, these insulin gels could only maintain low blood glucose concentrations for up to 12 hours (113, 177, 236).

Caliceti et al. reported preparation of hydrogels which change properties with change of environment such as temperature or pH. These gels could lower the blood glucose concentrations in diabetic rats after 2 hours after injection and maintained the effect for up to 100 hours. Kim et al.s reported the use of PLGA-PEG-PLGA polymer based gels (Relgel®) for preparation of insulin formulations (237). The gels are liquid at room temperature, but undergo a sol gel transformation at 37° C. Hence, an insulin containing solution of these gels when injected subcutaneously transforms to a solid and forms a depot, from which insulin can leach out for extended period of time. These formulations, upon subcutaneous injection in rats, could maintain low blood glucose concentrations for two weeks. Chen et al. made similar gels, but added zinc carbonate to them. The gel loaded with insulin Ultralente with zinc carbonate could keep the blood glucose concentrations in the normal range for 10 days in ZDF type 2 diabetic rats (113).
CHAPTER 2. PROLONGED EFFECT OF INSULIN IN PRESENCE OF WATER SOLUBLE AND WATER INSOLUBLE SOLUBLE ZINC SALTS

2.1 Introduction

Insulin is a metabolic hormone synthesized, stored and secreted from beta cells of pancreas. In beta cells of pancreas insulin is stored as granules consisting of insulin hexamers. These hexamers are composed of six units of insulin loosely connected by hydrophobic forces and further stabilized by presence of zinc ions (238).

In 1937, Scott and Fisher used zinc for the first time in insulin formulations to stabilize protamine zinc insulin formulation (31, 34). In 1951, Hallas-Moller et al. came up with an idea of Lente insulins. Semilente was a suspension of amorphous modification of insulin and zinc, and had somewhat of a longer duration of action than the normal insulin. Ultralente was a crystalline modification of insulin (insulin crystallized in acetate buffer) with a longer duration of action similar to protamine zinc insulin formulation (35-40, 82, 87, 239).

Insulin interacts with zinc ions to form hexamers, and the hexamers get further stabilized in the presence of zinc. When these hexamers are injected subcutaneously, because of their inability to cross the capillary barriers, they form a depot at the site of injection. To get absorbed into the blood stream, these hexamers have to dissociate first into dimers and then to monomers. When these hexamers are injected subcutaneously, because of their inability to cross the capillary barriers, they form a depot at the site of injection. To get absorbed into the blood stream, these hexamers have to dissociate first into dimers and then to monomers. This leads to delayed and extended effect of insulin (68, 124, 238, 240).

A novel injectable biodegradable gel system for controlled release drug delivery has been designed and characterized in our laboratory. It consists of a biodegradable polymer, PLGA, biocompatible plasticizer(s) and an active pharmaceutical ingredient (API). To date, several molecules, including antibiotics, opiate analgesics and narcotic antagonists, have been successfully incorporated into the gel system, and controlled release of the aforementioned APIs have been achieved in our laboratory.

The long term aim of this study was to develop an injectable biodegradable gel system which can be used to control the release of insulin for one week or longer, after a single subcutaneous injection in type-2 diabetic ZDF rats.

The objective of this part of the study was to investigate the effect of different water-soluble and water-insoluble zinc salts on blood glucose lowering effect of insulin in type-2 diabetic ZDF rats.
2.2 Materials and methods

2.2.1 Materials

Insulin (RHI) powder (26.9 IU/mg containing 0.4% zinc) was purchased from Diosynth France S.A. (Usine St. Charles, France) and Humulin® U (Ultralente insulin, 10 mL) (Elli Lilly and Company, Indianapolis, IN) was purchased from Cardinal Health. The HPLC analysis of insulin was performed on a Shimadzu HPLC system (Shimadzu Scientific Instruments, Inc., Columbia, MD). The primary column was a 250 x 4.6 mm Alltech Macrosphere RP 300 C18 5 μm column and the guard column (Alltech, Deerfield, IL) was 7.5 x 4.6 mm, packed with same material as the primary column. HPLC grade Acetonitrile (ACN), trifloroacetic acid (TFA) and distilled water were used for preparing the mobile phase. Male ZDF (fa/fa) rats (Charles River Laboratories, Inc., Wilmington, MA) weighing 300-400 grams were used for the in vivo studies. Blood glucose was measured using Bayer Ascensia® Breeze blood glucose meter (Bayer Corporation, Elkhart, IN). The pH of the formulations was determined using pH meter (Model IQ 240, IQ Scientific Instruments, Inc., San Diego CA).

2.2.2 Methods

2.2.2.1 Preparation of the insulin and zinc suspensions. An appropriate quantity of insulin (RHI) powder was suspended in normal saline to make 40 IU/ml suspensions. Tables 2-1 to 2-4 show the composition of insulin formulations containing varying concentrations of water-soluble (zinc sulfate or zinc chloride) and water-insoluble (zinc carbonate or zinc oxide) zinc salts. The ingredients were weighed into a 7-mL glass scintillation vial and water was added to bring the final weight to approximately 3 grams. An appropriate quantity of sodium chloride was added to each formulation to make the solution isotonic before adding insulin. Humulin® U suspension (100 IU/ml) was used as is.

2.2.2.2 High pressure liquid chromatography method of insulin. The HPLC analysis of RHI was performed using a Shimadzu HPLC system (Shimadzu Scientific Instruments, Inc., Columbia, MD) consisting of an SCL-10A vp system controller, a LC-10 AD vp pump, a DGU-14A degasser, an SIL-10 AD vp autoinjector with a cooling system, a CTO-10A vp column oven, an SPD-10 AD vp photo-diode array (PDA) detector and a computer loaded with the Shimadzu Class-VP 7.2 software. The primary column was a 250 x 4.6 mm Alltech Macrosphere RP 300 C18 5 μm column and the guard column was 7.5 x 4.6 mm, packed with same material as the primary column.

A gradient elution method was used for analyzing the insulin. Mobile phase A was 0.15% v/v trifluoroacetic acid (TFA) in water and mobile phase B was 0.13% v/v TFA in 95% v/v acetonitrile in water. Mobile phases were filtered through a Whatmann0.45 μm nylon membrane (Whatmann International Ltd, Maidstone, England) and degassed in an ultrasonicator (Model FS60, Fisher Scientific, Fair Lawn, NJ) for 10 minutes before pumping it through the HPLC system.
<table>
<thead>
<tr>
<th>Formulation #</th>
<th>Insulin (mg)</th>
<th>ZnSO$_4$ (mg)</th>
<th>NaCl (mg)</th>
<th>Adjust total weight with H$_2$O(g)</th>
<th>Amount of zinc in the formulation (mg/g)</th>
<th>Molar ratio of insulin to zinc</th>
</tr>
</thead>
<tbody>
<tr>
<td>OA-2-1</td>
<td>4.56</td>
<td>-</td>
<td>27.00</td>
<td>3.00</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>OA-2-2</td>
<td>4.58</td>
<td>3.15</td>
<td>26.52</td>
<td>3.00</td>
<td>0.24</td>
<td>1:14</td>
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<td>OA-2-3</td>
<td>4.53</td>
<td>6.33</td>
<td>26.04</td>
<td>3.00</td>
<td>0.48</td>
<td>1:28</td>
</tr>
<tr>
<td>OA-2-4</td>
<td>4.49</td>
<td>12.66</td>
<td>25.14</td>
<td>3.00</td>
<td>0.96</td>
<td>1:56</td>
</tr>
<tr>
<td>OA-2-5</td>
<td>4.50</td>
<td>25.35</td>
<td>23.19</td>
<td>3.00</td>
<td>1.92</td>
<td>1:112</td>
</tr>
<tr>
<td>Formulation #</td>
<td>Insulin (mg)</td>
<td>ZnCl₂ (mg)</td>
<td>NaCl (mg)</td>
<td>Adjust total weight with H₂O (g)</td>
<td>Amount of zinc in the formulation (mg/g)</td>
<td>Molar ratio of insulin to zinc</td>
</tr>
<tr>
<td>--------------</td>
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</tr>
<tr>
<td>OA-2-6</td>
<td>4.61</td>
<td>1.5</td>
<td>26.07</td>
<td>3.00</td>
<td>0.24</td>
<td>1:14</td>
</tr>
<tr>
<td>OA-2-7</td>
<td>4.51</td>
<td>3.0</td>
<td>25.14</td>
<td>3.00</td>
<td>0.48</td>
<td>1:28</td>
</tr>
<tr>
<td>OA-2-8</td>
<td>4.62</td>
<td>6.0</td>
<td>23.28</td>
<td>3.00</td>
<td>0.96</td>
<td>1:56</td>
</tr>
<tr>
<td>OA-2-9</td>
<td>4.59</td>
<td>12.0</td>
<td>19.56</td>
<td>3.00</td>
<td>1.92</td>
<td>1:112</td>
</tr>
<tr>
<td>Formulation #</td>
<td>Insulin (mg)</td>
<td>Zn CO₃ (mg)</td>
<td>NaCl (mg)</td>
<td>Adjust total weight with H₂O (g)</td>
<td>Amount of zinc in the formulation (mg/g)</td>
<td>Molar ratio of Insulin to zinc</td>
</tr>
<tr>
<td>---------------</td>
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<tr>
<td>OA-2-10</td>
<td>4.61</td>
<td>4.82</td>
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<td>1:56</td>
</tr>
<tr>
<td>OA-2-11</td>
<td>4.61</td>
<td>9.71</td>
<td>27.0</td>
<td>3.00</td>
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<td>1:112</td>
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<tr>
<td>OA-2-12</td>
<td>4.55</td>
<td>30</td>
<td>27.0</td>
<td>3.00</td>
<td>5.98</td>
<td>1:349</td>
</tr>
</tbody>
</table>
Table 2-4  Composition of insulin formulations containing zinc oxide

<table>
<thead>
<tr>
<th>Formulation #</th>
<th>Insulin (mg)</th>
<th>ZnO (mg)</th>
<th>NaCl (mg)</th>
<th>Adjust total weight with H₂O (g)</th>
<th>Amount of zinc in the formulation (mg/g)</th>
<th>Molar ratio of insulin to zinc</th>
</tr>
</thead>
<tbody>
<tr>
<td>OA-2-13</td>
<td>4.39</td>
<td>0.890</td>
<td>27.0</td>
<td>3.00</td>
<td>0.24</td>
<td>1:14</td>
</tr>
<tr>
<td>OA-2-14</td>
<td>4.60</td>
<td>1.79</td>
<td>27.0</td>
<td>3.00</td>
<td>0.48</td>
<td>1:28</td>
</tr>
<tr>
<td>OA-2-15</td>
<td>4.64</td>
<td>3.58</td>
<td>27.0</td>
<td>3.00</td>
<td>0.96</td>
<td>1:56</td>
</tr>
<tr>
<td>OA-2-16</td>
<td>4.44</td>
<td>7.16</td>
<td>27.0</td>
<td>3.00</td>
<td>1.92</td>
<td>1:112</td>
</tr>
</tbody>
</table>
The initial mobile phase composition of 70% solvent A and 30% solvent B was maintained at a flow rate of 1 mL/min for 2 minutes. Between 2 and 16 minutes, the percentage of solvent B was increased linearly to 50%. These conditions were maintained for 3 minutes from 16 to 19 minutes. Between 19 and 24 minutes, the percentage of solvent B was decreased linearly to 30%. The mobile phase was pumped at a flow rate of 1 mL/min. The injection volume of each sample, including the insulin standards, insulin controls, insulin samples (unknown concentration) and blank samples (0.01N HCl) without insulin was 50 μL. The temperature of the HPLC column was maintained at 30°C by using a column oven. The primary detection wavelength of insulin was set to 285 nm.

2.2.2.3 Preparation of standard, control and sample solutions. The standard solutions and controls of insulin were prepared from recombinant human insulin. Approximately 10 or 25 mg of insulin was weighed accurately using an analytical balance and transferred to a clean dry 10 or 25 mL volumetric flask. Either 1 mL or 2.5 mL of 0.01 N hydrochloric acid was added into the flask and the flask was swirled gently to dissolve the insulin powder. Isotonic phosphate buffered saline (PBS, pH 7.4) was used to make up the volume. The resultant solution contained approximately 1 mg/mL of insulin (working stock solution). Standard solutions were prepared from working stock solution by transferring the different amounts of working stock solution to five 10 mL volumetric flasks and making up the volume with PBS. The final insulin standard solutions contained 5, 10, 25, 50, 75, 100, 125, and 150 μg/mL of insulin, respectively.

The insulin controls were also prepared from the working stock solution. These controls were always prepared and dispersed into small aliquots. Each aliquot was stored in a 1 mL HPLC vial and sealed by Parafilm. The controls were frozen at –20°C and one vial was thawed before the analysis. Depending on the approximate concentration of the insulin samples in each batch of analysis, one or two insulin controls were prepared accordingly.

2.2.2.4 Process of chromatographic data. After each HPLC injection, the software generated the chromatograph of the sample and integrated the peak area of insulin. The peak area of the standard solutions was then plotted against the concentration of insulin.

2.2.2.5 Pharmacodynamic studies of insulin in ZDF rats. Pharmacodynamics of subcutaneously administered Humulin® U (Ultralente insulin, 10 mL) (Elli Lilly and Company, Indianapolis, IN) and insulin (Diosynth, Inc., Bensenville, IL) with and without varying concentrations of water-soluble and water-insoluble zinc salts was investigated in ZDF rats at a dose of 10 IU/kg.

ZDF rats were randomly divided into groups of 4 to 6 animals per group. Each rat was anesthetized by isoflurane (Halocarbon Product Corporation, River Edge, NJ) using a SurgiVet/Anesco ventilator (Waukesha, WI) prior to handling the animals for injecting the insulin dosage forms and glucose measurement. Body weight of each animal was then recorded to determine the dose of the insulin suspensions to be injected. Blood glucose concentrations of each animal was measured before injecting insulin, using the
Ascensia™ Breeze™ glucose monitoring system (Bayer Corporation, Elkhart, IN). Each group received one dose of a particular form of insulin formulation. An appropriate dose of insulin was injected subcutaneously on the dorsal neck of each rat using a ½ mL Becton Dickinson Lo-Dose™ U-100 insulin syringe with a permanently attached 28 gauge needle (0.5 inch in length) (Franklin Lakes, NJ). After injection, the rats were returned to the cages and allowed to recover from anesthesia. At pre-determined time intervals, the rats were anesthetized by the abovementioned method and blood glucose concentrations of the rats were measured. Blood glucose monitoring was stopped when the blood glucose concentrations reached their normally high concentrations.

2.2.2.6 In vitro studies. Tables 2-1-2-4 lists amounts of ingredients used to prepare various RHI formulations with and without zinc salts. After 16 hours at room temperature, the pH of these formulations was determined using a calibrated pH meter. The formulations were then centrifuged (Eppendorf Centrifuge 5415 C) and the supernatant were collected into HPLC vials. Insulin concentrations were determined in the supernatant using a HPLC method as specified before.

2.2.2.7 In vitro dissolution of insulin from insulin formulations with varying concentrations of zinc sulfate. The dissolution rate of RHI with and without zinc sulfate was studied using a dialysis method. Dialysis was conducted using the DispoDialyzer® ready-to-use tubes (500 μL capacity) purchased from Spectrum Laboratories (Rancho Dominguez, CA). The DispoDialyzer® tubes were made from cellulose ester, having a molecular weight cut-off of 50,000. The dialysis medium was isotonic PBS at pH 7.4.

Before using, the DispoDialyzer® tubes were rinsed with the dialysis media (PBS). The rinsed tubes were then kept, submerged in the dialysis media for 30 minutes prior to use.

An appropriate amount of insulin was suspended in PBS such that the concentration of insulin in the suspension was approximately 5 mg/mL (Table 2-4). In another set of samples, regular human insulin formulations with varying concentrations of zinc sulfate were prepared and used. Each formulation (0.5 mL) was then transferred to a DispoDialyzer® tube. Each DispoDialyzer® filled with insulin formulation was then placed into a 50 mL glass centrifuge tube, containing 25 mL of dialysis media. The DispoDialyzer® tube floated upright in the media. The centrifuge tube was capped, shaken at 100 rpm in an orbital shaker at room temperature.

At pre-determined time intervals, the centrifuge tubes were removed from the shaker and 1 mL of dialysis media was withdrawn from each tube. The concentration of insulin in the media was determined by the HPLC method. The rest of the dialysis media in each centrifuge tube was dumped and 25 mL of fresh media was refilled. The tubes were then returned to the shaker and shaking was continued. The cumulative percentage of insulin dissolved from each suspension was plotted as a function of time.
2.3 Results and discussion

2.3.1 Pharmacodynamic studies of insulin in ZDF rats

Figure 2-1 shows the effect insulin on blood glucose concentrations in ZDF rats after a subcutaneous injection of insulin and ultralente insulin. The blood glucose concentrations decreased from 535 mg/dL to 77 mg/dL within two hours after administering a subcutaneous injection of regular insulin. Normal blood glucose concentration (50 to 200 mg/dL) was maintained in the animals for approximately 2.5 hours, and within 6 hours after injection, the blood glucose returned to its initial high concentrations of 535 mg/dL. Therefore, the total duration of insulin activity in the animals was less than 6 hours. The blood glucose concentrations decreased from 431 mg/dL to 74 mg/dL within four hours after administering a subcutaneous injection of ultralente. Normal blood glucose (50 to 200 mg/dL) was maintained in the animals for approximately 7 hours, and within 14 hours after injection, the blood glucose concentrations returned to its initial high concentrations of 375 mg/dL. Therefore, the total duration of ultralente activity was less than 14 hours.

Figure 2-2 compares the mean and standard deviations of blood glucose concentrations in ZDF rats after a subcutaneous injection of insulin with varying concentrations of zinc sulfate. It is evident from the figure that adding varying amounts of zinc from 0.24 to 1.92 mg (obtained from zinc sulfate) to the regular insulin increased the Tmax (time at the lowest blood glucose concentration) from 2 hours (Tmax for regular insulin without any zinc) to 8 hours for formulation with 0.24 mg of zinc and 16 hours for formulation with 1.92 mg of zinc. The Emax (lowest blood glucose concentration) after a subcutaneous injection of RHI was 75 mg/dL.

The decrease in blood glucose concentrations from approximately 500 mg/dL to approximately 110 mg/dL was observed after subcutaneous injections of RHI with 0.24 or 0.48 mg of zinc (obtained from zinc chloride), respectively (Figure 2-3). Normal blood glucose concentrations (50 to 200 mg/dL) were maintained for approximately 6 hours in the animals, which were injected with both the formulations, and within 10-12 hours after injecting either formulation, the blood glucose returned to its initial high concentrations of approximately 500 mg/dL.

The drop in blood glucose concentrations after subcutaneous injections of insulin with 0.24 or 0.48 mg of zinc obtained from zinc chloride or zinc sulfate was similar to the drop in blood glucose concentrations after a subcutaneous injection of ultralente in the animals. The Emax after a subcutaneous injections of ultralente (Figure 2-1) was 140 mg/dL, whereas, the Emax values after subcutaneous injections of regular insulin with 0.24 or 0.48 mg of zinc obtained from zinc chloride or zinc sulfate was approximately 110 –160 mg/dL (Figures 2-2 and 2-3). The drop in blood glucose concentrations after subcutaneous injections of insulin with 0.96 or 1.92 mg of zinc obtained from zinc chloride or zinc sulfate was more gradual than that observed after an injection of insulin with either 0.24 or 0.48 mg of zinc obtained from zinc chloride or zinc sulfate. Moreover, the Tmax for regular insulin with 0.96 or 1.92 mg of zinc obtained from zinc chloride or zinc sulfate was approximately 3 hours, whereas, the Tmax for regular insulin with 0.24 or 0.48 mg of zinc obtained from zinc chloride or zinc sulfate was approximately 2 hours. The decrease in blood glucose concentrations from approximately 500 mg/dL to approximately 110 mg/dL was observed after subcutaneous injections of RHI with 0.24 or 0.48 mg of zinc (obtained from zinc chloride), respectively (Figure 2-3). Normal blood glucose concentrations (50 to 200 mg/dL) were maintained for approximately 6 hours in the animals, which were injected with both the formulations, and within 10-12 hours after injecting either formulation, the blood glucose returned to its initial high concentrations of approximately 500 mg/dL.
Figure 2-1  Blood glucose concentrations of ZDF rats after a single subcutaneous injection of insulin at a dose of 10 IU/kg (Mean ± SEM, n = 6)
Figure 2-2  Comparison of blood glucose concentrations in ZDF rats after a single subcutaneous injection of insulin with varying concentrations of zinc sulfate (Insulin dose-10 U/kg) (Mean ± SEM, n = 4)
Figure 2-3  Comparison of blood glucose concentrations in ZDF rats after a single subcutaneous injection of insulin with varying concentrations of zinc chloride (Insulin dose-10 U/kg) (Mean ± SEM, n = 4)
zinc sulfate was 12 or 16 hours, respectively (Figures 2-2 and 2-3). Hence, insulin release could be extended by varying concentrations of both the water soluble salts of zinc. Moreover, the effect of different concentrations of zinc obtained from two different water soluble salts on the Tmax and Emax were similar. However zinc chloride, due to its very hygroscopic nature, was not used further in these studies.

Figure 2-4 and Figure 2-5 compare the mean and standard deviations of blood glucose concentrations in ZDF rats after a subcutaneous injection of insulin with varying concentrations of water-insoluble zinc salts. Figure 2-4 depicts the blood glucose concentrations after subcutaneous injection of insulin with different concentrations of zinc, obtained from zinc carbonate. It is evident from the Figure 2-4 that adding varying amounts of zinc from 0.96 to 1.92 mg (obtained from zinc carbonate) to the insulin increased the Tmax (time at the lowest blood glucose concentration) from 2 hours (Tmax for regular insulin without any zinc) to only approximately 4 hours for formulation with 0.96 mg and 1.92 mg of zinc. However, the blood glucose concentrations reached back to high concentrations of 500 mg/dL within 8 hours.

Therefore, there was not much difference in the blood glucose concentrations of ZDF rats after subcutaneous injections of formulations of insulin containing zinc carbonate at two different zinc concentrations of 0.96 to 1.92 mg (Figure 2-1). Furthermore, the total duration of action in the animals for these formulations was shorter than that of Ultralente. Figure 2-5 depicts the blood glucose concentrations after a subcutaneous injection of insulin with different concentrations of zinc obtained from zinc oxide (another water-insoluble zinc compound). It is clear from the Figure 2-5 that
Figure 2-4  Comparison of blood glucose concentrations in ZDF rats after subcutaneous injection of insulin, ultralente and insulin with varying concentrations of zinc carbonate (Insulin dose-10 U/kg) (Mean ± SEM, n = 4)
Figure 2-5  Comparison of blood glucose concentrations in ZDF rats after a single subcutaneous injection of insulin with varying concentrations of zinc oxide (Insulin dose-10 U/kg) (Mean ± SEM, n = 4)
increasing the amount of zinc oxide in the suspension formulation of insulin does not prolong the blood glucose lowering effect of insulin.

Figure 2-6 shows a comparison of blood glucose concentrations in ZDF rats after subcutaneous injections of RHI with water-insoluble zinc carbonate and zinc oxide, and water-soluble zinc sulfate and zinc chloride. (Insulin dose-10 U/kg). It is evident from Figure 2-6 that the blood glucose lowering effect of insulin and maintenance of low blood glucose concentrations for a prolonged period of time in the ZDF rats was more than for the formulations prepared with water-soluble zinc salts than those prepared with water-insoluble salts. This could be due to complexation of zinc ions (obtained from the water-soluble salts) with the insulin and formation of an insoluble zinc-insulin hexamers. The hexamers could have been further stabilized to form higher molecular weight associates in presence of excess free zinc ions (86).

These high molecular weight insulin associates (72000-256,000 Dalton) and hexamers (36000 Dalton) cannot cross the capillary barrier, and can only reach the blood stream after dissociation to monomers (86). This explains the reason of slow and prolonged effect of insulin in presence of high concentration of water soluble-salts like zinc chloride and zinc sulfate. The water-insoluble zinc salts, on the other hand, could not dissolve in the aqueous medium and hence could not provide the zinc ions to complex with the insulin molecules.

2.3.2 In vitro release of insulin from insulin formulations with varying concentrations of zinc sulfate

To further investigate the effect of zinc on RHI formulations, in vitro release studies were performed on insulin formulations prepared with water soluble zinc sulfate. Figures 2-7 and 2-8 depict the percentage release of insulin from suspensions of insulin prepared without and with varying concentrations of zinc sulfate, respectively, across a semi permeable membrane (dialysis bags with a MWCO of 50,000). Figure 2-7 shows that approximately 100 percent of insulin was released within 12 hours through the semipermeable membrane from the RHI formulation without any additional zinc sulfate. However, as shown in Figure 2-8, the release of insulin from RHI formulations prepared with two different concentrations of zinc (0.24 and 0.48 mg/gram of formulation) obtained from zinc sulfate was only approximately three percent at 48 hour time point. Moreover, no insulin was released from these formulations until 4 hours, and no detectable quantity of insulin was observed from RHI formulations prepared with 0.96 and 1.92 mg/g of zinc (obtained from zinc sulfate) until 48 hours. The contents of dialysis bags containing RHI formulations were studied after 48 hours of release study.

Figure 2-9 depicts the percentage of insulin remaining in the dialysis bags after completion of the in vitro release study (i.e. 48 hours), with RHI formulations prepared with varying concentration of zinc. The results of the study also showed that after 48 hours of in vitro release study, approximately 3 percent of insulin was remaining inside the dialysis bags containing insulin formulations without any zinc. However, in the case
Figure 2-6  Comparison of blood glucose concentrations in ZDF rats after a single subcutaneous injection of insulin with water-soluble and water-insoluble zinc salts (Insulin dose-10 U/kg) (Mean ± SEM, n = 4)
Figure 2-7  Comparision of *in vitro* insulin release from insulin suspension formulations with and without zinc sulfate
Figure 2-8  Comparision of *in vitro* insulin release from insulin suspension formulations with zinc sulfate
Figure 2-9  Percentage of insulin remaining in the dialysis bags after completion of *in vitro* release study of insulin formulations with varying concentrations of zinc.
of insulin formulations prepared with varying concentrations of zinc sulfate, settled insulin powder could be seen at the bottom of the dialysis bags. Quantitation of insulin concentration in these dialysis bags showed that approximately 88 to 96 percentage of insulin was still present inside the dialysis bags even after 48 hours.

This further confirmed that fact that insulin forms hexamers and other complex associates of high molecular weight, which could not cross the dialysis bag with a MWCO of 50,000. To see the effect of pH on the concentration of insulin in the supernatant, insulin formulations with 0.24 mg of zinc/g of formulation (similar to Formulation OA-2-2) and without zinc (similar to Formulation OA-2-1) were prepared at different pH values of approximately 2.0, 3.0, 4.0, 5.0, and 5.6. Figure 2-10 shows the concentration of RHI in the supernatant obtained from these formulations at different pH values. It is clear from the figure that, as the pH decreased from 5.6 to 4.0, the amount of insulin dissolved in the supernatant increased from approximately 50 μg/mL to 1000 μg/mL. As the pH decreased further to pH 2.0, the amount of insulin dissolved in the supernatant increased up to 1500 μg/mL, i.e. insulin dissolved completely at pH 2.0. This is in agreement with the prior findings that insulin dissolves at lower pH values (238, 241).

Figure 2-10 also shows the concentration of insulin in the supernatant in formulations prepared with 0.24 mg of zinc/gram of formulation (similar to Formulation OA-2-2) at different pH values of approximately 2.0, 3.0, 4.0, 5.0, and 5.6. It is evident from the figure that the concentration of insulin in the supernatant in the presence of zinc was approximately 40 μg/mL and 100 μg/mL at pH 5.0 and 4.0, respectively. These insulin concentrations are considerably less than the insulin concentrations (approximately 400 μg/mL and 100 μg/mL) in the supernatant of the insulin formulations without zinc, at the similar pH values of 5.0 and 4.0. This may be because of interaction of insulin with zinc, leading to the formation and precipitation of insulin hexamers out of the solution. However at pH 3.0 and 2.0, the insulin concentrations in the supernatant of both the formulations prepared with and without zinc were similar. This also proves that insulin does not interacts with zinc at lower pH values (86).

Figure 2-11 also shows the pH and concentration of insulin in the supernatant obtained from formulations of insulin with varying concentrations zinc sulfate. The pH of insulin formulation without any additional zinc sulfate was approximately 5.6. As zinc sulfate was added to this insulin formulation, the pH decreased from 5.6 to approximately 5.05. With further addition of zinc sulfate, the pH decreased up to 4.85. The isoelectric point of insulin is 5.4 (238). Since insulin is more soluble in the acidic medium, as the pH went down towards the more acidic pH, i.e. away from the isoelectric point of insulin, more insulin dissolved in the medium. However, when these insulin formulations with varying concentrations of zinc sulfate were centrifuged and concentrations of insulin were determined in the supernatant, an opposite trend was observed.

Figure 2-11 also shows the concentration of insulin in the supernatant obtained from formulations of insulin prepared with varying concentrations zinc sulfate. As shown in Figure 2-11, as the zinc sulfate concentration increased in the formulations, the pH of
Figure 2-10  Concentrations of insulin in the supernatant obtained from formulations of insulin alone or insulin with zinc sulfate (0.24mg/mL) at different pH values
Figure 2-11  pH and concentration of insulin in the supernatant obtained from formulations of insulin (RHI) with varying concentrations zinc sulfate
the formulation decreased from 5.57 to 4.85. However, as the zinc sulfate concentration increased in the formulations, the insulin concentration in the supernatant of the formulation also decreased from approximately 50 μg/mL to 5 μg/mL. This approximately 10 fold decrease, in insulin concentration in the supernatant of the formulations may be due to an increase in the zinc ions in the formulations.

To investigate this phenomenon further, four different formulations were prepared at the final pH of approximately 5.05, 4.95, 4.90 and 4.85. These final pH values were selected based on the pH values of formulations of insulin with varying concentrations zinc sulfate. The insulin concentration was determined in the supernatant of these formulations after similar treatment as of formulations of insulin with varying concentrations zinc sulfate. As indicated in Figure 2-12 the insulin concentration, in the supernatant of the insulin formulations was much lower in the presence of zinc at similar pH values.

Figure 2-13 depicts the possible explanation of this finding. In the formulations of RHI with varying concentrations zinc sulfate, insulin was in suspension and zinc ions were in solution. With the addition of more zinc sulfate, the pH of the medium decreased and the medium became acidic. Since the solubility of insulin is more at the acidic pH, more insulin solublized at lower pH and hence was available to form hexamers in the presence of excess of zinc ions. As more hexamers and loose associates of hexamers were formed, they precipitated out of the solution thus resulting in a decrease in the net insulin concentration in the supernatant. When these insulin formulations with varying concentrations of zinc and containing hexamers were administered to the ZDF rats, the insulin hexamers and loose associates could not cross the capillary barrier; and thus formed a depot at the site of the injection. This depot formation may be the reason for the prolonged in vivo effect of the insulin formulations prepared with varying concentrations zinc sulfate, after subcutaneous injections in the ZDF type 2 diabetic rats.

2.4 Conclusions

The blood glucose lowering effect of insulin could be extended by varying concentrations of both the water-soluble salts of zinc. However, the blood glucose lowering effect of insulin and maintenance of low blood glucose concentrations for a prolonged period of time in the ZDF rats was more than for the formulations prepared with water-soluble zinc salts than those prepared with water-insoluble salts. This could be due to complexation of zinc ions (obtained from the water-soluble salts) with the insulin and formation of an insoluble zinc-insulin hexamers.
Figure 2-12  Concentration of insulin (RHI) in the supernatant obtained from formulations of insulin alone at different pH, or formulations of insulin with varying concentrations zinc sulfate
Figure 2-13  Possible mechanism of extended *in vivo* effect of formulations of insulin with varying concentrations zinc sulfate
CHAPTER 3. CONTROLLED RELEASE OF INSULIN FROM NOVEL BIODEGRADABLE INJECTABLE GELS

3.1 Introduction

Diabetes mellitus is a chronic disorder characterized by hyperglycemia. It is recognized as a worldwide epidemic by the WHO. The disease may be further classified as type-1 and type-2. Type-1 diabetes mellitus patients can survive only by regular administration of insulin, whereas type-2 diabetes mellitus patients can be treated with oral hypoglycemic agents and some other new classes of anti diabetic drugs. However, type-2 patients may also need insulin therapy at some point. Hence, insulin is pivotal in treating diabetes (242-244).

For successful insulin therapy, daily subcutaneous injections of insulin are required. Subcutaneous insulin has been satisfactory in terms of efficacy in majority of cases. However the use of these subcutaneous injections may cause hyperinsulinemia which may lead to hypoglycemia and diabetic micro and macro angiopathy. Further, these subcutaneous injections of insulin are also associated with pain, inconvenience, physiological stress, high cost, difficulty in handling insulin, localized insulin deposition leading to local hypertrophy and fat deposition at site of injection. Most insulin regimens require multiple daily injections which are associated with poor patient compliance. Reduction in number of insulin injections can significantly improve efficacy of insulin therapy (6, 18).

Insulin interacts with zinc ions to form hexamers. In the presence of zinc, these hexamers further get stabilized. When these hexamers are injected via subcutaneous route, they form a depot at the site of injection because of their inability to cross the capillary barriers. To get absorbed into the blood stream, these hexamers have to dissociate first into dimers and then to monomers. This leads to a delayed and extended effect of insulin in presence of zinc (124, 238, 245).

Various controlled release forms of insulin like microspheres, implants, gels, liposomes, and stereocomplexes have been investigated as extended release dosage forms of insulin. A novel injectable biodegradable gel system for controlled release drug delivery has been designed and characterized in our laboratory. It consists of a biodegradable polymer, PLGA, biocompatible plasticizer(s) and an active pharmaceutical ingredient (API). Chen et al. reported use of these gel systems for extended delivery of insulin for approximately one week (113). Chen et al. used ultralente crystals extracted from a commercial formulation (113). They also showed that adding a water-insoluble salt of zinc, zinc carbonate, further slowed down the release profile from these gels.

This work is continuation of the work done by Chen et al. and the aim of this study was to design insulin gel formulations, which could provide constant blood glucose lowering effect of the released insulin in diabetic rats for one week or longer, after a single subcutaneous injection. However in this part of study we intend to study the
incorporation of water-soluble salt, zinc sulfate, and its effect on prolongation of blood glucose lowering effect of regular human insulin, released from these gels.

3.2 Materials and methods

3.2.1 Materials

Insulin (RHI) powder (26.9 IU/mg, 0.4% zinc) used in this study was purchased from Diosynth France S.A. (Usine St. Charles, France). The polylactic-co-glycolic acid (PLGA) used in this study was Medisorb® 5050 DL 1A (i.v. 0.09 dL/g). It was donated by Alkermers Inc. (Cambridge, MA). According to the manufacturer’s technical information, PLGA 5050 DL 1A has a degradation time of 1-2 weeks (246). This grade of PLGA was selected because it has the shortest degradation time. Zinc sulfate used in this study was of laboratory grade and was bought from Sigma Chemicals (St. Louis, MO). Acetyl triethyl citrate (ATEC) and triethyl citrate (TEC) used in this study were generous donations from Morflex Inc. (Greensboro, NC). Male ZDF (fa/fa) rats weighing 300-400 grams used for the in vivo studies were received as donations from the Charles River Laboratories, Inc., (Wilmington, MA). Blood glucose was measured using Bayer Ascensia® Breeze blood glucose meter (Bayer Corporation, Elkhart, IN).

3.2.2 Methods

3.2.2.1 Preparation of insulin-zinc sulfate co-precipitates. An appropriate quantity of insulin (RHI) powder was suspended in normal saline to make insulin suspensions. Table 3-1 shows the composition of insulin formulations containing varying concentrations of zinc sulfate. The ingredients were weighed into a glass vial and water was added to bring the final weight to approximately 60 grams. Any lumps, if present, were broken using a stainless steel spatula. These formulations were kept at room temperature overnight (approximately 16 hours). The next day, the supernatant of the formulation was carefully removed and the precipitates were dried under vacuum at room temperature for 24 hours. The next day, the supernatant of the formulation was carefully removed and the precipitates were dried under vacuum at room temperature for 24 hours. The dried insulin-zinc sulfate co-precipitates were checked for any lumps, which if present were carefully triturated using a stainless steel spatula. Appropriate amounts of these insulin-zinc sulfate co-precipitates were loaded into the blank gel as shown in the Table 3-2 one day prior to subcutaneous administration to the ZDF rats. Zinc concentrations in the insulin-zinc precipitates were determined by atomic absorption spectrophotometer.

3.2.2.2 Fabrication of the gels. The gel formulations were prepared in two steps: dissolution of polymer in the plasticizer(s) to obtain a blank gel and loading of insulin alone or insulin zinc sulfate co-precipitates or insulin and zinc sulfate separately into the blank gel. The polymer, PLGA was allowed to come to room temperature in vacuum desiccators after it was taken out of the freezer.
Table 3-1  Formulations of insulin suspensions

<table>
<thead>
<tr>
<th>Formulation #</th>
<th>Insulin (mg)</th>
<th>Zn SO4 (mg)</th>
<th>NaCl (mg)</th>
<th>Adjust total weight with H2O (g)</th>
<th>Amount of zinc in the formulation (mg/g)</th>
<th>Molar ratio of insulin to zinc</th>
</tr>
</thead>
<tbody>
<tr>
<td>OA-3-3</td>
<td>90</td>
<td>63</td>
<td>530</td>
<td>60</td>
<td>0.24</td>
<td>1:14</td>
</tr>
<tr>
<td>OA-3-4</td>
<td>90</td>
<td>127</td>
<td>521</td>
<td>60</td>
<td>0.48</td>
<td>1:28</td>
</tr>
<tr>
<td>OA-3-5</td>
<td>90</td>
<td>253</td>
<td>501</td>
<td>60</td>
<td>0.96</td>
<td>1:56</td>
</tr>
<tr>
<td>OA-3-6</td>
<td>90</td>
<td>507</td>
<td>464</td>
<td>60</td>
<td>1.92</td>
<td>1:112</td>
</tr>
</tbody>
</table>
Table 3-2  Formulation composition of PLGA gel containing insulin and zinc

<table>
<thead>
<tr>
<th>Formulation #</th>
<th>ATEC (%)</th>
<th>TEC (%)</th>
<th>Insulin + ZnSO₄ co-precipitate (%)</th>
<th>Polymer (%)</th>
<th>Insulin + ZnSO₄ co-Precipitate of formulation</th>
<th>Actual Ratio Insulin: zinc</th>
</tr>
</thead>
<tbody>
<tr>
<td>OA-3-7</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
<td>OA-3-3</td>
<td>1:14</td>
</tr>
<tr>
<td>OA-3-8</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
<td>OA-3-4</td>
<td>1:28</td>
</tr>
<tr>
<td>OA-3-9</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
<td>OA-3-5</td>
<td>1:56</td>
</tr>
<tr>
<td>OA-3-10</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
<td>OA-3-6</td>
<td>1:112</td>
</tr>
</tbody>
</table>
The desired amount of plasticizer(s) was weighed into the scintillation vial containing the polymer. A plastic screw cap with aluminum liner was used to seal the vial. The vial was then vortexed on a vortex mixer to disperse the polymer particles in the plasticizer(s). The vial was placed and in a Lab-Line orbital shaker (Model 3527, Lab-line, Melrose Park, IL) maintained at 37°C and shaken at 150 rpm. It took 2 to 3 days for the polymer to completely dissolve.

The resulting blank gel was stored in a vacuum desiccator at room temperature until further use (typically 1 to 2 days). For preparing insulin-loaded gel, a desired amount of blank gel was transferred to a new glass scintillation vial, and an appropriate amount of either pure insulin particles (Table 3-3) or insulin-zinc sulfate co-precipitates (Table 3-4) was accurately weighed and transferred into the blank gel.

3.2.2.2 Preparation of physical mixtures of insulin and zinc sulfate. In the formulations containing insulin and zinc sulfate, the zinc sulfate was first passed through sieve number 400 (37.5 μm). An appropriate amount of the sieved zinc sulfate was weighed and mixed with insulin by the geometric dilution method. The resulting mixture of insulin and zinc sulfate was loaded into the blank gels. A stainless steel spatula was used to stir the mixture. Any visible agglomerates of insulin and/or zinc sulfate were broken using the spatula. Mixing was continued until a uniform mixture of insulin and zinc sulfate in the gel was produced.

3.2.2.3 Pharmacodynamic studies of in ZDF rats. All the experimental procedures for the animal studies were approved by the University of Tennessee Health Science Center’s Animal Resources Advisory Committee (Protocol 1622). The ZDF rats (Charles River Laboratories, Wilmington, MA) were at least 16 weeks old and weighed approximately 300 to 400 g. Hyperglycemia (> 400 mg/dL) in the ZDF rats was confirmed by blood glucose measurements.

The rats were group-housed (3 per cage) in the animal facility under a 12-hour light/dark cycle. They had access to food and water ad libitum throughout the studies. Before the animals were shipped from Charles River Laboratories, they were fed Purina diet #5008 until they fully developed diabetes (12 weeks old). The animals were on regular laboratory rodent food in our facility. The rats were acclimated to the environment for at least three days before they were used in the studies.

On each day of the study, the ZDF rats were transferred to the procedure room. The rats were randomly divided into groups, with 4 to 6 animals per group. Each group, except the control group, received one dose of an insulin formulation. The control group did not receive any insulin injection. Before injection, each rat was placed in an induction chamber and anesthetized by isoflurane (Halocarbon Product Corporation, River Edge, NJ) using a SurgiVet/Anesco ventilator (Waukesha, WI). Body weight of the animal was recorded to determine the volume of the insulin formulation to be injected. Blood glucose was measured using Bayer Ascensia® DEX® 2 or Ascensia Breeze blood glucose meter (Bayer Corporation, Elkhart, IN). The dose of insulin formulation was injected subcutaneously at the dorsal neck of the rat using a 1 mL syringe with an attached 23
Table 3-3  Formulation composition of PLGA gel containing insulin and zinc sulfate physical mixtures

<table>
<thead>
<tr>
<th>Formulation #</th>
<th>ATEC (%)</th>
<th>TEC (%)</th>
<th>Insulin (%)</th>
<th>Polymer (%)</th>
<th>ZnSO₄ (%)</th>
<th>Actual ratio insulin: zinc</th>
</tr>
</thead>
<tbody>
<tr>
<td>OA-3-11</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
<td>1</td>
<td>1:14</td>
</tr>
<tr>
<td>OA-3-12</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
<td>2</td>
<td>1:28</td>
</tr>
<tr>
<td>OA-3-13</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
<td>2.8</td>
<td>1:14</td>
</tr>
<tr>
<td>OA-3-14</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
<td>5.6</td>
<td>1:28</td>
</tr>
</tbody>
</table>
Table 3-4  Zinc contents in insulin-zinc co-precipitates

<table>
<thead>
<tr>
<th>Precipitate of Formulation #</th>
<th>Formulation composition of precipitate (mg/g)</th>
<th>Amount zinc (µm)/insulin (mg)</th>
<th>Ratio insulin : zinc</th>
</tr>
</thead>
<tbody>
<tr>
<td>OA-3-3</td>
<td>RHI-Zn-0.24</td>
<td>7.78 ± 0.69</td>
<td>06:04.1</td>
</tr>
<tr>
<td>OA-3-4</td>
<td>RHI-Zn-0.48</td>
<td>10.88 ± 0.96</td>
<td>06:05.1</td>
</tr>
<tr>
<td>OA-3-5</td>
<td>RHI-Zn-.96</td>
<td>13.77 ± 2.77</td>
<td>06:07.2</td>
</tr>
<tr>
<td>OA-3-6</td>
<td>RHI-Zn-1.92</td>
<td>11.67 ± 3.82</td>
<td>06:06.1</td>
</tr>
</tbody>
</table>
gauge 1 inch needle (Franklin Lakes, NJ). After injection, the rats were returned to the cages and allowed to recover from anesthesia. At pre-determined time intervals, the rats were anesthetized by aforementioned method and their blood glucose concentrations were measured.

A plot of blood glucose concentrations versus time was generated from the data to describe the glucose lowering effect of insulin, from which information such as time of peak action and duration of action could be obtained.

3.3 Results and discussion

Pharmacodynamic studies of different gel formulations of insulin in ZDF rats: Two different gel formulations consisting of 5 and 10 % PLGA were made by suspending insulin particles in the blank gels (Table 3-5). These insulin formulations were administered subcutaneously to ZDF rats to evaluate the biological effect of insulin released from these formulations. Specifically, the blood glucose concentrations and body weights of the rats were measured at periodic time intervals following the subcutaneous injection. Figure 3-1 shows changes in blood glucose concentrations of ZDF rats after a subcutaneous injection of PLGA (5 and 10%) gel formulation loaded with 4% insulin (Formulations OA-3-1 and OA-3-2). It is evident from Figure 3-1 that the blood glucose concentrations decreased from approximately 554 ± 29.4 mg/dL to 63 ± 10.6 mg/dL within 6 hours after a subcutaneous injection of Formulation OA-3-1 (5% PLGA) in the ZDF rats. The blood glucose concentrations were below 80 mg/dL in all the six rats, i.e. all the rats were hypoglycemic within six hours after injection. Further, the blood glucose concentrations of all the six rats were below 70 mg/dL within 12 hours after subcutaneous administration of the formulation. However, the blood glucose concentrations of all the six rats increased to normally high values of 443 ± 55.7 mg/dL within 24 hour after subcutaneous administration of the formulation, and remained high thereafter. A similar blood glucose concentration profile was obtained after subcutaneous injection of formulation OA-3-2 (10% PLGA). The blood glucose concentrations decreased from approximately 452 ± 27.38 mg/dL to 92.5 ± 4.9 mg/dL within 6 hours after a subcutaneous injection of the formulation.

The blood glucose concentrations further declined to below 75 mg/dL after 12 hours after administration of the formulation. As with 5 % PLGA formulation, the blood glucose concentrations of all of the six rats increased to normally high values of 331 ± 35.1 mg/dL, 24 hours after administration of the formulation and remained high thereafter. It is evident from these results that these insulin formulations consisting of 5 and 10% PLGA release insulin at a very rapid rate, leading to hypoglycemia in all the rats. It is evident from these results that these insulin formulations consisting of 5 and 10% PLGA release insulin at a very rapid rate, leading to hypoglycemia in all the rats. Further, the blood glucose concentrations could not be maintained at low concentrations (80 to 100 mg/dL) for a long period of time, and the all the rats were hyperglycemic within 24 hours after the subcutaneous injection of these formulations.
Table 3-5  Formulation composition of PLGA gel of insulin

<table>
<thead>
<tr>
<th>Formulation #</th>
<th>ATEC (%)</th>
<th>TEC (%)</th>
<th>Insulin (%)</th>
<th>Polymer (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>OA-3-1</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>OA-3-2</td>
<td>68.25</td>
<td>22.75</td>
<td>4</td>
<td>10</td>
</tr>
</tbody>
</table>
Figure 3-1  Changes in blood glucose concentrations of ZDF rats after a subcutaneous injection of PLGA (5% and 10%) gel formulations loaded with 4% insulin (Formulations OA-3-1 and OA-3-2) (Mean ± SEM, n = 5)
Figure 3-2 shows changes in the body weights of ZDF rats after a subcutaneous injection of the PLGA (5 and 10%) gel formulation loaded with 4% insulin (Formulations OA-3-1 and OA-3-2). The body weights of these ZDF rats 24 hours after subcutaneous injection of the PLGA formulations (5 and 10%) were higher than before the injection of the formulations. However, the body weights came back to approximate initial values within 48 hours after the injection of these formulations. This pharmacodynamic effect shown by the changes of the body weights after the subcutaneous injection is similar to the pharmacodynamic effect of the changes in the blood glucose concentrations. This shows that these formulations (Formulations OA-3-1 and OA-3-2) could not control the release of insulin for a prolonged period of time after a single subcutaneous injection in the ZDF rats, and the insulin effect lasted for less than 24 hours.

As shown in the previous study (Chapter-2) and reported by other groups, insulin zinc interaction can extend the in vivo duration of action of insulin (238). Further, as shown by Chen et al. (113) and Kim et al. (247), incorporation of water-insoluble zinc salt in a similar type of polymer based gels slowed down the insulin release and extended the in vivo duration of action of insulin. In this study, we purposed that incorporation of a water-soluble zinc salt, zinc sulfate, in these PLGA formulations of regular human insulin can prolong the blood glucose lowering effect of regular human insulin.

The insulin-zinc co-precipitates (Formulations OA-3-3, OA-3-4, OA-3-5 and OA-3-6) were loaded in 5% PLGA gels and four different formulations were obtained (Formulations OA-3-7, OA-3-8, OA-3-9 and OA-3-10). These formulations were injected subcutaneously in ZDF rats and change in blood glucose concentrations and body weights were monitored for a week. Figure 3-3 shows the changes in blood glucose concentrations of ZDF rats after a subcutaneous injection of 5% PLGA gel formulation loaded with 4% insulin alone (Formulation OA-3-1) or insulin-zinc sulfate co-precipitates (Formulations OA-3-7 and OA-3-8). It is evident from the figure that the blood glucose lowering effect of 5% PLGA formulation consisting of insulin alone (Formulation: OA-3-1) lasted less than 24 hours. After a subcutaneous injection of 5% PLGA formulation (Formulation OA-3-7), consisting of insulin-zinc sulfate co-precipitates of Formulation OA-3-3 (0.24 mg of zinc /gram of formulation), the blood glucose concentrations remained below 100 mg/dL until 24 hours before increasing to approximately 200 mg/dL after 48 hours after the subcutaneous injection of the formulation. The blood glucose concentrations steadily increased above 350 mg/dL and remained above this concentration thereafter. In the case of 5% PLGA formulation (Formulation OA-3-8), consisting of insulin-zinc co-precipitates of Formulation OA-3-4 (0.48 mg of zinc gram of formulation), the blood glucose concentrations decreased from approximately 426 ± 15.9 mg/dL to 75 ± 17.3 mg/dL within 12 hours after the subcutaneous injection. The blood glucose concentrations remained below 150 mg/dL for up to 3 days before increasing to approximately 330 mg/dL after 4 days after the subcutaneous injection of the formulation. The blood glucose concentrations increased steadily above 350 mg/dL and remained above this concentration thereafter. This shows that when the insulin-zinc co-precipitates were
Figure 3-2  Changes in body weights of ZDF rats after subcutaneous injections of PLGA (5 and 10%) gel formulations loaded with 4% insulin (Formulations OA-3-1 and OA-3-2) (Mean ± SEM, n = 5)
Figure 3-3 Changes in blood glucose concentrations of ZDF rats after subcutaneous injections of PLGA (5%) gel formulations loaded with 4% insulin alone (Formulation OA-3-1) or insulin-zinc co-precipitates (Formulations OA-3-7 and OA-3-8) (Mean ± SEM, n = 5)
loaded in the 5% PLGA gel formulations (Formulations OA-3-5 and OA-3-6) and injected subcutaneously in the ZDF rats, the blood glucose concentrations decreased slowly and remained at low concentrations (between 100 and 200 mg/dL) for a longer period of time. Addition of zinc also reduced the chances of occurrence of hypoglycemia in the ZDF rats after the subcutaneous injection of these PLGA formulations containing insulin. Further, with increase in the zinc, the in vivo effect of insulin was extended from 1 day to up to 3 days.

Figure 3-4 shows the changes in blood glucose concentrations of ZDF rats after a subcutaneous injection of 5% PLGA gel formulation loaded with 4% insulin alone (Formulation OA-3-1) or insulin-zinc co-precipitates (Formulations OA-3-9 and OA-3-10).

Figure 3-4 shows the changes in blood glucose concentrations of ZDF rats after a subcutaneous injection of 5% PLGA gel formulation loaded with 4% insulin alone (Formulation OA-3-1) or insulin-zinc co-precipitates (Formulations OA-3-9 and OA-3-10). After subcutaneous injection of 5% PLGA formulation (Formulation OA-3-9), consisting of insulin-zinc co-precipitates of Formulation OA-3-5 (0.96 mg of zinc /gram of formulation), the blood glucose concentrations decreased from approximately 492 ± 29.3 mg/dL to 105 ± 19.2 mg/dL within 12 hours after the subcutaneous injection. The blood glucose concentrations remained below 150 mg/dL for up to 2 days before increasing to approximately 300 mg/dL after 3 days after the subcutaneous injection of the formulation. The blood glucose concentrations increased steadily above 350 mg/dL and remained above this concentration thereafter. In the case of 5% PLGA formulation (Formulation OA-3-10 ) consisting of insulin- zinc co-precipitates of Formulation (Formulation OA-3-6 with 1.92 mg of zinc /gram of formulation), the blood glucose concentrations decreased slowly from approximately 467 ± 51.5 mg/dL to 99.8± 17.5 mg/dL within 24 hours after the subcutaneous injection. The blood glucose concentrations remained below 175 mg/dL for up to 3 days before increasing to approximately 300 mg/dL after 4 days after the subcutaneous injection of the formulation.

The blood glucose concentrations increased steadily above 350 mg/dL and remained above this concentration thereafter. This change in blood glucose concentrations shows that the effect of insulin does not last longer than 3 days after the subcutaneous injection of these 5% PLGA formulations containing insulin-zinc co-precipitates. Further, there was an increase in the duration of blood glucose lowering effect of insulin from Formulation OA-3-8 (5% PLGA gel containing 4% insulin-zinc co-precipitate with 0.48 mg of zinc/ gram of formulation) as compared to Formulation OA-3-7 (5% PLGA gel containing 4% insulin-zinc co-precipitate with 0.24 mg zinc / gram of formulation) and Formulation OA-3-1 (5% PLGA gel containing 4% insulin). However, there was no further increase in the in vivo duration of insulin when 5% PLGA gel formulations loaded with insulin-zinc co-precipitates having more zinc sulfate (obtained from Formulation OA-3-5 with 0.96 mg zinc/ gram of formulation and Formulation OA-3-6 with with 1.92 mg zinc / gram of formulation), were subcutaneously injected in the rats.
Figure 3-4  Changes in blood glucose concentrations of ZDF rats after subcutaneous injections of PLGA (5%) gel formulations loaded with 4% insulin alone (Formulation OA-3-1) or insulin-zinc co-precipitates (Formulations OA-3-9 and OA-3-10) (Mean ± SEM, n = 5)
After subcutaneous injections of 5 and 10% PLGA formulations of insulin (Figure 3-1) all the rats went to hypoglycemia within 6 hours. However, as the insulin-zinc co-precipitates loaded gels were subcutaneously injected in rats, the number of rats going into hypoglycemia decreased substantially. The blood glucose concentrations also decreased slowly after the subcutaneous injection of the 5% PLGA formulations of insulin-zinc co-precipitates.

Figure 3-5 shows the changes in the body weights of ZDF rats after a subcutaneous injections of 5% PLGA gel formulations loaded with 4% insulin-zinc co-precipitates (Formulations OA-3-7, OA-3-8, OA-3-9 and OA-3-10). It is evident from the figure that the body weights of ZDF rats increased 24 hours after the subcutaneous injections. However, in the case of Formulation OA-3-7, a very rapid weight gain was observed on day 1, after which it slowly declined. In the case of other formulations, the body weights of ZDF rats increased on day 1 and remained steady for up to day 3 followed by a slow decline after the subcutaneous injections. This change in body weights also showed that the effect of insulin does not last longer than 3 days after subcutaneous injection of these 5% PLGA formulations containing insulin-zinc co-precipitates. Since the blood glucose lowering effect of these 5% PLGA formulations containing insulin-zinc co-precipitates could not be extended for more than three days, the actual zinc content of these insulin-zinc co-precipitates was determined using atomic absorption spectrophotometer. As shown in Table 3-5, the actual zinc content of these co-precipitates was much lower. Two zinc ions are required to interact with six molecules of insulin to form a hexamer. These insulin-zinc co-precipitates have 4 to 7 zinc ions per six insulin molecules, which may be sufficient to transform all the insulin molecules into hexamers. Therefore, the initial blood glucose lowering effect was slow in formulations containing insulin-zinc co-precipitates. However, after subcutaneous injections, these co-precipitates were exposed to interstitial fluid, which dissolved and quickly removed the zinc ions from the injection site. Furthermore, the zinc ions present in these co-precipitates were not in excess to extend the effect of insulin for a long time.

To increase the amount of zinc in the formulations, zinc sulfate was directly loaded in the PLGA gels. Figure 3-6 shows the change in blood glucose concentrations in ZDF rats after subcutaneous injections of insulin loaded in 5% PLGA gel formulations with different concentrations of zinc sulfate (Formulations OA-3-11 and OA-3-12 containing 1 and 2% zinc sulfate, respectively). Formulation OA-3-1 prepared with 5% PLGA and having no zinc sulfate in it, showed a rapid decline in the blood glucose concentrations; reaching below 75 mg/dL in 6 hours. The blood glucose concentrations remained below 75 mg/dL for up to 12 hours, before climbing back to the normally high concentrations 350 mg/dL within 24 hours.

Formulations OA-3-11 and OA-3-12 containing 1 and 2% of zinc sulfate respectively, were prepared with 5% PLGA. Formulation OA-3-11 (containing 1% zinc sulfate) showed a slow decline in the blood glucose concentrations, reaching below 200 mg/dL in 12 hours and below 75 mg/dL in 24 hours. The blood glucose concentrations remained below 100 mg/dL for 3 days before slowly climbing up to 300 mg/dL concentrations at 5 days and remained above this concentration thereafter.
Figure 3-5 Changes in body weights of ZDF rats after subcutaneous injections of PLGA (5%) gel formulations loaded with 4% insulin-zinc co-precipitates (Formulations OA-3-7, OA-3-8, OA-3-9 and OA-3-10) (Mean ± SEM, n = 5)
Figure 3-6  Changes in blood glucose concentrations of ZDF rats after subcutaneous injections of insulin alone (Formulations OA-3-1) or physical mixtures of insulin (4%) and zinc sulfate (1 or 2%) loaded in PLGA (5%) gel formulations (Formulations OA-3-11 and OA-3-12) (Mean ± SEM, n = 5)
Formulation OA-3-12 (containing 2% zinc sulfate) showed a similar pattern as that of Formulation OA-3-11. When the zinc sulfate increased in the formulation, the blood glucose concentrations declined below 100 mg/dL even more slowly in 24 hours. The blood glucose concentrations remained below 200 mg/dL for 4 days before slowly climbing up to 300 mg/dL concentrations at 5 days and remained above this concentration thereafter. Thus, with addition of zinc sulfate in the 5% PLGA formulations, the blood glucose concentrations decreased at a slower rate. However, even with addition of 2% zinc sulfate, the blood glucose concentrations remained below 200 mg/dL only for 4 days. To investigate the effect of even higher concentrations of zinc sulfate, the amount of zinc sulfate was increased to 2.8 and 5.4% in the 5% PLGA gel formulations. Figure 3-7 shows the change in blood glucose concentrations in ZDF rats after subcutaneous injections of insulin loaded in 5% PLGA gels with higher concentrations of zinc sulfate (2.8 and 5.4%). Formulations OA-3-13 and OA-3-14 containing 2.8 and 5.6% of zinc sulfate, respectively were prepared with 5% PLGA. Formulation OA-3-13 also showed a slow decline in the blood glucose concentrations, reaching below 100 mg/dL in 24 hours.

Formulation OA-3-13 also showed a slow decline in the blood glucose concentrations, reaching below 100 mg/dL in 24 hours. The blood glucose concentrations remained between 100 to 250 mg/dL for 9 days before slowly climbing up to 300 mg/dL concentrations at 10 days. The blood glucose concentrations remained above 300 mg/dL thereafter. Formulation OA-3-14 (containing 5.6% zinc sulfate) showed a similar pattern as that of Formulation OA-3-13. However, the decline in blood glucose concentrations was slower (reaching below 100 mg/dL in 2 days) and lasted slightly longer after a subcutaneous injection of Formulation OA-3-14 compared to that of Formulation OA-3-13. Hence, the effect of insulin on the blood glucose concentrations was substantially increased with increasing zinc concentrations in the insulin-loaded gel formulations.

Figure 3-8 shows changes in body weights of ZDF rats after subcutaneous injections of 5% PLGA gel formulations loaded with 4% insulin and zinc sulfate mixture (Formulations OA-3-11, OA-3-12, OA-3-13 and OA-3-14). The changes in body weights of the rats repeated the pharmacodynamic trend that had been observed with the change in blood glucose concentrations after subcutaneous injections of these formulations.

### 3.4 Conclusions

A biodegradable injectable gel formulation prepared with zinc salts was able to sustain the release of insulin for up to 8 to 10 days following a single subcutaneous injection. The released insulin from the gel formulations was able to suppress the high blood glucose concentrations for the same period of time in the ZDF rats. However, a biodegradable injectable gel formulation of human insulin without zinc salts could maintain the blood glucose concentrations at low concentrations only for one day.
Figure 3-7 Changes in blood glucose concentrations of ZDF rats after subcutaneous injections of insulin loaded in PLGA (5%) gel formulations with different concentrations of zinc sulfate (2.8% and 5.4%) (Formulations OA-3-13 and OA-3-14) (Mean ± SEM, n = 5)
Figure 3-8  Changes in body weights of ZDF rats after subcutaneous injections of PLGA (5%) gel formulations loaded with 4% insulin and zinc sulfate mixtures (Formulations OA-3-11, OA-3-12, OA-3-13 and OA-3-14) (Mean ± SEM, n = 5)
CHAPTER 4. CONTROLLED RELEASE OF MODIFIED INSULIN GLARGINE FROM NOVEL BIODEGRADABLE INJECTABLE GELS

4.1 Introduction

Diabetes mellitus is a chronic metabolic disorder characterized by hyperglycemia, which may be due to deficiency or ineffectiveness of metabolic hormone insulin produced by Langharan’s beta cells of pancreas. The form of diabetes characterized by absolute insulin deficiency is known as type-1. The most prevalent form of diabetes is type-2 diabetes mellitus. It accounts for 90% of the cases and is due to ineffectiveness of insulin produced by the pancreas (6, 18, 242).

The major objective of diabetes treatment is to achieve normoglycemia by maintaining appropriate concentrations of insulin in the blood throughout the day. Insulin has been complexed with protamine or zinc to delay and extend the release \textit{in vivo} and prolong its effect for one day. However, these formulations have been associated with side effects such as hypoglycemia, and inter and intra-subject variability in absorption, and shorter duration of action than expected, thus leading to poor management of diabetes (6, 18, 248).

For intensive blood glucose control in people suffering from type-1 and type-2 diabetes, supply of basal insulin (a steady, low concentration of insulin that is constantly present in the circulation to cover post-prandial and overnight fasting periods) is very important. Protracted acting insulin preparations supply this basal concentration in clinical settings (128, 249, 250). One of the formulations that maintain basal concentration insulin is modified insulin called insulin glargine. It has been altered at two different positions: in A-chain, at position 21, where the asparagine has been substituted with glycine, thus imparting more resistance to deamidation in the acidic environment and providing more stability. The C-terminus of the B-chain has been elongated by the addition of two arginine molecules. This addition of two positive charges by the addition of arginines shifted the isoelectric point of the modified insulin glargine from pH 5.4 to 6.7. These modifications have made it possible to formulate insulin glargine into slightly acid stable solution, which is easy to inject and improves dose reproducibility. This insulin glargine solution forms stable hexamers which precipitate at neutral pH at the subcutaneous site of injection, thus forming a depot of microprecipitates of the modified insulin glargine, which then dissolve at a steady rate for a prolonged period of time (128, 249, 251).

Insulin and modified insulin like glargine interact with zinc ions to form hexamers. In the presence of zinc, these hexamers get further stabilized. When these hexamers are injected via subcutaneous route, they form a depot at the site of injection because of their inability to cross the capillary barriers. To get absorbed into the blood stream, these hexamers have to dissociate first into dimers and then to monomers. This leads to a delayed and extended effect of insulin and modified insulin glargine.
A novel biodegradable injectable gel system for controlled release drug delivery has been designed and characterized in our laboratory. It consists of a biodegradable polymer, PLGA, biocompatible plasticizer(s) and an active pharmaceutical ingredient (API). To date, several molecules, including antibiotics, opiate analgesics and narcotic antagonists, and insulin have been successfully incorporated into the gel system, and controlled release of the aforementioned APIs have been achieved in our laboratory.

Therefore, the aim of this study was to develop an injectable biodegradable gel system which can be used to control the release of insulin glargine for one week or longer, after a single subcutaneous injection in type-2 diabetic ZDF rats.

4.2 Materials and methods

4.2.1 Materials

Insulin glargine (Lantus®, Aventis Pharma, Parsippany, NJ) was purchased from Cardinal Health and insulin glargine particles were extracted using dialysis bags (Spectra/Por® Float-A-Lyzer®, Biotech grade Regenerated Cellulose (RC), 5 mL, 3.5K MWCO, Spectrum labs, Rancho Dominguez, CA). Labconco FreeZone® 12 liter Console Freeze Dry System (Labconco, Kansas City, MO) was used for freeze drying. Recombinant human insulin (insulin) powder (26.9 IU/mg, 0.4% zinc) was purchased from Diosynth France S.A. (Usine St. Charles, France). The HPLC analysis of insulin was performed on a Shimadzu HPLC system (Shimadzu Scientific Instruments, Inc., Columbia, MD). The primary column was a 250 x 4.6 mm Altech Macrosphere RP 300 C18 5 μm column and the guard column was 7.5 x 4.6 mm (Altech, Deerfield, IL), packed with same material as the primary column. HPLC grade Acetonitrile (ACN), trifluroacetic acid (TFA) and distilled water were used for preparing the mobile phase. The PLGA used in this study was Medisorb® 5050 DL 1A (i.v. 0.09 dL/g) donated by Alkermers Inc. (Cambridge, MA). Triethyl citrate (TEC) and acetyltriethyl citrate (ATEC) donated by Morflex Inc., (Greensboro, NC) were used as plasticizers. male ZDF (fa/fa) rats (Charles River Laboratories, Inc., Wilmington, MA) weighing 300 to 400 grams were used for the in vivo studies. Blood glucose was measured using Bayer Ascensia® Breeze blood glucose meter (Bayer Corporation, Elkhart, IN). The concentration of glargine in the plasma or serum was determined using a Mercodia human insulin ELISA kit (Mercodia AB, Uppsala, Sweden). Blood glucose and glargine monitoring in the animals was stopped when the concentration of blood glucose went back to the normally high concentration.

4.2.2 Methods

4.2.2.1 Extraction of insulin glargine particles. The Spectra/Por® Float-A-Lyzer® (dialysis bags) were removed from the refrigerator and allowed to equilibrate to room temperature. The cap was removed and the sodium azide solution inside the bag was discarded using a pipette. The dialysis bags were then cleaned with distilled water and placed in a one liter beaker containing 900 mL of distilled water for one hour to remove
any remaining sodium azide. The bags were removed from the beaker and again washed several times with distilled water. Insulin glargine (Lantus®) solution (5 mL) was loaded into each bag using a pipette. The insulin glargine solution filled bags were kept in the beaker containing 900 mL of distilled water and the beaker with the dialysis bags was placed in an ice bath and stirred continuously with a magnetic stirrer for 24 hours. The distilled water in the beaker was completely replaced with fresh water after 2, 4, 8 and 12 hours. After 24 hours, the cap of the dialysis bag was carefully removed and the sample was recovered using a pipette. The recovered sample was placed in a pre-washed 20 mL scintillation vial, which was covered with aluminum foil. Holes were punctured in the aluminum foil using an 18 gauge needle. This aluminum foil covered scintillation vial was then placed in a freezer at -80°C for 6 hours. The freeze dryer was then switched on and vacuum was allowed to come to the maximum concentration. The samples were removed from -80°C and freeze dried for 12 hours. The dried samples were pooled together in a clean 20 mL scintillation vial and sealed with Parafilm, capped and stored at -20°C until used for further studies.

4.2.2.2 High pressure liquid chromatography method of insulin glargine. The HPLC analysis of the freeze-dried insulin glargine was performed using a Shimadzu HPLC system (Shimadzu Scientific Instruments, Inc., Columbia, MD) consisting of an SCL-10A vp system controller, a LC-10 AD vp pump, a DGU-14A degasser, an SIL-10 Ad vp autoinjector with a cooling system, a CTO-10A vp column oven, an SPD-10AD vp photo-diode array (PDA) detector and a computer loaded with the Shimadzu Class-VP 7.2 software. The primary column was a 250 x 4.6 mm Altech Macrosphere RP 300 C18 5 μm column and the guard column was 7.5 x 4.6 mm, packed with same material as the primary column.

A gradient elution method was used for analyzing the insulin. Mobile phase A was 0.15% v/v trifluoroacetic acid (TFA) in water and mobile phase B was 0.13% v/v TFA in 95% v/v acetonitrile in water. The mobile phases were filtered through a Whatmann 0.45 μm nylon membrane (Whatmann International Ltd, Maidstone, England) and degassed in an ultrasonicator (Model FS60, Fisher Scientific, Fair Lawn, NJ) for 10 minutes before pumping it through the HPLC system.

The initial mobile phase composition of 70% solvent A and 30% solvent B was maintained at a flow rate of 1 mL/min for 2 minutes. Between 2 and 16 minutes, the percentage of solvent B was increased linearly to 50%. These conditions were maintained from 5 to 21 minutes. Between 22 and 25 minutes, the percentage of solvent B was decreased linearly to 30%. The mobile phase was pumped at a flow rate of 1 mL/min. The injection volume of each sample, including the insulin glargine standards, insulin glargine controls, insulin glargine samples (unknown concentration) and blank samples (0.01N HCl) without insulin glargine), was 50 μL. The temperature of the HPLC column was maintained at 30°C by using a column oven. The primary detection wavelength of insulin was set to 285 nm.

4.2.2.3 Preparation of standard, control and sample solutions. The standard and control solutions of insulin glargine were prepared from the commercially available
Lantus® solution with a concentration of 3.63 mg/mL as the working stock solution. Standard solutions were prepared from the working stock solution by transferring 50, 75, 100, 150 and 200 μL of the working stock solution to five 2 mL volumetric flasks and making up the volume with 0.01N HCl (pH 2.0).

The insulin glargine controls were also prepared from the working stock solution with a concentration of 100 μg/mL. For sample preparations, approximately 5 mg of freeze-dried insulin glargine particles were weighed on a calibrated Cahn C-31 microbalance (Cahn Instruments, Cerritos, CA) and transferred to a volumetric flask and dissolved in 0.01N HCl (pH 2.0). The final volume was made with 0.01N HCl (pH 2.0). The dissolved samples were immediately loaded on the HPLC tray.

4.2.2.4 Process of chromatographic data. After each HPLC injection, the software generated the chromatograph of the sample and integrated the peak area of insulin glargine. The peak area of the standard solutions was then plotted against the concentration of insulin glargine. A linear regression method was used to generate the equation from which the concentration of the unknown sample could be interpolated from the peak area. The concentration of the insulin glargine control was interpolated and compared to the theoretical value. The concentration of insulin glargine samples was determined using the standard curve.

4.2.2.5 Pharmacodynamic studies of insulin and insulin glargine in ZDF rats. All the experimental procedures for the animal studies were approved by the University of Tennessee Health Science Center’s Animal Resources Advisory Committee. The ZDF rats were at least 16 weeks old (weighing 300 to 400 g). Hyperglycemia (> 400 mg/dL) in the ZDF rats was confirmed by blood glucose measurements. The rats were group-housed (3 per cage) in the animal facility under a 12-hour light/dark cycle. They had access to food and water ad libitum throughout the studies. Before the animals were shipped from Charles River Laboratories, they were fed Purina diet #5008 until they fully developed diabetes (12 weeks old). The animals were on regular laboratory rodent food in our facility. The rats were acclimated to the environment for at least three days before they were used in the studies.

Insulin (RHI), insulin glargine and the commercially available formulation of insulin glargine, Lantus® were administered to the ZDF rats (type 2 diabetes) by a subcutaneous injection. An appropriate amount of insulin or insulin glargine powder was suspended in sterile normal saline to obtain a suspension with a concentration of 40 IU/mL (Table 4-1). Lantus® was used as a control and was administered to the animals as is (100 IU/mL).

Blood glucose concentrations in each animal were determined following a subcutaneous injection of each form of insulin at a dose of 10 IU/kg as follows: On each day of the study, the ZDF rats were transferred to the procedure room. The rats were randomly divided into groups, with 4 to 6 animals per group. Each group, except the control group, received one dose of a form of insulin. The control group did not receive any insulin injection. Before injection, each rat was placed in an induction chamber and
Table 4-1 Composition of gel formulations used for *in vivo* studies

<table>
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<th>Formulation #</th>
<th>Concentration of polymer <em>(% w/w)</em></th>
<th>Concentration of ATEC (% w/w)</th>
<th>Concentration of TEC (% w/w)</th>
<th>Insulin glargine (% w/w)</th>
<th>Zinc Sulfate (% w/w)</th>
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<td>0.10</td>
</tr>
</tbody>
</table>

OA-G-1 was prepared in phosphate buffer saline (PBS)

* Inherent viscosity of the polymer (dL/g) - 0.09
anesthetized by isoflurane (Halocarbon Product Corporation, River Edge, NJ) using a SurgiVet/Anesco ventilator (Waukesha, WI). The body weight of the animal was recorded to determine the volume of the insulin suspensions to be injected. Blood glucose was measured using Bayer Ascensia Breeze blood glucose meter (Bayer Corporation, Elkhart, IN). The predetermined dose of insulin (which was determined based on the animal body weight) was injected subcutaneously at the dorsal neck of the rat using a ½ mL Becton Dickinson Lo-Dose™ U-100 insulin syringe with a permanently attached 28 gauge x ½ inch needle (Franklin Lakes, NJ). After injection, the rats were returned to the cages and allowed to recover from anesthesia. At pre-determined time intervals, the rats were anesthetized by the aforementioned method and their blood glucose was measured. A plot of blood glucose concentrations versus time was generated from the data to describe the glucose lowering effect of insulin or insulin glargine, from which information such as time of peak action and duration of action could be obtained.

4.2.2.6 Fabrication of biodegradable gels. The gel formulations (Table 4-1) were prepared in two steps: dissolution of polymer in the plasticizer(s) to obtain a blank gel and loading of insulin glargine into the blank gel.

The polymer was allowed to come to room temperature in a vacuum desiccator after it was taken out from the freezer. Then an appropriate amount of polymer particles was accurately weighed into a dry, clean glass scintillation vial using a Mettler AE100 analytical balance (Mettler-Toledo, Inc., Columbus, OH). The desired amount of plasticizer(s) was weighed into the scintillation vial containing the polymer. A plastic screw cap with aluminum liner was used to seal the vial. The vial was then vortexed on a vortex mixer to disperse the polymer particles in the plasticizer(s). The vial was placed in a Lab-Line orbital shaker (Model 3527, Lab-line, Melrose Park, IL), which was maintained at 37°C, and shaken at 150 rpm. It took 2 to 3 days for the polymer to completely dissolve in the plasticizer. The resulting blank gel was stored in vacuum desiccators at room temperature until further use (typically 1 to 2 days). One day before the insulin glargine-loaded gel was required, a desired amount of blank gel was transferred to a new glass scintillation vial and an appropriate amount (approximately 80 mg) of insulin glargine particles was accurately weighed and transferred into the blank gel vial. A stainless steel spatula was used to disperse the insulin glargine particles uniformly into the blank gel. Any visible agglomerates of insulin glargine were broken using a spatula and the insulin glargine-loaded gel was ready to be used for the in vivo studies.

4.2.2.7 In vivo studies of insulin glargine-loaded gel formulations in ZDF rats. For the in vivo studies, blood glucose concentrations of the diabetic rats were measured before and after injection of insulin glargine-loaded gels. The concentration of insulin glargine of some of the formulations in plasma or serum was also determined and the data were plotted against time to characterize the release of insulin glargine from the insulin glargine-loaded gel formulations. For gel formulations consisting of insulin glargine and zinc sulfate, an appropriate quantity of zinc sulfate powder (< # 400 mesh or 37.5 µm) was accurately weighed and added to the insulin glargine-loaded gel.
Typically, a total of 2 g of gel (including insulin glargine and zinc sulfate) was prepared for each gel formulation. On the day of injection, the ZDF rats were transferred from the housing room to the procedure room. The animals were anesthetized by isoflurane. The dorsal neck of each rat (except the animals in the control group that did not receive any injection) was shaved to expose the injection site. Body weight of the rat was recorded. Blood glucose was measured prior to any injection using a glucose meter as described before. The insulin glargine-loaded gel was withdrawn into a barrel of a 1 mL syringe, without any needle. Then a 23 gauge needle (1 inch in length) was attached to the syringe, and the volume of the gel in the syringe was adjusted to the desired volume, depending on the dose of insulin for each animal. The shaved area on the dorsal neck of the animal was disinfected with isopropyl swab and a subcutaneous injection of the insulin glargine-loaded gel was made. The needle was inserted under the skin in such a way that the gel would be injected between the skin and the underlying muscle tissue. The actual procedure of injecting the gel was as follows: At the injection site, the skin was pinched and lifted by the finger and thumb. The needle was inserted under the skin between the finger and thumb. Approximately three-quarters of an inch of needle was pierced into the skin to make sure that the injected gel would not ooze out after the needle was removed from the injection site. After injection of the gels, at pre-determined time intervals, the rats were anesthetized and blood glucose concentrations were measured. Blood samples (300 to 400 μL) were taken from the tail vein and processed to collect plasma or serum. These plasma or serum samples were stored at -20°C until analysis.

4.2.2.8 Statistical analysis. Area under the effect curve (AUEC) which is defined as the area below baseline is representative of the effect of the formulation on lowering of blood glucose with greater AUEC implying greater effect. One way ANOVA within Excel 2007 was utilized to compare AUEC between treatment (N = 4) and control (N = 4) at 95% confidence level.

4.3 Results and discussion

4.3.1 Purity of insulin glargine particles

Figure 4-1 shows the chromatogram of placebo Lantus® formulation (without any insulin glargine). The placebo was made in house and each milliliter of the formulation contained 2.7 mg of m-cresol, 30 μg of zinc, (from zinc chloride), 20 mg of glycerol 85% w/w and water for injection. This formulation was similar to Lantus® formulation without insulin glargine. It is evident from the Figure 4-1 that the retention time of the peak of the placebo formulation is approximately 6 minutes.

Figure 4-2 shows the chromatogram of Lantus® formulation diluted in 0.01 N HCl. The chromatogram shows two peaks, one at 6 minutes, which corresponds to the placebo peak and another peak at 11 minutes. The peak at 11 minutes is the peak of insulin glargine or Lantus®. The diluted Lantus® formulations were injected at various concentrations and the areas under the peaks at 11 minutes were used to prepare the standard plot.
Figure 4-1    HPLC chromatogram of placebo of Lantus® formulation
Figure 4-2  HPLC chromatogram of Lantus® formulation
Figure 4-3 shows the chromatogram of purified (dialyzed) and freeze-dried particles of insulin glargine dissolved in 0.01 N HCl. The chromatogram shows a single peak at 11 minutes. It is evident from the Figure 4-3 that the placebo peak at 6 minutes is missing, thus indicating that the excipients in the Lantus® formulation were removed using the dialysis bags and pure insulin glargine particles were obtained after freeze drying. The calculated purity of these insulin glargine particles was found to be 94.6%.

4.3.2 Pharmacodynamic studies of insulin formulations in ZDF rats

Insulin and insulin glargine particles were suspended in sterile normal saline (0.9% NaCl) and administered subcutaneously to ZDF rats to evaluate the biological effect of different forms of insulin on the blood glucose concentrations of the rats. Figure 4-4 shows the blood glucose concentrations of these pharmacodynamic studies. It is evident from Figure 4-4 that the blood glucose concentrations decreased from approximately 452 ± 27.38 mg/dL to 92.5 ± 4.9 mg/dL within 3 hours after a subcutaneous injection of 10 IU/kg dose of RHI in the ZDF rats. Normal blood glucose concentrations (between 80 – 200 mg/dL) were maintained for approximately an hour before reaching the normally high blood glucose concentrations within 6 hours. Lantus® was subcutaneously injected at dose of 10 IU/kg (Figure 4-4). The blood glucose concentrations after Lantus® injection decreased from approximately 441 ± 17.02 mg/dL to 229 ± 45 mg/dL within 4 hours post injection. The blood glucose concentrations after subcutaneous injection of purified glargine particles suspended in saline decreased from approximately 481 ± 16.68 mg/dL to 136.23 ± 14.56 mg/dL within 2 hours (Figure 4-4).

Normal blood glucose concentrations (between 80 and 200 mg/dL) were maintained for approximately 6 hours before reaching the normally high blood glucose concentrations (350 mg/dL) within 12 hours. This pharmacodynamic effect of lowering of blood glucose concentrations shows that the insulin glargine particles (purified from the commercially available Lantus® formulation) and suspended in normal saline were bioactive. There was a significant difference between AUEC after subcutaneous injection of insulin glargine particles and control at 95% confidence level (P=0.003) indicating that insulin glargine particles causes significantly greater reduction in blood glucose when compared to control. Moreover, no significant difference was observed between AUEC after subcutaneous injection of insulin glargine particles and Lantus® (P>0.05) at 95% confidence level indicating that the Lantus® is no better than insulin glargine particles in reduction of blood glucose.

4.3.3 In vivo studies of gel formulations loaded with insulin glargine particles

Purified and freeze-dried particles of insulin glargine were suspended in phosphate buffer saline (PBS) at 4% loading, and 0.1 mL of this suspension was injected subcutaneously in the ZDF rats. Figure 4-5 shows the changes in blood glucose concentrations and body weights of ZDF rats after the injection. It is evident from the figure that blood glucose concentrations decreased rapidly below 100 mg/dL within 6 hours after injection and remained at that concentration until day 3 before rapidly climbing to above 300 mg/dL on day 5.
Figure 4-3  HPLC chromatogram of insulin glargine particles dissolved in HCl
Figure 4-4  Blood glucose profiles in ZDF rats after subcutaneous injections of insulin (RHI), Lantus® and insulin glargine particles suspended in saline (dose-10 U/Kg) (Mean ± SEM, n = 5)
Figure 4-5  Changes in blood glucose concentrations and body weights of ZDF rats after subcutaneous injections of 4% suspension of insulin glargine particles in PBS (Mean ± SEM, n = 5)
The rapid decline in blood glucose concentrations is not desirable. Moreover, the rats were nearly hypoglycemic for three days. Hence, there was a need to further slow down the release of insulin glargine, which produced a slower and extended effect in the animals. However, from Figure 4-5, it is clear that as the blood glucose concentrations decreased, the body weights of the rats increased rapidly. Moreover, the body weights of the animals started to decline as the blood glucose concentrations climbed back to the normally high concentrations above 350 mg/dL. This showed that the glargine particles not only produced a lowering in blood glucose concentrations, but also affected the body weights, thus proving that the purified and freeze-dried particles of insulin glargine were bioactive.

A similar effect in the change in blood glucose concentrations and body weights of ZDF rats was observed after subcutaneous injection of 0.1 mL of 4% suspension of insulin glargine particles in a 3:1 blend of ATEC: TEC plasticizers (Figure 4-6). Hence, it is clear from this study that loading insulin glargine particles in a combination of plasticizers alone does not provide extension of biological effect of insulin glargine. Figure 4-7 depicts change in blood glucose concentrations and body weights of ZDF rats after a subcutaneous injection of PLGA gel formulation (containing 5% PLGA) loaded with insulin glargine particles. The drug loading in the gel formulation (Formulation OA-G-3) as shown in Table 4-1 was 2% w/w. The drug loading in the gel formulation (Formulation OA-G-3) as shown in Table 4-1 was 2% w/w. The blood glucose concentrations reduced from 425 mg/dL to approximately 100 mg/dL within 24 hours after the injection and remained between 100 and 200 mg/dL for 4 days before steadily climbing back up to above 300 mg/dL on the day 9. This showed that addition of 5% PLGA to the plasticizer combination of ATEC:TEC (3:1) not only slowed down the in vivo effect of insulin, but also extended the duration of action of the drug for approximately 6 days before climbing back up to the normally high concentrations of above 350 mg/dL.

Figure 4-8 shows changes in blood glucose concentrations and body weights of ZDF rats after subcutaneous injection of gel formulation prepared with 5% PLGA and loaded with 4% w/w of insulin glargine particles (Formulation OA-G-4) (Table 4-1). It is evident from the figure that the effect of this gel formulation (Formulation OA-G-4) was similar to Formulation OA-G-3 with 2% drug loading. The blood glucose concentrations dropped from 445 mg/dL to about 100 mg/dL in 24 hours, and remained below 150 mg/dL for 9 days, before rising steadily above 300 mg/dL on day 13. Hence, doubling the drug loading from 2% (Formulation OA-G-3) to 4% (Formulation OA-G-4) extended the maintenance of normal blood glucose concentrations from 6 to 9 days, respectively. Moreover, the effect of insulin glargine was also seen on the body weights, which steadily increased as the blood glucose concentrations dropped. The body weights remained high and dropped slightly when the blood glucose concentrations began to rise steadily.

To explore the possibility of slower and longer duration of effect of insulin glargine in the ZDF rats, the polymer concentration in the gel formulation was increased
Figure 4-6  Changes in blood glucose concentrations and body weights of ZDF rats after subcutaneous injections of 4% suspension of insulin glargine particles in blends of ATEC:TEC (3:1) (Mean ± SEM, n = 5)
Figure 4-7 Changes in blood glucose concentrations and body weights of ZDF rats after a subcutaneous injection of gel (5% PLGA) formulation loaded with 2% insulin glargine particles (Mean ± SEM, n = 5)
Figure 4-8  Changes in blood glucose concentrations and body weights of ZDF rats after a subcutaneous injection of gel (5% PLGA) formulation loaded with 4% insulin glargine particles (Mean ± SEM, n = 5)
from 5 to 10% w/w. Figure 4-9 shows changes in blood glucose concentrations and body weights of ZDF rats after subcutaneous injection of PLGA gel (prepared with 10% PLGA) formulation loaded with 4% insulin glargine particles (Formulation OA-G-5) (Table 4-1).

The changes in blood glucose concentrations and body weights of ZDF rats were similar to Formulation OA-G-4 containing 5% PLGA gel (Figure 4-8). Hence, increasing the PLGA from 5% to 10% did not increase the in vivo duration of activity of insulin in the ZDF rats. Figure 4-10 depicts the comparison of blood glucose concentrations in ZDF rats after subcutaneous injections of different formulations of insulin glargine with 4% loading with (Formulation OA-G-4) or without PLGA (Formulation OA-G-1 & Formulation OA-G-2). There was a significant difference between AUEC after subcutaneous injection of formulation of insulin glargine with 4% loading with PLGA (Formulation OA-G-4) and control at 95% confidence level (P = 0.00001) indicating that insulin glargine with 4% loading with PLGA causes significantly greater reduction in blood glucose when compared to control. Furthermore, There was a significant difference between AUEC after subcutaneous injection of formulation of insulin glargine with 4% loading with PLGA (Formulation OA-G-4) and formulation of insulin glargine with 4% loading without PLGA (Formulation OA-G-1) at 95% confidence level (P = 0.0004) indicating that insulin glargine with 4% loading with PLGA causes significantly greater reduction in blood glucose concentrations, when compared to insulin glargine with 4% loading without PLGA.

Moreover, no significant difference was observed between AUEC after subcutaneous injection formulation of insulin glargine with 4% loading without PLGA i.e insulin glargine in PBS (Formulation OA-G-1) and insulin glargine in plasticizers (Formulation OA-G-2) (P>0.05) at 95% confidence level indicating that the insulin glargine in plasticizers (Formulation OA-G-2) is no better than insulin glargine in PBS (Formulation OA-G-1), in reduction of the blood glucose concentrations. It is clear from the figure that addition of 5% w/w PLGA not only slows down the lowering of blood glucose concentrations, but also increased the duration of action from 3 to 9 days. This extension in the blood glucose lowering effect of insulin glargine particles is due to presence of PLGA. This effect may be due to formation of a depot of insulin glargine particles encapsulated in the PLGA matrix. This PLGA matrix released the insulin slowly as it underwent erosion with time.

To study the concentration concentrations of glargine and its effect on blood glucose concentrations, Formulations OA-G-1 and OA-G-4 were injected into two different groups of 7 ZDF rats. The blood glucose concentrations and concentrations of glargine in serum were determined. Figure 4-11 shows the concentrations of glargine in serum and blood glucose concentrations after a subcutaneous injection of Formulation OA-G-1. The concentration of glargine in serum of the ZDF rats increased very rapidly to 3563 ± 1228 mIU/L within 6 hours after injection. The concentration of glargine remained above 1500 mIU/L until day 3 and then slowly declined below 100 mIU/L within 6 days after injection. The blood glucose concentrations dropped to a low concentration of 93 ± 31 mg/dL within 6 hours after injection thus corresponding to the
Figure 4-9  Changes in blood glucose concentrations and body weights of ZDF rats after subcutaneous injections of gel (10%) formulations formulation loaded with 4% insulin glargine particles (Mean ± SEM, n = 5)
Figure 4-10  Comparison of blood glucose concentrations in ZDF rats after subcutaneous injections of insulin glargine (4% loading) in different formulations (Mean ± SEM, n = 5)
Figure 4-11  Blood glucose concentrations and concentrations of insulin glargine in serum of ZDF rats after a single subcutaneous injection of Formulation OA-G-1 (Mean ± SEM, n = 7)
burst release of glargine from the formulation. The blood concentrations remained below 100 mg/dL for up to 4 days before it gradually went back up. The blood glucose concentrations went back up to more than 400 mg/dL on day 6 due to the decline in glargine serum concentrations.

Figure 4-12 shows the concentration of glargine in serum of the ZDF rats and blood glucose concentrations after a subcutaneous injection of gel Formulation OA-G-4. The concentration of glargine in serum of the ZDF rats increased to approximately 750 mIU/L within 12 hours after injection, and then declined to approximately 500 mIU/L within 24 hours after injection. The concentration of glargine was then maintained between 260 ± 134.9 mIU/L and 188 ± 55.9 mIU/L until day 10. Glargine concentrations decreased to 61.5 ± 16.5 mIU/L on day 11 and then gradually decreased further thereafter. The blood glucose concentrations were well correlated to the concentration of glargine in the serum. The blood glucose concentrations dropped to a low concentration (75 ± 14.5 mg/dL) on the first day after injection and were maintained at the low concentration (< 200 mg/dL) for 9 days before it gradually went back up. Blood glucose went back up to more than 400 mg/dL on day 13 and then was constantly maintained at normally high concentrations as seen before the injection.

Figure 4-13 shows a comparison of serum glargine concentrations in ZDF rats after subcutaneous injections of 4% insulin glargine loaded) in PLGA (5%) gel formulation (Formulation OA-G-4) and PBS (Formulation OA-G-1). There was a significant difference between serum glargine concentrations after subcutaneous injection of formulation of insulin glargine with 4% loading with PLGA (Formulation OA-G-4) and formulation of insulin glargine with 4% loading without PLGA (Formulation OA-G-1) at 95% confidence level (P = 0.001) indicating that insulin glargine with 4% loading with PLGA causes significantly extension in serum glargine concentrations when compared to insulin glargine with 4% loading without PLGA. It is evident from the figure that the serum glargine concentrations rose rapidly after subcutaneous injection of formulation of insulin glargine with 4% loading without PLGA (Formulation OA-G-1). The serum glargine concentrations decreased to near zero levels on day six. However, the serum glargine concentrations rose slowly after subcutaneous injection of formulation of insulin glargine with 4% loading with PLGA (Formulation OA-G-4). Furthermore, the serum glargine concentrations were above 2500 mIU/L until day 10.

To further study the effect of presence of zinc sulfate in the PLGA gels loaded with insulin glargine on the biological effect of these gels, three formulations at different concentrations of zinc sulfate were prepared. Figure 4-14 shows the comparison of blood glucose concentrations in ZDF rats after subcutaneous injections of different formulations loaded with 4% insulin glargine and various zinc sulfate concentrations. It is evident from the figure that addition of zinc sulfate in the formulations lowered the rate of blood glucose lowering effect. At the highest zinc sulfate concentration of 0.5% (Formulation OA-G-6), the blood glucose concentrations never dropped below 300 mg/dL. Moreover, the blood glucose concentrations remained between 300 to 400 mg/dL for 14 days.
Figure 4-12  Blood glucose concentrations and concentrations of insulin glargine in serum of ZDF rats after a single subcutaneous injection of Formulation OA-G-4 (Mean ± SEM, n = 6)
Figure 4-13  Comparison of serum glargine concentrations in ZDF rats after subcutaneous injections of 4% insulin glargine loaded in PLGA (5%) gel formulation (Formulation OA-G-4) and PBS
Figure 4-14  Comparison of blood glucose concentrations in ZDF rats after injections of insulin glargine (4% loading) in different formulations with various zinc sulfate concentrations (Mean ± SEM, n = 5)
There was a significant difference between AUEC after subcutaneous injection of formulation of insulin glargine with 4% loading with PLGA and zinc sulfate concentration of 0.5% (Formulation OA-G-6) and control at 95% confidence level (P = 0.02) indicating that insulin glargine with 4% loading with PLGA and zinc sulfate concentration of 0.5% causes significantly greater reduction in blood glucose when compared to control. Furthermore, there was also a significant difference between AUEC after subcutaneous injection of formulation of insulin glargine with 4% loading with PLGA (Formulation OA-G-4) and formulation of insulin glargine with 4% loading with PLGA and zinc sulfate concentration of 0.5% (Formulation OA-G-6) at 95% confidence level (P = 0.002) indicating that insulin glargine with 4% loading with PLGA causes significantly greater reduction in blood glucose when compared to formulation of insulin glargine with 4% loading with PLGA and zinc sulfate concentration of 0.5% (Formulation OA-G-6).

As the zinc sulfate concentration was reduced from 0.5% to 0.25% (Formulation OA-G-7), the blood glucose concentrations dropped from 520 mg/dL to about 178 mg/dL in 2 days, and remained below 200 mg/dL for 8 days, before rising steadily above 400 mg/dL on day 10. As the zinc sulfate concentration was further reduced from 0.25% to 0.1% (Formulation OA-G-8), the blood glucose concentrations dropped from 515 mg/dL to about 128 mg/dL in 24 hours, and remained below 200 mg/dL for 8 days, before rising steadily above 400 mg/dL on day 13.

Moreover, no significant difference was observed between AUEC after subcutaneous injection formulation of insulin glargine with 4% loading with PLGA (Formulation OA-G-4) and formulation of insulin glargine with 4% loading with PLGA and zinc sulfate concentration of 0.25% (Formulation OA-G-7) (P>0.05) and zinc sulfate concentration of 0.1% (Formulation OA-G-8) at 95% confidence level indicating that formulation of insulin glargine with 4% loading with PLGA (Formulation OA-G-4) is no better than formulation of insulin glargine with 4% loading with PLGA and zinc sulfate concentration of 0.25% (Formulation OA-G-7) and zinc sulfate concentration of 0.1% (Formulation OA-G-8), in reduction of blood glucose concentrations. Hence, zinc sulfate could be used as a release modifier for insulin glargine in the gels. However, the amount of zinc sulfate in the gel needs to be optimized, so that it can provide a prolonged effect in vivo.

4.4 Conclusions

Insulin glargine particles were purified and then freeze-dried from commercial Lantus® formulation. The purity of the freeze-dried particles of insulin glargine was determined using HPLC, and was found to be 94.6%. These insulin glargine particles were suspended in saline and subcutaneously injected in ZDF rats. The bioactivity, as measured by blood glucose lowering effect, was found to be similar to regular human insulin and the marketed formulation called Lantus® The PLGA gel formulations prepared with insulin glargine particles had duration of action of 10 days following a single subcutaneous injection. The concentration of insulin glargine was maintained
between 260 ± 134.9 mIU/L and 188 ± 55.9 mIU/L until day 10 after single subcutaneous injection of some of these formulations and the corresponding blood glucose concentrations were suppressed from above 400 mg/dL to below 200 mg/dL from day 1 to day 10 post-injection. The addition of zinc sulfate to the formulations prepared with purified insulin glargine particles further slowed down the drop in blood glucose concentrations.


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