THE ROLE OF INSULIN IN RELATION TO STRESS AND NUTRITIONAL STATE:

GLUCOSE HOMEOSTASIS AFTER SURGERY, IN OBESITY

AND OLD AGE

A thisls submitted in part fulfilment of the regularments for the Degree of

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Ьу

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ABSTRACT

Since the discovery of the hormone (Banting and Best, 1922), extensive research on insulin has been gaing an. Insulin is important not only in relation to carbohydrate metabolism, and thus in relation to diabetes metilitus, but also as one of the important hormones in protein and fet metabolism.

There is much evidence showing an impairment of insulin action in stress, whether the stress is physical or psychological. The way in which insulin resistance develops and its effects an matchalism may vary in different stress situations. This work investigates three forms of physical stress and attempts to show the role of insulin in each of these situations. The metabolic and clinical implications of this problem are discussed.

First, the effects of surgery were studied in cleven potents. Their mean age was 52 years, mean weight was 68 kg, and mean % ideal body weight was 94%. A nitrogen blance study showed that the negative balance after surgery coincided with elevated levels of plasma glucagon, non-estarified fatty acids, branched-chain amino acids, urlnary free cartisol, urlnary 17-OH-corricosteroids and with a decrease of total plasma amino acids. A temperary insulin resistance occurred in post-surgical patients, shown by hyperglycoemia and hyperinsulineamic during a two hour glucosa infusion.

The clinical significance of hyperglycoemia is discussed.

The second study was of hwenty nine obese patients (fasting blood glucose < 6.11 m mol/). Their mean age was 40 years, mean weight was 109 kg, and mean % ideal body weight was 166% (from 115% to 233%). These patients showed on average an imported response to the arel glucose tolerance test (aral GTT), intravenous glucose tolerance test (tv GTT) and intravenous glucose insulin

for insulin sensitivity. This impatment was related to hyperinculinaemia which followed glucese administration (oral or Iv). These abase patients seemed to fall into two groups: those with % ideal body weight < 160% showed impatred tolerance to glucese but relatively normal planes insulin responses; those > 160% showed marked hyperinsulinaemia.

[I le suggested that these responses represent those of 'active' and 'possive' obesity, but that the former may include a pre-clinical stage when insulin sensitivity is very high.

Thirdly, twenty three generic polients (festing blood glucose ζ 6.11 m mol/1) were studied. That mean age was 79 years and mean weight was 56 kg. These patients also showed an impairment in the oral GTT, in GTT and in GTT. The impairment was greater than that found in obese patients. Insulin response to glucose administration (areal or Iv) was sluggish, but the actual levels of insulin were not significantly lower than those found in young normal subjects (except for the peak value during in GTT). The major cause of impaired glucose tolerance was diminished insulin sensitivity, either in the peripheral tiesue, or, more probably in the liver, resulting in relative inability to switch off glucose output. An introvenous alanine tolerance test was carried out in eight elderly subjects (mean age was 78 years, mean weight was 52 kg), to assess gluconeogenic capacity of the liver, and again indicated the relative inability of endogenous insulin to suppress glucose production.

PARTI

INSULIN AND GLUCOSE HOMEOSTASIS

CHAPTER 1

INTRODUCTION

A. History:

Insulin is a hormore which is traditionally associated with diabetes mellitus.
Although this hormone was discovered not more than fifty years ago (Banting and Best, 1922), the discovered not more than fifty years ago (Banting and Best, 1922), the discovered not present the second of the work of the work of the second of Cappadocia, as "a moist and cold wasting of flesh and limbs into the urine" (A.D. 30 to A.D. 90). "Diabetes' is a Greak word meaning "siphon", and is descriptive of the body siphoning away through the urine.
It was a disorded discose and renained so for many years to come. Avicenna (Ibn Sina) a femous Anab physician (930-1033) (Fig. 1), gave a very complete description of the disorder, including some of the complications, such as diabetic garner, furunculosis, phthisis, and also the presence of a honey-like substance in the patient's urine. But many contucles before Aretaeus and Avicenna, old Chinese, Japanese and Hindu writings indicate that the disease had long been known to these peoples also.

The observations of Thomas Willis (1682) of glycosuria in unina however, marked the beginning of a new era when glycosuria was becaming an accepted diagnostic test for diabetes mellitue. Brunner (1683) observed in animal experiments that polyuria and polydipala occurred after removal of the pancreas, but II was Cawley (1788) who probably was the first to associate diabetes with the pancreas when he found multiple calculi and destruction of pancreatic tissue at an autopsy of a patient who had died from diabetes.

Figure '. An old woodout of Avisenna (The Sina) (930 - 1055).

(Courtesy of the Noyal Society of Medicine, London).

(Chartend . 1908)



. . .

Insulin is produced in the Islatt of Langerhans, named after Paul Langerhans (1869) who fins discovered them and described their structure embedded in the tissue of the pancreas. Their function and significance were then however still unknown.

The metabolic relationship between diabetes mellitus and the pancreas was clearly shown by Von Mering and Minkowski (1890). They found that pancreatectomy on dags produced hyperglycaemia and glycosuria and the dags finally died in ketasis and come. Their conclusion was that the pancreas eleborates a substance that keeps the blood sugar low and restores the metabolism to normal.

The whole world owes its thenks to Frederick Bonting and Charlos Best (1922), who discovered "the blood sugar lowering substance" now known as insulin. Their discovery led to the therapeutic use of insulin, thus saving thousands of lives, and has opened the doors to the possibility of its ultimate synthetic production.

It was not long before Abel (1926) achieved the crystallization of the hermans, but over hearty more years passed before Songer and his co-workers successfully pioneered the study of the sequence of amino acids in the insulin molecule (Songer, 1949; Ryle, Songer, Smith & Kitel, 1955).

Since its discovery and especially at present, extensive research on insulinhas been going on in relation not only to diabetes, but also to wider expects of body metabolism. Insulin is, no cloubt, the principal hormone in carbohydrate metabolism, but it is also an equally important hormone in protein (Manchester, 1970) and in fat metabolism (Avruch, Carter and Martin, 1972). This gives a significance much wider than the specialised limits of diabetes. In particular, since insulin has a central rate in the disposal of energy and in synthesis of protein, its relationship hashort and long term nutritional status is receiving increasing attention.

8. Insulin and Its Structure:

Insulin is produced and stored in the poncreatic islets of Langerhons (Mclaod, 1922). There is no evidence that it is produced normally elsewhere in the body (Beat, Japhcott and Scott, 1932). Although in some very rare cases it is produced in non-pancreatic tumous (Shames, Dhurandar and Blackard, 1968). Omenn, 1970).

Insulin is a polypeptide, and consists of two parallel chains of amine acids.

They are the A (acid) and B (basic) chains, and are joined to each other by two disulphide bridges. The third disulphide bridge is connecting two cystein-molecules within the A chain (Steiner, Kemmier, Clark, Oyer and Rubenstein, 1972). Although the destalled amino acids composition at insulin differs semewhat from one species to another, this two-chain structure and the relative positions of the three pairs of disulphide bridges are constant structured features (Fig. 2). Several kinds of experimental evidence indicates that these bridges are assented to the normal structural integrity and biolegical activity of the hormone (Humbel, Bosshard and Zahn, 1972).

Insulin is derived from a larger single polypeptide precessor, 'proinsulin' or "big insulin". In proinsulin, the A and B chain of insulin are joined in series by a further sequence of amino acids (the C-peptide), thus forming a continuous chain, with the same three pairs of disulphide bridges as in Insulin (Fig. 3). (Over, Cho. Figure 2. Primary structure of buman insulin (from Steiner, et.al., 1972).

dy. Ha. Val. Gla. Gla. Gya. Cys. Ser. Ha. Cys. Ser. Lev. Syr. Gla. Lev. Gla. Asn. Tyr. Cys. Asn. 9 10 11 12 13 14 15 16 17 18 19 1

Figure 3. Primary structure of human proinsulin (from Oyer, et.al., 1971).



Some proinsulin is released with insulin into the circulation and eithough cross reacting with insulin entitledies, it does not appear to be a serious source of error in insulin immunoassays. However, proinsulin is also biologically ective, although its effectiveness it only between 2 to 20 per cent of the biological ectivity of insulin (Rubenstein and Steiner, 1971).

CHAPTER II

A. Energy Balance:

Energy balance in man is a complex and highly integrated system of supply and utilization of energy, derived mainly from carbohydrate, fot and protein. The metabalizable energy of these nutrients is linked with requirements through energy couplers such as high energy phosphate compounds, e.g. adenotine triphosphate (ATP) and reduced forms of coenzymes, particularly reduced nicetinomide adenine dinucleatide phosphate (NADP 4 H*). In muscle, creatine phosphate has a specialized short term role in energy storage. This system of intermittent supplies and continuous but veriable demands involved the deposition of nutrient stores during periods of excess fuel inteke, and conversely, their mobilization in periods of distory nutrient deprivation. Narmonal control has an important function in this regulation of fivel supplies. Under normal circumstances, insulin action predominates during exagenous fuel excess (just after ingestion of food), whereas contra-regulatory hormones become aperative during the fasting phase, when energy has to be mobilized from endocenous stores.

Carbahydrate, fat and protein from exogenous sources are hydrolysed and absorbed within the gastra-intestinal system. The major endogenous fuel stores are in the forms of gycogen in the muscle and liver, triglyceride in adjace tissue, and if all fails, protein in the peripheral muscle. The main function of protein however, is to form the structural, contractibend enzymatic component of cells.

II. Glucose as an Energy Source:

Most tissues in the body use non-esterified fatty acids (NEFA) as their main source of energy to generate ATP. But there are several tissues which are dependent on glucose. They are divided into two groups.

- 1. Timums which exidize glucose completely (rich in mitochondria):
 - The brain and narrous tissues: The reason why these tissues do not use NEFA as their energy source is not yet clearly known. In these tissues, glucose is aridized to pyruvate and some ATP is generated, then pyruvate is axidized further to CO2 in the mitochandria with much greater ATP generation.
 - Bed muscle
 Although NEFA comprise the main fuel for red
 muscle, this tissue is able to use glucose also, and oxidises it
 completely to CO₂, thus generating the maximum yield of ATP.
- 2. Tissues dependent on glycolysis (mitochandria absent or deficient):
 - e. Red blood cell: This tissue does not have mitochondria, and glucose can, therefore, be exidized only to pyruvete. This exidation is coupled with the reduction of pyruvete to lockate. Lactate is then transferred to the fliver where it is converted back to glucose. Only small amounts of ATP are generated (2 males per one male of glucose) in oxidizing glucose to pyruvate, but recycling of pyruvate through lactate and the Carl cycle increases the energy yield of glycolysis considerably (see Chapter III C. on Carl cycle).

White muscle during exerction. This tissue does not have enough mitochondria for the complete evidation of glucose.

During exercise, therefore, factore is formed in the red blood cells, and this lactore is then also transferred to the liver to be converted back to glucose (see Chapter III C. on Carl cycle).

C. Fate of Exogenous Glucose added to the Blood Circulation:

That "carbohydrate given by mouth cen be converted into fat by the matebolic processes of the body is new an accepted fact." (Macdonald, 1967). Whether the tissues are glucose as NEFA dependent or whether mobilization of endogenous fuel stores involves glycogen or triglyceride, all may therefore, indirectly or directly be derived from distary carbohydrate. However, the <u>immediate</u> fate of dietary glucose or of the glucose load during and or introvenous glucose tolerance tests is much less certain.

In man there is a little comprehensive evidence on this question. Some suggested that up to 50 per cant of the ingested carbohydrate (arei) is taken up by the liver (Ensinck and Williams, 1972). This figure, however, is very likely to be too high, since after a glucose load, gluconeogenesis is suppressed in the liver (Madison, 1960) and studies on the rate of disappearance of glucose after oral or intravenous glucose load without using radioactive glucose tracer could be misleading. Thus experiments on rate, in which the orally administered glucose included a tracer dose of ¹⁴C-glucose, showed that after 180 minutes only 15 to 18 per cent of the ingested glucose had been taken up by the liver (Curtis-Prior, Trethewey, Stewart & Hanley, 1969; Jelicoote and Moody, 1969). However, even 15 per

cent uptake by the liver means that this organ is disproportionately active (relating to weight). It is instructive to look at some of the other organs in the same way and to compare introvenous and aroll administration of glucose (Table 1). The date are from an experiment of Curtis-Prior at. at. (1969). In this experiment on introvenous dose (750 mg/kg) and an introgestric dose (1500 mg/kg) of glucose were given to rais, in each case together with a tracer dose of (U - ¹⁴C) D-glucose. The results are summarized in Table 1.

The data in Table 1 show the amount of radio-active counts in each organ, therefore, they may not necessarily be that of glucose. It could be in the farm of glucose products (e.g. glycogen, pyruvate, locate). A correction figure for recycling is not included. Nevertheless, they provide us with a fair picture of the fate of exogenous glucose once it enters the circulation.

The body camposition of a normal man may not necessarily be the same as that of a ret. Table 2 shows the normal body composition of a normal male, weighing 70 kg.

Table 1

Distribution of radioactivity in argans of 200 g rats following an introvenous (750 mg/kg) or introgratric (1500 mg/kg) load of glucose, together with a tracer dose of (U = ¹⁴C) D-glucose (derived from Curtis-Prior et al., 1969)

load
180 min.
17.8
5.4
2.5
15.0
3.5
8.4
8.0
31.3

Normal body composition of a 70 kg male

	Tissue	% of body weight
	Skeletal muscle	45.0°
2.	Blood	7.6b
з.	Adlpose Hissue	19.6b
4.	Liver	1.90
5,	Allmentary tract	5.80
á.	Brain	2.00

- a. Darlved from Munra (1969)
- b. " " Olesen (1965)
- # " Johnston and Whillis (1954)

D. Maintenance of Blood Glucose Level:

1. Basal glucose levels

During festing, exogenous glucose is not available, and since there is always a certain demand for glucose (see Chapter II 8.), the body provide it by mobilization from stores (glycogen) and by gluconeogenesis. Cahili, (1970), summarised the substrate and harmone changes during a 24 hour less in a normal man, 70 kg in body weight, with a wity energy control of 7.5 https://day (1800.calorle/day) in Fig. 4.

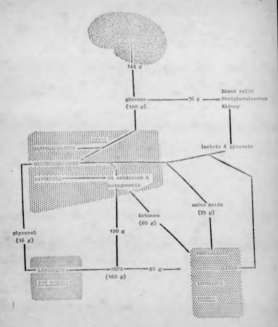
During this period of facting, about 180 g of glucose is produced by the liver and enters the circulation. Most of this is 'new' glucose which is derived from emino acids (75 g) from the muscle. While recycled rather than 'new' glucose is derived from pyruvate and locate (36 g) from muscle, nerve and blood cells, and kidney medulle. Glucose is also formed from glycerol (16 g) derived from fet

NEFA from the hydrolysis of triglyceride are oxidized in the liver, and also in the muscle. Some of the energy generated in this process is used in maintening gluconeogenesis (see Chapter III on gluconeogenesis) 80 per cent of the glucose produced by the liver will be diverted to the brain and the remainder to the cellular elements of blood, peripheral nerves and kidney medulla. Where through the process of glycolysis, it is converted back to pyruvate and lactate, which is then transported to the liver for reconversion to glucose (Fig. 4).

Pigura 4.

Scheme of fuel disposition in normal man, fasting for 24 hours (70 kg, detry energy) expenditure 7.5 MJ/day). Cluoses directed mainly for corebral consumption is released from hepstic glyocom, and new glucose is generated in liver from precursors derived from fat, masele, blood cells, nerve, and recal madulla. NEWS from triglyceride byfarelysis are smidited in muscle and liver as alternate energy sources.

(from Cabill, 1970).



Cabili (1970) further suggested that in prolonged starvation there is a decline in the amount of new glucose generated in the liver with a concomittant decrease in protein catabolism. In this situation, the levels of anti-insulin hormones increase in the circulation and they counteract the insulin action. (See Part II, Chapter II. at this situation, the protein counteract the insulin action. (See Part II, Chapter II. at this situation, the partic glucon-cogenesis.)

2. After Ingestion of food:

After Ingestion of load, the levels of glucose, embre acties, some gut hormones (e.g. gastrin, pancreagymin-chalacystakinine, socretin and entero glucogon) are increased in the blood. These substrates and hormones stimulate increased release of insulin (Porte & Bagdade, 1970). It is known that glucose alone is a potent stimulater of insulin release, but the combination of glucose and amino acids with the hormones (entero-insular axis) is responsible for the greater release of insulin during area glucose tolerance test (and GTT) campared to that found during intrevenous glucose between test (fund GTT), where glucose loading is given directly into the blood circulation. These changes of substrates and hormonel levels during the period following the legaciton of food are summerized in Fig. 5.

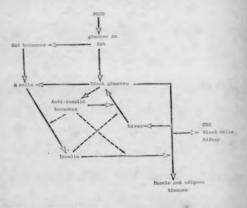
When carbohydrate is led, blood glucose rises, but saidom exceeds 8.88 m mol/1 [160 mg/100 ml] and usually subsides to preprondial levels by two hours. The insulin released during nutrient absorbtion promotes storage of the metabolic fuels in appropriate compartments, (e.g. as glycogen in musch and liver, and as triglyceride in adipose tissue). (Eminck and Williams, 1972). In muscle and fat, insulin enhances

Figure 5.

A subsect of glucose disposition after ingestion of food (earbohydrate) in Has.

Cluster is absorped in the gut and enter the circulation. Out hormones which are increased by the presence of food in the gut, stimulate 8 sells. On the other hand, blood glueces alone could also stimulate 5 sells. It calls produce insulin which prosets glucoss uptake by mascle and adjaces timunes. Blood glueces inhibits the secretions of sett-insulin hormones. These bormones spees insulin actions in liver (inhibition of gluecesopassie) and in muscle and adipose timuses (glueces uptake, inhibition of lipolysis and inhibition of protein breakdown).

But all anti-insulin hormones exhibit every action shows in this diagram.



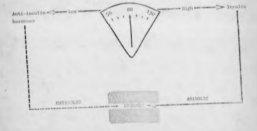
<-- stimulate or promote.

---- inhibit.

glucose autidation. It also enhances glycogen synthesis, potaselum and amino acid influx and protein synthesis in muscle (Reiser, 1967). Conversely, insulin inhibits the release of amino acide from muscle (Cohill, 1970). Insulin not only facilitates glucose entering adipose cells and thereby lipid synthesis, it also inhibits lipolysis. This is reflected in the decline in the circulating NEFA concentration which coincides with hyperglyceamic and elevation of insulin concentrations in blood, (Reiser, 1967). Rote and Bagdada, 1970). After ingestlen of carbohydrate, glucose is taken up by the liver, and most of the carbohydrate estimilated in the liver is deposited as glycogen, under the action of glycogen synthetase, an enzyme regulated by insulin (Reiser, 1967), and concomitantly, hepatic glycogenolysis and glyconeogenesis are obruptly reduced (Medison, 1969).

Fig. 6 summerised and simplifies the control of blood glucose level by insulin and anti-insulin hormones; e.g. growth hormone, apinephrine and glucogen. Figure 6. The control of blood glucose concentration by insulin and anti-insulin barmones.

(mg/100 ml)



aticalate or premate.

CHAPTER III

A. Definition

Glucomagenesis is strictly the "synthesis of new glucose fram non-corbohydrate pracursors", e.g. amino acid residues. But this process overlaps considerably with that whereby glucose is resynthesized from lactate and pyruvate. The term "gluconeogenesis" is therefore used here to include all these processes. The liver is the main organ where gluconeogenesis tokes place, although in prolonged starvation, the kidney becomes an important gluconeogeneic organ as well. In this situation the kidney takes up amino acids to produce "new" glucose as well as to produce NH3 to counteract the kebsis which is developed during starvation, (Owen, Felly, Morgan, Wahren and Cahiti. 1969).

Gluconeogenesis is important during starvetion and other altuations when carbohydrate intake from the alimentary canal is limited and the body glycogen stores are depleted. Pyruvate, lectate, glycerol and glucogenic amino acids are converted to glucose and glycogen. These amino acids, either from alimentary canal obscription or from protain breakdown in muscle, through gluconeogenesis, become an important source of energy.

B. Liver and Gluconeagenesis:

Amino acids enter the liver cell by a membrane transport system. Lectale, alonine, serine and glycine are converted into pyruvate in the cytosol (Fig. ?).

Pyruvate enters the mitochondria, is converted into axaloacetate by pyruvate carbo-

Phinter T.

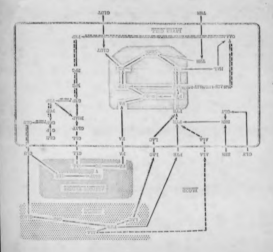
PATRWAY OF GLOODINGGENESIS.

Abbreviations are: LAC, lactate; PTR, pyruvate; ALA, alanine; SER, merines GLY, glycines På, fatty meids AcCoa, anetyl-Coas CIT, citrates at-EO, of-ketoglutarate; EFCC, maccinute; FUM, funarate; MAL, malate; GAA, ormicacetate; ASP, separtate; GLUF, glutamate; TRR, threonine; PEP, P-enol pyravate; 2PG, 2-P-glycerate; 3FG, 3-P-glycerate; GSP, glyceraldshyde-5-F; DEAP, dihydroxyacetone-P; GLYP, glycerol-1-P; GLL, glycerol; FDP, fructose-1, 6-41-P; F6P, fructose-6-P; G6P, glucose-6-P; CLD, glucose; C1P, glucose-1-P; GLH, glycogen.

(modified from Exton, 1972).

.enimele but forceging e.g. from glycerol and slanine.

. olog des cycle.



xylase or to acetyl coenzyme A by pyruvate dehydrogenose. Oxaloacetate is converted into malate and aspartate which leave the mitochondria or to citrate which is mainly metabolized in the Krebs cycle. Malate and aspartate are converted back to exalinacetate, and exploacetate is then converted by phospha-analinyruvate carboxykinase into phospho-malpyruvate (PEP) (Extan, 1972). Two males of high energy phosphate (ATP or GTP) are needed to convert one molecule of PEP. PEP is converted Into fructose - 1,5 diphosphate by a reversal of glycolysis. A further male of ATP and a male of (NADH + H*) are needed for each male of PEP utilized. Fructore = 1,6 diphosphote is hydrolysed to fructose - 6 phosphote by fructose - 1.6 diphosphotese. This enzyme is specific to tissues corrying out aluconeogenesis. Fructose - 6 phosphate Is converted to glucose - 6 phosphate and glucose - 6 phosphate is converted to glucose by another enzyms specific to gluconeogenesis, glucose - 6 phosphotose. The averall conversion of two moles of lactate to one mole of glucose, therefore, requires six moles of ATP (or equivalent as GTP) and 2 males of (NADH + H+). The former is provided by the axidation of NEFA and the latter by the reduction of NAD+ to (NADH + H*) In the convention of lactate to pyruvate. It has been suggested that pyruvate entry Into fiver mitochandria is a control point for glucaneogenesis which is influenced by epinephrine, certisol and allocation (Adam and Haynes, 1969).

Clycerol has a small but significant contribution in gluconeogenesis. It enters gluconeogenic pathway at the level of triose-phosphate by reacting with glycerokinase to form glycerol - 1 phosphate which is then exidised to dihydroxyacetone-phosphate by #C-glycero-phosphate dehydrogenes.

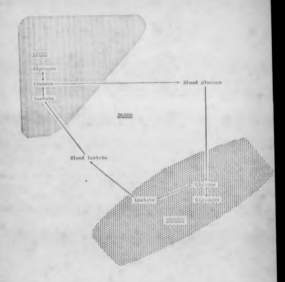
C. Corl Cycle:

In muscle the complete axidation of glucose involves production of pyruvate by a reaction in the cytosol (enacrobic) and the combustion of pyrovate by reactions In mitochondria (gerobic). Red muscle, (e.g. skeletal muscle), which is designed to work over long periods, depends upon the complete axidation of glucose and fatty acids. Such muscle contains more mitochondria to sustain a constantly high rate of axidation of acetyl conzyme A and has a higher content of myoglobin to deliver the required axygen for exidation in the mitochondria. Whereas in white muscle (e.g. breast muscle of a chicken) which is designed for short bursts of heavy activity, utilizes ATP during such exercise at a much greater rate than could be sustained by mitachondrial axidation. It therefore depends more upon a readily available high-energy phosphate store (viz. creatinine phosphate) and upon the rapid generation of ATP from the conversion of glucose to lactate in the cytosal. This kind of muscle has fewer mitochandria and less myaglabin. In reality, though, there is no clear cut division between sed and white muscles. In exercising white muscle, glucose is canverted into 2 molecules of lactate with the generation of 2 ATP in the cytosol. Lactate diffuses into the blood circulation and is taken up by the liver where it is converted back into alucase. (Fig. 8). This cyclic process of alycolysis (e.g. in muscle) and glucaneogenesis (in fiver), using glucase and lactate as transport material is known as the Carl cycle (Fig. 7 and Fig. B).

It is important to note that operation of Corl cycle does not result in a net

Increase in glucose formation for the body since lactate itself is derived from glucose.

Pigure 8. CORI CYLE.



However, there is a transfer of energy, since NEFA from adipose tissue are axiditized in the liver to provide ATF in glucon-eogenesis in the liver. Glucose recycling from loctate formed by glycelysis in blood cells, broin and other tissues may, in normal man, correspond to 10 to 33 per cent of total glucose tumover (Cahill, Harrera, Margan, Soeldner, Steinke, Levy, Reichard and Kipnia, 1966).

CHAPTER IV STRESS AND GLUCOSE METABOLISM

A Definitions

The term 'streen' has a wide range of meanings depending upon the context or the situation in which this word is being used. The Concise Oxford Dictionary (Fifth Edition, 1963), gives a definition of 'strees' as: a) a constraining or impalling force, b) an affort or demand upon energy, c) an emphasis on accentuation, d) (in mechanics) a force exerted between continuous bodies or parts of a body. However, in medicine in general, 'strees' is considered as "any stimulus of such magnitude as to tend to disrupt the homeostasis of the argentum" (Doell, 1966).

B. Classification of Stress

Stress may be classified into two types; first is physical and the other is psychological stress.

Table 3 summarized some of the different kind of stress folling into these two broad groupings.

It is very rare that one type of stress operates in Isolation. It is more usual for the primary stress to lead to a secondary stress of some other type, e.g.,

- An operation (mechanical) is followed by a decrease of food intoke (nutritional).
- An Infection (physical) is followed by a decrease in food intake (nutritional).
- An old person is feeling and and lonely (psychological), eats less food (nutritional, physical) and is likely to succumb more easily to disease and infections (physical).

Teble 3

Classification of Stress

- 1 Psychological
- 1. Environmental (e.g. unfriendly neighbours)
- 2. Endogenous (e.g. feeling sad)

- II Physical
 - 1. Nutritional (e.g. festing, starvation)
 - Mechanical (e.g. fracture, eperation)
 - Climatic (e.g. temperature, humidity)
 - 4. Infection and disease
 - 5. Physiological (e.g. pregnancy)
 - d. Exercise

It is obviously impossible to study all these interactions in a finited period of time. This thesis, therefore, attempts only to study three different kinds of stress, and concentrates only on measurable physical factors.

As indicated in the examples above, nutritional stress is often a secondary consequence at other forms of stress. This secondary nutritional stress usually has a component protein energy mainutrition (PEM), whatever other deficiencies may occur. The following chapter (Chapter V), therefore, does with PEM and its effect on glucose homeestasis and insulin production and effectiveness, as a background to observations after surgery (Part III), in abasity (Part III) and in ald age (Part IV).

CHAPTER V PROTEIN ENERGY MALNUTRITION (PEM)

A. Antiology:

PEM is a major problem in the developing world today. Distary deficiency of peatein and energy cen occur acutely from a sudden failure of food supply e.g. after a natural disaster such as flood or drought, or as a result of anorexia in III-ness or other acute stress, at even as a result of valuntary starvation in support of same political protest. Acute and prolonged starvation have both bean studied by Cabiti and his co-workers (Cabiti et. et., 1966; Owen et. et. 1969; Cabiti, 1970). In reality, however, in meet case nutritional deficiency is obviously not one of total starvation but various degrees of chronic ar seasonal shartage of food. Same of the earlier, simple concepts of protein and calorie (energy) deficiency in the earliagy of kwashlorkor (Williams, 1933; Platt, 1958) and meresmus (Platt, 1958; Jettiffe, 1966), respectively, have become difficult to sustain (Gerrow, 1966; Gapatan, 1968). Therefore, many workers prefer to use the general term of protein calorie membrutrition or pretain energy melanutrition (PEM) (Waterlow and Alleyne, 1971).

However, protein, energy supplies and matabolism cannot be separated into isolated compartments. Thus PEM is related not only to decongements of protein matabolism (Waterlow and Alleyna, 1971), but also to a decrease in the ability of the body to regulate blood glucose levels owing to associated andocrine changes (Heart, Plai) and Stewars, 1956; Stewars and Heard, 1959; Heard and Stewars, 1971).

B. PEM and Blood Glucase Levels:

Malnutrition is usually associated with hypoglycaemia. Balg and Edozien (1965) reported hypoglycaemia in kwashlorkow and Hadden (1967) also found hypoglycaemia in both kwashiarkar and marasmus. The actual levels of blood glucose reported very from one investigator to another, probably awing to different degrees of severity of the disease. The levels of fasting blood glucose In marasmic children in Hungary (when same of the children later died was between 0 to 1.4 m mol/j (0 to 25 mg/100 ml), (Kerpel-Frontus and Kalser, 1967), while reports from Ugande on children suffering from kwashloskor showed fasting blood glucose levels less than 2,2 m mal/| (40 mg/100 ml) (Whitehead and Harland, 1966). However, James and Coore (1970) found that In series of 26 main ourished children the mean fasting blood glucose was 3.1 m mat/j (55 mg/100 ml) initially and became 3.9 m mol/s (70 mg/100 ml) after recovery. Although the degree of hypoglycaemia was significant it was perhaps not as great as might have been expected from earlier reports from Africa and Hungary. Later, workers in Jamaica in a study designed to investigate the hypoglycaemia of PEM were somewhat follod by the absence of any primary nutritional hypoglycaemia. Hypoglycaemia was found anly when PEM had been superImposed on congenital defect and It had a tendency to persist after recovery, (Kerr, Stevens, Robinson and Picau, 1973). Most workers agreed that when a low fasting blood glucose concentration does occur, it improves with recovery from malnutrition unless sufficient energy is not provided with the rehabilitation diet (Balg and Edozien, 1965; Hadden, 1967).

There is an Impairment in glucose talerance in parients suffering from kwashlarker (Sloane, Teltz and Gilichrist, 1961; Baig and Edazien, 1965) and mercennus (Ozman, Maccioni, Zunige, Spade and Manckeberg, 1968), although some reported that in mercennus GTT could still be within normal limits (Hadden, 1967; Bowle, 1964). Thus the blood glucose homostatic mechanism seems to be disturbed in both directions, I.e. in diminished ability to deal with hypo- and hyperglycaemia.

C. PEM and Insulin Lavels:

It has been shown in maintained children that the festing plasma insulin level is usually low (James and Coore, 1970; Milner, 1971), and that the rise of plasma insulin concentrations in response to introvenous glucose load is usually small or obsent. Milner (1971) further could not show any increase in plasma insulin concentration after intravenous injection of glucogon. During recovery, insulin response to glucose is significantly improved, although it is still lower than in normal children (James and Coore, 1970).

However, the impairment in involin ensistivity rather than the actual deficiency of involin is probably the main contributor of poor glucose tolerance (Turner, 1966; Heard and Turner, 1967; Heard and Henry, 1969). Insulin sensitivity was measured in dags either by injecting involin clone (0.1 unit/kg 8w) or by injecting insulin (0.1 unit/kg 8w) tage ther with glucose (0.4 g/kg 8w). Insulin sensitivity is defined here as the effectiveness of insulin in lowering blood glucose concentration, and is expressed as a percentage rate constant for the fall in glucose concentestion. Dogs fed with few protein diet showed abnormalities in carbohydrate matebolism, but glucase telerance was correlated significantly with insulin sensitivity and not with circulating insulin levels (Heard and Henry, 1969).

CHAPTER VI

There is no doubt, therefore, that dietary stress produces an impairment in glucose homeostasis, but the mechanism and the role of insulin and other related homeones may differ from one type of stress to another. It is particularly important to understand the extent to which changes in glucose homeostasis, insulin sensitivity, etc. are adaptive and protective to the "stressed arganism. Almost inevitably phrases like "impairment" in glucose tolerance or in insulin sensitivity will be used in this thesis, as in many other reports, to indicate a quantitative change, but it may not necessarily mean "impairment" in the sense of being 'hormiful.

More information is needed on the complicated and sametimes obscure relationship between stress and insulin sensitivity. A study, therefore, was planned to investigate the effects of three types of physical stress on the role of insulin and other related harmones and substrates. This work concentrates on glucose homeosees in postents undergoing surgery, in obesity and in old age.

PARTII

GLUCOSE HOMEOSTASIS IN PATIENTS AFTER SURGERY

FART II

GLUCOSE HOMEOSTASIS IN PATIENTS AFTER SURGERY

CHAPTER I

A. Trauma and Catabolism:

Surgery is a type of physical trauma which consists of a mechanical stress (the actual operation) and is then often followed by dietary restriction (nutritional stress), (see Part I, Chapter IV on stress). Execuse it is usually elective, it is also preceded by other stresses derived from the condition which needs the operation and creates anxiety. As will be seen later, nutritional status before an operation is, therefore, not necessarily normal. Severe trauma, as in surgery, is a negative beforce of body protein associated with negative nitrogen belonce, and this is reflected by the increase in urinary nitrogen excretion (Cuthbertson, 1964). However, a negative nitrogen balance could also be found in immobilized, otherwise healthy subjects (Schenhelder, Heilskov and Olesen, 1954).

It is perhaps natural to assume that this period of negative nitrogen balance is due to increased careballism, i.e. Increased trackdown of protein. However, the loss of body nitrogen is associated with a fall in protein synthesis and no real evidence of an acute rise in the breakdown rates of body protein has been found in leambilized normal persons (Schenhayder, et. et., 1954), or in patients undergoing experations (O'Keefe, Sender and Jenes, 1974; Crans, Picou, Smith and Waterlow, 1976). This change in synthesis may result from an altered flow of substrates (i.e. amino acids and high energy phosphates and/or specific change in the rate of initiation and elongation. Each of these changes is probably mediated by endecrine balance. The regulation of protein synthesis has been extensively reviewed by Munro (1970, 1976).

This part of the thesis concentrates on endocrine changes following surgery and in particular on the relationship in glucose homeostasis between insulin and its counterregulatory hormones, contisol and glucoson (De Bado and Altzuler, 1958; Ensinck and Williams, 1972). But these same hormones that affect carbohydrate metabolism so that findings in one area usually have relevance to the other.

B. Anti-Insulin Hormones:

Anti-insulin harmones are defined as harmones which as physiological concentrations show anti-insulin effects. The anti-insulin effects could be in carbehydrate, fet as protein metabolism. However, this report limits itself only to harmones which have anti-insulin effects on carbohydrase metabolism. These harmones are:
Advenacosticotrophic harmone (ACTHI, glucocosticoids (e.g. costisol), glucagon,
growth harmone and cetecholomines (e.g. epinephtine).

Stress is usally associated with increased levels in blood of circulating
ACTH (Cooper & Nelson, 1962), cortisol (Yates and Urquhert, 1962; Ross, Welborn, Jehnston and Wright, 1966; Cuthbertson and Tilstone, 1969; glucagon
(Bloom, 1973; Linebey, Santeusania, Breaten, Falcona and Unger, 1974; Wilmere,
Mayland, Pruitt, Lineby, Falcona and Unger, 1974; Russell, Walker and Bloom,
1975), growth harmone (Greenwood and Landon, 1966), and catecholamines
(Walker, Zilell, Rautter, Schoemaker, Friend and Moore, 1959). ACTH stimulates the secretion of adrenocartical harmones (e.g. cortisol), while machanism
of the actions of the after harmones in muscle, liver and adipose tissue is summarized
in Table 4.

Table 4

The mechanism of action of some anti-insulin harmones in muscle, liver and adipose these. (Darlved from Ensinck and Williams, 1972).

	Hormones	Muscle	Liver	Adipose tissus
1.	Epinaphrina	Glycogenolysis Inhibits glucoes utilization	Glycogmolysis Gluconeogenesis	Glucoss upteks Lipolysis
2.	Glucagon	1. Protein breakdown	Glycogenolysm Gluconeogenesis	1. Lipolysis
3.	Growth hormone	Protein synthesis Inhibits glucoss utilisation		Lipolysis Inhibits glucom utilization
4,	Cortisol	1. Protein breakdown	Providing precursors for gluconeogenesis	1. Lipolysis
		Glycogenolysis Inhibits glucose utili setion		

This report fimits itself to the measurement of contisol and its metabolites and glucagon. Epinephrine, ACTH and growth hormone were not measured.

C. Trauma and Endagrine Balance:

Increased circulating levels and excretion of cortical are accepted as the usual consequence of many forms of stress (including surgery). Raw, et. el., 1966; Curthertson and Tilstone, 1969; and recently it has become increasingly avident that the same is true of glucagon (Bloom, 1973), Linday, et. el., 1974; Wilmore, et. el., 1974; Russell, et. el., 1975. The situation with regard to insulin, seems at first glance, rether doubtful. Some have claimed that severe trauma results in depression of plasma insulin levels in relation to blood glucose values (Wilmore, et. el., 1974; Linday, et. el., 1974), while others reported eleveted values and other signs of insulin resistence (Rass, et. el., 1966; Cuttherston and Illistone, 1969). However, the evidence suggests that during the accute phase, pleame insulin levels in relation to blood glucose values are indeed law (Allison, et. el., 1968; Wilmore, et. el., 1974; Linday, et. el., 1974), and they became alevated in the later phase of trauma (Ross, et. el., 1966; Allison et. el., 1966.

Resolution of these problems has considerable practical importance in providing the rationals for affective distary and possibly hormonal therapy after surgery and other forms of trauma. The present investigation seeks, therefore, to delineate the time and extent of hormonal and metabolic changes after surgery, to estempt to correlate these with nitrogen balance and in particular, with the patient's insulinganic aspacity in a two hour glucous infusion test, carried out one day after the operation (day 1) and on 'recovery' (Chapter III, D. on glucous infusion.)

CHAPTER II MATERIAL AND METHODS

A. Subjects

1. Patlants:

Eleven patients undergoing abdominal operations were studied. They were admitted to the surgical word, University College Mospital, London. Written consent was obtained from each patient for his participation in this study. None of the patients was diabetic. Their mean age was 52 4 3.4 years, mean height was 173 ± 2.3 cm and mean body weight was 68 ± 3.9 kg. The body weights ranged between 93 per cent and 118 per cent of the ideal weight for a given height and age (the mean was 94 ± 4.8 per cent). The ideal weight for a given height and age used for comparison was from the data of everage weights of adults in Gelgy Scientific Tables, 1970; besed on the data of "Insured Persons in the United States" (Seclety of Actuaries, 1979), (Table 5).

A complete filtrogen belance was done on three of the eleven patients,

while serial 24 hour urine collections were carried out an each patient. Fasting

blood samples were taken from each patient during pre-operative, post-operative
and 'recovery' periods. In addition to this, one day after the operation (day 1) and
on 'recovery' (days 9 - 21), a two hour glucose infusion test was carried out an each
patient. 'Recovery' is defined here as the time when the surgeons considered that
the patients were fill enough to be sent home, and this varied from patient to patients.

2. Cantrols

Four healthy young subjects were used as controls. Their mean age was 27.4.2.3 years, mean height was 170.2.6, 3.6, and mean body weight was 62.2

Sex, age, body weight and type of operation of patients who participated in this study

	Patient	Sex (M/F)	Age (years)	Weight [lig]	Height (cm)	% of ideal	Type of operation
1,	HG	М	34	85	172	118	Proximal gentric vagotomy
2.	HT	F	41	90	168	74	Repair previous gastrectomy Rouse-en-Y conversion
3.	NT	М	60	75	170	95	Vagatomy, pylerap esty, fundaplication and dilatation of ossophagus
4,	MC	М	57	89	179	111	Prostate hypertrophy, prostatectomy
5.	MD	M	32	56	175	75	Spleenomegally, spleenectomy
6.	NB	F	50	74	158	118	Repair of previous gretrectory
7.	BS	м	ól	73	179	92	Repair of previous colostomy
8.	EC	M	65	61	178	77	Carcinama colon, calectomy and colontomy
9.	CL	F	59	49	160	93	Proximal gastric vagatomy. Finney type pyloropiasty
10,	WD	М	53	70	179	87	Proximal gestric vagatomy
11,	WT	м	55	71	174	93	Proximal gentric vagotomy
	n (* S.E.		S2 ± 3.4	68 = 3.9	173 2 2,3	94 ± 4,8	

^{*}Compared with the data from Goigy Scientific Tables (1970)

4.2 kg. The body weights ranged between 86 per cent and 100 per cent of the ideal body weight for the given height and age (the mean was 92 ± 3.2 per cent), (Table 6). There was no significant difference between either the mean obsolute body weights (p>0.05) or the mean percentage ideal body weights of the controls and the patients (92 per cent and 94 per cent). These controls received the similar two hour glucose infusion test as given to the patients.

B. Nitrogen Balance Study

Three patients participated in this study.

1. Sample preparation:

a. Food samples:

The patients were asked to eat or dishk only food or liquid which was given by the hospital. The amount of food offered was recorded and protein content was calculated from Food Tables (McCance and Wilddowson, 1940). Nitrogen content of the food was calculated from the protein content, using the equation: 1g nitrogen equal to 6.25 g protein. Any food which was left over was collected for each 24 hour partial. Special plastic bags (weights known) were used for the food collection. The left over food was then weighted, mixed with water and was homogenized in a "Kanwood" mixer. Then 11 was made to 500 ml with more water, and re-hamogenized. Alliquots were put into universal containers and stored at -20° C until analysed for nitrogen.

Table 6
Sax, age, height and body weight of control subjects

	Subjects	Sex (AVF)	(years)	Height (cm)	Weight (kg)	% of ideal weight"
1.	PW	F	22	1.59	54	100
2.	WS	M	32	1 59	55	86
3.	PG	м	30	178	68	57
4.	GG	м	24	183	70	93
	(25.E.M.)		27 # 2.3	170 ± 6.31	62 ± 4.2	92 ± 3.2

^{*}Compared with the data from Geigy Scientific Tables (1970)

b. Feecal samples

Daily faecal excretions except for a few days following the operation when there were none, were also collected. The same plastic bags (weights known) were used for these collections. These bags fitted to special tin consulters which enabled the potients to perform the collections easily. The faeces were also weighed, mixed with water and were homogenized in a "Kenwood" mixer. They were also made to 500 ml with more water and re-homogenized. All quots were also put into universal containers and stored at -20° C until analyzed for nitrogen.

e. Urine samples:

Urine callections were done to all of the potients. Urine was preserved with 6N HCI (20 ml /24 hour callection). After measuring the volume of each 24 hour callection, samples of urine were put into universal containers and stored at -20° C until analyzed for nitrogen. Separate aliquots for steroid estimation, (see Section E, on analytical mathed), had a few drops of chloroform added.

d. Calculations:

Allquint of food and facces and of unine were analyzed for nitrogen content. (See Section E. on analytical methods) the daily nitrogen balance was given as: $ND = (Ng + N_{\frac{1}{2}} + N_{\frac{1}{2}})g \text{ nitrogen}$ where ND = food nitrogen(g) of faced in 24 hours

Ng = total nitrogen content (g) of food residue and waste in 24 hours

Np = total faecal nitrogen (g) in 24 hours

Nat = total urinary nitrogen.

ND was calculated from food tables, while $N_{\rm g}$, $N_{\rm g}$ and $N_{\rm U}$ were obtained by direct analysis. Attempts were made to have duplicate meals, one was given to the patients and the other was analyzed for nitrogen content instead of getting the values of nitrogen from food tables. But these attempts proved to be too much time and energy consuming, and the results were more or less similar to the results done through food tables and did not give any more accuracy as it was first expected.

C. Blood Samples:

1. Callection of blood and separation and storage of plasma

Fasting blood samples were taken almost every morning on the first three patients, but in the others they were taken two days before the operation, the first three days after the operation, day 8 and on 'recovery'. Fasting blood was taken at 8,30 a.m. after the patients had been fasted from midnight. Patients who were still having intrevenous fluids, had normal soline substituted for glucose at midnight.

For glucegon estimation, 4.5 ml of blood was added to a heparin tube conteining 0.5 ml cold salution of Trasylol (10,000 KILI/ml Bayer) and after rapid centrifugation the supernaturi was frazen immediately. The rest of the blood was put into
another heparin tube and an altquot of 0.05 ml was taken for blood glucess estimation.

The tube was then centrifuged and the plasma separated. 0.05 ml of plasma was used
for plasma glucese estimation and the rest was put into a polythene specimen tube and
stored at -20° C until the day of estimation. Stored plasma samples were used later
for estimations of plasma insulin, plasma NEFA, plasma amino acids and plasma corthoi.

2. Whole blood or plasma for glucase estimations

It has been reported that plasma glucose concentration is usually about 4 per cent higher than blood glucose (ingram, ingram, Turtle, Sturrock and Applegatith, 1971). Plasma is the 'carrier system' which carries glucose either from gestraintestinel treat or from the liver to verticus tissues, including the blood cells. However, most results are usually reported in terms of blood glucose (e.g. W.H.O. definition of diabetic, etc.). In the present work, both blood and plasma glucose were measured (see Section E, an analytical methods.) Although our values for plasma glucose were mostly higher than values for blood glucose, the difference was not always 4 per cent as reported by Ingram et. al. (1971). But we also found that with fasting values are in situations when the subjects were given insulin (see Pert III and Part IV of this thesis and the glucose concentration fells below fasting values, the blood glucose concentration is usually higher than that af plasma. These differences in the values of plasma and blood glucose did not give any significant difference to the results of the various tests. Another benefit of measuring

plasma glucose is that it could be used as a check if there was a technical error in the blood glucose estimation (e.g. using a wrong pipette, stc.), since when the blood is centrifuged and plasma separated, we were not obta to repeat the blood glucose estimation. In this thatis, to avoid pointless duplication, plasma glucose values are not included in the results.

D. Glucose Infusion:

This test was used rather than the luGTT used in the obese (Part III) and gariatric patients (Part IV) because it was thought to be the least burden to the patients, especially on the day immediately after the operation (day I). And because continuous intravenous glucase infusion is the usual "distary regime" for most patients during the post-apprelive pariod, therefore, the substrates and hormonal changes would be measured in the most usual circumstances.

All glucose infusion tests were carried out in the morning. Fasting blood was taken at 8.30 a.m. as usual. A Holter Roller pump (Extra Corporeal Medical Specialistics Ltd.) was used for infusing glucose through a butterfly needle inserted into an ante-cubital valu of one arm. The rate of glucose infusion was 0.33 g kg⁻¹ h⁻¹ (Reaven and Farquhar, 1969), and the test lasted for two hours. Another butterfly needle in the colateral ante-cubital valu was used to withdrawing blood samples. Normal saline was used to fill the laster butterfly needle to prevent the blood from clotting. Sometimes the blood did clot despite the regular flushing of the butterfly needle with saline, and we had to find another value to insert another.

needle. This was later solved by mixing haparin (Haparin Injection 8P, 1000 units/ml) with the normal saline (2000 units haparin/500 ml saline). In praliminary work this concentration of haparin appeared to be the optimum for producing constant flow of blood without clatting, while still not affecting the NEFA concentration. Stronger solution of haparin (2000 units/20 ml saline) gave very high and creatic values for plasme NEFA concentration. (The values could be as high as 3000 μ mol/1 to 4000 μ mol/1).

After the Initial fasting blood sample had been taken, further blood samples were taken at 15°, 30°, 60°, 90° and 120° after the start of the Infusion. The blood samples were treated as described in the previous section (see Section C.1. In this chapter).

E. Analytical Methods:

1. Nitrogen (N):

a. Food and faeces:

Aliquan of 0.5 g of food and forcal homogenate were digested with 3 ml concentrated sulphuric acid and selenium catalyst in a Kjeldahl flosk. Glass beads were used to prevent humping. After the samples had cleared, digestion was continued for a further one how. The samples were then allowed to cool and ware diluted with distilled water to 100 ml. They were then analyzed for nitrogen by the

sodium phenate method for NHg In the Technicon autoanelyser, using (NHg12 \$Oq as standards. The nitrogen content of the standards ranged between 5 \$Ig/mt to 150 \$Ig/mt. Semples which fall autiliae the range of the standards were redigested if the final nitrogen concentration was too low, and made to more appropriate volume or the final solutions were diluted, if the concentration was too high.

Samples were run in the euroanelyser at a rate of 36 seconds sampling and 48 seconds washing (Technicon).

b. Urine:

Allquots of 0.1 ml urine were digested with 0.5 ml concentrated sulphuric acid and selentum catalyst in a Kjeldahl flask. Gless beads were also used to prevent humping. After the samples had cleared, digestion was continued for a further half hour. They were then allowed to cool and were diluted with distilled water to 10 ml. The rest of the procedure was similar to that of nitrogen analysis for food and faecal homogenote. From time to time, analysis for nitrogen content of random samples of food and faecal homogenate and urine were done manually using the Markham method (Wootton, 1969) a compare results with those done on outcomplyer.

2. Creatinine:

Creatinine was estimated in the urine. Urine samples were diluted with distilled water (1/20 dilution) and were put into the antonolyser (Technicon). Creatinine solutions ranged between 1.0 mg/100 ml to 15.0 mg/100 ml were used as standards. Any samples which fell outside the range of the standards had the urine radiiluted and the assay

repeated. Samples were run through the automalyser et a rate of 36 seconds sampling and 48 seconds weeking (Technican). Some urine samples chasen at random were enalysed manually as a comparison to results obtained from automalyser.

3. Glucose:

Glucose in blood and plasms was measured enzymatically using the glucose axidase Perid method and kit of Boehringer Carporation (London) Ltd. Aliquatr of 0.05 ml of blood or plasma was mixed with 1.0 ml uranylacetate satution (160 mg/ 100 ml of normal satine) to pracipitate the protein. After centrifugation, 200 µ1 of supernations were mixed with 5 ml glucose oxidase solution. The blank (200 µ1 of water) and standard (200 µ1 of diluted Boehringer standard) were treated similarly. After standing 30 minutes at room temperature, the samples and the standard were read against the blank at 420 nm in a Unican Specirophotometer (Unican SP600), using 10 mm glass curette. For the standard, the solution supplied with the Boehringer kit (100 mg glucose/100 ml) was diffused 1+1 with water.

4. NEFA:

The Boshringer kit for NEFA estimation was used (Boshringer Corporation, Lendon Ltd.). Aliquots of 200 µt of plasma and standard (500 µt out/1) were mixed with 5 ml of chloroform and 1 ml of solution comprising a mixture of 0.27 M capper nitrate and 0.45 M triethanolamine buffer. For the blank, a tube containing all of the reagents but no plasma or standard was treated similarly. After 10 minutes shaking and 5 minutes centrifugation at 3000 RPM, the supernatural tagether with interfaction protein layer were removed by aspiration. A 2 ml aliquot of the remaining

chloreform extract was then added to 0.2 ml of 9 mM diethyl-dithlacarbamets. The samples and the standard were read against the blank in a Unicam spectrophotometer (Unicam SP600) at 436 nm, using 10 mm glass curette.

In the later part of this work, we observed that the reading of the standard in the spectrophotometer was getting lower and lower, resulting in apparent higher values of NEFA concentration in the plasma samples we were analysing. It was found out later that the standard supplied with the kit, contained less than the amount indicated in the label (500 µmol/1). We had to repeat some of our last few NEFA assays using polimitic acid (488 µmol/1) as the stendard. A complaint was sent to Boshringer Corporation (Landon) Ltd. Their head office (Mannhaim, West Germany) confirmed our findings. They apologised, promised to withdraw the faulty standard, and also promised us some compensation. They did send the compensation, unfortunately there were only few new NEFA kits. From then on, we always used polimitic acid (488 µmol/1) as the standard.

5. Insulin:

Plasmo insulin was estimated by the radioimmuno assay method of Holes and Randle (1963), using the kit supplied by the Radiochemical Centre, Amerikam, England. The method is based on a principle that the insulin in plasma and in standard solutions compete with the added radioactive insulin (125 |-insulin) for reaction with an antibody specific to insulin. The insoluble insulin-entibody complex which is formed, is filtered aut and measured for radioactivity. The level of radioactivity in the filter paper is related in an inverse manner to the amount of insulin

present in plasma or stondard solutions.

The emount of unlabelled insulin present in plasma is calculated as follows:

Where I - the concentration of unlabelled insulin

Co = the radioactivity of insulin-antibody complex when the concentration of unlabelled insulin is zero

CI = the radioactivity of insulin-entibody complex when the concentration of unlabelled insulin is I

Ca/CI is linearly related to 1. In practice, the slope of the line is obtained by having a series of standard solutions made from human insulin standard provided by the kit. The standard solution ranged between 5 Munit/ml to 244 Munit/ml, and any samples which fell outside this range had to have the assay repeated. The plasma had to be dilluted if the concentration was too high, at a larger amount of plasma sample was needed if the concentration was too law. An aliquot of 0.05 ml of plasma was used in this assay. In addition to this, a salution of unlabelled human insulin (40 Munit/ml) was made, put into a series of polythene lubes and stored at -20° Centigrade. This unlabelled human insulin solution was always included in each insulin assay and served as a quality control, and one batch of quality control overlapped with the next batch.

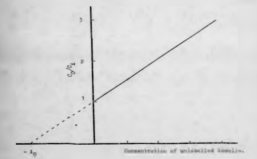
The counting of radioactivity was done using a well typed&-scintillation counter (ECKO Electronics Ltd., England).

- Figure 9. Hadia-immine assay of Insulin with Insulin-antibody Precipitate.

 Theoretical relationship between the ratio of the radioactivities
 in the insulin-antibody complex in the absence and presence of
 unlabeled insulin. (See details given in the text).

 Co- radioactivities in the insulin-antibody precipitate in the
 absence of unlabelled insulin.
 - \mathbf{C}_{j^m} redicactivities in the insulin-antibody precipitate in the presence of unlabelled insulin.

(from Hales and Randle, 1963) .



4. Plasma Amina Acids:

Plasma amino acids were estimated using the ninhydrin method (Spackman, Stein & Moore, 1958) in an emino acid analyser (Locarte, England).

7. Glucagon:

Plaina glucagon was measured by radialimmuno-assay using a pancreatic alucagon specific antiserum (Russell et. al., 1975).

8. Carticosterolds:

a. Plasma cortisol:

Flama contisol was measured by the competitive binding method of

b. Urinary free cartisal:

Urinory free contisol was measured similarly to plasma contisol but the extracted curtisol was purified by paper chromotography and lesses monitored by the addition of ^{1,4}C-certisol

g. Urlnery 17-OH-corticosteroids:

These staroids were measured by the sodium borohydride reduction, sodium periodate axidation method (Grey, Baron, Brooks and James, 1869).

F. Assessment of the Data:

Palrod 1 lost (Armilogo, 1971) was used in assessing changes within the patients.

The ordinary student's 1 lest was used when comparing data from the patients with those of the controls.

CHAPTER III

A. Nitragen Balance:

A complete nitrogen balance study in three patients indicated the extent and duration of negative beforce which was equivalent to about 50 g protein loss peridar, for a period of 5 to 6 days after the operation. Since during the first part of this period the patients neither received food nor had any local losses, urinary (nitrogen) excretion adequately reflected the changes in the (negative) balance. However, this figure did not include the amount of blood transfusion (if any) or blood loss. The mean altropen balance in the three patients is shown in Fig. 10.

By day 10, a positive nitrogen balance had been re-established with a full in urhary nitrogen excretion and an increase in food intake.

8. Urinary Nitrogen Excretion:

The 24 hour unlarry nitrogen excretion showed an increase during the period following the operation. The mean values for eleven patients showed that the increases were significant compared to the mean pre-operative levels, on days 1, 2 and day 3 (p < 0.05, paired t test), and a significant decrease on day 6 (p < 0.01, paired t test) (Fig. 10).

C. Fascal Nitrogen Excretion:

During the days immediately following the operation, there was no foecal excretion. Table 7 shows the delity values of foecal nitrogen excretions in the three patients studied.

Plaire 10. Mean (* 588) 24 hour wrinary aitrogen excretions is 11 patients (vertical lines), and mean nitrogen balance is 3 patients (vertical blocks) before and after operations

There were significant increases (from preoperative values) in urinary mitrogen exerctions on day 1, 2 and 3 (p < 0.05), and a significant decrease on day 6 (p < 0.01) (paired t test).

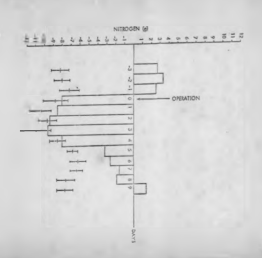


Table 7

Daily foscal nitrogen excretion in 3 patients (g)

Subjects	Day (g)												
	-3	-2	-1	Operation	1	2	3	4	5	6	7	8	9
HG		2,2		4								2.2	
нт	3,8	6.3	0.4	14	-	-	-	-	-	3,4	1.4	0.1	0.2
NT		1,3	0.7		-		-		•	-	-	-	0.1

- = na faecal excretion

blank = not measured

D. Urinary Creatinine Excretion:

The 24-hour urinery creatinine excretion in eleven patients followed closely the pattern of urinary nitrogen loss. There were significant increases from the mean pre-operative levels on days 1, 2 and 3 (p< 0.05, paired i test), and significant decreases on days 5 and 6 (p<0.03, paired t test). Fig. 11 shows the delly values of wrinery creatinine excretion in the eleven patients studied.

E. The Relationship between Nitrogen Balance and Dally Harmonal and Substrate Values:

1. Plasma Insulin:

The period of negative nitregen balance coincided with increased festing plasma insulin level, although it was only on day 1 that the difference between the values reached a significant level (p < 0.05, poired t test) (Table 8).

2. Plasma glucagoni

The fasting plasma glucegon level was also significantly raised from preoperative levels in the first few days after the operation (paired t test) (Table 8).

3. Plasma cortisol

The fasting plasma cortical level, (the insulin, was significantly relised from gree-operative levels only on day 1 (p < 0.05, points) to table 8).

4. Urinary steroids:

The 24-hour excretion of urinery 17-OH-conflicationality, however, was significantly relised for the first few days ofter the paration (point it set) (Fig. 12), as also was the 24-hour uninary free continol excretion (paintd (test) (Fig. 12).

There were significant increases (from preoperative values) on day 1, 2 and 5 (p < 0.05), and significant decreases on day 5 and 6 (p < 0.01 and p < 0.05 respectively).

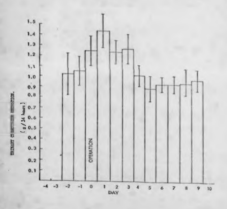


Table 8

Mean fasting levels of blood glucose, plasma insulin, plasma glucegon, plasma certisel and NEFA in patient and centrals (\$2.55M). Faired is test was used for comparing pre-operative values with the values on the days following the appention.

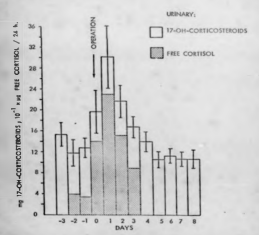
POST-OPERATION

	No. of	Pre- operation	Day 1	Day 2	Day 3	Day 8	Recovery*	Controls
Blood glucoss (mmol/1)	10	5.320.14	5.920.23	5,870.37	5.620.30	5,620,15	5.120.15	5,1=0,35
Plasma insulin (uunit/ml)	10	1620.8	25*2.2	24 [±] 4.0	1,643,1	2012.5	1621.8	15#2.5
Plasma glucagon (ng/l)	10	60=10.0	173±35,1		136#24,1*	64±19.6	6621 6.3	35±12.1
Plasma cartisal	9	359#26.5	486±57,1	282#30.1	339±18,2	386436.2	315#19.6	306/13.5
NEFA (amal/I)	9	357258	625275	4322541	432756°	465-82	4592103	500=197

Significant difference: "p<0.05; ""p<0.01; ""p<0.001. (Paired t test

Meure 12. Rean values of 24 hour urinary secretions of 17-08-corticosteroids in 10 (4 SDM), and free corticol in 5 patients:

There were significant increases (from preoperative values) in urinary 17-08-corticosteroids on day 1, 2 and 3 (y < 0.05), and is urinary free cortisol on day 1 and 2 (y < 0.05) (paired t test).



5. Blood glucose:

The fasting blood glucase level, like insulin, was significantly raised from pre-operative level only on day 1 (p < 0.05, pained # test) (Table B).

6. Plasma NEFA:

The fasting plasma NEFA concentration was significantly raised from preaparative level during the first few days following the operation (paired t test)

(Table 8).

7. Plasmo omina acids:

There was a decrease in the levels of fasting total planes amine acids.

However, fasting fevels of plasma feucine, iso-leucine and valine were raised in

concentration (Table 9).

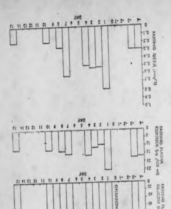
These substrate and hormonal changes in relation to nitrogen balance are shown in Fig. 13 for one patient (HG).

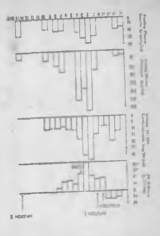
The overell picture is one of an acute rise in blood glucose, and in plasma insulin and cortisol on day 1 after the operation. This may be compared with the sustained changes lasting for about the period of the negative nitrogen balance, in plasma glucogen, in urinery glucoriticolds excretion in plasma amino acid and NEFA concentrations.

Table 9

	"Recovery"	2412 2673 2800 3469 3469	22222	99233	222.23.23.23.23.23.23.23.23.23.23.23.23.
During the of total plasma valine.	Day 8	2139 2115 2486 1991	2283	2552	107 230 230 207
g plasma emino acids in patients and centrals (\$ SEM). During se operation, there was a decrease in the traiting levels of total increase in the fasting levels of tactucion, levelne and valing	Doy 4	1810 2884 3108	8 = =	222	327 327
its and confre sees in the fi fluctavelous	Day 2 Day 3 Day 4 AMINO ACIDS (mol/1)	1431 1838 2567	25 77 25 25 25 25 25 25 25 25 25 25 25 25 25	22.52	270
ds in patien was a decre tog levels o	Day 2	1614 1973 2222	SO-LEUGINE (p.mal/1) 32 66 82 81 33 81	101 163 167	156 219 256 256
1 8 c	Day 1 TOTAL PLASMA	1545 2045 2240 2501 2501 2689	150-L	25 7 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2 1 2	22 22 22 22 22 22 22 22 22 22 22 22 22
of festing after the	Pre-operation TO	2152 2581 2391 3678 2333	32332	133 126 148 135	205 231 238 258 274
Daily values of first few days omino acids,		4	17 - VI - 21 - 21 - 21 - 21 - 21 - 21 - 21 - 2		11.8 (n = 4) 17.7 (n = 4)
	Subjects	E IN SE	AT NT NT NT NAC	ē ·	Controls 114 # 11.0 (n = 4) HT NO NO NO Controls 195 # 17.7 (n = 4)

Pierre 13. Daily values before and after the operation for one patients (MD) showings changes in nitragen belance, in urinary 17-08-corticasteroids excretion, and in fasting concentrations of glucagon, insulin, corticol, glucose and MEFA in plasms.





F. Substrate and Hormonal Changes during Glucase Infusions

Except for the lesting levels (p + 0.05), there was no significant difference between the glucose values at any one time in infusion 1 (day 1: and at the same time in infusion 11 ('tecovery'), but there was a trend to a higher 2-hour level on day 1 (12.08 m mal/1 against 10.36 m mal/1). However, both curves were significantly higher than in normal subjects where the mean blood glucose levels reached a plateau at 6.07 m mal/1, Fig. 14).

Glucose infusion caused a substantial fall in plasma glucagon concentration (Fig. 14) but because of the higher faiting glucagon levels, plasma glucagon levels continued to be significantly higher during the Infusion on day I than at similar times during the Infusion done on "recovery" (p < 0.05). Similarly, the mean glucagon curve during infusion on "recovery" was higher than in controls, but the difference was not smitstically significant (p > 0.05).

Infusion of glucose also decreased the levels of plasma NEFA, total plasma amino acids and plasma cortisol on both occasions in the patients and also in the controls (Table 10), but the changes in plasma cortisol during glucose infusion in controls were not significantly different from those found in normal subjects at rest (J.D. Few, unpublished date).

Plasma insulin levels during glucose infusion were significantly higher on day 1 than on "recovery" both in respect to the fasting levels (p.<.0.001) and the levels reached diving glucose infusion, in which a significant difference was reached at 30 minutes (p.<.0.05). At both times the patients had levels which were significantly higher than those of the normal controls. A significant difference between the patients' Pigure 14.

Heam (2 SSM) blood glucose, plasma insulin and plasma glucosym concentrations during 2 hour glucose infusion $(0.55~g~g^{-1}b^{-1})$ in patients! infusion I (Δ), patients! infusion II (O) and infusion of soutrols (\Box),

Significance of differences :

Pasting volume: See Table 6.

Glucova concentration: Infusion I z infusion II, non significant.

Infusion I and II s sontrols , from 60° p < 0.01.

Insulin concentration: Infusion I x infusion II, from 30° p < 0.05.

Infusion II z Controls, from 901 p < 0.05.

Glucason concentration: Infusion I z infusion II, all times p < 0.05.

Infusion II x controls, non significant.

The data were from 10 patients and 4 healthy control subjects. The paired t test was used for comparing values in patients' infusions I and II, and Student' t test for comparison of patients and control subjects.

ě,

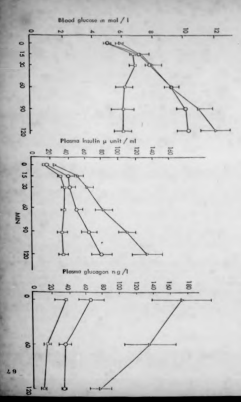


Table 10

The effect or 2 hour glucose infusion (0.35g kg⁻¹ h ⁻¹ ten NEFA, total amino acids and control levels in plasma of patients and controls (2 SEM. Student's t test was used for comparing mean values between controls' and patients' infusion (()) and between controls' and patients' infusion (()) and between controls' and patients' infusion (()). Paired t test was used for comparing values within individuals.

	Flama NEFA (umal/1)			Total plasma amino acida (-mot/1)			Plasma cortisol (resul/1)			
	Fast		Post Infusion	Fost		Post Infusion (140 J	Fost (©)		Post Infusion (1801)	
Infusion I	624 2 75	***	132 = 28	2208 2 192		1701 2 84	486 ± 57.1		403 ± 47.2	
No. of parients		9		***	5			9		
Infusion II	4.59 ± 103	***	121 ± 36	2879 ± 179		2360 ± 186	315 = 19.6	*	275 4 29.3	
				*(t)		*** (1)			a(II)	
Controls	500 a 197	***	208 ± 67	2893 ± 175	*	2408 2 36	306 ± 13,5	9	160 ± 74.5	
No. of controls		4			4			4		

Significant difference: "p .05; ""p < .01; """p < .001

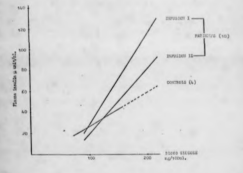
Infusion I and controls' Infusion was reached at 30 minutes (p=0.01), and between pattents' Infusion II and controls' Infusion at 90 minutes (p<0.05) (Fig. 14).

Although the high levels of plasma Insulin during the glucose infusion were essociated with a higher level of blood glucose, the insulin levels rended to be higher for a given glucose value in the immediate post-operative period. Thus after 1 hour the mean blood glucose levels in the patients' two infusions were identical (Fig. 14) but the corresponding insulin values in the immediate post-operative state were almost twice those found in 'recovery'.

Fig. 15 shows the regression lines between plasma insulin and blood glucose concentrations during plucose infusion of controls and during infusion I and II of patients. A given glucose level tended to be associated with a higher insulin level in infusion I than in infusion II of the patients, and higher in infusion II of the patients.

Picture 15. Regression lines between plasma installin and blood glucose concentrations during patients' infusion I, patients' infusion II and infusion of controls.

Data from 10 patients and 4 healthy control subjects.



CHAPTER IV

When surgery is followed by a period of dietery deprivation, alleviated only by the use of intravenous glucose, patients obviously suffer from both acute protein and energy deliciency. This is manifested by negative nitragen balance and an increase in the levels of those harmones and metabolites known to change in response to starvetion.

increased levels of glucegon, which promote rapid mobilization of fuel from carbohydrate, fat and protein (Foa, 1972) and of cartiaci, which affects glucense-genesis (De Bado and Altzuler, 1958) are features of the body's response to fuel shartuge. The increased output of urinary nitrogen in the face of a felling protein intake illustrates the effectiveness of energy mobilization and gluconeogenesis, which have a priority in the short term over protein retention.

The major difference between the effects of surgery and those of fasting in otherwise healthy subjects may till in the insulin response. The immediate acute response to surgery is the same as to starvation, i.e. reduced levels of plasma insulin. After surgery this is followed by a period in which basel plasma insulin levels are raised (Lindsey, et. al., 1974) and we have found that glucose infusion results in an elevated insulingfucose ratio compared with ratios found on recovery or in normal control subjects (Fig. 15). Therefore, the apparent discrepancies between reports of low insulingfucose ratios and ours of high ratios, are indeed related to the time of observation.

During the initial acute phase of trauma, insulin secretion is probably blocked by the high circulating levels of catechalamines (Cersal, Luft and Efendie, 1971).

Our measurements were made, however, 24 hours after surgery at a time when the nutrient supply may well dominate the hormonal response.

The elevated levels of basal plasma insulin and high values for the insuling glucom ratio in our patients can be interpreted as a compensation in insulin-output as a result of peripheral insulin resistance (Parte, 1975). This resistance may be an innete response to troums since patients after surgery when infused with amino acid solution alone, though having lower absolute levels of insulin and glucose, have the some Insuling lucose ratio as patients receiving parenteral glucose (Blackburn, Flatt, Clowes and O'Donnel, 1973). In the early stages after surgery, analysis and the degree of surgical trauma certainly play a rais in the metabolic response of hepatic and peripheral timues (Long, Spencer, Kinney and Geiger, 1971; Wikland and Jorfeldt, 1975). High circulating levels of catecholomines can induce lipplysis (Rosell, 1966), blockade the release of insulin (Carasi et. al., 1971) and stimulate the output of glucore from the liver (Beam, Billing and Sherlack, 1951), appropriate analysis in the most-operative period can reduce the circulating NEFA and alucose levels and the rate of release of glucose from the liver (Wikland and Jorfeldt, 1975). Elevated levels of NEFA may themselves be expected to induce a reduction in the rate of alucose outflow from the circulation with the glucose intolerance and insulin resistance (Randle, Hales, Garland and Newsholms, 1963; Balasse and Neef, 1974) although our NEFA results are not particularly high. Thus, averproduction of glucose by the liver and slower uptake by the periphery may both tend to increase blood glucose levels and an afteration in either process may be responsible for evidence of insulin resistance (fellig and Webren, 1975).

This resistance seems to take some time to return to normal. There was no evidence of obesity in the perients, a condition which would explain a peristently low glucose tolerance and high insulinglucose ratio (see Pert II of this thesis). Furthermore, the age of the patients would load one to expect lower insulinglucose ratios (see Part IV of this thesis). The persisting increase in plasma glucogon in the patients studied in the 'recovery' phase argues against their hormonal status having resumed completely to a pre-aperative level despite being studied two weeks after surgery.

The demonstration of high circulating levels of glucagam after surgery is in keeping with the results of other studies of patients in the post-apparative phase (Lindsey et al., 1974). Russell et al., 1975). The elevation of glucagan pensists despite the mildly raised blood glucase level but the glucase in fusion led to a suppression of glucagan towards normal. This impaired suppression presumably reflects the realistance in the response of the al-cell to glucase and may be part of the more generalized insulin resistance at a cellular level (Samats, Tyler and Marker, 1972).

While glucogon itself may play an important role in the control of gluconeagenesis and glucose production, there is increasing evidence that I is role in producling hyperglycoemia depends an insulin deficiency itself and the glucogon-insulin
ratio as being of less importance in the pre-diabetic state (Sharwin, Fesher, Hendler
and Felig, 1976). Thus enhanced hapatic glucose output after surgery may reflect
either a direct short term effect of catecholomines on glycogonal vals or the combined

effects of insulin resistence and hyperglucagoneemia on gluconeogenesis. The additional role of enhanced corticosteroid secretion in stimulating gluconeogenesis probably operates ever the 4-5 days after surgery as judged from the urinary data (Fig. 12).

It is not possible from the present evidence to quantitate the relative roles of hepatic averproduction of glucose and periphend resistance to glucose uptake and axidation despite the use of an infusion of glucose which might be expected to repress hepatic glucoseogenesis. Our infusion rate was approximately five times the basel glucose tumover rate in normal man (Bower and Moorhouse, 1973) and it seems vary untikely in healthy controls that this quantity of glucose did not suppress glucon-agenesis (Moditon, 1909). In fewour of the suppression of hepatic glucose production in our patients, as well as in the controls during glucose infusion were the similar falls in glucegon, NEFA and amino acid levels, but we have no direct evidence this substrate supply for gluconeogenesis become rate limiting, or that the peripheral tissues were more or less sensitive to insulin than the hepatic cell.

A diminished protein synthetic rate post-operatively (O'Keefe et. al., 1974; Crene, et. al., 1976) might be the result of insulin resistance, but this espect of protein metabolism requires further investigation. The net increase of protein breakdown is shown by the increased fasting plasma levels of branched-chain amino acids (Tobie 9). The rise of these emino acids, however, may not be specific to this phenomenon since prolonged starvation eline may also give a similar or even a greater rise (Adibl., 1968; Felig, Owen, Wehren and Cahill, 1969).

Therapeutic attempts to compensate for insulin resistance, assuming the resistance is undestrable, have sometimes taken the form of massive insulin administration (Hinton, Allison, Littlejohn and Llayd, 1971). Insulin sentitivity may differ between various organs and clasues and even between different actions of insulin within the same time. This success caution in the use of insulin. Furthermore, it could even be around that insulin resistance protects the subject from the less desirable feature of elevated Insulin levels. An alternative approach would be to give parenteral amino acids rather than glucose (Blackburn, et. al., 1973). The lower levels of insulin induced by amina acid infusions would permit more ready mobilization of lipids and, therefore, allow the conservation of protein and glucase. Interestingly, plasma NEFA levels in our patients given glucose were not particularly high. If however, the development of Insulin resistance places the patient at a disadvantage, this could be countered not only by giving insulin but also by blacking the release of insulin antagonists. It has been shown that surgery under morphine anaesthesia is not accompanied by the usual elevation of plasma cortisol and growth hormone concentration, and that the lack of edreng-cortical stimulation had no adverse clinical affect (George, Raier, Lanese and Rower, 1974).

It is shown, therefore, that after surgery, metabolic responses have some elements in common with stervation, but exacerbated and modified by hormonal stress responses. Similarly, there are resemblences with the diabetic, e.g. in hyperglucogonaemia, but in surgical patients the suppressibility of glucagon with glucose, possibly via alevated insulin levels, marked a clear distinction between the two PART III
GLUCOSE HOMEOSTASIS
IN
OBESITY

CHAPTER I

A. Definition:

Obesity is defined as a state in which an excessive amount of fet accumulates and where the body weight exceeds by at least 20 per cent the normal or desirable weight for a given height and age (i.e. 'ideal body weight') (Starm and Mirsch, 1972; Davidson, Passmore, Brock and Truwell, 1975). Others take 10 per cent above the ideal weight as the upper limit for normal weight, (Craddack, 1973). In the present work 'ideal' body weight for a given height and age was derived from the data of everage weights of adults in Gelgy Scientific Tables (1970). These are based on the data of insured persons in the United States (Society of Actuaries, 1959). Some may not agree with the use of relative excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since the excess of body weight as an indication of obesity, since the excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity, since this excess of body weight as an indication of obesity.

B. Problem of Obesity:

Millions of people in the developing world suffer and die from inadequate food intake, while transcally, millions of their counterparts in the western countries are in the opposite situation. They eat a lot and some even too much, and have too little physical exercise, which inevitably leads to obesity.

To some, chastry simply means an aesthetic problem, yet to many it comes as a sarious health hazard. Obsetry is known to be related to several diseases, a.g. diabates (Jaslin, 1921); Smith and Levine, 1961); hypertension (Marks, 1970); cardio-respiratory fellure (Barlayne, 1958); and many others. Not suprisingly, therefore, abosity is also related to an increase in the rate of mortality and marbidity (Marks, 1970).

C. Aetlalogy of Obsiliys

The question is inevitably asked whether obstity is always a simple problem of overeating and lack of physical exercise. The entwer obviously would not be as simple as that.

Under normal conditions, the body is capable of regulating its energy belonce quite accurately and the body weight of a healthy adult remains relatively steady. The human body has the ability to interrupt its food intake. Therefore, it is capable of having results at cartain times of the day, rether than as repeated small snacks, and thus it is able to do creative work during the day and have long intervals of these at right. All this is possible only because of the body's ability to store energy when it is evaluable in excess of immediate requirements, and use it lates when external energy supply falls below its requirements. However, interestingly in other higher primates such as gotiles and orang-utens, they spend most of their day time foreging for food and this is interrupted only by brief pations of rest and sex. This may be due to the type of food they east rather than a major physiological difference the apas and man. The eges do not set high energy density foods as humans do, therefore, they have to eat a lot to be able to meet their energy requirements.

For some curtous reasons, aur deficate energy before a con be disrupted resulting in an excessive amount of energy being stored, 1.e. more than will be used in the interval before the next meal. This extra energy is stored mostly in the form of fat in adjose itsue. This may be the result of simple over eating, either caused by social pressure or other psychological stress, or may be due to something much more complicated and have an actual organic cause resulting in metabolic malfunctions.

The latter could be of genetic, hyporhalamic or endocrine origin (Mayer, 1957).

This work does not attempt to investigate these complicated forms of obesity, eithough from the range of patients we studied, such forms of obesity cannot be excluded.

D. Insulin's Role in Obesity:

Excessive energy is stored mostly in adipose stasse as triglyceride. In fact 80 to 85 per cent of adipose tissue in an adult consists of triglyceride, (Craddock, 1973). In the presence of insulin, for synthesis is enhanced, while when insulin is obsent, glucose enters adipocytes less easily and the supply of lipogenic precursors is diminished. At the same time I polysis in adipose tissue is accelerated.

It is known that obesity is usually related to Insulin resistance (Franckson, Malaisse, Amauld, Rasia, Ooms, Balesse, Canrad and Bastenie, 1966; Chlauverekis, and White, 1969), and is associated with hyperinsulinaemia (Back, Kaumans, Winterling, Stein, Daughaday and Kipnis, 1964; Frankson at al., 1966; Perley and Kipnis, 1966; Chiles and Tzagaumis, 1970). Insulin resistance could be defined here as the relative inability of insulin either endogenous or exagenous, to lower the blood gluces concentration. The alucase balenace test in obesity may sametimes still be

within the normal limits (Perley and Kipnis, 1966) but most aften it is impaired (Paullin and Sauls, 1922; Beck et. al., 1964; Perlay and Kipnis, 1966; Chiles and Taggournis, 1970). Whether the degree of insulin resistance correlates with the degree at obesity and whether the amount of circulating insulin may in a way be the major factor in contributing to obesity, are being investigated in this work. This work concentrales mainly on the role of insulin in obese subjects faced by a glucose load, either aral or intravenous (Iv), and compares them with the data collected from young normal adults. This work was not planned as an attempt to unravel the mysteries of obesity. Rather the studies were planned to extend the range of conditions exhibiting altered glucose tolerance, insulin sensitivity and plesma insulin levels. Insofar as the obesity is known to be associated with varying degrees of diminished glucose tolerance and increased insulin resistance and enhanced insulinogenic response to glucose, the biochemical signs of abegly are similar to those of the post-surgical patients. Yet such patients suffer from undemutrition while the obese struggle with evernuturition. It was hoped to find some clues why these very different nutritional states and up with a similar metabolic picture.

CHAPTER II

A. Subjects:

1. Patlanta:

twenty nine patients were studied (Table 11). Their mean age was 40 ± 2.9 years, their mean height was 165 ± 6.5 cm, their mean body weight was 100 ± 5.5 kg and they ranged from 115 per cent to 233 per cent of their ideal body weight (the mean was 166 ± 6.7 per cent). These patients were either from the abesity clinic, University Callege Hospital, or patients admitted from this clinic to the methodic ward (Manson Ward, Hospital for Trapical Diseases), both of the University Callege Hospital Group. The patients were asked whether apart from their course of treatment, they would be willing to consider participating in our study. It was then explained in detail what kinds of tests were going to be involved in their study, and consent (oral or written) was obtained. None of the patients were known diabetic and all had fasting blood glucose concentrations less than 6.11 mmol/1 (110mg/100 ml). 6.11 mmol/1 is the upper limit for normal feating blood glucose recommended by a working party appointed by the Callege of General Practitioners, England (1963). None of the patients had allowed allowed to the callege of General Practitioners, England (1963). None of the patients had allowed.

The delly energy intake of the out-patients prior to the tests was not studied, but from information given by the patients it was estimated at between 8.4 and 12.6 MJ /day (2000 to 3000 Celoriet/day) including a reasonable carbohydrate intake.

Patients who were admitted to the ward, received a diet of 8.4 MJ/day (2000 Celoriet/day) with a carbohydrate content of 200 g/day, far 3 to 4 days before they underwent

List of Obese Patients Table 11

Tests

											Insuffi		Insulfi		
iv GTT and Iv GITT	0 Z	No	No	°Z	No	o Z	oN.	°Z	No	No.	Ne	No	2	Yes	
Oral GTT (ZO potients)	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	745	Yes	Yes	Yes	Yes	Yes	
% ideal body weight	141	218	115	140	142	156	130	124	95	218	202	207	21.8	192	Sles (1970)
Weight (kg)	0	139	R	16	71	105	22	10	11	22	124	122	178	12	jeniffe To
Helght (cm)	126	175	168	163	179	162	22	155	157	156	25	168	178	169	*Compared with the date from Geigy Scientiffe Tables (1970)
(year)	25	28	17	45	922	53	N	62	11	46	100	99	8	24	of the fron
SE X	u.	14.	u -	*	W	16.	16.	i.	Me	ú.	u	ш.	×	W	of with t
Subject	b	r,	8	af	u. o.	03	MS.	DR	25	No.	KOM	50	*	NC	*Compare

Subject	Sex	Age (years)	Height (cm)	Weight (kg)	% ideal body weight*	Oral GTT (20 parlants)	(To patients)	
DR	F	33	166	110	175	Yes	Yes	
	F	21	173	135	213	Yes	Yes	
HG		17	163	90	160	You	Yes	Insulin not measured
MR	F		160	128	217	Yes	Yes	
WB	F	47		86	116	Yes	Yes	Insulin not measured
8C	F	42	175		122	Yes	Yes	
FH	F	22	162	75		No	Yes	
WT	F	14	161	94	177		Yes	
GR	28	20	176	180	233	No	Yes	
KY	F	27	161	107	179	No	Yes	
CW	F	53	162	80	119	No		
AD	F	68	158	79	123	Nà	Yes	
AR	м	45	168	120	164	No	Yes	
	F	52	156	96	156	No	Yes	
VR		51	155	100	164	No	Yes	
GG PR	F	31	168	123	191	No -	Yes	Insulin not measured

^{*}Compared with the data from Gelgy Scientific Tables (1970)

any treatment or any further dietary restriction. The tests on these in-patients reported in this work were done during the end of this period.

2. Normal subjects:

Fourteen young non-obese normal subjects were used as controls (Table 12). Their mean age was 27.6 ± 1.3 years, their mean height was 169.2 ± 2.72 cm. their mean body weight was 61 ± 2.21 kg, and ranged between 85 per cent and 108 per cent of their idnot hody weight (the mean was 94 ± 1.9 per cent). All the control subjects were told to eat sufficient load for at least three days prior to the test, including a reasonable carbohydrate intake.

B. Tests:

Most subjects, patients as well as the controls, received three kinds of test-Each was done in the morning.

1. Oral glucose talerance test (oral GTT):

A butterfly needle attached to an ontecubited vali on one arm was used for drawing blood samples. The fasting blood sample was taken at 9.30 a.m., after the subjects had been fasted from midnight. A drink of 50 g glucose in 250 ml water was given and serial blood samples were taken at 15°, 30°, 60°, 90°, 120° and 150° after the oral glucose. Happiin tubes were used to prevent the blood from clotting. An alliquot of 0.05 ml of blood was taken for blood glucose estimation. The blood bubes were centrifuged and the plasma separated. An alliquot of 0.05 ml of plasma was them used for plasma glucose estimation, and the rest of the plasma was stored. Plasma samples were stored at -20° C until analyzed for plasma insulin and NEFA (see Part II). Chapter II, Methods).

List of Normal Subjects

6513	in GIT and in GITT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	ů.	Ž	e Z	2	Yes
9-	Oral GTT (T3 subjects)	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes.	Yes	Yes	Yes	Yes	ž
	% ideal	25	06	88	108	25	100	100	100	28	42	98	96	06	13
	Weight (kg)	8	89	99	\$	22	23	E	15	13	8	23	78	85	13
	Height (cm)	165	178	178	165	185	158	10	160	159	18	160	100	169	168
	Age (years)	26	8	27	25	24	24	23	22	23	25	9	B	R	11
	X-D	W	×	u.	u.	×	SS .	34.		×	×	×	×	11-	2
	Subjects	NS	50	MG	03	DA	e H	R	NS NS	115	99	d	00	SM	23

If the whole glucose (or insulin) curve in an oral GTT is thought of as being made up of 15 minute units (Fig. 16), then the 0' - 15' period is one unit, 15' - 30' is one unit, but 30' - 60', 60' - 90' etc. are each 2 units. The total number of 15 minute units is 10.

The average values of glucose or insulin concentrations during an oral GTT were, therefore calculated as:

Average value =

$$\frac{c_0 + c_{15}}{2} + \frac{c_{15} + c_{30}}{2} + 2 = \frac{c_{30} + c_{60}}{2} + 2 = \frac{c_{60} + c_{120}}{2} + 2 = \frac{c_{120} + c_{150}}{2}$$

10

where C₀, C₁₅, C₃₀, C₆₀, C₉₀, C₁₂₀, and C₁₅₀ were blood (or Insulin)concentrations at 0°, 15°, 30°, 60°, 90°, 120° and 150°.

The molar concentration of insulin was calculated from the concentration measured in the assay (µ unit/ml), on the basis of 1 unit of insulin being equivalent to 0.04167 mg (1 mg = 24 units) and the molecular weight of human insulin being 5807 (Scientific Tables, Geigy, 1970).

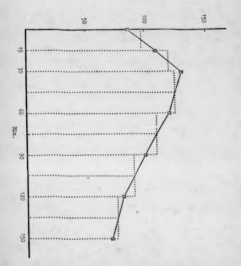
The insulin-glucase malar ratio during the oral GTT was calculated for each subject as follows:

motor ratio= Average insulin value (in mol/1)
Average glucose value (in mol/1)

Calculation of the average value of blood glucose concentration during oral glucose telerance test (oral CTT).

(m) are observed concentrations of blood glucoss at various times during the test.

(See details given in the text).



2. Intravenous glucose tolerance test (Iv GTT):

This test was also done using a butterfly needle attached to an antecubital velo of one arm. A fasting blood sample was applied taken as 9,30 a.m., and through the same needle a solution of 50 per cent alucose was injected. In the first obese patients an attempt was made to insert a butterfly needle in each arm, one for the injection of glucose and the other for blood sampling. However, it is difficult enough to find one suitable vein in prossly overweight subjects, without subjecting patients and operator to the added traums of trying to do it twice. There was no evidence that the injected glucose in any way interfered with subsequent glucose sampling, as, acially as the tubing and needle were flushed in the normal saline after each operation (see Part II, Chapter II, Methods). The subjects received 0.33 g glucose/kg body weight (maximum 25g per person) (Franckson et. al., 1966). Glucose was injected within 4 to 5 minutes and serial blood samples were taken at 5°, 10', 20', 30', 40', 50' and 60' after glucose injection. The procedure for handling blood samples was similar to that in the oral GIT. The blood glucose concentration (mg/100 ml) was platted against time in minutes on Log-Linear graph paper and the best fitting straight line drawn through the points (Fig. 17). The percentage removal rate of glucase (Kn) was calculated from this line using the equation;

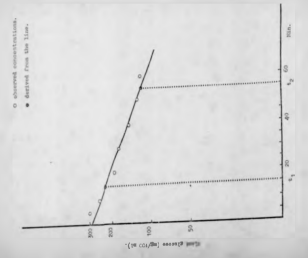
$$\kappa_G = \frac{2.303 (\log C_1 - \log C_2)}{i_2 - i_1}$$
 | 1 100% min

where C₁ and C₂ were glucase concentrations read from the graph at any convenient times t₁ and t₂ minutes after glucase injection (likkes and Luft, 1957). Heard, 1966).

The validity of this test will be discussed in Part V (General Discussion).

Calculation of the % removal rate of glucome during intravenous glucome tolerance test (iv GTT) (No value).

(See details given in the text).



It was found that plasme NEFA concentration, during the Iv GTT, decreased in the same way as did blood glucose concentration, so that when the plasma NEFA concentration (μ mol/I) was plotted against time on Log-Linear paper, a straight line could be drawn through these points also. From the best fitting line, the percentage removal rate of NEFA (K_F) was calculated using the same equation as for alucose.

3. Intravenous alucasa insulin talerance test (Iv GITT):

This test was devised as a test of Insulin sensitivity (with respect to blood glucose homeostasis), (Heard and Henry, 1969 a). The principle is that having derived a value for K_G from iv GTI described above, the test is immediately repeated, but on this occasion with insulin added to glucose at a concentration which would avamp the body's endogenous circulating insulin. The differences between subjects in the percentage removal rate of glucose in this test (K_{G+1}) cannot, therefore, be due to differences in plasma insulin concentration, and must be due to variations in responsiveness to insulin.

The validity of the IV GITT when carried out immediately after the IV GIT depends on the differences in results being due solely to the added insulin and not to any priming affect of the first glucase load on the disposal of the second.

Conard (1955) has reported that If a second IV GIT is carried out immediately after the first, they give no difference in the KG. Similar results have been reported by Sansola and Marks (1965), Heard and Henry (1969 a.).

The procedure in the Iv GITT was similar to that in the Iv GITT. Subjects were given a second equal date of glucase at the end of the first test, but this sime insulin (insulin 8P, Wellcome, 20 units/ml) was added to the glucose (10 units of insulin in 50 ml of 50 per cent glucose; 0.133 units insulin/kg body weight). Therefore, the maximal dose of insulin per person was 10 units. Blood samples were taken at 10°, 20°, 30°, 40°, 50° and 60° after the glucose and insulin administration, and $K_{\rm G}$ + 1 was calculated in the same way as $K_{\rm G}$. The procedure for handling blood samples was similar to that followed in the two previous tests,

C. Analytical Methods:

Analyses of blood or plasma glucose, plasma insulin and plasma NEFA wers
corried out according to the procedures reported in the previous study in surgical
patients (see Part II, Chapter II, Analytical Methods.

D. Dietary Treatment

Dietary treatment, general word management, clinical investigations and all tests other than oral GTT, Iv GTT and Iv GTT, were under the clinical control of Professor J.C. Waterlow, Dr. Andrew Tompkins, the late Dr. Peter Sender, Dr. Suson Ell, Dr. Graeme Glugston, Mrs. Inger O'Moere and Miss Elizabeth Roe (dieticlone), and all of the nurses in Manson Ward, Hospital for Trapical Disease.

For the first 3 to 4 days in the ward, all the hospitalized obese subjects received 8.4 M.1/day (2000 Calories/day) before they were submitted to any further dietary restrictions. The dietary restrictions were not always the same from patient to patient. It depended on the clinical situation and the ability of the patients to

cape with the treatment. Basically the restriction consisted of two parts:

1. Initial restriction:

The patients received a dist of either

- A non-protein diet of 1.3 MJ/day (300 (etoriet/day es Hycel (63 per cent glucose syrup, Beecham Products, England) or es
- b. A dist of 2.1 MJ/day (500 Catories/day) which contained 50g protein. The initial restriction was given for a period of 2 to 3 weeks.

2. Maintenance dieti

This diet followed the Initial restriction and consisted of either-

- a. A diet of 2.1 MJ/day (500 Calories/day) which contained
 50 g protein, or as:
- b. A diet of 3.4 MJ/day (800 Calories/day) which also contained 50 g protein, or:
- c. A diet of 4.2 MJ/day (1000 (alories/day) which also contained 50 a protein.

Patients were sent home on this maintenance diet and were followed for periods of time which varied from patient to patient, depending on their co-operation.

An extempt to correlate the type of dietary restriction and weight loss, insulin sensitivity and weight loss in these patients is discussed in the latter part of this report.

E. Assessment of the Data:

The student's I test was used for comparing the patients date with those of the controls.

CHAPTERIII RESULTS

A. Oral GTI:

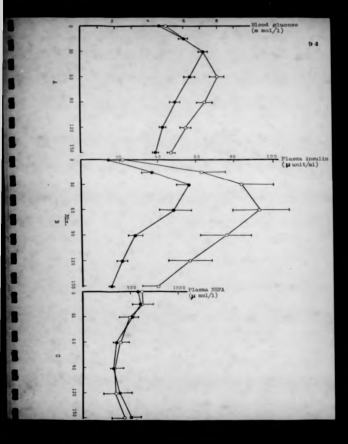
Oral GTT was done in 20 obese patients and in 13 normal control subjects (Tables 11, 12), 16 of these patients had their plasme insulin measured. To avaid unnecessary repetition, results reported below were only the results of 15 patients who had their insulin levels measured. There was no difference in the mean glucose curve between these 16 patients and the mean curve of all the 20 patients.

1. Blood glucose:

The mean fatting blood glucose concentration was 4.89 ± 0.16 m mol/1 (88.1 ± 2.93 mg/100 ml) in the obese patients against 4.62 ± 0.16 m mol/1 (83.2 ± 2.92 mg/100 ml) in controls (Table 13). The difference was not significant (p > 0.05). However, during the test the concentrations of blood glucose in the patients continued to rise after 30° and the peak was at 60° , whereas the peak in the controls was at 30° . Blood glucose concentrations in the patients were higher than in control subjects, throughout the rest of the test, and the difference reached significant levels after 60° (p < 0.05) (Fig. 18A).

The average value for blood glucose concentration during the arel GTT was $4.73 \pm 0.28 \text{ m mol/} (121.2 \pm 5.10 \text{ mg/} 100 \text{ ml})$ in the obese pattents against $5.71 \pm 0.18 \text{ m mol/} (102.8 \pm 3.25 \text{ mg/} 100 \text{ ml})$ in control subjects. The difference was significant (p < 0.01) (Table 13).

- Pieure 18. Oral OTT in 16 obese patients (O) and 13 normal control subjects ().
 - A. Mean blood glucose values during oral GT?.
 There were mignificant differences in the mean fasting values (p < 0.05); and in the mean values at 60°, 90°, 120° and 150° (p < 0.05) between the two groups.</p>
 - Mean planess insulin values during oral CFF.
 There were significant differences in the sean fasting concentrations (p < 0.05); and in the sean values at 60', 90', 120' and 150'
 (p < 0.05) between the two groups.
 - Mean plasma MEFA values during oral GFF.
 The differences between the mean values at various times during the test were not significant.
 (mean values of a plasma see 7 normal control subjects).



2. Plasma Insulin:

The mean fasting plasma insulin level was $21.4 \pm 2.88 \, \mu$ unit/ml. In the abota patients against $12.8 \pm 1.31 \, \mu$ unit/ml. In control subjects. The difference was significant (p < 0.05) (Table 13). During the test the levels of plasma insulin in the patients continued to rise ofter 30' and the peak was at 60', whereas the peak in the controls was at 30'. Plasma insulin levels in the patients were higher than in control subjects throughout the rest of the test and the difference reached significant levels after 60' (p < 0.05) (Fig. 188).

The everage value for plasme insulin concentration during oral GTT was $69.1^{\pm}10.63 \mu$ unit/ml in the obese patients against $33.5^{\pm}3.51 \mu$ unit/ml in control subjects. The difference was significent (p < 0.01) (Table 13),

During the selection of patients, any obvious diabetic (fasting blood glucose concentration higher than 6.11 m mol/1) were excluded. (See Chapter IIA, Patients). WHO criteria for globetes (1965) states that in normal and GTT, the upper limit of 120 minute value for blood glucose concentration is 6.11 m mol/1 (110 mg/100 ml). According to this criteria, among 16 obese patients studied, 7 were 10 labetic and 8 were han-diabetic. Therefore, we could divide that 16 obese patients into two groups. The han-diabetic abese groups did not show any significant differences in the mean blood glucose concentrations at various times during and GTT compared to the normal control subjects, although they showed differences in mean plasma insulin concentrations. The differences between these two groups of obese patients and between these two groups and normal control subjects are summarized in Table 14.

2. Plasma insulla:

The mean festing planne Insulin level was 21.4 \pm 2.88 μ unit/ml in the obese patients against 12.8 \pm 1.31 μ unit/ml in control subjects. The difference was significant (ρ < 0.05) (Table 13). During the test the levels of planne insulin in the patients continued to rise after 30° and the peak was at 60°, whereas the peak in the controls was at 30°. Planne Insulin levels in the patients were higher than in control subjects throughout the rest of the test and the difference reached significant levels after 60° (ρ < 0.05) (Fig. 1 Eq.).

The everage value for plause insulin concentration during and GTT was 69.1 $^{\pm}$ 10.65 μ unit/ml in the obese patients against 33.5 $^{\pm}$ 3.51 μ unit/ml in control subjects. The difference was significant (ρ < 0.01) (Table 13).

During the selection of patients, any abvious diobatic (festing blood glucose concentration highes than 6.11 m mol/1) were excluded. (See Chapter IIA, Patients). WHO criteria for diobates (1965) states that in normal and GTT, the upper limit of 120 minute value for blood glucose concentration is 6.11 m mol/1 (110 mg/100 ml). According to this criteria, among 16 obese patients studied, 7 were slobatic and 8 were han-diobatic. Therefore, we could divide the 16 obese patients into two groups. The han-diabatic' abese group did not show any significant differences in the mean blood glucose concentrations at various times during and GTT compared to the narmal control subjects, although they showed differences in mean plasme insulin concentrations. The differences between these two groups of obese patients and between these two groups of obese patients and between

Toble 13

The mean values for feeting blood glucose and plasma insulin, everage concentrations of blood glucose and plasma insulin, and insulin-glucose moler retio during one GTT, in patients and control subjects (2 SEM). Number of observations in parenthesis.

	Mean feeting	concentrations	Mean of the	average velue fo	1 50 min .	10 ⁶ s mean insulins glucoss malar ratio
	Blood glucoss	Plasma insulin	Blood glucom	Plas	na insulin	
	(m mal/1)	(µ unit/ml)	(m mol/1)	(μ unit/ml)	(106 x = mol/1)	
Patients (16)	4.89 ± 2.93	21.4 = 2.88	6.77 ± 0.28	AP.1 = 10,65	495.8 2 76.4	75.3 ± 11.89
	- 4		-07	**	140	
Controls (13)	4.62 ± 0.16	12.8 ± 1.31	5.71 ± 0.18	33.5 ± 3,51	240.4 ± 25.2	42.2 2 4.50

[&]quot;g < 0.05; "°g < 0.01; ns = not significant

7 9

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	164	21		ğı	B1	<u>R</u>		8 i	Δı		91	ğΙ	M	$\underline{s}_{\parallel}$
-	5.51 0.26	3-8	8,4	g 4 g	g = 6	0.3 0.3		N . G	3-2		76.	3 - 5	5-3	17.65
0,00	,				:	3								
To the figure (9)	- EZ'0	6.27 2 0.30	7.22	3-3	2 B	1 2	8 g	20,4 4,38	8 .8	$\lim_{n\to\infty} m = \frac{1}{n!}$	21,09	3 - E	3 - 5	F - F
11 - 11	:			÷	1	1		1			1		:	. 0
Control (12)	3+5	5 4 6	E-8	3 - 21	515	5.5		3.5	N . 8	# - # # # # # # # # # # # # # # # # # #	10.3	3.65	8 · 2	3-5

z

Since there was no clear cut difference between the two groups of these obese patients, they will be reported in this work as one population.

The differences between non-diabetic and diabetic obese patients will be discussed later in Part V (General Discussion).

3. Insuffin:glucose molar ratio:

The mean insuling fuces malar ratio in the obserpations was (75.3 \pm 11.89) \times 10⁻⁶ egainst (42.2 \pm 4.50) \times 10⁻⁶ in control subjects. The difference was significant (ϕ <0.05) (Table 1.3).

4. Plasma NEFA:

There was a great range in the levels of plasma NEFA both in the obese parliants and in the controls. This applied to tasting levels as well as the levels during the test. The mean fasting plasma NEFA level in the obese patients was 627 ± 120 μ mol/l against 590 ± 68 μ mol/l in control subjects. The difference was not significant (p> 0.03). In keeping with the elevated glucose and insulin values, there was a tendency to a lower 120' plasma NEFA value in the patients, but throughout the test, the difference in the mean plasma NEFA levels between the two groups was not significant (p): 0.05) (Fig. 18C).

8. Iv GTT and Iv GITT:

Iv GTT and Iv GTTT were done in 14 obese patients and in 10 normal controls (Table 11 and Table 12).

1. Kg values:

The mean KG values in the obese patients was 1.11 \pm 0.18 %/minute against 1.81 \pm 0.28 %/minute in control subjects. The difference was significant (a < 0.05) (Table 1.5).

There was no significent correlation between KG and fasting plasma insulin values during to GTT, in the obese patients nor in the control subjects (Fig. 19A). A higher KG value was usually related to a lower fasting plasma insulin (avel.

There was also no significant correlation between K_G and peak plasma insulin values, in the obese patients not in the control subjects (Fig. 198). However, again higher K_G was usually related to a lower peak plasma insulin level.

2. Peak Insulin levels during oral G TT and Iv GTT:

There were 5 above patients and 9 normal control subjects who had both oral GTT and iv GTT dane.

The mean peak insulin levels during and GTT were higher than the mean peak insulin levels during to GTT both in potients and in the control subjects. The differences, however, were not significent ($\rho > 0.05$) (Table 1 Δ).

The mean peak insulin level during and GTT in the obese patients was 160.6 \pm 24.98 μ unit/ml against 72.0 \pm 7.26 μ unit/ml in the control subjects. The difference was significant (p < 0.01) (Table 16).

Yable 15

The mean values for KG , KG + 1 and KF in the obese patients and in serial control subjects (= SEM). Number of observations in parenthesis.

		Mean Values	
	KG (%/mbn)	Kg +1	(%/min)
Ohese patients	1,11 \$ 0.18	2,81 ± 0.25 (16 ²	2.86 ± 0.90 (4)
			ns.
Controls	1.81 ± 0.28 (10)	4_55 ± 0.51 (10)	3.10 ± 0.73 (6)

< 0.05; ns = not significant

Pieure 12 4 Correlation between No and feating plasma insulin values during iv GTT,
in the obese patients (O) and normal control subjects (0).

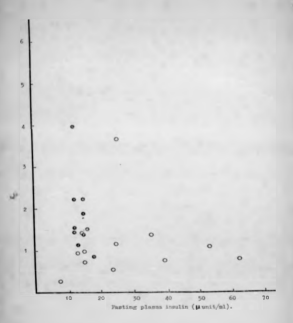
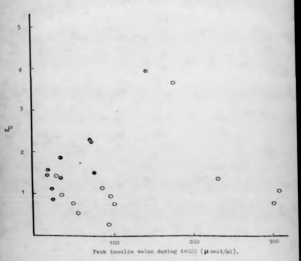


Figure 19 h. Correlation between K_Q and peak plasms immalin values during iv GPF, in the obese patients (O) and normal control subjects (O).



The mean peak insulin level during to GTT in the above patients war 138.4 $^{\pm}$ 11.17 μ unit/mi against 54.0 $^{\pm}$ 13.60 μ unit/mi in the control subjects. This difference was also significant (μ < 0.05) (Table 14).

3. KG + 1 velues

The mean K_{G+1} value in the obserpations was 2.81 2 0.35 %/minute against 4.55 2 0.31 %/minute in control subjects. The difference was significant (p < 0.05) (Table 15).

There was no correlation between K_{G+1} and fasting plasma insulin values in the obese patients nor in the control subjects (Fig. 20A). However, the lower K_{G+1} values were usually related to the higher fasting plasma insulin levels. This, however, is really just saying that lower K_{G+1} values and higher fasting plasma insulin levels are both characteristic of obesity.

There was also no correlation between K_{G+1} and peak plasma insulin values during by GTT in the obese patients nor in the control subjects (Fig. 208). A higher K_{G+1} was usually related to a lower pook plasma insulin level

There was a significant correlation between K_G and K_{G-4-1} values in the obese patients and in the control subjects (p < 0.001) (Fig. 21).

4, Kg valuns

The mean KF value in the obese patient was 2,86 ± 0,90 %/minute against 3,10 ± 0,73 %/minute in the normal subjects. The difference was not significant (Table 15).

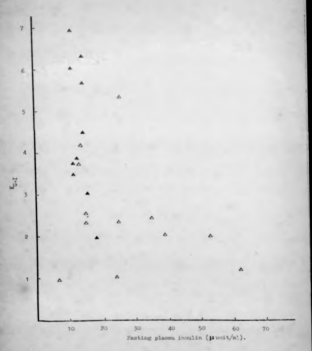
Table 16

The mean peak insulin levels (\pm SEM) during oral GTT and iv GTT in patients and narmal control subjects. Number of observations in parenthesis.

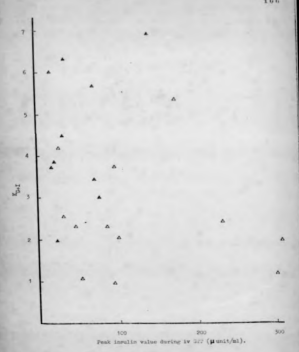
	A	lean Pack Insulin Level	
		(µ unit/ml)	
Oral GTT		ly GTT	Ratio of oral: le
160.6 2 24.98	ns	138.4 ± 11.17	1,16
**			
72.0 ± 7.26	ns	54.0 2 13.60	1,33
	160.6 2 24.98	Oral GTT 160.6 2 24.98 ms	Orel GTT by GTT 160.6 2 24.98 ms 138.4 2 11.17

[&]quot;p < 0.05; ** p < 0.01; ns = not significent

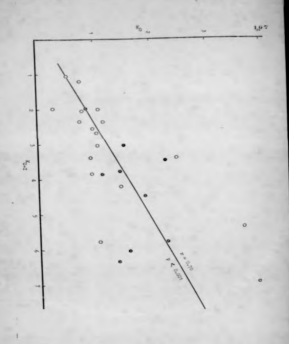
Distra 20 A. Correlation between the % removal rate of glucome during the intravenous glucose insulin telerance test (iv CTTP) ($E_{\Omega+\Sigma}$ value) and fasting planes insulin connectration, in the chase patients (Δ) and mornal control subjects (A).



Pinure 20 l. Correlation between $\frac{1}{2}$ and peak plasma insulin values during iv OTT, in the obest patients (Δ) and normal control subjects (Δ).



Regression line between the values of K_{Q} and k_{Q} in the : obest present obest patients (O) and normal control subjects (O).



C. Carrelation between Blochemical Results and Degree of Obesity:

1. Blood glucom:

Since subjects with a feeting blood glucose greater then 6.11 m mol/1 (110 mg/100 ml) were excluded from this study, the range in the feeting blood glucose concentrations in gettient as well as in controls was very nerrow, fying between 3.39 m mal/1 (61 mg/100 ml) and 6.05 m mol/1 (107 mg/100 ml). In contrest to this, there was a very wide spread in the range of % ideal body weight in these subjects.

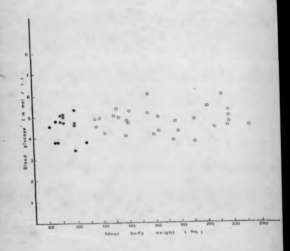
The % ideal weights varied between 85 per cent (the lowest of the controls) and 233 per cent (the highest of the obese group) (Fig. 22A). There was no correlation between feeting blood glucose values and body weight as % ideal body weight.

2. Plasma Insulin:

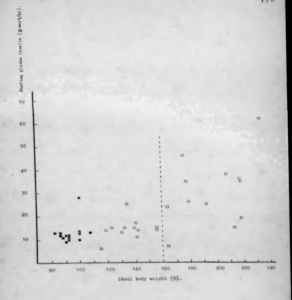
There was a positive correlation between the logarithm of the value of fasting plasmo insulin concentration and % ideal body weight (p < 0.001). (Fig. 228).

There was no statistical correlation between the peek planed insulin concentration during and GTT (at the logarithms of these values and % ideal body weight. However, the lower values of peek plasma insulin concentration during and GTT were usually related to lower values of % of ideal body weight. If we take 160 per cent of ideal body weight as a dividing line, the obese patients and normal control subjects could be divided into two groups. One group was made up of the subjects who had weight less than 160 per cent of their ideal body weight. Among 22 subjects (13 normal and 9 obese) in this group, all except 5 (1 normal and 4 obese) had peek plasma insulin concentrations test than 100 ps unity/ml. The other group contrated of subjects

Correlation between fasting blood glucose concentration and % ideal body weight in the obese patients (O) and normal control subjects (•).



Plaura 22 h. Carrelation between fasting plasma insulin concentration and % ideal body weight in the obese patients (C) and normal control subjects (C).



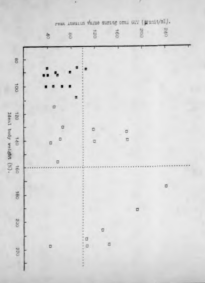
who all had weights higher than 160 per cent of their ideal body weight. Among 7 subjects (all obese) in this group, all except one had peak plasma insulin concentrations higher than 100. In units/ml. However, only 5 out of 9 obese subjects who had weights less than 160 per cent ideal body weight, had low peak insulin concentrations as found in the normal control subjects (Fig. 23A).

There was no statistical correlation between the peak plasma insulin concentration during to GTT and % of ideal body weight. However, here again, the lower values of peak plasma insulin during to GTT were usually rate ted to lower values of % ideal body weight. And if once more we take 1 40 per cent ideal body weight as a dividing line, there were 14 (10 normal and 4 obese) whose body weight was less than 140 per cent of their ideal body weight, and all except 2 (1 normal and 1 obese) had peak plasma insulin concentrations less than 80 µunity/ml. However, 4 of the obese who had weights less than 140 per cent ideal weight, 2 still had low peak plasma insulin concentrations as in normal control subjects. There were 9 athers subjects (all obese) who had body weights higher than 140 per cent ideal body weight, and all except 1 had had peak plasma insulin concentration inligher than 80 µunits/ml. We chose 80 µunits/ml instead of 100 µunits/ml, because usually there is a lower peak plasma insulin concentration inligher than 80 µunits/ml instead of 100 µunits/ml, because usually there is a lower peak plasma insulin value for ity GTT compared to orat GTT, either in obese or in normal control subjects (Fig. 238).

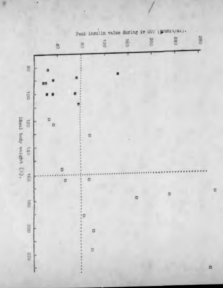
2. KG values

There was no statistical correlation between KG value during to GTI and % ideal body weight. However, the lower values of KG were usually related to higher values of K ideal body weight. If, here again, we take 160 per cent ideal body weight as a dividing line, 15 subjects (10 normal and 5 above) who had their weights less than 160

Figure 23 A. Correlation between peak plassa insulin value during oral GFT and
% ideal body weight in the obese patients (D) and normal control
subjects (.).



Please 1 B. Correlation between peak plasma insulin value during iv OTT and % ideal body weight is the obese patients (D) and normal control mubicots (D).



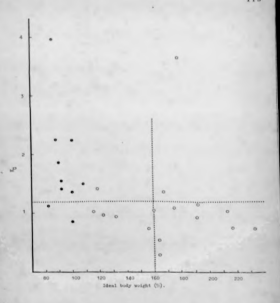
per can't their ideal body weight, 9 (8 normal and 1 obess) had KG values higher then 1.20 %/ain and only 6, (2 normal and 4 obess) had KG values lower than 1.20 %/min, 5 of the obess subjects who had weights less than 160 per can't ideal body weight, enly 1 had KG value higher than 1.20 %/min, as in normal control subjects. And emong 11 subjects who had their weights equal or higher than 160 per cent their ideal body weight (all obess), all except 2 had KG values lower than 1.20 %/min. (Fig. 24).

4. Kg + L values

There was no statistical correlation between $K_{G-\theta-1}$ value during in GTT and % ideal body weight. However, the lower values of $K_{G-\theta-1}$ were usually related to higher values of % ideal body weight. If, here also, we take 160 per cent of ideal body weight as a dividing line, 15 subjects (10 normal and 5 obese) who had weights less than 160 per cent their ideal body weight, 10 (8 normal and 2 obese) had $K_{G-\theta-1}$ values higher and only 5 (2 normal and 3 obese) had $K_{G-\theta-1}$ values lower than 3.20%/min. However, only 2 out of 5 obese subjects who had weights less than 160 per cent ideal body weight had $K_{G-\theta-1}$ values higher than 3.20%/min as in normal control subjects. 11 subjects had their weight higher than 160 per cent their ideal body weight (all obese). All except 3 had $K_{G-\theta-1}$ values less than 3.20%/min. (Fig. 25).

D. The Effect of Dietary Restriction on Weight Loss in the Obese Patients:

As mentioned above, dietery management of patients was related firstly to the need of the patients and secondly to those of the investigations a. the investigation of protein tumover by Professor J.C. Waterlow, Dr. P. Sender, Dr. Susan Ell. Pieure 24. Cerrelation between a value and % ideal body weight in the obess patients (O) and normal central debjects (•).



<u>Pigure 25</u>. Correlation between K_{G+I} value and % ideal body weight in the obese patients (\odot) and normal control subjects (\bullet).

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.(%) suggested foot leads

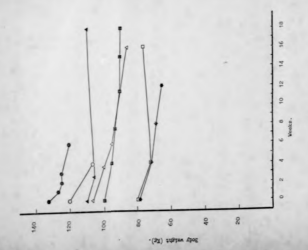
Figure 26. Weight loss during distary restriction in some obese patients.

(■) NC. (■) GG.

(O) AR. (D) CV.

(A) DR. (+) A.

(A) KY.



Dr. P. Garlick and Dr. G. Clugston (using realescrive tyresine and leucine), and for this reason drug therapy was not usually amplayed except for a very small number who received thyronine; b. our study of gluces homeosteris.

There was no statistical correlation between the weight loss in the patients (kg/day) and the type of diet during dietary restriction. There was also no statistical correlation between weight loss and KG or KG + | values. There was also no statistical correlation between weight loss and fasting values of plasma insulin or peak plasma insulin during oral GTT and iv QTT (Table 17).

Fig. 26 shows body weight changes during dietary treatment in some parlants showing different patterns of weight loss.

The effect of various dietar restrictions in relation to K. . . and plasma issuits values in more cheen patients. The patients surburs are the care as those given in Table 11.

Patients	Age (years)	Kg, K		MJ/day	INITIAL Duration (days)	DIET Weight loss (Kg/day)		Duration (days)	E DIRT Weight loss (Kg/day)	INITI:	% ideal weight*		LIN (puni Peak in oral GTT	
23. KY	37	3.65	5.35	1.3	26	0.27	4.2	90	0.14	107	178	26		175
24. CW	53	1	4.20	1.3	33	0.21	2.1	33	0.11	80	119	15		28
25. AD	68	0.97	2.56	1.3	27	0,22	2.1	73	-0.05	79	123	15		34
26. AR	45	0.00		1.3	30	0.43	ILEO-	JEJUNAL B	Y-PASS	120	164	24		55
27. VR	53		2.31	1.3	14	1.14	2.1	17	0.18	96	156	15		48
14. MC	29		2.34	1.3	18	0.44	4.2	18	0.22	133	192	25	193	86
15. DR	33	1116	2.00	1.3	10	0.40	4.2	153	-0.03	110	175	46	241	308
17. MR	17	100	3.01	2.1	13	0.46			ght again	90	160	Not :	easured	
18. MB	47	100	2.04	2.1	30	0.37	2.1	fter dis	0.13	138	217	36	144	100
19. BC	42	1000	2.67	2.1	20	0.20	2.5	30	0	86	116	Not :	easured	
20. FH	22	100	3.75	2.1	30	0.23				75	132	13	118	96
28. GG	51	1	0.97	3.3	30	0				100	164	7		94
29. PR	31		3.38	3.3	30	0				123	191	Not :	easured	
16. HG	21	100	5.68	3.3	20	0				135	213	15	107	102
6. DB		Not me		120	60	0.07	3.3	45	0.04	105	156	14	57	

CHAPTER IV

The results show the familiar findings of an impairment in glucose telerance in abesity (Paulin and Saul, 1922; Back et. al. 1966; Parley and Kipnis, 1966; Chiles and Taggaurnis, 1970, Rabinowitz, 1970), and that this impairment is vauelly accompanied by hyperinsulingemia.

A decrease in insulin sensitivity in respect to cerbohydrote metabolism was present in these patients as shown by high blood glucose in the presence of high plasmo insulin concentrations during the oral GTT (Fig. 18A; Fig. 188 and Table 13). Similarly, the Iv GTT in the obese patients showed that lower K_G values were usually related to higher insulin concentrations (Fig. 19A and 1981.

During tv G1TT, the obese patients thoward a significantly lower mean value of KG + \parallel than did the normal controls (p < 0.01°, (Table 15). The value of KG + \parallel is a measure of insulin sentitivity (in respect to carbohydrate metabolism), (Franckson et. al. 1966; Heard and Henry, 1969 a). Glucose tuterance (KG value) was significantly correlated to insulin sensitivity (p < 0.001) but not to the actual concentration of plause insulin (Fig. 21, Fig. 19A and Fig. 19B). These results confirm similar findings reported in melinourished dogs on the significance of insulin sensitivity rather than insulin concentration in determining glucose tolerance (Turner, 1966; Heard and Turner, 1967; Heard and Henry, 1969 a).

'Insulin sensitivity' or 'Insulin resistence' are terms which have usually been used in the context of cerbohydrate metabolism, (Franciscon et. al. 1966; Heard and Turner, 1967; Heard and Marry, 1969 at.

Insulin resistance in respect to fat matabolism probably also occurred in the obese potients. Most reports show that obese subjects have slightly elevated plasma NEFA. In the present unail series, however, there were no significant differences between the obese potients and control subjects in the values of festing plasma NEFA (Fig. 18C), not in the fall of plasma NEFA concentrations during oral GTT (Fig. 18C) and Iv GTT (Kg; Table 15). In all cases, however, the observed 'normal' values of NEFA in the observable to the subjects were accompanied by higher concentrations of plasma insulin (Fig. 188), implying an overest resistance of fat metabolism to insulin.

Insulin inhibits (Ipolysis (Kipnis, 1965)), on the other hand, insulin promoted glucose uptake and Ilipogenesis (Jennianud and Renold, 1969). Either will result in the fell in pleame NEFA concentration. Only a very low level of insulin is needed to inhibit ilipolysis, and ance this level has been exceeded (Kipnis, 1965) any further effect of Insulin on NEFA levels would be expected to operate via enhanced glucose uptake and resetrification of NEFA. The insulin resistance reflected in NEFA: insulin resistanchips might therefore be in large part another measure of insulin resistance in respect to glucose materialism. Kipnis (1965) also showed that growth hormone administration resulted in an apparent dissociation between muscle and adjoore tissue in response to insulin, and he discussed the question of whether this is due to different argen sensitivity to insulin and different responses of glucose uptake and Inhibition of Ilipolysis.

Peasons, Malkiejohn, Dewer and Thow (1955) observed that weight gain in thin young man was less than expected when they were subjected to an overfeeding for 14 days. Whereas overfeeding in two fet young women resulted in a greater weight gain than that which was found in these thin man (Passmare, Swinndal and El Din, 1963). Not only are obses people still susceptible to increased energy in the same way as thin people do. It is interesting to note that Rolly (1921), more than Rity years ago showed that the specific dynamic action of food stuff was reduced to practically zero after the development of obesity. "Specific dynamic action" is the term used to describe the affect of food in relsing the metabolic rate above the value when firsting.

All obese patients must at some time out in excess of finely requirements in order to tig the energy bolance and put on weight. This early stage could be described as an "active" obssity, and reflects a "normal" subject who ears to excess. He early, therefore, still have a normal insulin sensitivity (in respect of cerbohydrate and fet). "Active" obssity is represented here by the similarity between controls and some obssessablects under 160 per cent ideal body weight in the values of fasting insulin (Fig. 228) and peak insulin during arel GTT (Fig. 23A) and it GTT (Fig. 22B). Some of these patients, however, had started to heve higher values of fasting insulin, peak insulin during onel GTT and it GTT. Interestingly, most of them had elready shown a decrease in its glucose tolerance (KG, Fig. 24) and insulin sensitivity (KG + (r. Fig. 25). This group may represent a transition period before they go into a further stage, i.e. "pastive" obesity. "Passive" obesity could be represented here by

the obase subjects above 140 per cant ideal body weight showing predominant pictures of high fasting insulin concentrations (Fig. 228), high peak insulin during oral GTT (Fig. 23A) and Iv GTT (Fig. 238), low $K_{\widehat{G}}$ values (Fig. 24), and low insulin ansitivity ($K_{\widehat{G}} + 1$) (Fig. 25'.

Some apparently normal* subjects showed very high values for K_G and $K_G+\{(Figs. 21, 24 \text{ and } 25).$ It is speculated that these subjects may be in an early phase of 'activa' obesity. This speculation is based on the similar pattern of changes in K_G and K_{G+} found in experimental animals fed diets deficient in protein (Heard and Turner, 1967). Heard and Stewart, 1971). In all cases, early signs of hypersensitivity to insulin were later followed by an impatrment of glucose tolerance and diminished insulin sensitivity. However, such speculation requires further investigation.

Our results are in tine with the view that 'gorging' should be avoided, aspecially if the dist is rich in carbohydrate. 'Gorging' leads to a greater increase in :;pagenesis (Gwinup, Byran, Routh, Kruger and Hanwi, 1973; Bray, 1972). This affect would be accentuated in those who are already obese and in whom any carbohydrate intake (oral or iv) leads to vary high levels of plasma insulin (Yable 12, Figs. 23A, and 239).

It is very difficult to assess the affect of dietery restriction on body weight in these patients. It depends more on the will power to stick to the diet and the mental attitude of the individual. Some followed the diet restriction properly, some did not, resulting in an inability to show any weight loss. As the patients primarily came for treatment, the type of diet may, therefore, differ from subject to subject.

None of the blochemical parameters measured in this work showed any gradictive value in respect of effectiveness of dietary therapy.

This may seem surprising II obesity is to be regarded not as a 'discose', but rather as a 'glift'. The genetic development of obesity is not at all well understood in man. The genotype for obesity was assumed to be present since early human population (Montegu, 1966). To encient people in the poleolithic times, or perhaps even at present, in creas where food could comestimes be scarce, the ability to become obese, would have provided greater reserves of food supply upon which to draw in times of scarcity.

PART IV

GLUCOSE HOMEOSTASIS

IN

OLD AGE

CHAPTER I

A. Ageing Process:

In the past century, man's life expectancy has increased considerably (Department of Health and Social Security, England, 1976), through the advencing knowledge of modern medicine. This has been in two main areas: In preventive medicine, through improvement of the environmental sentration, better management not only in maternal and child health care, but also the health of the population in general; and in the field of curetive medicine, especially following the discovery of entitlations. While the medical successes are considered a good thing, the results expose us to different kinds of new problems. The increasing number of old people in the community emphasizes the importance of seeking batter knowledge of againg processes, and better entities in caring for and understanding old people.

The word 'againg' means to 'grow old' (Oxford Conclus English Dictionary).

However, in biological studies, this term could have many different connotations.

Bellomy (1970) gave a definition of againg as a "decline in the ability of the homeostatic system of the body to cope with fluctuations in the external world". In the process of growing old, there is a steady decline of physiological functions. In conjunction with a decline in anatomical, blochemical and hormonal features and performance (Narris and Shock, 1966). A decrease in physical activity with an without painful feelings in the hones and joints, a decrease in food intake, emotional disturbances and many other physical or psychological inadequacies show that old age and stress are closely associated. Old age, therefore, is not merely the pseuing of

years, but many malfunctions added together become a generalized picture of old age, and, whatever the original cause, some elements of stress either physical or psycholealcal or both are always present in old age (see Part I, Chapter IV, Stress).

8. Ageing and Carbohydrate Metabolism

Increasing age has always been especiated with a decline in glucose tolerance, either oral GTT (Brent, 1960). Burch and O'Meelly, 1967). Chiauverakis, Jarrett and Keen, 1967), or the Iv GTT (Streeten, Gentein, Mermar and Doisy, 1965). Franckson et. al. 1966; Cackford, Harbeck and Williams, 1966). The decline in Iv glucose tolerance is associated with a decrease in insulin production (Cockford et. al. 1966). However, in the oral GTT, some have claimed that the prolonged hygietylycamia resulted in higher levels of plasma insulin (Chiauverakis et. al. 1967). Andres, Poxefsky, Swerdloff and Tobin, 1970).

It is important to try to decide whether we should apply the same criteria in diagnosing and treating diabetes in the alderly as are used for young adults. Most authorities would suggest making "allowences" for age (WHO Expert Committee on diabetes mellitus, 1965; Andres, 1973). This seems reasonable if any because elderly people probably have diminished food intake and sametimes have impaired absorptive capacity (Webster and Leaming, 1975), and, therefore, do not, under normal circumstances, load their systems to the same extent as young people. In telemona tests on the other hand, the same load is given to young end old allike.

Andres (1974), in his review claimed that impaired glucose tolerance (Intravenous test) in the elderly was due, not to impaired smallfully to insulin, but to deficient insulin production. However, he also acknowledged that there was not yet any clear consensus botween various workers about plasma insulin levels during the oral GTT.

In all the other structions we have studied, insulin sensitivity played as large a part (if not more) in determining glucose talerance as did insulin levels.

Therefore, we sought to differentiate between insulin levels and insulin effectiveness in the olderly.

Because of the gastro-intestinal treat changes in old people (Agate, 1963), including possible impairment in intestinal insulinogenic capacity, and also the inconclusive reports on insulin response during oral GTT, it was also desirable to compare the effects of oral and by glucose loads on blood glucose and plasme insulin fevels.

The decrease in glucose tolerance in old age could be due either to the decreased uptake of glucose by the liver and peripheral tissues, or due to the hability of insulin to stop liver gluconeogenests. Therefore, we also compared the gluconeogenic capacity of the liver between the geriatric patients and normal control subjects. This was carried out by introvenous injection of 1-alenine (Wise, Handler and Fallg, 1973). Mestyan, Schultz and Harvath, 1974) and measuring the rise in blood glucose concentration, the changes in plasma insulin and glucogon, and the disappearance of injected 1-alenine from the circulation. The % removal rate of injected 1-alenine (K_A) was calculated the same way as those calculations for K_G and K_{G+1} . (See Part III, Chapter II, Methods).

C. Intravenous I-glanine Tolerance Tast and a Measure of Glucaneogenesis:

Gluconeogenesis is very difficult to measure. A number of different "tolerence" tests have been used in which substrates capable of serving as glucoss precurent wave been administered. These include dihydroxyecotone or glycarol (Femendes, Koster, Gross and Sorgedroger, 1974), fructose (Pogliare, Kerl, Keeting, Brown and Kipnis, 1972), pyruvate (Moorhouse, 1964), Lectate (Kreibeeg, Penninghon and Boshell, 1970) and alanine (Felig, Mariliss, Owen and Cohill, 1969). Whis et. al. 1972; Mastyfin et. al. 1974). Alanine is of special importance. It has been suggested that alanine uptake by the liveramounts to 50 per cent of the total amino acid uptake by this organ (Felig, Wehren, 1971). This is accounted for by the fact that alanine is also released by the muscles in amounts for exceeding the alanine content of muscle protein. Protein contents only 5 to 7 per cent alanine (Kominz, Hough, Symond and Laki, 1754), yet alanine accounts for about 30 per cent of total amino acids released from muscle (Londen, Feley and Webb, 1963).

This production of alanine requires pyruvate to serve as an amino group acceptor in alanine amino transferase reactions. Pyruvate is derived from glucose by glycolysis in muscle. In the liver, pyruvate is regenerated from alanine, and then serves as a gluconeogenic precursor to replanish the glucose removed from the blood by muscle and other tissues, and so complates the 'glucose-alanine cycle' (Felig, 1973). The success of this process, therefore, depends upon the availability of glucose precursors and the afficiency of the liver in converting alanine into glucose.

Alasina is known to promote both gluconsogenesis and glycogenalysis through stimulation of glucogen production (Wiss, Hendler and Felig, 1972). Thus, the rise in blood glucose cencentrations after introversus alanine administration connot be attributed to gluconeogenesis alone, unless liver glycogen stores have been deplated by festing. Ideally, the subject should be lested 24 to 48 hours to be sure of more at less complete exhaustion of liver glycogen stores (Hullman and Nilsson, 1971). However, such a procedure was not possible on geriatric patients. With this limitation taken this occount, geriatric patients and normal control subjects were given an intravenous elegine test after fasting for 8 hours from midnight.

CHAPTER II

A. Subjects:

1. Butfante:

Patients from the geriatric wards, St. Pancres Hospital, were studied. They were asked "-thes they would be willing to consider participating and the kinds of tests involved were explained in detail. Consent (oral or written) was thus obtained from each patient and control subject participating in the study. These patients were presented to us as non-diabetics, and any subjects who had a festing blood glucoss ascentration higher than 6.11 m mai/1 (110 mg/100 ml) (working party, College of General Practitioners, 1963) were excluded. Their medical records showed that none of these patients had any glycosuria, and all received normal hospital diet.

The first three tests (and GTT, Iv GTT and Iv GTT) were carried out on 23 patients (Table 18). Their mean age was 79 ± Ll years, their mean body weight was 56 ± 2.2 kg. For three to four days before the tests, each of them received an extre 150 g cart-chydrate/day. This was to ansure that all the subjects had a reasonable carbohydrate Intoke before the tests were done. This extre carbohydrate was in the torm of 150 g/600 ml solution, consisting of 50 per cent orange juice and 50 per cent.

Calorsen (Scientific Hospitel Supplies Ltd., England).

Unfortunately, some of the patients were not able to participate in the fourth

Table 18

List of gariatele patients who participated in oral GTT, Iv GTT and Iv GITT

Subject	Sax	(years)	Rody weight (kg)	Previous diseases
GV	F	88	50	Gangrene of the toes,
WT	F	87	51	Myocardial Infarct
СН	м	74	69	Ajherosclerosis
OV	M	83	69	Myocardial Infaret
VL.	M	83	60	Left hemiplogia
CR	M	74	73	Prostatectomy
ED	F	77	71	Kyphosis
EL	F	18	50	Rheumatold arthritis
PS	F	77	60	Myocardial Infarct
SH	F	76	50	Mitral stanosis
FD	M	73	62	Myocardial inferct
BJ	F	78	57	C.V.A., tight hemiplegia
BX	F	82	43	Demantia
OT	M	79	37	Dementie
FL	F	82	47	C.V.A.
MR	M	78	67	C.V.A., paraplegia
EN	м	86	61	Prostate hypertrophy
MG	F	74	62	Carvical spandylitis
MC	м	78	55	C.V.A., right hemiplagia
PG	м	77	53	Alld left ventricular fallure
DW	F	73	67	Dermoid tumour in the paivis
PN	F	78	34	Carcinoma cervix
ST	м	68	49	C.V.A., right hamiplagia

unable in do so because of their clinical conditions. Therefore, a few more patients had to be added to make up a reasonable number. Fig. 19 shows the first of 8 gertetric patients who had the 1v alonine tolerance test. Their mean age was 78.2.1, 6 years and their mean body weight was 52.2.3, $9 \, kg$.

Due to the advanced age of all gerietric patients studied, we were unable to get a reliable measurement of their heights, thus also unable to calculate their % ideal body weights, and these measurements were not included in this report.

2. Normal subjects:

Normal control subjects for the anal GTT, Iv GTT and Iv GTTT, were the seme 14 young, non-abose subjects used as controls in the study on abosity (see Part III, Chapter II, Normal Subjects), (Table 12).

As with the patients, so also with the control subjects; some of the original volunteers were unwilling as not available to participate in further tests. More new volunteers were, therefore, recruited to make a reasonable number of control subjects for the IV alanine tolerance test Table 20 shows the list of B control subjects for the IV alanine tolerance test. Their mean age was 28 ± 1.5 years, their mean height was 172 ± 3.5 cm, their mean weight was 63 ± 4.2 kg, and their mean %

Table 19
List of gerlatric patients who participated in the intravenous alamine televince test

Subject	Seese	(years)	Body weight
JG	м	78	67
FL	F	82	47
EL	F	gi	46
CL	F	62	41
AD	F	77	56
HE	M	71	63
BJ	F	78	57
JL	F	71	35

Table 20

List of control subjects who participated in the intravenous signing tolerance test

Subject	Sex	Age (perca)	Halght (cm)	Weight (kg)	% Ideal weight
AT	м	34	181	76	94
SM	F	32	1 69	58	90
RA	M	31	170	70	97
А.	м	27	188	83	99
WS	м	32	1.59	55	86
50	F	24	165	54	95
VP	м	23	178	74	102
ON	F	24	162	52	95

^{*}Compared with the data from Gelgy Scientific Tables (1970)

ideal body weight was 95 ± 1.8%.

8. The Tests

1. Oral GTI, iv GTI and iv GITTs

The methods and procedures used in these tests, the calculation of the average glucose ar insulin concentrations and insulin: glucose malar ratio during onal GTI; the calculation of K_G and K_F in the Iv GTT, and K_{G-b-1} in Iv GTT, have all been described (see Part III, Chapter II, Methods).

Although all garietric patients showed low values of K_G and K_{G-d-1} , a few of the first patients tested showed hypoglycoemic signs a few minutes after the end of the iv GUT. For that reason, all later patients received an extra 20-30 mL of 50 per cent glucose solution introvenously at the end of the iv GUT, and the number 300 per cent glucose solution introvenously at the end of the iv GUT, and the number 300 per cent glucose solution introvenously at the end of the iv GUT, and the number 300 per cent glucose solution introvenously at the end of the iv GUT, and the number 300 per cent glucose solution in 300 per cent glucose solution.

2. Intravenous alanine talerance test (Iv ATT):

This test was devised as a measure of the gluconeogenic capacity of the liver.

The subjects were fasted from midnight and the test was carried out in the marning.

A butterfly needle attached to an antecubital value of one arm was used for drawing blood sample. Two fasting blood samples (10 minutes apart) were taken at about 9.30 a.m. Through the same needle, a solution of 10 per cent 1-alanine (0.15 g/kg body weight) (Wise et. al. 1973) was injected. The atlanine solution (see section 8.3. below) was injected within 5 minutes, and serial blood samples through the same needle, were taken at 51, 101, 201, 301, 601 901 and 1201 after

the injection. The process of for handling blood complex was similar to those in the arel GTT, Iv GTT or Iv GTT. Except that for glucegon estimation, that 4.5 ml of blood was also taken and added to another haparin tube containing 0.5 ml cold solution of Trasylal (10,000 KIU/ml, Bayer) and after immediate centrifugation the supernatort was frazen rapidly in solid CO₂ and stored at -20° C until the day of analysis.

Although name of the subjects complained of having any clinical complications from this test, a few of them however, experienced a slight stomach discomfort for several minutes following the injection of alanine.

3. 10% I- alanina solution:

Alanine solutions were made by distaining sterile I-elanine (Kabi-Vitrum Pharmacceutical, Sweden) in sterile distilled water as a 10% solution. This solution was buffered to a ph between 7.0 and 7.3 with sodium hydroxide. It was then passed through a millipare filter and autoclaved at 120° C for 10 minutes (Mestyan et. al., 1974).

This elanine solution was made by the Department of Pharmacy, University Callage Hospital. We exempted to make our own solutions, but the Hospital refused to allow us to do the test unless the solution was made by the Hospital's own Pharmacy and from time to time the quality control of the solution was checked.

C. Analytical Methods:

Analysis for blood or plasma glucase, plasma NEFA, plasma insulin and plasma glucagen were carried aut according to the procedures reported in the previous study (see Part II, Chapter II, Mathods'. Plasma clanine was analysed by an enzymatic

method, using 1-alanine dehydrogeness (Reilly, 1975) with reagents supplied by the Bookinger Corporation, London Ltd.

D. Assessment of the Date:

Student's t test was used for comparing the potients data with those of the controls.

CHAPTER III

RESULTS

A. Onl Gil:

The oral GT1 was carried out in 23 garietric patients and in 13 normal control subjects (Tables 12 and 23).

1. Blood glucose:

The mean blood glucose concentration was 4.92 ± 0.11 m mai/1 (88, 6 ±2.02 mg/100 mg) in the partiatric patients against 4.62 ± 0.16 m mai/1 (80, 2 ±2.92 mg/100 ml) in control subjects (Toble 21, Fig. 27). The difference was not eignificant (p > 0.35), but during the test the concentrations of blood glucose in the patients continued to rise after 30° and the peak was at 90°, whereas the peak in the controls was at 30°. Blood glucose concentrations in the patients were higher than in control subjects throughout the rest of the test. And the difference reached significant levels after 60° (p < 0.01, and from 90° p < 0.001) (Fig. 27A).

The average value for blood glucose concentration during the onal GTT was 7.72 \pm 0.28 m moi/1 (139.1 \pm 5.05 mg/100 mill in the gariantic settlents against 5.71 \pm 0.18 m moi/1 (102.8 \pm 3.25 mg/100 mill in control subjects. The difference was significent (p < 0.01) (Table 21).

2. Plasma Insulin:

The mean testing plasma insulin concentration was 12.3 \pm 0.54 gaunit/mi in the parietric patients against 12.8 \pm 1.31 μ unit/mi in control subjects. The difference was not significent (p> 0.05) (Table 21). During the test the concentrations

- Oral OTT im 25 geriatric patients (O) and 13 normal control mabjects ().
 - Mean blood glucose values during oral GTT.

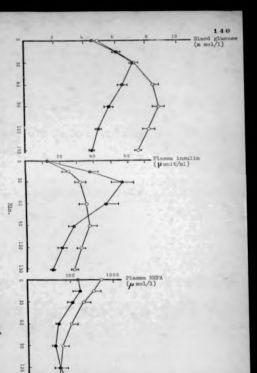
 There were significant differences in the mean values at 60° (p < 0.01), and at 90°, 120° and 150° (p < 0.001) between the two groeps.
 - Mean plasma insulin values during oral GFF,

 There were mignificant differences in the mean values at 30'

 (p < 0.01), at 120' (p < 0.05) and at 150' (p < 0.01) between these
 two groups.
 - Mean plasma NKFA values during ovel OTT.

 Except for the mean fasting levels (p < 0.05), the differences in
 the mean levels at various times during the tast between the two
 groups were not significant.

 (mean values of 12 swrittic and 7 mormal control subjects).



as plasma insulin in the generative patients did not rise as high as in the control subjects, but continued to rise after 30°, and the peak was at 90°. Whereas the peak in the centrols was at 30°. Plasma insulin concentrations in the patients were lewer than in centrol subjects, up to 60° (significant difference at 30°, $p \le 0.01$). But, while the insulin concentration after 60° decreased in the control subjects, the insulin concentration in the patients remained high, and was significantly greater than in the controls at 120° and 150° ($p \le 0.05$ and $p \le 0.001$ respectively). (Fig. 278).

The average value for plasma insulin concentration during and GTT was $31.6 \pm 2.52 \,\mu$ unit/mi in the gerietric patients against $32.5 \pm 2.51 \,\mu$ unit/mi in control subjects. The difference was not significent (p > 0.05) (Table 21).

As also in the study of obesity (Part III, Chapter II, Patients) any overt diabetics (fasting blood glucose higher than 6.11 m mol/1) were excluded during the initial selection of patients. WHO (1965) further gave as a criterion for a "narmal" oral GTT, that the 120' blood glucose value should be less than 6.11 m mol/1 (110 mg/ 100 ml) and gave the lower limit for diabetes as a 120' value of 7.22 m mol/1 (130 mg/ 100 ml). Having regard to the advanced age (mean 79.2.1,1 years) of these patients we arbitrarily chose 7.77 m mol/1 (140 mg/ 100 ml) for an upper limit for "non-diabetic" geriatric subjects. According to file new criterion, among 23 geriatric patients studied, 15 could be classified as "diabetic" and 8 as "non-diabetic" geriatrics (Table 22).

Fortuitously, the dividing line of 7.77 m mol/1 corresponded closely with the mean

The mean all of festing blood glucoss and plasms insulin, average concentrations of blood glucose and plasms insulin, and insulin glucose mater ratio during and GTT, in gerletric patients and control subjects (\$ \$EM). Number of observations in parenthesis.

	Mean feating	cancentrations	Menn of I	glucose moler ratio		
	Blood clucose Masma insulin		Blood glucoss	Plane	a Insulin	
	(m mgl/11	(µ unit/ml)	(m mol/1)	(µunit/mi)	(10 ⁶ x m mol/1)	
Patients (23)	4,92 2 0.11	12,3 ± 0.56	7.72 2 0.28	31.6 ± 2.52 ns	227.0 ± 18.1	29.8 ± 2.27
Cantrols (13)	4,62 2 0.16	12,8 2 1,31	5.71 ± 0.18	33,5 2 3,51	240.4 ± 25.2	42.2 ± 4.50

100 - more levella-

^{*} p < 0.05 ** p < 0.01 ns = not significant

120' blood alucose value for all 23 patients (8.12 m mg//1).

3. Insulingfucose malar ratios

The mean insulingluces moler ratio in the geriatric patients was $(29.8 \pm 2.27) \times 10^{-6}$ against $(42.2 \pm 4.50) \times 10^{-6}$ in control subjects. The difference was significant (a = 0.01) (Table 21).

4. Plasma NEFA:

There was a great range in the levels of plasma NEFA both in the geriatric patients and in control subjects. This applied to the fasting concentrations as well as to the concentrations during the test. The mean festing plasma NEFA concentration in the geriatric patients was 866 $^{\frac{1}{2}}$ 86 μ mol/1 against 990 $^{\frac{1}{2}}$ 68 μ mol/1 in control subjects. The difference was significant (p <0.05). Although the plasma insulin concentrations during the arei GIT were lower in the geriatric patients, there were no significant differences in the mean NEFA concentrations at any time during the test between the geriatric patients and the normal controls. The NEFA curve during the arei GIT in the patients mitrored their singgish insulin curve, showing a daleyed and prolonged reaction. Thus at 150° the mean NEFA concentration was lower in the patients than in the controls, since in the latter the NEFA concentration had started to rise again (Fig. 27C).

8. Iv GTT and Iv GITT:

The Iv GTT and Iv GITT were carried out in 23 geriatric patients and in 10 normal controls (Tables 23 and 12).

1. Kg values:

The mean KG value in the periatric patients was 0.98 # 0.09 %/min against

Mean blood glucose and plasma insulin concentrations during and GTT in the "diabetic" and "non-diabetic" garietrics and in normal cantral subjects (2 SEM). Number of observations in parenthesis.

Oal GIT

			Ma	as blan	d alucas	e concil	n tration	ıs		Med	n plasn	us insulin	concen	ra Hone	
			Meen blood glucase concentrations (m mel/1)					(µ unit/ml)							
		0-	15'	304	60°	901	1201	1.50*	01	1.9	301	604	90"	120°	150'
1	'Diabetic' gerietrics (15)	5.07 ± 0.15	6.30 0.22	7.66	9.24	9,74 ± 0.52	9.07 ± 0.46	0.43	12.3 ± 0.72	21.5 2 2.27	31.9 2 4.64	37.1 2 5.28	35,7 2 4,03	34.1 2 4.01	28.8 1 2.46
	Licil					**	***	in.							
П	'Non-dishetic' geriatrics (8)	4,63 1 0.08	5.77	6,43	7.08	7.04 0.37	6.35	5.78	0.82	31,4 2 10,18	12.0 2 4.92	32.1 2 5,37	40.0 ± 11.60	28.1 2 2.45	25.9 2 3.96
	1 x 11) 1(x 11)				***	***	***	***		**				*	***
н	Control subjects	4.62 ± 0.16	6.15	7.19 0.30	6.40	5.50 2 0.29	4.81 2 0.20	4,36 2 0,19	12,8 2 1,31	37.3 ± 5.39	56.3 1 6.82	47.1 \$ 6.97	27.8 ± 3.65	20.6 2 2.94	14.8 2 1.34

⁺p<0.05 ** p<0.01 *** p<0.001

1.81 \pm 0.28%/min in the control subjects. The difference was significant (p <0.001) (Table 23),

There was no statistical correlation between K_G and festing plasma insulinvalues during Iv GTT in the combined group of garletric pateInts and the normal controls (Fig. 28A).

There was also no correlation between K_G and peak plasma insulin values, in the geriatric potents and control subjects (Fig. 281). However, a higher K_G was usually related to a lower peak plasma insulin concentration.

2. Peak Insulin concentrations during one GTT and in GTTs

There were 23 gertatric and 9 normal control subjects who had both oral and Iv GTT done.

The mean peak insulin concentrations during the oral CTT were higher than the mean peak insulin concentrations during to CTT both in the geriatric patients and in the control subjects. The differences, however, were not significent (p > 0.05) (Table 24). Whereas in the control subjects the peak insulin concentrations during the oral GTT were always higher than the peak during the IV GTT, in the geriatric patients however, B out of 23 patients had peak insulin concentrations during the IV GTT higher than the peak during the oral GTT. This suggested that in these 8 geriatric patients there was some degree of small intestinal malabsorption. The mean peak insulin concentration during the oral test in the geriatric patients was 48.0 \pm 5.34 μ unit/ml against 72.0 \pm 7.26 μ unit/ml in the control subjects. The difference was significant (p <0.05) (Table 24).

Table 23

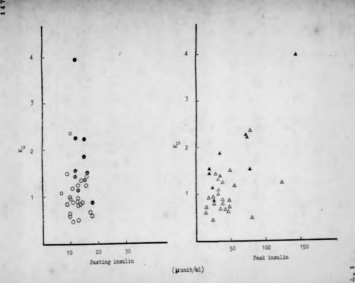
The mean values for KG, KG + μ , Kc in the switchis patients and in normal control subjects (2 SEM). Number of observations in parenthesis.

Meon Volues

	KG	KG +1	Kp	
	(%/mln)	(%/min)	(%/win	
Geriatric patients	0.98 20.09	2.29 ± 0.16 (23)	2.21 ± 0_34 (10)	
	***	***	76	
Control subjects	1,81 = 0.28	4.55 ± 0.51 (10)	3.10 ± 0.73 (6)	

transfingle ten = an | 100.0 > a ***

- Figures 28 A. Correlation K_Q and fasting plasma insulin values during iv GFT in the geriatric patients () and normal control subjects (◆).
 - 28 B. Correlation between K_G and peak plasms insulin values during iv OFF in the geriatric patients (A) and normal control subjects (A).



The mean peak insulin concentration during tv GTT in the gariatric patients was 41.6 * 5.13 μ unit/ml against 54.0 \pm 13.60 μ unit/ml in the control subjects. This difference was not significant (p>0.05) (Table 24).

3. EG+T values:

The mean $E_{\rm OST}$ value in the geriatric patients was 2.29 \pm 0.46%/mimte against 4.55 \pm 0.51 in the control subjects. The difference was significant (p.(0.001) (Table 25).

There was no correlation between the and fasting plasma insulin values in the geriatric patients nor in control subjects (Fig. 29 A).

There was also no correlation between and peak plasma insulin values during iv OTI in the geriatric patients nor in control subjects (Fig. 29 B).

There was a highly significant correlation between x_i and x_i values in the combined geniatric patients and normal control subjects. (x = 0.71, p < 0.001) (Fig. -30).

4. Ky values :

The seas $\frac{1}{2}$ value in the geriatric patients was 2.21 \pm 0.34 %minute against 3.10 \pm 0.75 %minute in the normal control subjects. The difference was not significant (Table 25)

Table 24

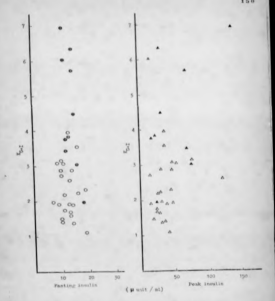
The mean peak insulin concentrations during are IGTT \equiv iv GTT (2 -LM), geriatric patients and normal control subjects. Number of observations in parenthesis.

Mem Peak Insulin Concentrations

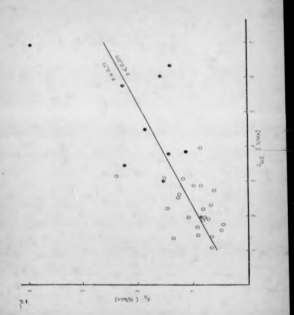
	Oral CTT	Iv GTT	Ratio of oral; ly
Gariatric parlants (23)	48.0 ² 5.34 ms	41.6 ± 5,13	1,15
	*	166	
Controls (9)	72,037,26 19	34.0 = 12.40	1,50

^{*} p < 0.05; ns = net significant

- Flowers 29 A. Correlation between \mathbb{K}_{U+1} and fasting places insulin values during iv OTF in the gariatric patients (O) and normal control subjects (O).
 - 23.2. Correlation between E_{C+1} and peak planes insulin values during iv GTP in the geriatric patients (A) and normal sectrol subjects (A).



Regression line between the values of $K_{\mathbb{Q}}$ and $K_{\mathbb{Q}+1}$ in the combined group of switching prients (\mathbb{Q}) and normal control subjects (\mathbb{Q}).



C. Iv Alanina Telerance Test (Iv ATT):

1. Blood glucose:

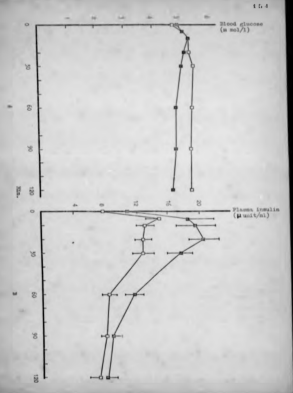
The mean fasting blood glucose concentration was 4.77.2.0.17 m mol/1 (85.9.4.3.02 mg/100 ml) in the garietric patients against 4.92.2.0.07 m mol/1 (86.6.2.1.32 mg/100 ml) in the central subjects (Fig. 31A). The difference was not significent (p > 0.05). During the test in the garietric patients, blood glucose concentration continued to rise after 10° and reached a plateau only at 30°, whereas the blood glucose concentration in the central subjects only rose and reached a pook at 10° before felling again. Blood glucose concentrations in the generatric patients were significantly higher than in the control subjects from 30° onwards (p < 0.01) (Fig. 31A).

2. Plasma Insulin:

The mean fasting plasma insulin concentration was 7.8 \pm 0.86 μ unit/ml in the gestatric patients against 10.9 \pm 1.22 μ unit/ml in the control subjects. The difference was significant (μ < 0.01) (Fig. 318). During the test the levels of plasma insulin in the control subjects continued to rise and the peak was at 30°, whereas the peak in the gestatric patients was at 5°. Throughout the test the plasma insulin concentrations in the control subjects were always higher than in the gestatric patients. The differences were significant at 10° and 20°, (μ < 0.001), at 30°

- Plante 11. Intravenous Alanias tolerance test (0.15 g l-alanias/Kg 36)
 in 8 gariatris patients (0) and 8 normal control subjects
 (a).
 - Mean blood glusces concentration fallowing the iv injection of slanine.

 There was no significant difference in the mean fasting blood glusces was raised and remained elevated throughout the teet. The difference in the mean levels remained elevated throughout the teet. The difference in the mean
 - F. Mean places insulin values at various times during the test. There was a significant difference in the fasting values between the two groups (y < 0.01); at 10° and 20° (y < 0.001), at 30° and 60° (b < 0.01).</p>



and 60' (p < 0.01) (Fig. 318).

3. Plasma alanina:

The mean fasting plasma alanina concentration was $398.2.64~\mu$ mol/l in the geriatric patients against $387.9.62~\mu$ mol/l in the control subjects. The difference was not significant (Table 25).

The peak plasma alanina concentration was 5536 2 362 µ mai/1 in the garletric patients against 5456 2 661 µ mai/1 in the control subjects. Both were at 5' effor an injection of 1-clanina and the difference was not significant (Table 23).

The mean % ramoval rate of clanine (K_A) was 2.93 $^{\pm}$ 0.18 %/min in the generatoric patients against 5.31 $^{\pm}$ 0.71 %/min in the control subjects. The difference was highly significant (p \leq 0.001) (Table 25).

4. Plasma glucagan:

Radialimmune essay of glucagon is carried out in Dr. S.R. Bloom's loboratory, The Rayal Post Graduate Medical School, Landon. At present, they are having difficulty with the essay technique, and at the time of writing this thesis, results of plasma glucagon levels during the livistancine test have not been received. The glucagon results cannot therefore be included in this report, but it is hoped to odd them as a separate appendix as soon as they are available.

Table 25

The mean feeting and peak concentrations and the mean percentage removal rate of plasms atenine $(K_{\widehat{\mathbf{A}}})$ during to atenine tolerance test (2 SEM), in the gerietric patients and in the control subjects. Number of observations in parenthesis.

	Plane o (p re	(%/min)		
	Mean fasting concentration	Mean pack concentration		
		(at 5°)		
Gariatelo patients 4)	398 ± 64	5534 2 362	2.93 # 0.18	
	ni	ns	***	
Control subjects 4)	3g7 ± 62	5456 2 661	5,31 ± 0 71	

p (0.05; ns = not significant

CHAPTER IV

Our results show the familiar findings of an impairment in oral GTT in old age (Srant, 1960; Burch and O'Meally, 1967; Chlauverakis, et. al., 1967; Smith and Hall, 1973! (Fig. 27A). Insulin secretion showed a sluggish response to the glucose food, with a delayed peak at 90! (Fig. 278). The peak of plasma insulin corresponded with the peak at the glucose curve. There was no significent difference in the mean fasting concentration and the mean average concentration of plasma insulin during the oral GTT, between the geriatric patients and normal control subjects. However, the insulin-glucose malar ratio was significantly lower in the geriatric patients (patients) (p. < 0.01) (Table 21).

This work has demonstrated that looking only at a single time for insulin during the anal GTT, e.g. the 60° value as reported by Chlouverakis, et. at. (1967), could be very misleading, since at that stage (60°) insulin concentration in geriatric patients was still rising (peak at 90°) whereas in the control subjects it was in the process of going down (peak at 30°). Therefore, at this time (60°), there was an apparently higher mean plasma insulin value in the geriatric subjects (Fig. 278). Our results are in line with those reported by Smith and Hall (1973), where they also had serial measurements of plasma insulin during the oral GTT.

It is interesting to note that the decrease in the level of plasma NEFA during the oral GTT was not significantly different. between the geniatric patients and normal controls (Fig. 27C), despite the fact that the geniatric patients had a delayed insulin curve. This suggests that insulin levels in the periatric patients were still.

able to prevent NEFA release from adipose three. Plasma NEFA concentrations, like those of plasme insulin also showed a sluggish response to glucose administration in the elderly.

Our results also continued earlier findings that old people have an impairment in Iv GTT (Streeten, et. al., 1965). Franckson et. al., 1965; Cockford, et. al., 1966). There were algoriticant differences in the mean K_G ($p \leqslant 0.001$) and $K_G + 1$ ($p \leqslant 0.001$), but not in the mean KF values, between the generational and the normal controls (Table 23). This way suggest that old age offacts carbohydree more than for metabolism (represented by enti-lipsiytic action of insulin).

The patients had suffered at veriable times before the test from the usual range of clinical conditions found in geriatric wards, e.g. cerdiavascular accident (C.V.A.), myocardial infarct, gangrams, etc. Same still had the results of their previous illnesses such as hemiplegia, poraplegia, kyphosis, etc., at the time of the test (Table 18), However, all were in reasonably 'good health'. Name of these previous illnesses (e.g. myocardial infarct), show any correlation with the results of glucose tolerance. This was seemingly surgrising for isheamic cordiovescular diseases show correlation with imported glucose homeostasts in the 50 to 60 year ald adults (Wohlberg and Thompson, 1968). Possibly in the younger people, law KG values as offer indices of impatred glucose homeostasts may be signs associated with those more prone to heart attacks. But, by the advanced age of the patients in this work, all seem to have reached such a degree of impatrent, that differences due to factors other than age are difficult to distinguish.

Orag treatments received by these patients (It any) were discontinued for at least 12 hours before the test.

The mean peak insulin level during the area GTT was significantly lower (p <0.05) in the gestatric patients compared with normal control subjects. The mean peak insulin level during the IV GTT was also lower in the elderly but not significantly to (p> 0.05) (Table 24). These findings of low peak levels of insulin response to glucose load, appear to support the hypothesis of Andres (1973), who gave grime importance to the decreated # -cell response to glucose stimulus in equitability to the importance to the decreated # -cell response to glucose stimulus in

As in the study in obsolity, the results for geriatric patients show that intuitine exercitivity (K_{G-4}) valual is a more important factor in determining glucose tolerance then the actual plasma insulin concentration. This is shown first by the highly significant correlation between K_G and K_{G-4-1} values in the combined group of geriatric parlants and normal control subjects (r=0.71, p<0.001) (Fig. 30), and secondly by the fact that there was no statistical correlation between K_G and fasting or peak plasma insulin values during the I_F OTT (Figs. 28A and Fig. 28B). Nor did sentitivity to insulin relate to bosel or peak insulin levels during the I_F OTT, since there was no statistical correlation between K_{G-4-1} and fasting or peak insulin values during its overlation between I_{G-4-1} and fasting or peak insulin values during its

Glucose absorption by the small intestine (during and GTT) must also play a part in determining the shape of the curve. Webster and Leasing (1973), using a modified

xylasa falerence lest and the method of Ros and Rice (1948) for determining xylass concentration, showed that 26 per cent of gerlatric subjects showed some degree of malebsorption. If this is true, the delayed entry of glucose into the circulation would add its effect to those of other factors responsible for prolonged hyperglycaemia, and increase its severity (Fig. 27A). He patic glucose production is the most obvious non-dietary condidate for the cause of delayed hyperglycaemia in the oral GTT. It was happed that this could be essessed by I-alanine administration, but in order to separate the happetic from gut factors, alanine had to be given introvenously.

Injection of I-elanine produced a significantly higher concentration of blood glucose in the geriatric patients than in control subjects (Fig. 31A). It seemed that in the geriatric patients, plasma insulin (which was low compared to normal control subjects) was unable to suppress gluconeogenesis by the liver (Fig. 31B). Therefore, the glucose level remained high, whereas in the control subjects it want down after 10°. Unfortunately glucogon values during th's test are not yet available at the time of writing.

Glycogen content in the liver must have been low if not depleted altogether (at least in the geriatries) as shown by low fasting values of blood glucose in both groupt (Fig. 31A), (Hultman and Nilsson, 1971), so that the new glucose must in large part be due to the conversion of clanine rather shan glycogenolysis in the liver.

The fact that fasting must have been reasonably adequate is shown also by the fact that elanine produced, even in control subjects, anly a modest increase in Insulin secretion (Fig. 318). If the glucagon results had been available, they would

probably have shown a marked rise (Unger, 1972). Only if glucose is available does elanine have a marked silmulation effect on the /2 -cell and under such circumstances the effect on glucegon release is minimal.

Within the limitations of this simple test, 1-atonine injections seemed to indicate some inability of andogenous insulin to suppress glucose production by the fiver in the geriatric subjects. However, the clonine results on their own do not really prove that the higher glucose values in the geriatric patients are not simply due to the shortage of insulin.

As in the study in obesity, the geriatric patients were also divided into two groups based on 120° blood glucose value during and GTT. The 'diabatic' group has 120' blood glucose value higher than 7.77 m mol/1. [140 mg/100ml] and the 'non-diabatic' group values lower than 7.77 m mol/1. Fostultously, the dividing line (7.77 m mol/1), which had been attitratily chosen, corresponded closely with the mean 120° blood glucose value for all 23 patients (8.12 m mol/1) and with the 50th percentile value for the 120° blood glucose value of 80 year olds given by Andres (1973). The degress of impairment in and GTT in these two groups was not related to the degree of hypoinsulinaemia (Table 22). In fact, the average insulin values during and GTT were almost identical for the 'diabatic' and 'non-diabatic' groups, each being slightly less than for normal controls. The only difference lay in the shape of the curves. This confirms the argument advanced earlier for the geriatric patients as a whole, that insulin deficiency was not the main cause of impaired glucose tolerance.

The question really is, if the elderly do indeed produce less insulin in response to a glucose challengs than young people, does insulin 'deficiency' constitute a limiting factor in glucose homeostasis? Another way of looking at this problem is to pick only control subject who had low insulin-glucose molar ratio during and GTT (low-malo controls). In these low-ratio controls (LRC) (n = 3), their mean insulin-glucose molar ratio was 27.8 ± 1.33 against 29.8 ± 2.29, in the elderly (p > 0.05).

The mean average plasma insulin concentration during and GTT in the LRC was 20.8 ± 0.55 punit/ml against 31.6 ± 2.52 punit/ml in the geniatric patients. The difference was also not significant. However, the LRC, despite the lact that they had slightly lower mean value (n 'normal') of average blood glucose concentration during the and GTT. This mean value was 5.38 ± 0.25 m mal/l in these LRC against 7.72 ± 0.28 m mal/l in the geniatric patients. The difference was highly significant (p < 0.001) (Table 26). These results again show that hypoinsulineemia was not the main cause of impaired glucose loisence in the elderty.

Our results show significant differences in carbohydrate metabolism between aid and young people, and we are convinced that the liver plays an important rale. Future work should attempt to define this role more clearly. For instances, if a tracer does of labelled glucose accompanied the test load, measurements of specific activity of planne alucose at vertous times during the test would throw some more light on

Table 26

The mean values for everage concentrations of blood glucose and plasma insulin, and insuling lucose maler ratio during and IST; in controls, 5 controls with low insuling glucose maler ratio (low-ratio controls) and in gerietric potients (4 SEM). Number of observations in parenthesis.

		Mean of the avera	glucam malar ratio		
		Blood glucose	Plasma insulin		
		(m mal/1)	(µunit/ml)		
I	Controls (13)	5.71 ± 0.18	33.5 ± 3.51	42.2 ± 4.50	
	1 x II	RS	ns	ns	
	1 × 10	**	ns	**	
II	Low-Ratio Controls (5)	5.38 ± 0.25	20.8 ± 0.55	27.8 4 1.33	
	H×III	***	ns	ns	
III	Geriatric patients	7.72 2 0.28	31 ,6 ± 2 ,52	29.8 2 2,27	

^{***}p < 0.01; *** p < 0.001; ns = nat significant

the significance of liver gluconeogenesis.

As with carbohydrate metabolism, It is not surgrising that protein synthesis and protein turnover is also reduced in the alderly (Young, Steffee, Pencharz, Winterer and Schrimshaw, 1975; D.J. Millword, personal communication). Our results show that added alanine disappeared significantly wore slawly (lew Kg) from the blood of the elderly than from normal young people (p < 0.001; Table 25). This may indicate that transfer rates for amino acids like alanine become distributed in old age. Dr. D.J. Millward (personal communication) reports that infusion of lubelled amino acids takes considerably longer to reach plateau levels in old rate than in young rate. Possibly, Impaired glucose tolerance may even be a part of this general phenomenon.

PART V
GENERAL DISCUSSION

INTRODUCTION

The common thread running through this work is the rate of insulin in glucose homeostasis in three different situations, each of which may be regarded as presenting same degree of stress viz. surgery, obesity and aid age. Each of these topics has already been discussed individually in some detail. The object of the present discussion is to comment briefly on a few general points which have excepted earlier reference or which are of general importance across the whole area of this investigations.

CHAPTER II

A. Chalce of Subjects:

Patients participating in the study of glucose homeostests after surgery were chosen by the surgical staff, Department of Surgery, University Callege Haspital, Landon (Head: Professor C.G. Clerk), based on our specification (see Part II, Chapter II, Subjects).

This study was planned in such a way that each patient would be his own central by comparing tests carried aut on day 1 post-operatively, and an 'recovery'. Four young normal subjects were also included in the study and were referred to as 'controls'. However, they could not serve as 'controls' in the pure sense, because, although their mean body weights and mean % ideal body weight were not significantly different from those of the potients, they were significantly younger. The mean age of the normal subjects was 27.3.2.3 years againt 52.3.3.4 years, in the patients (F < 0.01). (See Part II, Chapter II, Subjects). If work with glucose infusion were to continue, It might be useful in the future to recruit more volunteers to make up the number, and extend the age range, so that II covers the age range of the patients. However, in the present work, any differences between the sungical patients and the 'control' subjects could, therefore, not only be due to the effect of surgery and food deprivation, but also the effect of age.

If age (52 years) were a significant factor in determining the retuins of glucase infusion, one would expect at least a trend towards impaired rather than heightened insulin vectorion is response to glucose (cumpare geriatric patients, Part IV). Therefore, the affect of age in the surgical study is likely to have been minimal.

In the choice of patients in the study in obesity, we had 29 abese subjects who attended the Nutritian Clinic, University Callege Hospital and willing to participate in our study, but not all agreed to have the three tests carried out on them. Some agreed to having the arci GTT while the others agreed only to participate in the ly GTT and in GTT (these two tests were carried out one after another on the same maming). And only 7 subjects agreed to have all three tests carried out (Yobio 11).

It is hoped that in future work a larger number of obese patients will be studied, so that they can be classified not only according to % ideal body weight alone (as in the present study), but to be extended to the possible classification of patients by combined age and % ideal body weight.

Fertunately, in the study in the elderly, we were able to persuade the volunteers (gestatric patients) to participate in all of the three tests (arel GTT, ly GTT and Iv GTT). Besically, most of these patients were lonely, and they loved to have someone show an interest in them. In general they were less concerned about the traumo (pain) of the needle used during the test, compared to the attitudes of the sungical and obese patients, and even more, the healthy controls.

However, we also faced difficulties when we asked the elderly patients to participate in the IV alanine tolerance lest. This may be due to the fact, apart from some of the patients not being available, the IV alanine balerance test gave a slight stomach discomfort to some, for 10 to 20 minutes after the injection of I-alanine. As this became known, the difficulty in obtaining valuntaers increased, not only in the elderly but also among potential "control" subjects.

B. Chalce of Tests

It would be ideal if we could have the same test for all three studies (surgery, ebesity and old egs). However, each test has its own marits and disdunntages.

1. Glucose Infusion tests

This test was considered the best for surgical patients, because it represents only a minor change (a pump instead of the usual infusion drips) in the method of infusing glucose which most patients received as usual post-operative 'dietery' therapy. It was, therefore, of minimal burden to these immediate post-operative patients (day 1); and included those who post-idental infusion yeard most of them did not seem to mind this change. However, during the 'recovery' period the presence of the operator together with his infusion set and infusion pump was not a welcome sight.

The emount of glucose infused (0.35 g kg⁻¹ h⁻¹) would certainly have suppressed hepatic glucose output (Madison, 1906). Therefore, when a plateau of blood glucose cancentration had been reached, we could discount the hepatic factor, and the rate of infusion at this stage represents the peripheral glucose uptake, unless blood glucose concentration exceeds the renal barrier, in which case trans would be lost through urine.

Since the rate of infusion was the same on all occasions, so also was the total glucose uptake. Now total glucose uptake = concentration x fractional rate af uptake (Francison, et. et. 1966). Therefore, those patients who showed the highest plateau concentrations of blood glucose (day 1, past-operatively) must have had the lowest fractional rates of glucose removal (equivalent to the lowest KG values).

We also tried similar glucose Infusions on some obese and geriatric patients.

They also shared the view of those recovering surgical patients (during the "recovery" period), of not being keen on having this test carried out. This, together with other technical problems such as the difficulty in getting two nice values, one in each arm.

In some obese subjects, meant that there seemed no good reason to persist with glucose infusion as apposed to the other tests.

2. Oral GIT:

This test is straightforward and has not been associated with any complaints of discomfort from patients or subjects. Its main obvious advantage is that it is physiological, in that the glucose enters by mouth and is subject to the full range of factors before it reaches the tissues. Its main disodvantage is that one is presented with a curve or a set of results rather than a single enswer. This may necessitate quite difficult value judgements, e.g. in deciding the weight to be estached be festing or 120' blood glucose values in diagnosing 'diabetes'. (Working Party, Royal College of General Practitioners, 1963, WHO, 1965).

The fact that the oral GTT has been so widely used under fairly standardised conditions in a large variety of subjects, makes it a valuable reference test in comparing data from one source with another.

3. Iv GTT:

The text is relatively unfamiliar to most clinicians compared with the oral GTT, and its advantages are the short duration of the text (60 minutes) and the fact that the test has a result which can be expressed as a single figure (Kg). This parmits statistical comparison between individuals or gas ups and between the same subject at different periods of time. Moreover, since it bypasset the gut, the ly GTT eliminates the compilation of having to consider the absorption and gut insulnagenic factors. However, it also has some alterivantages.

Lefferty, Gliddings and Mangnell (1975) argued that % disappearance rates (KG values) for glucose were without value and absolute removal rates alone have any meaning. It can, however, be argued that "normality" in absolute glucose removal rates atone tells is very fittle about the state of the subjects. For instance, in the earlier discussion (8.1.) on singlical patients, it was pointed out that all had the same absolute glucose removal rates, but because blood glucose concentrations differed, KG values must have differed similarly (but inversely). The KG values gave a measurer of insulin sensitivity (they significantly correlate with KG + | values; gave a measurer of insulin sensitivity (they significantly correlate with KG + | values; and insulin levels reveal the extent and manner of adaptation to this changing sensitivity to insulin. It is in fact difficult to see how absolute glucose removal rates could be impaired, unless there was a severe failure in the Acali or wastage of glucose via the unless.

4. Iv GITT:

This test is extensively used in animals (Turner, 1966; Heard and Turner,

Table 27

Correlation between KG and KG $_{\Phi\,I}$ values in control subjects, obese and geniatric patients, and in the combined three groups.

	No.	Regression Line	<u>r</u>	E
Control subjects	10	y = 0.33 x + 0.32	0,60	> 0.05
Obese patients	16	y = 0.34 x + 0.15	0.65	< 0.01
Garlatric patients	22	y = 0.23 x + 0.44	0.43	< 0.05
Combined group	49	y = 0.33 x + 0.21	0.70	< 0.00

1967, Heard and Henry, 1969, a, b) resembles others which have been used in man (a.g. Franckson, et. et., 1966) and is in part, identical to that of Silverstone, Branfonbrener, Shock and Ylengss, (1957). The key difference is that in the present test the iv GITT should be coupled with tv GITT and measurements of plasme insulin, to give simultaneous essessment of glucose tolerance and insulin status and sensitivity. The ky GITT is then, a good measure of insulin sensitivity (Table 27).

The administration of a relatively large dose of exogenous insulin in this test should certainly switch off glucose output from the lives, and, therefore, any differences in the KG + 1 values can only be due to the differences in peripheral tissue responsiveness.

The 10° values of plasma insulin concentration after the injection of glucose and insulin, could be as high as 1.50 to 300 $\,\mu$ unit/ml in the normal subjects, and even higher in some of the obese patients.

C. Skinfold Measurements:

Skinfold measurements were certied out in some obese, elderly and control subjects, as a measure of their nutritional status. These measurements, however, were so unreliable in the obese, especially the extremely obese (reproducibility very poor). They seemed to have better reproducibility in the elderly, but the present data for comparison (Durain and Wamersley, 1974) did not cover very old subjects (the highest was 50 + years). The measurements of the elderly subjects (mean age 79 years) if compared with their date, gave apparently lower values of body fat centent. This may be due to the fact that in the very old the elasticity of the skin had become very poor, and the measurements gave a false, lower reading of skinfold thickness.

CHAPTER III STUDY IN SURGICAL PATIENTS

A. Insulin Resistance:

The patients showed a "temporary" insulin resistence, manifested in infusion 1, by high planeau values for blood glucose (1.e. low K_G values; see Chapter II, 8.1), in the presence of levels of insulin which were higher than normal. In infusion II blood glucose plateau concentrations were almost the same as in infusion 1, and therefore K_G values had not changed significantly, but plasma insulin values were lower. Therefore, presumably insulin resistence had diminished, but had not returned to normal (compare with young 'control' subjects).

In Infusion 1, the development of insulin resistance was probably due to the high circulating levels of cortisal and glucagan and probably other anti-insulin hormanes (e.a. catecholamines and growth hormane).

B. Significance of insulin Therapy:

As discussed earlier, insulin administration has been advacated and practised to avercome the insulin resistance of trauma (Hinton, et. al., 1971) (see Part II, Chapter IV). It must be acknowledged that insulin treatment is probably necessary for very serious traums, like burns. The very high catabolic state in hums would deplate glycogen stores and enhance lipolysis. This increase in lipolysis causes raised levels of NEFA, which were shown to cause massive deposition of fat in the liver (Felgelson, Pfatfs, Karman and Stienberg, 1961). On the other hand in moderate or minor types of surgery (or other physical trauma, lipolysis is probably desirable, to enable the body to use its energy reserves of for rather than protain.

Giving emino acids, instead of the usual glucose, would then permit the improvement of negative nitragen balance and by not encouraging release of high levels of insulin in the circulation would permit lipolysis (Blackburn, et., al., 1973).

C. Nitrogen Balance

As judged by the degree of the uninary nitrogen exception (mean between 9 and 11 g nitrogen loss per day), the degree of trauma in the present series of surgical patients appears low, compared with, for instance, massive uninary nitrogen excretion reported by Cuthbertson (1964), from patients following fracture. However, his patients received a diet with 70 g protein/day, unlike the surgical patients who received none for the first few days fallowing the operation. Another factor to be borne in mind is the influence of previous diet. If low in protein, this might lead to minimal increased loss of urinary nitrogen following trauma (Munro, 1964). Conclusions drawn from urinary nitrogen figures alone are dangerous. A proper nitrogen balance must be estimated.

CHAPTER IV STUDY IN OBESITY AND ELDERLY SUBJECTS

A. Impaired Glucase Tolerance

The abese and elderly subjects showed a chronic or permanent form of Insulin resistance.

Both groups showed impaired glucose colerance, shown by the significently higher mean values for average alood glucose concentrations during oral GTT (Table 28), higher and delayed peak of glucose levels during and GTT (Fig. 18A and Fig. 27A), and significently lower values of KG during by GTT (Table 29), compared with the mean values in the normal subjects.

The Impaired glucose talerance in the abose subjects was related to higher mean levels of everage plasma insulin concentrations and peak insulin response during area GTT (Tables 28 and 30°, and to significantly higher mean peak plasma insulin concentration (Table 20) during IV GTT, than in the normal control subjects.

In contrast to the obese, Impaired glucose tolerance in the elderly was associated with normal or marginally diminished insulin levels during oral GTT and Iv GTT (Tables 28 and 30).

B. Possible Role of the Liver:

The main feature of the oral GTT in the alderly was the delayed and exaggerated peak glucose values. The main cause of which probably the Impaired ability of the liver (in the elderly) to switch off glucose production in response to endogenous insulin. This impairment resulted in prolonged hyperglycaemia. But one must also

Toble 28

The mean values for festing blood glucose and plasma inselin, excess concentrations of blood glucose and plasma inselin, and inselin; glucose actor ratio during oral GTI in control, where and gentatric subjects (\$ SEM). Number of observations in parenthesis.

	Mean fasting concentrations		Mean of the Average Value for 150 min.		10 ⁶ x mean insulin:	
	(m mol/)	Plasma insultin	Blood glucose (m mol/l)	Plean	a Insulin	
	(as many .	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		(µ unit/ml)	$(10^6~\mathrm{m~mol/l})$	
I Controls (13)	4,62 = 0.16	12,8 2 1.31	5.71 ± 0,18	33.5 ± 3.51	240.4 ± 25.2	42.2 2 4.50
l at H		٠		64	**	+
I x III			**			**
II Ohese (16	4,89 1 2.93	21.4 ± 2,88	6.73 ± 0.28	69.1 ± 10.61	495,8 2 76.4	73,5±11.8F
11 x 111		ere:	4	100	100	
III Gerlettic (23)	4,92 ± 0,11	12.3 ± 0.54	7.72 ± 0.28	1 31.6 = 2.52	227.0 ± 18.1	29.8 ± 2.27

^{*}p < 0.05; ** p < 0.01; *** p < 0.001

The mean values for KG , KG , 1 = KF in the control, obese and white subjects (2 SEM). Number of observations in parenthesis.

			Mean Values	
		KG (SQmin)	KG +1	(%/min)
I	Control	1.81 ± 0.28 (10°	4.55 ± 0.51 (10)	3.10 ± 0.73 (6)
	1xH		+	ns
	Ex III	***	100	ns
II	Obese	1,11 ± 0,18 (16i	2.81 ± 0.35 (14)	2.86 ± 0.90 (4)
	11 × 111	ns	nt r	nă.
III	Gerianic	Q.98 ± Q.86 (23)	2.29 ± 0.16 (23)	2,21 ± 0,34 (10)

^{*} p < 0.05; *** p < 0.001; ns = not significant

acknowledge that the Insulin response to oral glucose administration was sluggish in the alderly as shown by the delayed peak insulin level during this test.

It is unlikely that the deleyed peak Insulin level was the result of delayed ebeorption rate of glucose, as suggested by Smith and Hall (1973), but proof is lacking. The evidence which is against this argument is a. the increase in blood glucose level was equal in the elderly and in young people during the first 30°; b. after 30° the elderly still increased their blood glucose level nearly as much again as they did in the first 30 minutes. The peak level of blood glucose corresponded with the peak insulin level at 90°; and c. the simple is alonine tolerance test showed that the elderly has increased levels of blood glucose which were not suppressed by their endagenous insulin secretion.

The Impairment of IV GTT was also probably due to the decreased ability of endogenous insulin to suppress gluconeogenesis. The mean K_G value was significently lower in the elderly (Table 29), while the mean peak insulin level, ethough lower, was not significantly so (Table 30).

The logarithm values of K_G were significantly correlated with 120° blood glucose values during and GTT in the combined group of the abase, elderly and control subjects (p=0.001; Fig. 22). This suggests that iv GTT would be just as good (or as bad) as the anal GTT in detecting impairment in glucose solarance.

C. Insulin Sensitivity:

1. Blood glucose and NEFA levels:

While interpretation of the Iv GTT might be ambiguous, the Iv GITT seems

Pictre 3?. Correlation between K₀ value (%/min.) during iv GTT and 170° value of blood glucose concentration during oral GTT, is the obese, geriatric and control subjects.

(•) obese patients.

(O) overt disbatio obese patient.

(.) geriatric patients.

(A) overt diabetic geristric patient.

(m) normal control subjects.

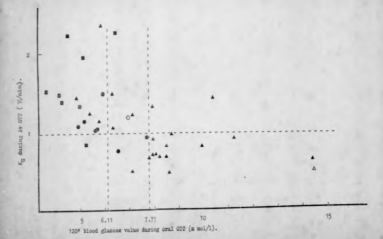


Table 30

The mean peck insulin concentrations during ere! GTT and iv GTT (2 SEM), in control, obese and geriatric subjects. Number of observations in perenthesis.

Mean Peak Insulin Concentrations (II unit/ml)

				(In and 6 are	
		Oral GTT		Iv Gft	Ratio of oral: 1v
I	Controls (9)	72.0 ± 7.26	ns	54.0 ± 13.60	1,33
	1 x II	10		14	
	t a til			ns	
11	Obess patients (5)	160.6 24.98	ne	138.4 ± 11,17	1,16
	Lx II	***		***	
777	Geriatric paties	th 48.0 2 5.34		41.6 = 5.13	1,15

[&]quot;p <0.05; "" p <0.01; "" p < 0.001; ns = not significant

quite clearly to measure peripheral tissue responsiveness. It is very unlikely that liver gluconeogenesis was not suppressed (even in the obese and the elderly) by the relatively high concentration of exagenous insulin which was given. The circulating levels 10 minutes after injection of insulin was found to lie in the range of 150 unit/mil to 400 unit/mil, i.e. about 5 to 10 times higher than peak levels of andogenous insulin measured during to GTT. Both the obese and the elderly showed significantly decreased mean KG +1 values (Table 29), indicating that peripheral tissue response to insulin (as judged by glucose uptake) is reduced in these obese and elderly people.

In the elderly, it has been argued that this impaired peripheral tissue sensitivity extends also to suppression of hapatic glucose output (see Chapter II 8. Oral GTI). There seems no good reason why these arguments should not also apply, though perhaps to a lesser degree, to the interpretation of the oral GTI of the obese-

Both In the study in obstity and ald age, the evidence showed that this decrease in insulin sensitivity in relation to carbohydrote is more pronounced than in relation to far metabolism. This is shown by the absence of any significant difference in the NEFA curves during and GTT (Figs. 18C and Fig. 27C) or K_F values during by GTT (Table 291, suggesting that the lipolytic action is not yet offected by obesity or old suggest

This is perhaps not surprising since insulin concentrations which waximally situation glucose uptake by insulin sensitive itsues are considerably higher than concentration required to inhibit lipolysis.

2. Correlation of glucose talerance (KG ! with insulin sentitivity (KG + 1)

This work demonstrated highly algorificant correlations between K_G and K_{G-+1} values (p. 0.001, Table 27, Fig. 55). It is interesting that this significant correlation was present in even the obese group alone (p. 0.01) or the garietric group alone (p. 0.05), and given a larger number of control subjects, they would certainly have shown the same correlation. Another interesting (e.c. is that in such individual group, the shape of the regression line was almost identical, except in the garietric group which was slightly lower (Table 27). Perhaps this is a sign of some degree of $\frac{1}{2}$ cell failure.

3. Hypersensitivity to insulin in "normal" subjects:

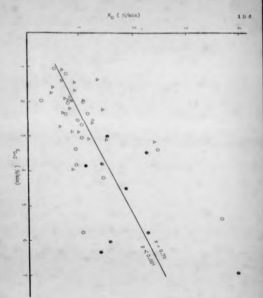
Some apparently normal subjects showed vary high values for K_G +1 and therefore, also for K_G. It has been suggested in Part III, Chapter IV, that these subjects may be in an early phose of 'active' obesity. A longitudinal study of such people may or may not reveal that this speculation is true.

This speculation is based on the similar pattern of changes in K_G and $K_{G\to f}$ found in experimental enimals. These animals were fied on diets marginally deficient in protein (Heard and Turner, 1967; Heard and Stewarr, 1971). In all cases, hypersensivity to insulfn was the earliest phase and fed on to impaired glucose tolerance and diminished insulin sensitivity.

It would remain also to estempt to distinguish genetic from dietary influences.

In the development of obesity.

Pigure 35. Regression line between the values of K_G and K_{G+1} in the combined group of obese (O), geniatric (A) and normal control subjects (•).



The phrains 'active' and 'passive' in regard to obesity, may have been independently used by other authors in a different context; e.g. Craddock (1973) used these terms for the ability or inability of obess subjects to gain more weight. While this work uses these phrains in relation specifically to insulin manufactivity (in respect to carbohydrate mesobolism). (See Part III, Chapter V).

As it was shown (Part III, Chapter III), that treatment of obesity is very difficult and unreliable, a longitudinal study of insulin sensitivity eight detect makes in the factive pre-abose phase or floor who are prone to obesity.

Prevention of abesity rather than treatment is likely to offer greater success.

4. What determines insulin sensitivity?:

Two main groups of factors must have been likely to be important. They are: a. anti-insulin hormones, insulin antagonist, etc. (see Part II, Chapter I), and b. callular factors including a valiability of receptor sites. Early anthulasm for the concept that insulin sensitivity and receptor evallability would always correlate exactly, has been rather dampened (Cuatraceses, 1974; Kern, 1975) and more work on this aspect is still needed.

CHAPTER V DIAGNOSTIC SIGNIFICANCE OF THE TESTS

Obviously a combination of tests would be ideal in determining an impairment in glucose tolerance. For practical purposes, however, using only one particular test, and even in extreme cases, only looking at fasting blood glucose values, would probably be sufficient; although, a few borderline cases may escape from detection.

Additional elimination of subjects who had festing blood glucose concentrations higher than 6.11 m mat/1 (thus all having normal festing blood glucose), 120' value for blood glucose during and GTT revealed that some of the obese (\geq 6.11 m mot/1) and some of the generatic subjects (\geq 7.77 m mot/1) were 'diabetic' (Fig. 32). Furthermore, although it was shown earlier that the logarithm values for KG during to GTT were significantly correlated with 120' values of blood glucose during and GTT, there were some subjects who would be normal by one test but 'diabetic' by the other. If one's aim to to get early detection of a disease, or an impairment, then a combination of tests is very necessary. (Fig. 32).

(v GTT) is best carried out immediately after iv GTT, because it could serve as a measure of peripheral tissue responsiveness to insulin, since the affect of liver aluconeogenesis has been eliminated.

CHAPTER VI

A. Surgical Patients:

A nitrogen belance study showed a negative belance emounting to 50 g pretain loss per day for a period of 5 to 6 days after the operation. This period of negative nitrogen belance coincided with elevated faiting levels of plasma glucegon, NEFA, branched-chain amine acids, urinary free cortisal, urinary 17-OH-conticosteroids and with a decrease of total amine acids. Festing levels of plasma insulin and plasma cortisal were only significantly elevated on day i post operatively.

A two hour Infusion of glucose (0.35 g kg⁻¹ h⁻¹) and ay 1, resulted in hyperglyceemia and hyperinsulinasmia, suggesting a "temporery" insulin resistance. Glucose also suppressed glucogon secretion to the same extent anday 1, and an "recovery". It also decreased plasmaNEFA, total amino acids and plasma cortisol in both infusions. But, the fall in plasma cortisol was probably due to the effect of mating rather than the actual effect of glucose.

Several eyestlans remained unanswered:

Insulin resistance seems to protect the subjects from the less destrable features of elevated insulin levels. Then perhaps amino acid administration rather than glucose would be more beneficial, since it improves the nitrogen belance without having to increase plasma insulin levels and permits mobilization of feet (Blackburn, et. al. 1973). More work, however, is still needed to establish the best post-operative regime for patients undergoing surgery.

 It is known that protein synthesis is decreased following surgery (O'Keafe, et. al. 1974) Crane, et. al. 1976). Whether insulin resistance is also responsible in causing the decrease of protein synthesis should also be investigated.

B. Obese Subjects

This work demonstrates that above subjects showed Impoled resource to and I, and I of I. The impolement was associated with hyperinsulineania. The results showed that K_G or K_G+I values were not statistically correlated with fasting or pack insulin levels during in GTI. However, higher values for K_G or K_G+I were usually associated with lower values of fasting or pack insulin. There was a significant correlation between K_G and K_G+I values in combined groups of patients and control subjects.

The logarithm values for feating intuitin were correleted with % ideal body weight (p<0.001). K_G or K_{G+1} were not statistically correlated with % ideal body weight however, lower values for K_G or K_{G+1} were usually associated with higher % ideal body weight.

Unlike that of carbohydrate metabolism, this present work was unable to show any significant impairment of insulin sensitivity in regard to fat metabolism (ludged by the difference in NEFA curves during and GTT). Ky during Iv GTT).

Oral GTT, Iv GTT and Iv GTT did not give any pradiction of the possible success or failure of any distany treatment of an obese patient.

This work speculates that there are two types of obesity (based on their insulinsansitivity): a. 'active' abesity, [udged by the relatively low % ideal body weight (<160%) low values for K_G , K_{G+1} and low fasting insuling in. 'possive' obesity [udged by high % ideal body weight, low values for K_G , K_{G+1} and high fasting insuling and c. apparently normal subjects who had vary high values of K_G and K_{G+1} are probably in an early stage of 'active' obesity (see Chapter IV, C).

There are a few aspects of this study which need further investigations

- 1. more 'normal' subjects with high K_G and K_{G+1} values to be investigated. Lengitudinal study of such people may or may not reveal that the above speculation is true.
- II. More obesit patients are needed, so that they can be classified by the combination of age and % Ideal body weight. This is hoped to throw some light on the development of obesity.

C. Elderly Subjects:

This work demonstrates the familiar impairments in and GTT and iv GTT in the alderly. Although the impairment of glucose talerance was associated with hypolasulinasmia and decreased paripheral tissues insulin sensitivity, the decreased ability of andogenous insulin to self-chiff liver gluconeogenesis seemed to be the major cause of hyperglycosenie. After introvenous injection of indianine, the elderly showed increased levels of blood glucose which were not suppressed by their endogenous insulin secretion.

As in the study on obesity, this work confirms earlier findings in animal experiments (Turner, 1966; Heard and Turner, 1967; Heard and Henry, 1969 a), about the significance of insulin sensitivity (K_{G-q-1}) in determining glucose tolerance rather than the actual levels of circulating plasma insulin. This was shown by the significant correlation between the K_{G} and K_{G-q-1} values in combined groups of elderly potients and control subjects, and also in the combined group of obess, alderly and normal control subjects.

Also, as in the obese, this work was unable to show any significant impairment of insulin sensitivity in regard to fee metabolism (NEFA curves during and GTT).

Ke values during by GTT).

Few aspects in the study in the elderly need further investigations

- As old age is not merely the passing of years, but a generalized picture of many multurestons added together, the question now arises, whether the againg phenomenon should be prevented. And If it should, in what way?
- 11. As skinfold measurements are unreliable (see Chapter II, C) and % Ideal body weight is unable to be measured, what is a good, simple criterion in judging the nutritional status of the very old?
- III. A possible role of the liver in producing hyperglycoemia needs further investigation. By including a tracer dose of labelled glucose in the ardinary glucose load, and measuring the plasma glucose concentrations and the specific activity of labelled glucose at various times during the test, one would be able to get more information about the impaired suppressibility of liver gluconeogenesis, in the elderly (perhaps also in the above).

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MERICALE AND HORNOTAL CHANTES ANTES SUBCISIVE HYPERINGLINABILA DOMING GLOCORE INPURIOR.

V.E. EDENOMIENOVO, C.B.C. HEAND, W.P.T. LANS, J.FRA & G.B. MICON. London Ombosi of Hygisme and Tropical Medicine and Hoyal Peat-Graduate Medical School, Rendon.

Hereo patients were desired; Justin cheer had a part administration of the second seco



This work confirms that hyperpluonoponemia full-son marginal appearation () and demonstrates that this as morphomethic still pluodes, although low plants insulint gluone wakes that we reported the every home and garden generation (). A three operation will be supported to make the operation who insuling resistance.

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Glucose tolerance, plasma insulin levels and insulin sensitivity in periatric patients, By C. R. C. Hyann, W. S. Sossjodinnoro and Sylvia M. France, Chnical Natrition and Metabolism Unit, Department of Human Natrition, Lundon School of Hypiene and Trapical Medicine, London WC1E HT, and A. N.

ERYON-SMITH, Department of Gariatries, University College Hospital, London Vennus blood glucose values, 120 min after an oral load of 50 g glucosa (G.m) are used to diagnose diabetes (WHO, 1965). Values 16:11 mmol I are classified as normal and >7-22 mmol/l sa diabetic. However, Gine increases steadily with age and this raises problems in the elderly both of clinical significance and of physic-

logical interpretation This communication reports the results of an investigation of twenty-four gerintric patients (mean age 79 years) in whom both availability of insulin and insulin semaitivity were sessessed.

Each patient received two tests on successive days; (s) and glucose tolerance test (GTT) (50 g glucose) lasting 150 mair, (2) intravenues (i v.) GTT (0 33 g glucose, kg) leating to min followed immediately by i.v glucose i insulin (0:33 g glucose and 0-193 units insulin kg) with blood sampling for a further 60 min (Heard & Henry, 1969). Half the petierts had the oral test on the first day, half had the i.v. test. Patients were fasted from midnight. Blood was sampled and glucuse and insulin injected via an indwelling husterfly needle in the sim. Glucose (glucose axidase method) and insulin (Radinchemical Centre method) were estimated in plasma. Gluense disappearance rates (k; % per min) in the i.v. tests were calculated to give An for glucose alone and has for glucose + issulin Normal values are about a and 5% per min respectively (Franckson, Malaise, Arnould, Russo, Balasse, Conrad & Boutenie, 1956).

Although only one patient was overtly dishetic (fasting glucose 8 to mmol/l), mixteen of the twenty-four had Gam values >7.77 mmol l. Of these, fourteen also had neverely impaired i.v. gluents tolerance (Ag <1000 per mis). Only two other patients had Aq <1 0% per min. Clear signs of glucose malabourption occurred in three patients, who therefore showed very low plasma insulin levels during the oral GTT compared with values during the i.v. GTT. Another five patients without malabaseption also showed low plasma insulin levels during the oral GTT suggesting Imperiment of insulinogenic gut factors. Although plasma insulin levels during both tests were lower than those reported for young normal subjects, insulin sensitivity (Acres) was also low in all the patients (mean a 4% per min). The shape of the oral GTT curves auggested that failure to auppress hepatic glucose release was a feature of this inculin intensitivity.

The extent to which these changes are typical of old age per se and whether deterioration in nutritional status contributes to the effects remain to be cotablished.

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