AN ECTOPIC SYNTHESIS OF THE MELANIN IN THE ADIPOCYTES
OF THE MORBIDLY OBESE SUBJECTS.

by

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A dissertation submitted in partial fulfillment of the requirements for the degree of Doctor of Philosophy at George Mason University

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DEDICATION

This thesis is dedicated to my parent in laws  
Prem Singh and Harbhajan Kaur Randhawa  
who have been a source of encouragement and inspiration thorough out my studies.

This thesis is dedicated to my parents  
Gurdev Singh and Narinder Kaur Dhillon  
who introduced to me the joy of reading from birth,  
enabling such a study to take place today.

This thesis is dedicated to my dear husband  
Harpal Singh Randhawa  
Thanks for providing me strength, courage and support that urged me to strive to achieve  
my goals in my life.

This thesis is dedicated to my lovely sons  
Akashdeep Singh Randhawa  
Swaraj Singh Randhawa  
There were many times when this thesis took me away from precious moments with  
them. They teach me about unconditional love.

This thesis is dedicated to other family members and friends who have contributed in  
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ETBR..........................Endothelin-B receptor  
FBS............................Fetal bovine serum  
FGF............................Fibroblast growth factor  
GSH..........................Glutathione  
GSK3β .....................Glycogen synthase kinase-3β  
GTP...........................Guanosine-5’-triphosphate  
HIF-1..........................Hypoxia inducible factor-1  
HPLC.........................High-performance liquid chromatography  
IBMX..........................Isobutylmethyl xanthine  
ICAM-1..........................Intercellular cell-adhesion molecule-1  
IgE..............................Immunoglobulin E  
IGF.............................Insulin-like growth factor  
IL..............Interleukin  
LAM............................Lymphangioleiomyomatosis  
LDL.............................Low density lipoprotein  
L-DOPA…………………..3, 4-L-dihydroxyphenylalanine  
LEF 1/TCF..................Lymphoid enhancer-binding factor 1  
LF..............................Lipofuscin  
LPH.............................Lipotropin  
LPL..........................Lipoprotein lipase  
LPS..........................Lipoplysaccharide  
MAPK................Mitogen-activated protein kinase  
MCH............................Melanin concentrating hormone  
MCP-1........................Monocyte chemoattractant protein 1  
MCR............................Melanocortin receptor  
MIF................Macrophage migration inhibitory factor  
MIF..........................Migration inhibitory factor  
MITF..........................Microphthalmia-associated transcription factor  
MM..............................Malignant melanoma  
MMP..........................Matrix metalloproteinases  
mRNA..........................Messenger ribonucleic acid  
MS.................Metabolic syndrome  
NCC...........................Neural crest cells  
NF-kB..........................Nuclear Factor κB  
NGF............................Nerve growth factor  
NM..............................Neuromelanin  
ob/ob..........................Obese/obese  
OCA........................Oculocutaneous albinisms  
OCT..........................Optimal cutting temperature  
ONECUT-2......................One cut domain 2  
p90RSK......................p90 ribosomal S6 kinase  
PAI-1..........................Plasminogen activator inhibitor-1  
PAM..........................Peptidyl α-amidating mono-oxygenase  
PAR2..........................Protease acitivated receptor 2

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PAX3..........................Paired box gene 3
PBS..........................Phosphate Buffered Saline
PC..........................Prohormone convertase
PCR..........................Polymerase chain reaction
PD..........................Parkinson disease
PKA..........................Protein kinase A
PKC-β.....................Protein kinase C-β
PNGaseF....................Peptide: N-glycosidase F
PNS..........................Peripheral nervous system
POMC........................Proopiomelanocortin
PPAR/RXR...............Peroxisome proliferator-activated receptors /retinoid X receptor
PTCA........................Pyrrole-2,3,5-tricarboxylic acid
PUFA........................Polysaturated fatty acids
QTL..........................Quantitative trait locus
ROS..........................Reactive oxygen species
RPE..........................Retinal pigment epithelium
RSK..........................Ribosomal S6 kinase
SCF..........................Stem cell factor
SDS-PAGE.....................Sodium dodecyl sulfate polyacrylamide gel electrophoresis
SN..........................Substantia nigra
SRY..........................Sex determining region Y -box 10 (SOX 10)
STAT3......................Signal transducer and activator of transcription 3
TF................................Transcription factor
TGF-β........................Transforming growth factor- β
TH..........................Tyrosine hydroxylase
TIMP........................Tissue inhibitor of metalloproteinases
TNF-α........................Tumor necrosis factor-α
TRP..........................Tyrosinase related protein
TYR..........................Tyrosinase
TYRP1........................Tyrosinase related protein 1
TYRP2........................Tyrosinase related protein 2
UV..........................Ultraviolet
VEGF........................Vascular endothelial growthfactor
VLDL........................Very Low-Density Lipoprotein
α, γ, β-MSH ............α, γ, β-melanocyte stimulating hormone
ABSTRACT

AN ECTOPIC SYNTHESIS OF THE MELANIN IN THE ADIPOCYTES OF THE MORBIDLY OBESE SUBJECTS

Manpreet Randhawa, PhD.

George Mason University, 2008

Dissertation Director: Dr. Ancha Baranova

Melanocortins produced from the post-translational processing of pro-opiomelanocortin (POMC) are regulators of pigmentation in the hair and skin and are also critical for proper maintenance of energy balance. Particularly, α-MSH is a melanocortin ligand involved in the control of both pigmentation in skin and energy homeostasis through various subtypes of the melanocortin receptors present both in the skin and the adipose tissue. Expression profiling of 40,173 individual cDNA clones with RNA from visceral adipose samples showed statistically significant overexpression of genes encoding tyrosinase-related protein 1 (TYRP1), dopachrome tautomerase (DCT/TYRP2), melanosome transport protein RAB27a and Melan-A (MLANA). These findings lead to the following two hypotheses:
1) Melanin biosynthesis pathway is functional in adipose tissue and is excessively stimulated in morbid obesity.

2) Melanogenesis may be stimulated in morbidly obese individuals due to high levels of α-MSH.

Fontana-Masson staining performed on the adipose tissue registered the presence of a black pigment in the periphery of the adipocytes. Granules of the pigment were found in higher quantities in visceral adipocytes from obese individuals as compared to samples from lean subjects. An LC/MS/UV analysis of the chemical composition of the pigment proved that these granules contain melanin.

The expression of the TYR, TYRP1, TYRP2, MITF and MC1R genes involved in melanogenesis was studied by real time PCR in adipose tissue samples obtained from morbidly obese and from lean subjects. The expression level of TYR, encoding the rate-limiting enzyme required for the conversion of L-DOPA to dopachrome, was found to be expressed at much higher levels in obese subjects as compared to lean subjects whereas no expression was registered in liver and gastric tissues. The other genes showed the same pattern as TYR, but the differences were less pronounced as compared to TYR. The expression of TYR was further localized to the adipocytes as determined by in situ hybridization of adipose tissue slides, where TYR was found only in the periphery of the cell.

The study of the expression of tyrosinase protein in adipose tissue by Western blotting revealed properly folded and mature tyrosinase homodimer of 140kDa. The presence of the tyrosinase as well as TYRP1 and TYRP2 proteins were confirmed in the
human adipocytes by immunohistochemistry. A substantial difference has been seen between adipose samples of obese and lean subjects, with more tyrosinase in adipocytes from obese samples. The biological activity of TYR was evaluated by $^{14}C$ assay that showed increased enzyme activity in the adipose tissue from morbidly obese subjects as compared to lean subjects, whereas no activity was found in gastric and liver tissue samples.

Both murine 3T3-L1 adipocytes and primary human adipocytes at different stages of differentiation were exposed to different concentrations of $\alpha$-MSH for different time periods. Real-time PCR performed on mRNA extracts obtained from murine 3T3-L1 cells and human adipocytes provided no consistent expression data for melanogenesis related genes. The enzymatic activity of a tyrosinase from protein extracts obtained from 3T3-L1 adipocytes was evaluated by L-dopa assay. A gradual decrease in the rate of L-dopa oxidation was observed spectrophotometrically during the differentiation of adipocytes. $^{14}C$ assays indicated the presence of minimal residual activity of tyrosinase in cultured human cells. Western blotting performed on extracts from human adipocytes showed the presence of a specific band characterized by a smaller molecular weight than normal tyrosinase. The glycosidase digestions confirmed that this band corresponds to an inactive, nonglycosylated form of tyrosinase.

These collective findings indicate that the melanin synthesis pathway is functional in intact human adipose tissue while further work on the appropriate cellular model of the adipocytic melanogenesis is warranted.
**Introduction**

Pigmentation is one of the most variable phenotypes in humans. Skin color varies not only between races, but also between individuals of the same ethnic group. Differences in the human color traits in hair, skin, and eyes are primarily due to a pigment called melanin. Other chromophores present in the human body such as hemoglobin and carotenoids play only a minor role in skin pigmentation. The wide variation in constitutive pigmentation among humans is caused by enormous differences in the rate of the synthesis of the two forms of melanin, eumelanin and pheomelanin, and the rate of the transfer of the melanosomes to keratinocytes. These pigments are synthesized in lysosome-like organelles called melanosomes in the melanin-producing specialized cells known as the melanocytes. Pigmentation differences mainly arise due to variation in the number, size, composition and distribution of the melanosomes; whereas melanocyte numbers typically remain relatively constant.

Loss or gain of pigmentation due to genetic and environmental reasons leads to various pigmentary disorders. Darkening of the skin may result from abnormal distribution of melanin due to an increase in the number of melanocytes, to production of excessive amounts of melanin or to increased amounts of melanin supplied by a normal population of melanocytes. This results in various hyperpigmentary disorders like
epidermal melanosis, freckles, epidermal melanocytosis, lentigines etc. Diminished skin color most commonly results from the decrease in epidermal melanin content, e.g. leukoderma and hypopigmentation, caused by defects in the formation of the melanin (Nordlund et al., 2006). Vitiligo, a skin disorder characterized by white, pigment-free skin, mostly gradually growing patches with a hyperpigmented edge, is attributed to absence or loss of melanocytes that finally leads to a loss of the melanin synthesis. In addition to cutaneous pigmentary disorders, loss of pigmentation also affects the other organs of the body, such as eyes, ears and hair, thus, suggesting that these pigments have functions in addition to providing skin color. For example, loss of pigmentation in eyes, which might be due to aging or other reasons, results in macular degeneration, whereas in ears it results in ototoxicity.

Skin physiology has also been shown to be altered in individuals with high Body-Mass Index (BMI). Particularly, obesity affects the skin barrier function, development of the sebaceous glands and sebum production, sweat glands, lymphatics, collagen structure and function and wound healing, micro- and macrocirculation, and the distribution of the subcutaneous fat. Speaking generally, obesity aggravates existing skin disorders, for example, elevation of the levels of androgens and other hormones frequently seen in obese patients activates function of the sebaceous glands (Deplewski & Rosenfield, 1999). The most common dermatological manifestation of obesity is an acanthosis nigricans that is characterized by symmetric, velvety, hyperpigmented plaques mostly seen in the axilla, groin and posterior neck areas. Obesity is also associated with alteration of the collagen structure and function and impaired wound healing. A study of
obese mice demonstrated slower wound healing and decreased wound collagen deposition (Goodson & Hunt, 1986), which could be related to the structural changes in subcutaneous adipose tissue.

This is the first time the presence of melanin granules have been documented in the adipocytes of adipose tissue obtained from morbidly obese subjects. In this study the enzymes required for pigmentation was shown to be present in human adipose tissue. The presence of melanin in adipocytes implies that the changes in adipose tissue associated with obesity may be due to peripheral action of α-MSH, the main melanocortin peptide required for pigmentation and which also plays a critical role in energy homeostasis. Study of the regulation of this pathway in adipose tissue may elucidate the pathogenesis of the obesity and the secondary consequences of this metabolic disorder.
Chapter I

Obesity

Obesity is a multifactorial chronic disease involving environmental, genetic, physiologic, metabolic, behavioral, and psychological components. Obesity is defined as a condition of excess body fat, and is associated with a large number of debilitating and life-threatening consequences, particularly, cardiovascular, metabolic and other noncommunicable diseases. Obesity has been increasing at an alarming rate world-wide over the past two decades to the extent that it has become pandemic. In the United States, obesity is considered as the second leading cause of preventable death and affects every segment of the population, including men, women and now even children.

Obesity is defined as an increase in the amount of adipose tissue in the body. For the past fifty years both visceral and subcutaneous adipose tissues were considered as simple depots of triglycerides. Recently, the complex metabolic endocrine functions performed by adipose tissue has become increasingly clear. In other words, adipose tissue can be defined as a very active, multifunctional endocrine organ which is involved in intense signaling crosstalk with other peripheral tissues (Kershaw & Flier, 2004). It acts as a site for mediation of chronic inflammation through the action of known pro-
inflammatory adipocytokines (Degawa-Yamauchi et al., 2003; McTernan et al., 2002).

Obesity can be characterized by a state of chronic low-grade inflammation (Yudkin et al., 1999; Das, 2001; Festa et al., 2001).

**Morphology of Adipose Tissue**

Adipose tissue could be described as loosely connected tissue that functions as the major storage site for the fat in the form of triglycerides. The major bulk of adipose tissue is a loose association of lipid-filled cells called adipocytes, which are held in a framework of collagen fibers. In addition to adipocytes, adipose tissue contains stromal vascular cells, fibroblastic connective tissue cells, leukocytes, macrophages, and pre-adipocytes (not yet filled with lipid), which contribute to structural integrity. The size of adipose tissue is a function of both adipocyte number and size (Johnson & Greenwood, 1988).

The adipose tissue can be differentiated into two types depending on the structure and composition of adipocytes: white adipose and brown adipose tissue. Brown adipose tissue is present in human newborns and in hibernating mammals; its primary purpose is to generate body heat. The white adipose tissue makes the omental fat, present in the deep abdomen, and serves many other functions besides acting as storage of triglycerides. Brown adipose tissue is more vascularized than white adipose tissue. Furthermore, brown adipose tissue differs from white adipose tissue in terms of composition of adipocytes: adipocytes in brown adipose tissue have multilocular lipid droplets whereas adipocytes of white adipose tissue have one big unilocular lipid droplet. Morphologically, nuclei and
the cytoplasm of adipocytes of white adipose tissue are pushed to the periphery of the cells whereas the central vacuole is filled with a big lipid droplet.

Adipocytes originate from mesenchymal multipotent stem cells that develop into adipocyte precursor cells (usually termed preadipocytes) by largely unknown mechanisms. *In vitro*, isolated preadipocytes can be stimulated to differentiate into mature adipocytes by treatment with specific adipogenic factors that activate PPARγ/RXR receptors (Ailhaud & Hauner, 1997; Spiegelman & Flier, 1996). After differentiation from fibroblast-like preadipocytes, adipocytes start accumulating lipids once they reach critical size. Increases in adipocyte turnover rate, differentiation and apoptosis can result in the weight gain or loss. Substantial weight gain, or an increase in body mass index (BMI) to ≥ 30, is known as obesity - a metabolic disorder, associated with multiple secondary complications.

**Adipose tissue changes in obesity**

During obesity, adipose tissue undergoes drastic changes besides an increase in size and number of adipocytes. Histologically, adipose tissue of obese individuals is characterized by infiltration by mononuclear cells, relative rarefaction of blood vessels and neural structures (Wellen & Hotamisligil, 2003). Structurally, adipose tissues are less vascularized under diseased conditions as compared to normal conditions. Obesity is also often accompanied by chronic inflammation; this results in reduction of the cross sectional area of blood vessels per unit weight of adipose tissue which leads to the reduction in blood flow as compared to non-obese conditions (Summers *et al.*, 1996;
Summers *et al.*, 1999). Surprisingly, in terms of energy homeostasis, obese subjects found to be hyperglycemic and hyperinsulinaemic as compared to lean group show the same glucose uptake by the adipose tissue (Coppack *et al.*, 1996). So, instead of being absorbed or transported to the adipose tissue, obesity leads to an increase of the concentration of glucose in the blood.

In addition to all the factors discussed above, various paracrine and endocrine factors which include several hormones, cytokines and adipokines are differentially expressed under the obese conditions. Various epidemiological studies have established a direct association between the increased expression and secretion of cytokines and the development of a variety of pathological processes, particularly, insulin resistance, glucose intolerance, dyslipidaemia, elevated blood pressure, impaired fibrinolysis and endothelial dysfunction (Bjorntorp, 1991). Obesity is also associated with a constellation of risk factors that are nowadays recognized as components of what is now called Metabolic Syndrome (MS) (Reaven, 1988; Bjorntorp, 1991; Hauner, 2002). Metabolic syndrome is commonly accompanied by changes in the secreted adipokines that are regulated both on the mRNA and protein levels. This indirectly indicates that adipokines could serve as major players in pathogenesis of metabolic syndrome, as they modulate complex networks of homeostatic, immunological and inflammatory processes. Changes in cytokine levels produced by immune cells infiltrating adipose tissue also have deleterious consequences on such processes. For example, Esposito *et al.* found that levels of some anti-inflammatory cytokines, such as IL-10, are decreased in metabolic syndrome patients, but remain normal in healthy obese controls (Esposito *et al.*, 2003).
Table 1: Classification of BMI according to World Health Organization:

<table>
<thead>
<tr>
<th>Terminology</th>
<th>BMI, Kg/m², Range</th>
<th>WHO classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Underweight</td>
<td>&lt;18.5</td>
<td>Underweight</td>
</tr>
<tr>
<td>Normal</td>
<td>18.5-24.9</td>
<td>Normal range</td>
</tr>
<tr>
<td>Overweight</td>
<td>25.0-29.9</td>
<td>Preobese</td>
</tr>
<tr>
<td>Obesity class 1</td>
<td>30.0-34.9</td>
<td>Obese class 1</td>
</tr>
<tr>
<td>Obesity class 2</td>
<td>34.9-39.9</td>
<td>Obese class 2</td>
</tr>
<tr>
<td>Obesity class 3</td>
<td>&gt;40.0</td>
<td>Obese class 3</td>
</tr>
</tbody>
</table>

Correlation between obesity, levels of the melanocortins and metabolic disorders

To maintain energy homeostasis by balancing caloric intake to caloric expenditure, mammals have developed highly adaptive and intriguingly redundant mechanisms. Loss of this balance may lead either to anorexic dystrophy or to obesity. Obesity is characterized by an excess of fat mass that predisposes one to the development of insulin resistance and other secondary complications like non-alcoholic fatty liver disease, coronary artery disease and hypertension. Obesity resulting in the alteration of energy homeostasis is rapidly reaching epidemic proportions all over the world.

The central melanocortin system is the best-characterized neuronal pathway that is involved in the regulation of energy homeostasis (Ellacott & Cone, 2004). It is
considered a promising target for the treatment of eating disorders and obesity (MacNeil et al., 2002). The melanocortin system refers to a set of hormonal, neuropeptidergic, and paracrine signaling pathways that are defined by five G protein-coupled melanocortin receptors (MC-Rs), peptide agonists derived from the proopiomelanocortin (POMC) preprohormone precursor and their endogenous antagonists, agouti and agouti-related protein. The location of POMC neurons (arcuate nucleus of the hypothalamus, nucleus tractus solitarius of the brain stem) and AgRP neurons (arcuate nucleus of the hypothalamus) suggests their role in the regulation of satiety and energy expenditure (Ellacott and Cone, 2004).

**Obesity and inflammation:**

Adipose tissue can be described as an endocrine organ that produces various cytokines, aromatized steroid hormones and protein adipokines like leptin and adiponectin. More than 100 soluble factors have been identified as produced by and released from adipose tissue; these adipokines exert multiple effects at both the local and the systemic level by modulating many different metabolic processes (Hauner, 2005). Any change in the mass and the composition of adipose tissue associated with Metabolic Syndrome are accompanied by changes in secreted adipokines that are regulated both on the mRNA and protein levels. This indirectly indicates that adipokines could serve as major players in pathogenesis of metabolic syndrome, as they modulate complex networks of homeostatic, immunological and inflammatory processes. For example, levels of some anti-inflammatory cytokines, such as IL-10, are decreased in metabolic
syndrome patients, but remain normal in healthy obese controls (Esposito et al., 2003), while the production of the inflammatory peptides is increased in the sickest cohort of the morbidly obese patients (Jarrar et al., 2008). In addition, there is a link between low levels of circulatory adiponectin and high systolic blood pressure (Huang et al., 2003). Recent studies revealed that obesity induces ER stress and that the latter in turn activates an inflammatory response, thus contributing to insulin resistance in the liver and adipose tissue (Ozcan et al., 2004; Nakatani et al., 2005). The general assumption is that inflammation is consequent to obesity, but obesity has also been suggested to be infact, a result of inflammatory disease (Das, 2001).
Fig (1): Expansion of the adipose tissue during weight gain leads to the recruitment of macrophages through various signals. Both macrophages and adipocytes are involved in the secretion of soluble proteins. LPL, lipoprotein lipase; PAI-1, plasminogen activator inhibitor-1; CRP, C-reactive protein; IL, interleukin; NGF, nerve growth factor; TGF-β, transforming growth factor-β; TNF-α, tumor necrosis factor-α; MCP-1, monocyte chemoattractant protein 1; VEGF, vascular endothelial growth factor.

Leptin

Leptin, the most studied hormone secreted by adipocytes, helps to maintain a balance between calorie intake and calorie expenditure. It is the best characterized peripheral signaling molecule that interacts with both orexigenic and anorexigenic
pathways (Trayhurn et al., 1999). The leptin receptor is expressed in several regions of
the central nervous system (CNS), including the hypothalamus (Tartaglia et al., 1995), as
well as in a number of peripheral tissues. The hypothalamus is considered as primary
location where leptin acts to inhibit feeding (Jacob et al., 1997; Satoh et al., 1997; Tang-
Christensen et al., 1999b). This energy regulation is mediated through hypothalamic
neuropeptide-POMC neurons in the hypothalamus where leptin induces catabolic effects
(Mizuno et al., 1998, Woods et al., 1998; Schwartz et al., 1999). Thus, leptin, along with
insulin which also has direct actions in the CNS (Schwartz et al., 1994; Woods et al.,
1996), functions as a negative feedback signal to the CNS to regulate energy balance.
Particularly, leptin inhibits the actions of neuropeptide Y, melanin concentrating hormone
(MCH), orexin A, agouti-related peptide and cannabinoid systems (Meister, 2000; Di
Marzo et al., 2001), whereas POMC and cocaine- and amphetamine-regulated transcripts
(CART) are upregulated by leptin (Schwartz et al., 1996; Kristensen et al., 1998; Meister,
2000). Mutations in the genes encoding leptin (Montague et al., 1997; Strobel et al.,
1998) or the leptin receptor (Clement et al., 1998) results in an obese phenotype both in
mice (ob/ob) and humans.

α-MSH has been shown to inhibit leptin secretion in differentiated rat adipocytes
cultured in vitro (Hoggard et al., 2004). Leptin administered to ob/ob mice increases the
release of α-MSH into the circulation, suggesting a possible feedback loop between the
sites of α-MSH release and the release of leptin from the adipose tissue (Hoggard et al.,
2004; Norman et al., 2003). The physiological significance of this putative feedback
probably depends upon the underlying state of energy balance, since low plasma levels of α-MSH in fasting animals are paralleled by low plasma leptin (Hoggard et al., 2004).

**Adiponectin**

Adiponectin is one of the most abundant protein hormones secreted into the bloodstream by adipocytes. It plays a role in the suppression of inflammation-associated metabolic disorders, particularly, of type II diabetes, obesity, and atherosclerosis. Serum levels of the adiponectin show a distinct decrease in circulating levels in obesity and an inverse correlation with BMI. Adiponectin possesses antidiabetic properties due to its insulin-mimetic and insulin-sensitizing characteristics, its anti-inflammatory and anti-atherosclerotic effects have also been consistently reported (Hotta et al., 2000). Additionally, a number of epidemiological studies have implicated adiponectin in modulating angiogenesis, and cardiac remodeling (Spranger et al., 2003; Yamauchi et al., 2001; Maeda et al., 2002). These effects are thought to be mediated by adiponectin binding to its cell surface receptors (Yamauchi et al., 2003; Hug et al., 2004) and subsequent activation of signaling pathways within the target cell (Yamauchi et al., 2003). Anti-inflammatory properties of the adiponectin play a critical role in reducing TNF-α production, as it antagonizes several of the inflammatory effects of TNF-α (Ouchi et al., 2003). In contrast, the pro-inflammatory agents produced in the obese subjects attenuate the production of adiponectin (Ouchi et al., 2003). Clinically, plasma adiponectin levels inversely correlate with levels of C-reactive protein in humans, which
is another inflammatory agent secreted by adipose tissue of individuals with high BMIs (Ouchi et al., 2003).

Adipocyte precursors have potent phagocytic capacity and can be transformed into macrophage-like cells in response to appropriate stimuli (Charriere et al., 2003). Adiponectin plays a critical role in this process, as it promotes the clearance of early apoptotic debris (Takemura et al., 2007). In macrophages, adiponectin stimulates clearance of early apoptotic bodies by their engulfment and suppresses TNF-α production. Slower clearance of such debris might exacerbate symptoms of inflammation and autoimmunity (Savill et al., 2002; Potter et al., 2003), as the delay in the engulfment of apoptotic bodies leads to the production of proinflammatory cytokines (Savill et al., 2002).

Levels of adiponectin in the serum are inversely correlated with the size of the central (visceral) depot of the fat. Once established, these levels cannot be altered by lowering of leptin level via fasting or external administration of leptin (Gavrila et al., 2003). Peripheral administrations of melanocortin receptor agonist-like peptides, for example, melanotan II has been shown to upregulate the expression of adiponectin receptors 1 and 2, whereas no effect was seen on the level of adiponectin in the serum (Blüher S et al., 2004). Serum level of α-MSH was found to be higher in obese individuals as compared to lean subjects (Hoggard et al., 2004), but serum levels of adiponectin remained low.
**Resistin**

Human studies have also highlighted substantial expression of resistin in adipose tissue (Savage *et al.*, 2001), particularly in abdominal depots (McTernan *et al.*, 2002). Furthermore, a positive correlation between serum resistin and body fat content has also been reported (Zhang *et al.*, 2002). Resistin expression is positively correlated with the BMI and visceral fat area (Azuma *et al.*, 2003; Vozarova de Courten *et al.*, 2004), which is mainly due to an increase in adipocyte number and differentiation (Asensio *et al.*, 2004). According to one hypothesis, adipose tissue in obese subjects is under continuous infiltration pressure by macrophages that also contributes to resistin expression. The infiltration itself further propagates the recruitment of more macrophages, which perpetuates an increase in resistin expression. Alternatively, macrophage recruitment may arise from trans-differentiation of pre-adipocytes into macrophages-like cells (Lehrke *et al.*, 2004; Kaser *et al.*, 2003), which may also result in high resistin expression.

Resistin is postulated to contribute to insulin resistance and other inflammatory responses; some correlations may exist between levels of resistin and other inflammatory factors produced by obese subjects. One such study has shown that in obese patients with obstructive sleep apnea syndrome, resistin levels are significantly correlated with levels of IL-6 and ICAM-1 (intercellular cell-adhesion molecule-1) (Kaser *et al.*, 2003). Correlations between expression and secretion of resistin and other inflammatory cytokines, including IL-6, Leptin, TNF-α and CRT have also been reported in patients with severe inflammatory disease, obesity and type II diabetes (Lehrke *et al.*, 2004).
Resistin expression in humans has been shown to be stimulated by LPS, IL-1, TNF-α and IL-6 (Mc Ternan et al., 2003; Vendrell et al., 2004; Wellen & Hotmisligil, 2003).

**Cytokines**

In addition to adipokines, white adipose tissue secretes cytokines as well as other soluble proteins directly involved in lipid metabolism, the complement system and vascular homeostasis. The main consideration is the regulation of the cytokine production by adipokines (Ahima & Flier, 2000). The wide variety of cytokines secreted by white adipose tissue includes TNF-α, IL-6, IL-1β, IL-8, IL-10, IL-18, IL-17D and TGF-β. Circulating levels of all these cytokines except IL-10 increases with obesity and decreases with weight reduction (Esposito et al., 2003) and can be correlated to the amount of adipose tissue, especially the visceral fat.

The earliest identified secretory protein with autocrine effects recognized as the product of adipocytes was lipoprotein lipase responsible for the breakdown of circulating triacylglycerols (chylomicrons and VLDL) to fatty acids. White adipose tissue is also a major source of angiotensinogen, which regulates blood pressure and stimulates the production of prostacyclins acting as a signal for the differentiation of preadipocytes to adipocytes (Zorad et al., 1995; Ailhaud et al., 2000). The circulating levels of angiotensinogen are higher in obesity is correlated with an increase in adipose tissue mass.
TNF-α was the first inflammatory cytokine described as expressed in and secreted by adipocytes of white adipose tissue (Hotamisligil et al., 1993). Besides its critical role in host defense, it also plays a major role in lipid and glucose metabolism (Pennica et al., 1984). Even though TNF-α mRNA is expressed by the adipocytes of white adipose tissue, its secretion by other infiltrating cells of adipose tissues, for example, macrophages and vascular fraction, has also been documented (Weisberg et al., 2003; Fain et al., 2004a). The serum levels of TNF-α are significantly correlated with visceral adiposity (Hotamisligil et al., 1993). Administration of TNF-α to experimental animals has been shown to induce insulin resistance (Lang et al., 1992) whereas inhibition of TNF-α by a blockade of TNF-α bioactivity in vivo (Hotamisligil et al., 1993; Ofei et al., 1996; Cheung et al., 1998) or genetic knockout of the TNF receptor/ligand (Uysal et al., 1997; Schreyer et al., 1998; Ventre et al., 1997) improves insulin sensitivity and glucose metabolism. Moreover, TNF-α acts on murine adipocytes in vitro to suppress expression of many adipose-specific genes, including the enzymes involved in lipogenesis (Kawakami et al., 1986). Inactivation of the lipogenesis enzymes leads to hyperlipidemia by increasing the level of triglycerides (Semb et al., 1987; Sherman et al., 1988), and can be the main cause of hyperlipidemia and hyperinsulinemia in obese subjects.

TNF-α also plays a pivotal role in the production of several cytokines and other adipokines (Coppack, 2001). Nowadays, TNF-α, in addition to its role in inflammation, is recognized as a multi-functional regulatory cytokine, implicated in other cellular processes, particularly, in cell apoptosis and survival, cytotoxicity, production of acute-phase protein, haptoglobin and other cytokines, such as IL-1 and IL-6 (Chiellini et al.,
TNF-α strongly stimulates the expression of IL-18, IL-8 and IL-1β by fat cells (Wood et al., 2005).

Recent developments in the area of the role of central nervous and neuroendocrine systems in host responses (Besedovsky & del Rey, 1992; Goetzl & Sreedharan, 1992) have provided some new insights about the niche of the neuronal and soluble mediators in these systems that influence inflammation. One of these mediators is the endogenous neuropeptide α-MSH which is believed to modulate host reactions by inhibiting the actions of cytokines (Lipton, 1990; Catania and Lipton, 1993). Both α-MSH and its synthetic analogues exert potent anti-inflammatory activity when administered systemically (Deeter et al., 1988; Hiltz & Lipton, 1989, 1990; Catania & Lipton, 1993; Hiltz et al., 1992). For example, one recent study demonstrated that systemically administered α-MSH inhibits TNF-α concentrations circulating in the brain. TNF-α production by human peripheral blood mononuclear cells is inhibited by α-MSH, which indicates that a direct effect of the peptide is mediated through melanocortin receptors on monocytes. In mice, the level of TNF-α induction after intraperitoneal injection of lipopolysaccharides (LPS) was found to be reduced by the simultaneous injection of α-MSH or its analogues (Gonindard et al., 1996). In addition, α-MSH has the properties to modulate the cytokine regulatory properties of TNF-α. Recently α-MSH has been shown to inhibit TNF-α induced expression of the matrix metalloproteinase-13 in human chondrosarcoma cells by modulating NF-κB signaling (Yoon et al., 2008).
IL-18 gene expression was registered in human subcutaneous and visceral adipose tissue, both in mature adipocytes and the stromal-vascular fraction. In cultured human adipocytes, IL-18 mRNA levels dramatically rise in response to TNF-α (Wood et al., 2005). When IL-18 operates at a local level, its proinflammatory action invokes a cascade of inflammatory events involving cytokines, chemokines and adhesion molecules (Gracie et al., 2003). IL-18 has been also shown to stimulate the expression of IL-6 and IL-8 in human lung carcinoma cell line (Kim et al., 2005). Interestingly, recent reports indicate that under specific conditions IL-18 may act as both an angiogenic activator (Qiao et al., 2004) and angiogenic suppressor (Cao et al., 1999).

IL-8 is another chemokine secreted by adipocytes. It has been implicated in artherosclerosis and coronary heart disease (Ross, 1999) as it plays a role in modulating an inflammatory response. IL-8 possesses chemoattractant properties as it serves to recruit the neutrophils, T lymphocytes, induces the adhesion of monocytes to endothelium and promotes the migration of vascular smooth muscle cells (Yue et al., 1993). All these events lead to intimal thickening resulting in artherosclerosis. Initially IL-8 was thought to be produced by macrophages and monocytes only (Baggiolini, 1995). Recently two in vitro studies by Bruun et al. revealed that IL-8 is produced and secreted by human adipocytes (Bruun et al., 2000; Bruun et al., 2001). Other studies have shown that oxidized low-density lipoprotein particles are also able to stimulate production and secretion of IL-8 by macrophages embedded in the adipose tissue (Liu et al., 1997). Therefore it can be speculated that a correlation between the severity of obesity and the
development of secondary complications of this disorder, particularly the atherosclerosis and cardiovascular disease, is partially due to an increase in IL-8 levels.

**Underlying pathology behind inflammation**

**Lipid peroxidation and reactive oxygen species**

Lipid peroxidation refers to the oxidative degradation of lipids. This free radical-generating process occurs on every membrane containing structure of the cell. Actually, the peroxidation of the lipids is a major contributor to the ROS production. Obesity can be considered as a major cause of increased lipid peroxidation including oxidation of cell membranes and proteins in conjunction with disturbances of cellular redox homeostasis (Shattock *et al.*, 1994; Girotti *et al.*, 1998; McDuffee *et al.*, 1997). Free radicals produced during lipid peroxidation are known to be involved in a number of human pathologies including atherosclerosis (Steinberg, 1997), cancer (Cerutti, 1994) and hypertension (Russo C *et al.*, 1998). Recent evidence has shown that obesity is also associated with increased myocardial lipid peroxidation and susceptibility to oxidative damage *in vitro* (Vincent *et al.*, 1998). Increased production of ROS has been documented in myocardium muscles in obese subjects, which may be due to an increase in mechanical and metabolic load leading to increased consumption of oxygen and finally resulting in the formation of hydrogen peroxide from mitochondrial respiration (Turrens *et al.*, 1997). In the heart, significant oxidative injury produced by lipid peroxidation can ultimately lead to cardiac arrhythmias, poor contractility, infarction, cardiac failure or sudden death (Yu, 1994; Shattock & Haddock 1991).
Low density lipoprotein oxidation has been reported as the prominent cause of atherosclerosis (Witztum et al., 1991). Lipid accumulation in steatotic liver is also associated with a phenomenon of peroxidation (MacDonald, 2001) analogous to that observed in LDL of the atherosclerotic lesion (Glass & Witztum, 2001). In both organs the oxidized LDL particles are taken up by macrophages which act as a chemotactic aid for monocytes and vascular smooth muscle cells, simultaneously inducing their proliferation, partially by stimulating expression of adhesion molecules, cytokines and growth factors. Besides this, oxidized LDL also induces humoral and cell-mediated immune responses (Witztum, 1994; Jialal & Devaraj, 1996).

An excess of the adipose tissue is associated with increased release of cytokines that in a number of ways stimulate the generation of ROS. Although cells might produce both enzymatic and non-enzymatic antioxidants to help remove ROS, in morbidly obese subjects the pool of these antioxidants gets rapidly depleted leading to an increase in cellular stress due to continuous production of ROS (Farshad Amirkhizi et al, 2007). The consequence of the ROS overproduction in human obesity is constant oxidative stress resulting in progressive tissue damage that attracts more ROS-producing cells of the immune system, thus, forming a vicious circle.

**Hypoxia**

Hypoxia occurs when oxygen availability does not match the demand of the surrounding tissue, resulting in decreased oxygen tension. Hypoxic stress plays a pivotal role in normal human development and physiology, including embryogenesis and wound
repair. Hypoxia has been well studied for its importance in the pathogenesis of several human diseases, including heart disease, stroke, diabetes, and cancer (Semenza, 2000). Hypoxia is known to increase the mRNA expression of HIF-1 (Hypoxia inducible factor-1) and a wide variety of other genes that stimulate erythropoiesis, angiogenesis, and glycolysis (Semenza, 2000). Besides engaging the HIF-1-dependent pathway, hypoxia also activates HIF-1 independent pathways that contribute to the cellular adaptation to this type of the stress (Bi et al., 2005).

Obesity can be morphologically described as hypertrophy and hyperplasia of adipocytes. Adipocytes under normal conditions have a limited capacity for hypertrophy; one reason for this is the diffusion limit of oxygen, which is at most 100 µm (Helmlinger, 1997). The resistance of adipose tissue to hypertrophic and angiogenic stimuli is one of the reasons why white adipose tissue is not richly vascularized. In obesity, adipocytes become hypertrophic as their size increases up to 140–180 µm in diameter. Therefore, hypertrophic adipocytes have to endure a less-than-adequate oxygen supply, which suggests the existence of hypoxic conditions in the adipose. Hypoxia has now been directly demonstrated in adipose tissue of several obese mouse models (ob/ob, KKAy, diet-induced) and molecular studies indicate that the level of the HIF-1 is increased, as is expression of the hypoxia-sensitive marker gene, GLUT1 [Ye et al., 2007; Trayhurn et al., 2008]. Hypoxia is known to have a great impact on the expression of genes encoding adipokines, including angiopoietin-like protein 4, IL-6, leptin, macrophage migration inhibitory factor and vascular endothelial growth factor (Trayhurn et al., 2008; Hosogai
et al., 2007). One of the well documented examples in hypoxia-dependent degradation is degradation of adiponectin mRNA in adipocytes (Hosogai et al., 2007).

Hypoxia in adipose tissue has at least two major effects on obesity-associated chronic inflammation. First, it induces the adipocyte expression of cytokines, for example, MIF and MCP-1 factors that promote macrophage infiltration in the adipose tissue. Second, hypoxic environment stimulates macrophages, which are much more active than adipocytes in the production of inflammatory cytokines (Kadowaki et al., 2006). The hypoxia may also increase lipolysis that leads to the release of free fatty acids (FFA) from adipocytes, which later on also join the cycle of the endocrine alteration and inflammation. Hypoxia also directly leads to an increase of the levels of mRNAs of ER stress marker genes, CHOP and GRP78 (glucose-regulated protein, 78 kD) (Hosogai et al., 2007). In conclusion, hypoxia can impair adipocyte function and secretion of adipokines. Therefore, hypoxia may contribute to the development of the secondary complications of obesity, particularly, insulin resistance and metabolic syndrome (Wellen & Hotamisligil, 2005).

**Cutaneous manifestations of obesity**

Obesity is associated with a variety of dermatoses, including acanthosis nigricans, acrochordons, keratosis pilaris, hyperandrogenism and hirsutism, striae distensae, and adiposis dolorosa, and as well with fat redistribution.
Acanthosis nigricans is the most common dermatological manifestation of obesity. It is traditionally characterized by hyperpigmented, velvety thickening of epidermis that primarily affects the axillae, posterior neck fold, flexor skin surfaces, and umbilicus, and infrequently is diffuse with involvement of the mucosal surfaces of body folds (Schwartz, 1994). This increased pigmentation may range from a brown to gray color depending on the underlying skin tone. It is typically found in obese persons, individuals with type II diabetes, or those with a family history of diabetes mellitus (Gibson, 2004), however, its prevalence and significance in an unselected adult obese population is unknown.

Acanthosis nigricans is a reliable cutaneous marker of hyperinsulinemia, a consequence of insulin resistance that is associated with obesity (Katz et al., 2000). Obese children with acanthosis nigricans have also been shown to have insulin resistance (J.F. Fu, et al., 2004). The blood glucose levels may or may not be elevated as the high level of insulin is often able to maintain glucose homeostasis in the presence of insulin resistance (Katz et al., 2000). The proposed mechanism of how hyperinsulinemia leads to this epidermal change begins at the cellular level. Increased levels of circulating insulin lead to a decrease in the number of functional insulin receptors (Rendon et al., 1989; Cruz & Hud 1992). These “classic” insulin receptors regulate glucose uptake, cell growth, DNA synthesis, and protein and fat metabolism via tyrosine kinase activity. Keratinocytes and fibroblasts both express insulin-like growth factor (IGF) receptors that are also capable of binding insulin and have growth-promoting effects (Hermanns-Le et
al., 2004). Decreased numbers of functional insulin receptors cause a shift to increased binding to IGF receptors contributing to the development of acanthosis nigricans.

As could be seen from what is written above, the current explanation for the connection between insulin resistance and an increase in the cutaneous pigmentation is rather vague. There is a possibility for other, not yet unidentified factors produced by adipose and influencing the skin. There are indications of significant crosstalk between adipocytes and the melanocortin system, which point to the existence of novel pathways in adipocytes. Study of these pathways may elucidate the pathogenesis of the metabolic disorders.

**Melanocortins:**

**POMC (proopiomelanocortin)**

*POMC* gene encodes a polypeptide hormone precursor that undergoes extensive and tissue-specific post-translational processing via cleavage by specific enzymes known as prohormone convertases. This gene is actively transcribed in several tissues, including the corticotroph cells of the anterior pituitary, neurons originating in the arcuate nucleus of the hypothalamus, and cells in the dermis and the lymphoid system (Bertagna, 1994). The POMC-related peptides ACTH and β-endorphin have also been documented in the cardiac tissue of rat. There is a possibility that POMC is synthesized by cardiomyocytes, but not by cardiac neurons (Forman & Bagasra 1992; Millington, 1993). Besides that, not
much is known about their cellular localization and function in heart (Saito 1983; Forman et al., 1989; Forman et al., 1994).

The primary site of POMC expression, the pituitary gland, consists of two POMC-expressing lineages: Corticotroph and Melanotroph. The single copy of POMC gene is expressed in both pituitary lineages but under different developmental and hormonal regulatory pathways. At the cellular level, this regulation is exerted by transcription factors that regulate both POMC transcription and cell differentiation. The corticotroph lineage constitutes the central link integrating signals from the brain and periphery that serves as a regulator of the production of glucocorticoids. In the cells of this lineage POMC is processed into adrenocorticotropicin (ACTH). In the melanotrophs the POMC is processed further, so α-melanocyte stimulating hormone (α-MSH) is produced instead of ACTH.

Apart from differential expression of POMC in several tissues as mentioned above, other mechanisms of POMC regulation exist. First, POMC could be regulated at the transcriptional level through the formation of truncated and non-truncated forms of POMC mRNA. The 5’ truncated POMC transcript lacks the POMC signal sequence encoded by exon 2 of the POMC gene, so it is unlikely to undergo translation (Jeannotte L, 1987; Lacaze-Masnoteil T, 1987; Clark AJL, 1990). In the testes, ovaries, pancreas and other peripheral tissues expressing the POMC gene this short POMC transcript is far more abundant that full-length POMC mRNA (DeBold CR, 1988; Hummel A, 1994;
Ivell R, 1994). In these tissues POMC peptide concentrations correlate with full-length, not 5’ truncated, POMC mRNA levels (DeBold CR, 1988).

Second, POMC production is regulated at the posttranslational level. POMC undergoes extensive and tissue specific posttranslational processing to yield a range of biologically active peptides. The POMC gene produces a 32 kDa propeptide, which is targeted via a specific signal into regulated secretory granules when passed through the Golgi stacks (Cool et al. 1997). The processing of POMC depends on the range of processing enzymes present in the tissue. For example, in pituitary, expression of prohormone convertase 1 (PC1), but not PC2, results in the production of only N terminal peptide, joining peptide, ACTH and β-lipotropin (White and Gibson 1998). In contrast, expression of PC2 in hypothalamus in addition to PC1 leads to the production of CLIP, β and γ-MSH due to its action on ACTH and gamma-LPH respectively (Bertagna et al., 1988).

In addition to prohormone convertases and their endogenous inhibitors in hypothalamus, other enzymes play a key role in generating mature POMC-derived peptides. The generation of mature α-MSH from ACTH1-17 is catalysed by carboxypeptidase E (CPE), peptidyl α-amidating mono-oxygenase (PAM) and N-acetyltransferase. ACTH is absent in the hypothalamus due to the presence of PC2, PAM and CPE enzymes which process the ACTH into smaller peptides. In this process, carboxy-terminal basic amino acids are trimmed from ACTH1-17 by CPE which is subsequently amidated by PAM to generate desacetyl α-MSH (DA-αMSH), which is
Further converted to mature form of α-MSH by N-acetyltransferase. Besides pituitary and hypothalamus, POMC and POMC derived peptides undergo extensive post-translational processing in cardiac tissue (Millington, 1993).

Peptides derived from POMC serve a wide range of functions such as skin pigmentation and energy homeostasis (Hadley & Haskell-Luevano, 1999; Cone, 2005). In addition to this, POMC and POMC-derived peptides regulate a remarkably diverse array of physiological functions including adrenocortical steroidogenesis, natriuresis, erectile responses, and exocrine gland secretion and also exerts immunomodulatory effects.

The biological effects of POMC-derived peptides are diverse. Mutations in POMC in humans result in metabolic disorders, such as early onset of obesity due to adrenal insufficiency (Krude et al., 1998) and hyperphagia (Krude and Gruters 2000). Furthermore, POMC also serves as a quantitative trait locus (QTL) in human obesity (QTL is a genetic factor that interacts with other genes and environmental factors to predispose to complex human phenotypes) (Comuzzie et al., 1998). Mutations in POMC have also been involved in the development of the different skin color phenotypes due to switch from eumelanin to pheomelanin resulting in the formation of pale skin and red hair pigmentation. Besides, mutations in enzymes required for POMC processing, for example, PC1, also disrupt melanocortin signaling, as lack of α-MSH generation in the hypothalamus ultimately results in obesity (Jackson et al., 1997).
Melanocortin receptors

The biological effects of POMC-derived peptides are largely mediated through melanocortin receptors (MCR). To date, five melanocortin receptor genes have been cloned and characterized. All five receptors are G-protein-coupled receptors with seven transmembrane domains, which mediate cell signaling by activating cAMP-dependent
pathways. These receptors differ in their tissue distribution and in their ability to recognize various melanocortins and their physiological antagonists.

MC1R was the first MC receptor cloned (Mountjoy et al., 1992; Chhajlani & Wikberg., 1992). The human MC1 receptor affinity is equally high for α-MSH and ACTH and lower for β-MSH and γ-MSH (Mountjoy, 1994.). Initially, this receptor was considered as peripheral receptor, and was thought to be expressed only by melanocytes, keratinocytes and melanoma cells, with its main function involved in the regulation of skin physiology and melanogenesis (Chhajlani, 1996; Eberle, 1988). However, the expression of MC1R has been documented by macrophage/monocytic cells (Star et al., 1995), lymphocytes with antigen-presenting and cytotoxic functions (Neumann et al., 2001), neutrophils (Catania et al., 1996), and dendritic cells (Becher et al., 1999), which suggests a specific role of MC1R in inflammation. It is also expressed at low levels in adipocytes (Boston & Cone, 1996) and endothelial cells (Vergoni & Bertolini, 2000). Recently, high expression of MC1R has been documented in the adipocytes of severely obese subjects as compared to adipocytes of lean subjects (Hoch et al., 2007). The authors suggested a functional role of MC1R in regulation of fat cell proliferation (Hoch et al., 2007). Function of MC1R has not been proven yet, so there is another possibility that MC1R expression in the adipocytes of morbidly obese subjects is involved in their pigmentation.

MC2R is mainly expressed in the zona fasciculata and zona glomerulosa of the adrenal cortex (Mountjoy et al., 1992). Recently expression of MC2R has been
documented in adipocytes of both subcutaneous and visceral adipose tissues (Schaffler et al., 2005). MC2R shows great affinity for ACTH peptides and mediates the production and release of steroids in the adrenal cortex (Xia & Wikberg, 1996; Buckley & Ramachandran, 1981). Hereditary isolated glucocorticoid deficiency, a rare autosomal disorder, is caused by mutations in the MC2R gene (Tsigos et al., 1993).

MC3R is expressed within the central nervous system, for example, within the hypothalamus, thalamus, hippocampus, anterior amygdala, and cortex. In the periphery it is expressed in the placenta, ovary, mammary gland, testis, gastrointestinal tract, and the heart (Chahajlani & Wikeberg, 1992; Grantz et al., 1993; Chhajlani, 1996). Recently expression of MC3R has been documented on adipocytes from both subcutaneous and visceral adipose tissues (Smith et al., 2003). The receptor shows same binding affinity for all MC peptides and plays an important role in the regulation of cardiovascular functions, thermoregulation, and feeding behavior (Getting, 2002; Low, 2004). Besides, MC3R is also involved in mediating some of the anti-inflammatory effects of α-MSH (Getting, 2002; Getting et al., 2003). Knockout mice lacking MC3R are obese, but not hyperphagic (Butler and Cone, 2003).

MC4R is expressed within the neuroendocrine system in the brain: those receptors are particularly dense in regions of hypothalamus. MC4R is reported in the rat osteosarcoma cell line, UMR-106 and in the growth plate in the rat fetus (Dumont, 2001; Mountjoy et al., 2003). Expression of MC4R has also been shown in adipocytes from both subcutaneous and visceral fat (Smith et al., 2003; Schaffler et al., 2005). Both
MC3R and MC4R have been identified as important downstream effectors regulating energy homeostasis in response to neuropeptides secreted by POMC- and AgRP-ergic neurons. MC4R shows the highest affinity for α-MSH and ACTH. This receptor is particularly involved in the control of the feeding and behavior, specifically in the paraventricular nucleus, the dorsomedial hypothalamus and the lateral hypothalamic area, which are (Mountjoy, 1994). Mutations in the MC4R gene in humans lead to hyperinsulinemia and obese phenotype (Vaisse et al., 1998), which confirms its role in appetite regulation. Even heterozygous MC4R mutations display an obese phenotype which signifies that appetite regulation is sensitive to quantitative variation in MC4R expression.

MC5R is localized to sebaceous glands, exocrine glands, hair follicles, and epidermis in human skin, adipocytes and skin mast cells as well as in cultured human sebocytes (Thiboutot et al., 2000, Slominski et al., 2000, Bohm et al, 2006). Mutations in the MC5R gene in mice result in defective water repulsion and thermoregulation as well as in reduced sebaceous gland secretion (Chen et al., 1997).

Therefore, adipocytes express all types of MC receptors. Additionally, all types of melanocortin receptors were found in bone cells (both osteoblasts and osteoclasts) and chondrocytes (Zhong et al., 2005). However, their exact function in bone tissue is not known yet.
α-MSH

α-MSH is derived from post-translational cleavage of POMC. This peptide was first discovered due to its role in pigmentation, but now is observed to be involved in various biologic processes (Lipton, 1998). In the brain, α-MSH is mainly produced by neurons in the arcuate nucleus of the hypothalamus. α-MSH could be produced by immunocompetent cells (Schiöth et al., 1997; Taylor et al., 1994; Taylor et al., 1992). Particularly, α-MSH has been discovered in lymphocytes, monocytes/macrophages (Rajora et al., 1996; Star et al., 1995), Langerhans cells and epidermal cells (Schauer et al., 1994), where it exerts immunomodulatory effects. The mechanisms by which its anti-inflammatory effects are executed are not fully understood, but they clearly involve direct action of the peptide on its receptors in peripheral inflammatory cells. Additionally, an indirect action on peripheral inflammation is thought to be mediated by stimulation of α-MSH receptors present within the brain. Genetic defects inactivating the receptors for α-MSH in experimental animals and in humans have been shown to result in obesity (Faroosti et al., 1999).

Central action of α-MSH

POMC-derived α-MSH peptides regulate the central control of feeding and energy balance (Pierroz et al., 2002) by decreasing body weight mediated by the decrease of the fat mass and improvements of glucose uptake. Intracerebroventricular administration of melanotan, a synthetic analogue of α-MSH, to morbidly obese rats has been shown to elevate serum leptin level independently of morphological changes in
white adipose tissue (Banno et al., 2007). Furthermore, an increase in serum adiponectin level and insulin sensitivity after central administration of Melanotan has also been documented (Banno et al., 2007). Therefore, activation of the central melanocortin systems with α-MSH or its analogues reduces adiposity by producing anorexigenic signals, which might result in increased insulin sensitivity through their action within the white adipose tissues. Besides an important role in appetite regulation, α-MSH has also been shown to reduce fever and acute inflammation in the skin when administered centrally. The above provides evidence suggesting that anti-inflammatory influences of neural origin that are triggered by α-MSH could be used to treat systemic inflammation.

**Peripheral action of α-MSH**

α-MSH has antipyretic and anti-inflammatory roles. Recently, peripheral action of α-MSH generated much interest, as this peptide and its synthetic analogues were shown to provide protection against ischemia/reperfusion injury in various tissues. In myocytes, α-MSH induces expression of cytoskeleton proteins and represses immune, inflammatory, cell cycle, and protein turnover mediators (Colombo et al., 2005). α-MSH modulates the production of proinflammatory cytokines like IL-1β, and IL-6 but do not block them completely (Hiltz & Lipton 1998; Lipton and Catania 1997). Similarly α-MSH also modulates the formation of NO and TNF-α, but can not completely stop the production completely, by inactivating the NF-kB pathway (Manna & Aggarwal, 1998). In senile plaques of Alzheimer’s disease patients α-MSH modulates inflammation (Galimberti et al., 1999). Another interesting consequence of α-MSH signaling is the
suppression of the collagen synthesis and deposition (Bohm et al., 2004) and modulation of the balance between MMP-1, MMP-8 and their inhibitors (TIMPs) (Wang et al., 2006), thus pointing at α-MSH as potential antifibrotic drug target.

Besides repressing the level of various inflammatory mediators, α-MSH also increases the production of anti-inflammatory cytokines like IL-10 by human monocytes (Bhardwaj et al., 1996). α-MSH also has immunosuppressive effects as it suppresses the synthesis of the endotoxin receptor CD14 present on the macrophages (Sarkar et al., 2003), induces neutrophil elastase (Manna et al., 2006), decreases the production of interferon gamma by human T-cells (Taylor, 2003) and modulates IgE synthesis by human B cells (Aebischer et al., 1994). In addition, intraperitoneal administration of a small concentration of α-MSH has been shown to modulate the circulating level of inflammatory agents like TNF-α, IL-6 and others in the serum (Hernández et al., 1999). α-MSH levels are elevated in plasma of obese men (Katsuki et al., 2000; Haggards et al. 1994) compared with non-obese subjects, suggesting possible peripheral resistance to anorexigenic action of this peptide, while peripheral administration of α-MSH to POMC-null mice leads to substantial weight loss, with direct effects on adipocytes (Yaswen et al., 1999 and Zemel & Shi, 2000). Therefore, peripheral action of α-MSH is equally important for the homeostasis of energy. Additionally, the broad anticytokine effects of α-MSH suggest that this peptide might as well regulate adipokine release, thus producing effects on a variety of obesity-related pathological phenotypes.
Chapter II

Pigmentation

Mammalian skin is pigmented due to the cellular accumulation of amorphous, opaque, insoluble mixtures of biopolymers called melanins. These pigments are synthesized in the melanocytes located in the basal layer of the epidermis, the hair bulb, and the iris. Melanins also occur naturally in the inner ear and choroid of eye (Prota, 1992). Synthesized melanin is stored in special lysosomal-like organelles called melanosomes (Raposo & Marks, 2002) present in the melanocytes. Later the melanosomes are actively exported to keratinocytes along the dendritic processes of melanocytes. Each functional epidermal melanogenic unit that produces and distributes melanin is composed of one melanocyte and approximately 36 neighboring keratinocytes (Jimbow, 1995; Ortonne, 1992). Melanin is produced from L-tyrosine, which acts as the major substrate, by special enzymes produced by the tyrosinase gene family. Melanin synthesis can be induced by exposure to ultraviolet radiation, inflammatory modulators such as leukotrienes and prostaglandins, and hormonal and growth factor stimuli such as α-melanocyte-stimulating hormone and endothelins.
Cutaneous pigmentation:

Skin, the main source of melanin, is one of the largest organs with vital function. It serves as an interface between the body and an external environment filled with stresses such as solar radiation, thermal and chemical insults. Cutaneous pigmentation plays a critical role in camouflage, mimicry and social communication. Furthermore, it also provides a basis for gender recognition, as in many mammals the epidermis of females is less melanized than that of males (Robins, 1991), which may be due to higher need of vitamin D during pregnancy and lactation (Wilson, 1990).

The amount of pigment in the skin depends on the quantity, quality, and distribution of the melanin. Cutaneous pigmentation results from exquisite interactions between various cell types in the skin. The best described of these are the interactions between the specialized melanin-synthesizing epidermal cells called melanocytes and the keratinocytes, and between the melanocytes and the dermal fibroblasts. To be more specific, skin pigmentation relies on melanocytes to provide, among other things, photoprotection and thermoregulation. Melanocytes produce melanin and transfer it to the recipient keratinocytes. This cellular interaction is subject to regulation by a wide array of factors, some of which are endocrine, while others are paracrine and/or autocrine. Many of these factors regulate both the constitutive pigmentation and the ultraviolet radiation (UVR) or inflammation-induced hyperpigmentation.
**Structure of the skin**

The human skin consists of three individual layers: epidermis, dermis and hypodermis. The epidermis is a continually renewing surface epithelial layer—the outermost layer of skin—which can be divided into stratum corneum, granular, squamous, and basal layer. Keratinocytes make up 90% of the epidermal cells. These cells arise in the basal layer of epidermis. Later on they migrate toward the surface of the skin and gradually lose their nuclei to form the stratum corneum (Slominski *et al.* 2004). The granular layer of epidermis is composed of tightly apposed keratohyaline granules and 5 to 7 layers of polyhedral cells makes the squamous layer, attached to each other by intercellular bridges that contain desmosomes. Langerhans cells are the type of dendritic cells present in squamous layer. The basal layer of epidermis is made up of cuboidal or columnar basal cells which lie along the dermoepidermal junction, attached to each other and to the overlying squamous cells by intercellular bridges with melanocytes scattered among them. Melanocytes are melanin pigment-producing specialized cells, with ample, pale, non-filamentous cytoplasm, and are not connected to each other. These cells possess large oval nuclei, a small nucleolus and pale dendritic possesses that extend between the intercellular spaces of keratinocytes. Melanocytes contain small sub-cellular membrane-bound endosomal organelles called melanosomes, where melanin is synthesized and later on packed into granules and targeted preferentially to the ends of the dendrites for transfer out to neighboring keratinocytes (Hearing 2005). A melanocytes itself rarely contains melanin granules unless the cell has been activated to produce increased
amounts of pigment. Besides producing melanin, epidermal melanocytes also maintain the skin homeostasis under hypoxic conditions (Stucker et al., 2002).

The dermis layer of the skin is a connective tissue-based structure made up of collagenous and elastic fibers, with variable number of fibroblasts scattered among them. It underlies the epidermis and is organized into a papillary and reticular region.

The hypodermis or subcutis is a loose connective tissue mainly composed of brown adipose tissue defined by fibrous connective tissue septa present underneath the dermis (Young et al., 2000; Fawcett et al., 2002).

Fig 3: Structure of skin
Types of melanin in mammals

Melanocytic melanin

Melanin is a ubiquitous biological pigment (Kollias et al., 1991). Most of the visible pigmentation in mammals results from the synthesis and distribution of two types of melanin, whereas most feather coloration in birds is due to carotenoid pigments. The two types of melanin pigments are eumelanin and pheomelanin. The eumelanin provides brown/black color phenotype and preferentially contains indolic monomers, while the pheomelanin provides contains yellow/red color phenotype and consists of benzothiazine subunits (Prota, 1988). These two pigments differ not only in color but also in the size, shape and packaging of their granules (Slominski 2004).

Both pigments provide body color, but their functions are very different from biological point of view. Eumelanin is photoprotective for the skin cells, as it possesses an intriguing and rather unique set of physicochemical properties, including strong broadband UV and visible light absorption (Meredith & Riesz, 2004); powerful anti-oxidant and free radical scavenging ability (Sarna et al., 1986); and, somewhat curiously, electrical conductivity and photoconductivity in the condensed phase (Jastrzebska et al., 1995; Rosei et al., 1996). Pheomelanin is believed to be carcinogenic, as it acts synergistically with UV radiation (Harsanyi et al., 1980). It serves as a potent UV photosensitizer that possibly contributes to increased sensitivity of fair-skinned individuals with yellow or red hair to sunburn, premature aging, and/or malignant transformation (Takeuchi et al., 2004).
Melanocytic extracutaneous melanin

Besides skin, extracutaneous melanin has been documented in eyes, ears, hair, mucosal tissues and other organs. Recently, extracutaneous production of melanin has been reported in heart (Ambani et al., 2005). These findings were supported by the discovery of the melanocytes in the adult heart, where they are localized primarily in the valvuloseptal apparatus (Mjaatvedt et al., 2005).

In the eye, melanin is found at a higher concentration than anywhere else in the human body (Potts, 1964). Ocular pigmentation is believed to be formed for lifetime during a brief period of the fetal and perinatal developmental window. In birds, the deposition of the melanin in the retina of the eye occurs primarily dural fetal development and continues for several days after hatching. The pigmented cells of the eye are non-dividing and no melanin renewal is known to occur at this location. Abnormalities in melanogenesis in eyes have been seen during the vision loss due to age-related macular degeneration (Young, 1987; Bressler et al., 1988).

The eye contains two pigmented cell types, the choroidal/uveal melanocytes and the retinal pigment epithelium (RPE) cells, which forms melanosomes indistinguishable from those in the skin melanocytes. RPE forms the outermost layer of retina and is composed of a monolayer of polygonal cells which are continuous cranially with the pigmented epithelial layer of the ciliary body and epithelium of iris. Both in RPE and other pigmented cells of the eye melanin shows very little turnover (Ings, 1984; Sarna, 1992). Besides melanin, the eye contains another pigment, called lipofuscin, which is also
black in color like melanin but differs in chemical composition, biosynthesis and function. Inconsistant with melanin, amount of lipofuscin increases with age.

Apart from the eye, extracutaneous melanocytes are also present in the cochlear and vestibular tissues, mostly in the cuticle, ampullae and parts of the semicircular canals of ear. They are also present near the vascular plexus of the modiolus (Franz et al., 1990). The highest number of melanocytes has been documented in the stria vascularis (Savin, 1965; Conlee et al., 1989; Cable & Steel, 1991), a well vascularized, three-layered epithelium where the pigment cells form the intermediate layer. In this layer, the pigment cells are interplaced with the secretory cells, which are responsible for the unique ionic composition of the endolymph and are in close contact with the capillary vessels. Little is known about the function of inner ear melanins, and no data are available about inner ear melanogenesis. However, an association between pigment abnormalities and hearing defects has been clearly established (Meyer zum Gottesberge, 1988), suggesting an important role for melanin and/or for the melanin-synthesizing cells. Melanocytes of the inner ear are involved in the regulation of the secretion of endolymph, the trans-epithelial ion transport, and the protection against ototoxic drugs and high-intensity noise damage (Barrenas & Lindgren, 1990).

Pigmentation of the hair is another example of extracutaneous formation of melanocytic melanin. The color of hair depends on the pigment content of the hair shaft, with melanin granules mainly found in the cortex, whereas no melanin is seen in the cells of the inner root sheath. The process of the pigmentation of hair follicles follows
sequences of events identical to those seen in epidermis and results in a terminally differentiated keratinized endproduct, the hair shaft, which eventually being shed off. Follicular melanocytes are derived from epidermal melanocytes during hair follicle morphogenesis. In contrast to epidermis, the hair follicle undergoes cyclical regeneration, and has a more complicated proliferative profile and architecture.

The "follicular-melanin unit" consists of one melanocyte for every five keratinocytes located in the hair bulb (Tobin et al., 1998). Hair bulb melanocytes are larger than epidermal melanocytes, and have longer and more extensive dendrites, contain more developed Golgi and rough endoplasmic reticulum, and produce two- to four-fold larger melanosomes (Bell, 1967; Toda et al., 1972). In hair follicles, active melanocytes reside in the wall of the pilary canal (infundibulum) and in the immune privileged proximal hair bulb, close to the upper part of the dermal papilla (Billingham & Silvers, 1971; Westgate et al., 1991), whereas the inactive melanocytes have been observed in the outer root sheath of middle and lower follicles (Staricco, 1963). The hair bulb, however, is the only site of pigment production for the hair shaft. It contains both highly melanogenic melanocytes and a minor subpopulation of poorly differentiated pigment cells (Tobin et al., 1995; Tobin et al., 1999) (Fig 4 A).
Fig 4: (A) Structure of hair. (B) Developmental stages of hair.
The hair follicle cycle consists of stages of rest (telogen), hair growth (anagen), follicle regression (catagen), and hair shedding (exogen) (Fig 4 B). The entire lower epithelial structure is formed during anagen, and regresses during catagen. Hair is formed by rapidly proliferating matrix keratinocytes in the bulb located at the base of the growing (anagen) follicle. Matrix cells eventually stop proliferating, and hair growth ceases at catagen when the lower follicle regresses to reach a stage of rest (telogen). After telogen, the lower hair-producing portion of the follicle regenerates, marking the new anagen phase. Generally, in parallel to anagen onset in humans, the hair shaft is shed during the exogen stage (Milner et al., 2002). Undifferentiated melanocytes are transferred in the epithelial column of the catagen stage follicle which later differentiates into active melanocytes during the telogen and anagen stage follicles of the hair germ. The differentiated melanocytes then transfer melanosomes to the follicular epithelial cells. The numbers of melanocytes in follicles during catagen and telogen are very low with little cytoplasm and less developed Golgi complex and rough endoplasmic reticulum, but their number increases during the early anagen stage. These observations suggest that the bulb population of melanocytes has a perpetuating system, that is, a melanocyte reservoir that contributes to normal hair pigmentation despite the shedding and replacing of hairs throughout the life (Sugiyama S, 1976).

**Extracutaneous non-melanocytic melanin**

Extracutaneous pigmented granules were first found in the human central nervous system (Scherer, 1939). Neuromelanin (NM) is a non-melanocytic extracutaneous
pigment found in the substantia nigra and locus coeruleus of the brain (Zecca et al., 1996). Synthesis of neuromelanin itself is understood poorly. Most researchers consider neuromelanin as an inert cellular by-product of dopamine synthesis produced via simple autoxidation (Double et al., 2000; Zecca et al., 2003). In humans, these pigments keep accumulating in the nigrostriatal dopamine neurons up to the age of 60 years and then decrease afterwards (Mann et al., 1977). Neuromelanins are insoluble brown/black macropolymer granular pigments composed of aminochromes and noradrenalinochromes, with the properties of both pheomelanins and eumelanins (Odh et al., 1994). The absence of NM in the central nervous system is possibly associated with pathogenesis of Parkinson disease (Zecca et al., 2003).

Reactivation of the melanocyte–specific differentiation program is also a characteristic of pulmonary lymphangioleiomyomatosis (LAM). LAM cells could be stained with antibodies against melanocytic markers gp100 and CD63 (tetraspanin) as well as with PNL2, an antibody against an as yet uncharacterized melanocytic antigen (Zhe & Schuger, 2004). Occasional findings of extracutaneous melanin were also described in non-melanoma neuroendocrine tumor of lungs (Iihara et al., 2002), pheochromocytoma (Bellezza et al., 2004), composite paraganglioma-ganglioneuroma of the urinary bladder (Dundr et al., 2003) and in black pigmented adrenal nodules found in association with Cushing's syndrome (Damron et al., 1985).

The liver of Rana esculenta is also a site of extracutaneous production of melanin. In the amphibian liver melanin is produced not by melanocytes or any other pigment cell
of neural origin, but by Kupffer cells of macrophage lineage (Pintucci et al., 1990, Sichel et al., 1997, Guida et al., 1998). The fact that the liver possesses its own endogenous melanogenic ability (Cicero et al., 1982, Sciuto et al., 1988) is supported by findings of dopa oxidase activity in the frog liver (Cicero et al., 1989) and in the cultured Kupffer cells, which do indeed express a tyrosinase (Guida et al., 2000).

**Non-melanin pigments**

Argentaffin pigments and lipofuscin are examples of extracutaneously produced pigments which resemble melanin in color but differ in chemical composition. Lipofuscin, also called age pigments, are fine yellow-brown granules composed of lipid-containing residues of lysosomal digestion that cannot be degraded by lysosomal hydrolases and also cannot be exocytosed. This intralysosomal polymeric materials are found mainly in the liver, kidney, heart muscle, adrenals, nerve cells, and ganglion cells and are specifically arranged around the nucleus. Lipofuscins originate from autophagocytosed cellular components that have become oxidized outside or inside the lysosomal compartment and continued to accumulate over time within metabolically active postmitotic cells such as cardiac myocytes, neurons and RPE (Boulton et al., 2004, Cuervo and Dice, 2000 and Yin, 1996).

An accumulation of lipofuscin (LF) pigments with age may be hazardous to cellular functions (Brunk & Terman, 2002a) as these pigments have metal binding capacity. In the case of cells of the RPE, most of the lipofuscin pigments are derived from the ingestion of photoreceptor outer segments. These photoreceptor outer segments are
mainly composed of polysaturated fatty acids (PUFAs) which are susceptible to attack by oxygen-derived free radicals. Due to the rich oxygen supply of the retina and its exposure to light of short wavelengths, PUFAs are peroxidized by iron-catalyzed intralysosomal oxidation and further transformed into fluorophores, which are known components of LF (Gutteridge, 1982). Since RPE cells do not divide and thereby cannot reduce their LF content, a substantial part of the RPE cell cytoplasm may finally be occupied by LF (Feeney-Burns et al., 1984). Some of the retinal disorders like age-related macular degeneration (AMD) and retinitis pigmentosa (Boulton et al., 2004) are accompanied by an abundance of RPE-derived lipofuscin whose composition is similar to age-related lipofuscin.

The content of lipofuscin in the liver, the heart and the neurons positively correlates with mitochondrial damage and the production of ROS. Most likely, the lipofuscin accumulation and the mitochondrial damage have common underlying mechanisms, particularly, imperfect autophagy and lysosomal degradation of oxidatively damaged mitochondria and other organelles (Terman et al., 2004). Several qualitative analyses provide supportive evidence that aging is accompanied by increases in the volume and/or in the number of dense bodies (secondary lysosomes, residual bodies, lipofuscin) in rodent and human hepatocytes (Schmucker, 1990 and Schmucker & Sachs, 2002).

The argentaffin-producing cells of the enteroendocrine system found in gastric glands secrete black pigment called argentaffin granules. The cells are located randomly
within the mucous membrane lining of the intestine and in tube-like depressions in that
lining known as the Lieberkühn glands. The argentaffin granules contain a chemical
called serotonin which stimulates smooth muscle contractions. Besides argentaffin
granules, these cells secrete a number of hormones, including gastrin, enteroglucagon,
serotonin and histamine.

Properties of melanin

In the human skin, melanin acts as a natural sunscreen and photoprotective
pigment. Melanin is capable of absorbing and scattering solar radiation, particularly, the
energetic UV and short wavelength visible photons. The pigment absorbs the harmful UV
radiation. The energy of the absorbed photons is rapidly utilized. However, the absorbing
potency decreases exponentially from the UV to the visible range (Sarna, 1992). Melanin
reacts continuously to impinging photons and either gets degraded or photoionized, or
produces free radicals on exposure to light. A very efficient non-radiative de-excitation of
melanin following the absorption of ultraviolet and visible photons has been reported in
several studies (Crippa et al., 1991; Forrest et al., 2000).

Environmental ultraviolet (UV) is known to cause a wide range of cutaneous
effects including skin cancer, immune suppression, photoaging and erythema (Phan et al.,
2006). These effects are a result of various physical and biochemical changes such as
proliferation of epithelial cells, skin hyperpigmentation, DNA damage, activation or
inactivation of a wide range of enzymes and proteins, and oxygen free radical production
(Lim et al., 2008). UVA-induced facultative pigmentation is limited to the basal cell layer
of epidermis whereas UVB-induced facultative pigmentation is distributed in keratinocytes throughout the epidermis, activated melanocytes and corneocytes of the stratum corneum (Rosen et al., 1990). Cutaneous melanin protects the skin, while ocular melanin protects the eye, specifically the retina, from chronic exposure to solar radiation, which reduces the incidence of age-related macular degeneration (AMD) (Young, 1988).

UV photons can mediate damage through two different mechanisms, either by direct absorption of UV via cellular chromophores, resulting in formation of an excited state and subsequent chemical reaction, or by phosensitization mechanisms, where the UV light is absorbed by endogenous (or exogenous) sensitizers that are excited and their further reactions lead to formation of ROS. These highly reactive species can interact with cellular macromolecules such as DNA, proteins, fatty acids and saccharides causing oxidative damage. Direct and indirect injuries result in a number of harmful effects such as disrupted cell metabolism, morphological and ultrastructural changes, attacks on the regulation pathways and alterations in the differentiation, proliferation and apoptosis of the skin cells. These pathological processes may lead to erythema, sunburn, inflammation, immunosuppression, photoaging, gene mutation, and development of cutaneous malignancies.

In dark-skinned subjects eumelanin acts as a filter against UV radiation and as an antioxidant that scavenges the ROS thus protecting the skin against UVB-induced DNA lesions and UVA-induced membrane damage (Montagna & Carlisle, 1991). On the contrary, pheomelanin, which is abundant in the light skinned individuals is not a good
scavenger of free radicals. This type of the melanin can produce free radicals and damage cellular material including DNA. Subtypes of eumelanin also differ in their properties: darker and highly polymerized DHI-eumelanin provides optimal photoabsorption but potentially increases cytotoxic effects (Jimbow, 1995; Schmitz et al., 1995) whereas DHICA-eumelanins have somewhat reduced photoabsorbing properties, but they are associated with decreased toxicity towards cells (Hochstein & Cohen, 1963; Pawelek & Lerner, 1978; Urabe et al., 1994).

Besides producing ROS and DNA damage, UV radiation also induces inflammatory responses. Prostanoids and eicosanoids play an important part in triggering melanin synthesis in UV-induced pigmentation (Nordlund & Askenase, 1983; Tomita et al., 1992; Morelli and Norris, 1993), in a manner similar to that of other melanogens such as endothelin-1 (Imokawa et al., 1995) and α-MSH (Hunt et al., 1994). Prostanoids and eicosanoids activate a wide variety of responses in the skin, including: increased vasodilation of blood vessels, increased vascular leakiness of fluid into the skin, the activation of nerve endings, an increased release of histamine, and an activation of the melanocytes. If the skin is continually exposed to solar radiation over many years, the constant production of inflammatory hormones ultimately results in a permanent activation of melanocytes. Another ubiquitous inflammatory factor, histamine, exerts its stimulation through H2 receptors. Histamine has a melanogenic effect on cultured human melanocytes (Tomita et al., 1988). In human skin, histamine is produced and released by dermal mast cells and keratinocytes, and an increased release of histamine by UV irradiation was reported by Gilchrest & Eller (1999). These findings strongly suggest the
involvement of histamine in UV-stimulated pigmentation. In conclusion, UV irradiation increases the amount of melanin directly and indirectly through different pathways. Antioxidant properties of melanin are responsible for scavenging of ROS, thus, reducing the DNA damage.

**Origin of melanocytes**

Melanocytes originate from a subpopulation of neural crest cells (NCC) called melanoblasts that emerge from the dorsal neural tube. The NCC migrates dorso-laterally to form melanocytes of the skin, hair and inner ear mainly within cochlea, including the stria vascularis (Schrott & Spoendlin, 1987; Peters *et al*., 1995; Conlee *et al*., 1994; Motohashi *et al*., 1994). These cells can also migrate ventrally and contribute to the formation of sensory and glia neurons and glia of cranial ganglia and mesenchymal cells capable of differentiating into tendons, adipocytes, cartilage and bone, connective tissue, neuroendocrine cells (Le Douarin, 1982; Le Douarin and Kalcheim, 1999).

The NCCs are derived from neural crest, a transient embryonic structure that arises from neuroectoderm and gives rise to many cell types and tissues of the adult vertebrate organism. NCCs originate dorsally at the tip, or ‘crest’, of the neural tube within developing embryo, and undergo a well-defined extensive migration following epithelial-to-mesenchymal conversion and detachment from the neural tube (Le Douarin & Kalcheim, 1999; Bronnner-Fraser & Fraser, 1988). At the onset of migration, the NCC are composed of a heterogeneous population of cells endowed with different proliferation and differentiation potentials, capable of forming pigment cells, neurons and glial cells of
the peripheral nervous system (PNS) and endocrine cells (Sieber-Blum & Cohen, 1980). NCC that differentiate into melanoblasts migrate by a dorso-lateral pathway between dermatome and the overlying ectoderm; from embryonic day (E10.5) they migrate ventrally through the developing dermis. At E14.5, they begin to invade the overlying epidermis and then migrate into the developing hair follicles, where they continue to proliferate and differentiate, before beginning to synthesize pigment at around postnatal day 4 (Mayer, 1973; Jordan & Jackson, 2000b).

The pigment cells of eye, retinal pigment epithelium (RPE), develop from a totally different source, the anterior neuroepithelium: optic cups of the brain (Bharti et al., 2006). These optic vesicles arise as paired bulb-like outpocketings of the forebrain. The vesicles indent to form a double-layered cup whose innermost layer develops into the highly complex neural retina while the outer one gives rise chiefly to the single-cell-layer pigment epithelium. Any defect in retinal pigmentation leads to visual impairment.

As shown by in vitro lineage analysis, both the cardiac and the trunk NCCs possess the ability to form pigmented melanocytes (Sieber-Blum and Cohen, 1980; Baroffio et al., 1991). There is also a possibility that melanocytes derived from NCCs may arrest during migration and may have undergone malignant transformation in situ. Non cutaneous occurrence of the malignant melanoma (MM) indicates a good example of above mentioned assumption. Furthermore, the mRNA transcripts of tyrosinase, the most specific marker of melanocytic differentiation, have been detected by PCR analysis in various non cutaneous tissues of normal human organs such as lymph nodes, antrum,
colon, kidney, lung, testis, ovary, breast, and peripheral nerves (Battayani et al., 1995). These finding is explained by the presence of the scattered fully-differentiated melanocytes, melanocytic precursors, or Schwann cells bearing potentialities of melanocytic differentiation within normal tissue. However, since the non-melanocytic human cells can express tyrosinase but can not form the functional enzyme, these PCR experiments cannot be viewed as the ultimate proof of the “scattered melanocytes” theory.

On the other hand, the presence of the melanocytes has also been reported in the adult heart where these cells are localized mostly in the valvuloseptal apparatus (Mjaatvedt et al., 2005; Ambani et al., 2005). Heart melanocytes were revealed in the transgenic mouse model that expresses the LacZ reporter gene under the control of the Dopachrome tautomerase (Dct) promoter (abstract in PASPCR "The origin of melanocytes in the murine heart"). The contribution of NCCs to heart development has been established earlier (Kirby & Waldo, 1995). The analysis of embryos from E10.5 through E12.5 showed that the melanocytes destined to the heart initially follow the same migratory route as those moving towards the skin and originate from the same region as the cardiac NCCs. At E11.5 they leave the skin, travel inwards and reach the heart by E12.5. By E14.5, their numbers have increased. The location of the melanocytes within the embryonic heart and within the adult heart is the same.
**Melanosomes and their biogenesis**

Eumelanin and pheomelanin synthesis takes place in specialized, pigment-containing, lysosome-related organelles called melanosomes. The melanosomes are present within the melanocytes located in the skin and other places of the body such as the eye, the ear, the meninges, the hardarian gland, and accessory lacrimal glands on the inner side of the orbit. In birds, melanosomes are found in the melanocytes of the feather-follicles as well as other tissues. Melanosomes are also present in the connective tissue of the portal canals, in the space of Disse and around the sinusoids of amphibian and reptilian livers (Eberth, 1866; Oppel, 1990).

Melanosomes, like other cellular organelles, are membrane-bound structures containing a variety of molecules including lipids, proteins and melanin. The structure and contents of melanosomes depend on their origin and correlate with the type of melanin produced. Eumelanin-producing eumelanosomes are of elliptical shape and contain fibrillar matrix, pheomelanosomes have a more rounded contour and contain vesiculoglobular matrix (Ozeki *et al.*, 2007). Neuromelanin-producing neuromelanosomes have a complex structure. Their melanin component is bound to metals, peptides, and lipids, which at present have unknown structure (Zecca *et al.*, 2000; Zecca *et al.*, 1992; Zecca *et al.*, 1996).

The biogenesis and the maturation of melanosomes in melanocytes and their transfer to adjacent keratinocytes provide the basic processes underlying the mammalian pigmentation (Marks and Seabra, 2001). Melanosomes originate from the trans-Golgi
apparatus and mature within the melanocyte or the developing RPE through four morphologically distinct stages (Marks and Seabra, 2001). Stage I melanosomes, also called premelanosomes, are formed by outpouching of a smooth membrane from the RER; these organelles are characterized by the early matrix organization, while in stage II the matrix is already organized but melanin is not formed yet. The structural foundation of the fibrils in stage I and II melanosomes is PMEL 17 protein. In stage III, there is a deposition of melanin. In stage IV, melanosomes are fully melanized. Pigment-loaded melanosomes of stage IV are translocated bidirectionally to their recipient keratinocytes along the dendrites of melanocytes.

Melanin biosynthetic enzymes tyrosinase (TYR), tyrosinase-related protein 1 (TYRP1) and tyrosinase-related protein 2 (TYRP2) are integral membrane proteins that are enriched in stage III and IV melanosomes (Raposo et al., 2001; Kushimoto et al., 2001; Theos et al., 2005). The process of the cellular sorting for these integral membrane proteins is unique to these specialized cells. When expressed ectopically in non-pigmented cells, most melanosomal proteins are localized to late endosomes and lysosomes (Bouchard et al., 1989; Vijayasaradhi et al., 1995; Berson et al., 2001).

Microtubule-based transport of mature melanosomes to keratinocytes is facilitated by dynein and kinesin motor proteins. These mature melanosomes are captured by actin-based motor proteins which are later on dispersed into the cytoplasm of keratinocytes (Seabra & Coudrier, 2004; Barral & Seabra, 2004). Capture on actin filaments is mediated by a protein complex that consists of myosin Va, the small GTPase RAB27a.
and the linker protein melanophilin (also known as SLAC2a) (Seabra & Coudrier, 2004). Genetic, morphologic and biochemical studies of three mouse coat color mutants, *dilute*, *ashen* and *leaden*, provided the understanding of the mechanisms that regulates the binding of melanosomes to myosin Va and helps their (Barral & Seabra, 2004; Goud, 2002; Hammer & Wu, 2002). A protease activated receptor 2 (*PAR2*) that is present on the keratinocytes also plays an important role in melanosome transfer (Boissy, 2003).

In contrast, RPE melanosomes are synthesized *in utero* and remain unchanged thereafter. Unlike the cutaneous melanocytes, the melanosomes of the eye never transfer to adjacent cells, but are retained inside the cells of origin. Premelanosomes are rarely found in the normal retina, which suggests a slow turnover of melanosomes in mature RPE. The melanosomes in the retina are thought to function in the absorption of stray light that enters the eye, thus, minimizing light scatter, and in the detoxification of oxidative intermediates (Schraermeyer & Heimann, 1999).

Defects in the formation, maturation and trafficking of melanosomes within melanocytes result in several genetic disorders, particularly, Hermansky–Pudlak syndrome, Chediak–Higashi syndrome, Griscelli syndrome and Elejalde syndrome. Individuals with one of these syndromes show defects in skin and/or hair pigmentation. In cells of these patients melanosomes appear to be clustered around the perinuclear area due to improper transportation, instead of being scattered at the periphery, as seen in normal melanocytes.
Melanosomal pH has been documented to control the rate of melanogenesis, eumelanin/phaeomelanin ratio and melanosome maturation in melanocytes (Ancans et al., 2001). Catalytic domains of the enzymes required for melanogenesis face the intracellular environment of melanocytes, so proper pH activates these enzymes. Particularly, melanosomal pH regulates multiple stages of melanin production in human pigment cells and the enzymatic activity of TYR (Ancans et al., 2001). In fact, the intact melanosomes are reported to have a proton gradient similar to that of lysosomes (Klaus, 1971; Hori et al., 1968), which makes the melanosomes acidic.

Melanocytes from different skin types differ in the pH of their melanosomes (Fuller 2001). Generally, more highly pigmented melanosomes are more acidic than less-pigmented melanosomes (Bhatnagar et al, 1993; Ramaiah, 2002). Higher enzymatic activity of TYR at low pH values has been reported (Tabachnick, J. 1970; Burka, R.C. 1965), probably because of the activating change in the conformation of TYR at low pH. Tyrosinase enzyme activity can be also boosted by a single dose of UV irradiation (Pathak et al., 1965). An inducible increase in the activity of TYR is explained by the hypothesis that it mainly exists in an inactive or partially inhibited state and is activated “on demand” by a decrease in pH of the intact melanosomes or by an increase of the ROS (Fitzpatrick, 1965).

Genetics of pigmentation:

Melanogenesis is a very complex, enzymatically regulated process controlled by more than 120 distinct genes (Bennett and Lamoreux, 2003). These genes work to
determine the development and distribution of melanocytes, their differentiation and their regulation by physiological factors. Protein products of these genes act as enzymes, structural regulators, transporters, receptors, and growth factors.

Melanogenesis starts with the oxidation of monophenols and/or ortho-diphenols that yield the corresponding o-quinones, which evolve through coupling enzymatic reactions toward the formation of melanins (Prota, 1995). The whole process is mediated by a group of enzymes uniquely expressed in melanocytes, called the tyrosinase-related protein (TRP) family. The TRP family has three known members: TYR, TYRP1 and dopachrome tautomerase (DCT or TYRP2) (Burchill et al., 1991; Hearing and Tsukamoto, 1991). These three enzymes are type I membrane glycoproteins targeted to and located in the melanosomal membrane.

Before reaching the melanosomal membrane in their final catalytically active conformational state, the proteins of TYR family undergo a complex post-translational processing. Correct processing and intracellular trafficking of tyrosinase and TYRPs are very critical to normal pigmentation. This processing involves N-glycosylation, movement from the endoplasmic reticulum (ER) to Golgi apparatus, copper binding and finally sorting to the melanosome. In mice, tyrosinase processing involves N-glycosylation at six glycosylation sites (Branza-Nichita et al., 2000) whereas the human TYR displays seven potential glycosylation sites (Ujvari et al., 2001), including one located in the copper binding site CuB. The glycosylated protein moves from the ER to the Golgi apparatus, binds to copper at designated sites CuA and CuB (Spritz, 1999), and
then is finally sorted to melanosomes. The ER acts as a quality control checkpoint for the posttranslational processing. The ER retains the incorrectly folded tyrosinase that is not further processed for copper binding.

Mutations in \textit{TYR} gene, including those leading to aberrant processing and/or trafficking defects, result in pigmentary disorders, including oculocutaneous albinisms type 1 (OCA1) and type 2 (OCA2). OCA1 is a genetic autosomal recessive disorder due to mutation in the \textit{TYR} gene. It is characterized by a loss of pigmentation in hair, skin, eyes and reduced vision (Oetting & King, 1999). OCA2 disorder is due to mutations in the pink-eyed dilution (\textit{p}) gene (Lee \textit{et al.}, 1994), which contributes to the correct processing of \textit{TYR} and its trafficking to the melanosome (Chen \textit{et al.}, 2002).

All the melanogenesis enzymes act in a series of reactions in the melanogenic pathway to control melanin production in melanosomes (Hearing, 1999; Hearing, 2005). Melanin biosynthesis depends on the amount of L-tyrosine present in the cell. It starts with hydroxylation of L-tyrosine to L-3, 4-dihydroxyphenylalanine (L-DOPA), an obligatory and rate-limiting step carried by tyrosine hydroxylase activity of tyrosinase. L-DOPA produced by tyrosine hydroxylation serves as a precursor to both melanins and catecholamines being subsequently oxidized to dopaquinone (Hearing & Jimenez, 1987; Hearing & Jimenez, 1989; Prota, 1988; Hearing, 1999). Eumelanogenesis involves the further transformation of dopaquinone to leukodopachrome, followed by a series of oxidoreduction reactions with production of the intermediates 5, 6-dihydroxyindole (DHI) and 5, 6-DHI carboxylic acid (DHICA) that is catalysed by TYRP2. This reaction
is followed by polymerization catalysed by TYRP1 and resulting in eumelanin formation. Pheomelanogenesis also starts with dopaquinone; which is conjugated to cysteine or glutathione to yield cysteinyldopa or glutathionyldopa, for further transformation into pheomelanin. The formation of eumelanin or pheomelanin is directly determined by the presence or absence of cysteine, of GSH in fully reduced thiolate state, and by the redox potential. When concentration of sulfhydryl compounds is low, dopaquinone is converted to dopachrome, initiating the eumelanogenic pathway. High concentration of cysteine or glutathione leads to the formation of pheomelanin.

Melanogenesis commences when the TYR enzyme is cleaved. Initiation of activity depends on an acidic environment provided by proton pumps (Mani et al., 2001; Puri et al., 2000). Formation of intermediates in the melanin biosynthesis pathway is also dependent on pH, which makes melanogenesis all together a pH-dependent process (Francisco García Canovas, 1982). Other direct regulators of melanogenesis include PMEL17 protein encoded by HMB-45/gp100/silver locus, catechol-O-methyltransferase (COMT), peroxidase, and macrophage migration inhibitory factor (MIF) (Hearing, 1999). PMEL17 has been proposed to catalyze the polymerization of DHICA to melanin, and that PMEL17/GP100 acts as scaffold for the deposition of melanin in melanosomes, and for stabilizing the melanin intermediates. COMT is responsible for O-methylation of DOPA and for the formation of DHI intermediates, whereas peroxidase catalyzes the oxidation of DHI and DHICA. MIF possesses D-dopachrome tautomerase activity as it transforms D-dopachrome, dopaminechrome or its derivatives to their indole compounds. Enzymes that indirectly affecting melanogenesis include glutathione reductase and
glutathione peroxidase, which regulate glutathione reduced/oxidized levels respectively, and catalase which regulates hydrogen peroxide removal (Schallreuter et al., 1998a).

Fig 5: Scheme illustrating common pathway for follicular and epidermal melanogenesis.

Less is known about the biosynthesis and function of extracutaneous melanins. In the eye, the biosynthetic pathway most likely involves the same enzymes as in the skin, and thus is relatively similar to the cutaneous pathway (Orlow et al., 1994; Benedito et
al., 1995). However, the biosynthetic pathway involved in the formation of neuromelanin in brain is still uncharacterized and, apparently, involves neither TYR (Barden, 1981), nor a peroxidase activity (Rabey & Hefti, 1990). Neuromelanin in the substantia nigra is assumed to be formed by catalyzing the conversion of the amino acid L-tyrosine to the dopamine (DA) precursor of dihydroxyphenylalanine (DOPA), by tyrosine hydroxylase (TH). DA and other catecholamines later on undergo non-enzymatic autoxidation to form neuromelanin (Breathnach, 1988). Alternatively, TH-mediated oxidation of dopamine has been proposed (Havvik, 1997). In addition to this, enzymatic activity of macrophage migration inhibitory factor was also suggested for neuromelanin synthesis, because it converts catecholamines into dihydroxyindole derivatives, the potential precursors of neuromelanin (Matsunaga et al., 1999). Tyrosine hydroxylase has been used as a marker of putative dopaminergic neurons and the existence of this enzyme in the human midbrain neurons is considered to be an evidence of DA synthesis in this area. A recent study, however, has found a low level of TYR mRNA expression in the human SN (J. Haavik, 1977). Some authors proposed that TYR could be involved in neuromelanin biosynthesis, because TYR mRNA (Haavik, 1977) and TYR promoter activity (Tief et al., 1998) have been detected in the substantia nigra. However, TYR protein was not detected in substantia nigra by immunohistochemistry (Ikemoto et al., 1998). Moreover, albinos, who lack TYR gene, display normally pigmented substantia nigra (Foley & Baxter, 1958).

In human substantia nigra, the neuromelanin content increases until age 60 and decreases afterwards (Fenichel & Bazelon, 1968), probably, due to loss of catecholeaminergic neurons during the process of aging. This neurodegeneration results in
a loss of dopaminergic neurons in the substantia nigra, which results in the loss of TH containing nerve endings in the striatum, and diminished levels of the striatal dopamine production causing abnormal motor behavior, finally leading to degenerative disorder called Parkinson’s Disease (PD) (Booij et al., 2001).

**Regulatory pathways in melanogenesis**

Melanogenesis is a tightly controlled process that can be regulated by posttranslational, translational, and transcriptional mechanisms (Eberle, 1988). A key role in the control of pigmentation in mammals is played by melanocortins. Melanocortins are a group of pituitary hormones that include ACTH and α, β and γ-MSH generated by the posttranslational cleavage of prohormone proopiomelanocortin. These peptides act through melanocortin receptors. In addition to their well-known roles in pigmentation and in neuroendocrine pathways, melanocortins serve as regulators of immune function, lipid metabolism and feeding behaviors in rodents and humans (Wintzen et al., 1996; Huszar et al., 1997; Luger et al., 1999).

α-MSH peptide is the main product of POMC acting upon the skin (Slominski et al., 2000). The other products, particularly, ACTH, γ-MSH and β-endorphin also serve as important mammalian cutaneous mediators (Inoue et al., 2003; Johansson & Wang, 1993; Kauser et al., 2003). In cultured human melanocytes, nanomolar concentrations of α-MSH, β-MSH, and ACTH stimulate melanogenesis, cell proliferation, dendrite formation, and cAMP production (Abdel-Malek et al., 1995; Rousseau et al., 2007; Hunt, 1995; Suzuki et al., 1996), whereas γ-MSH stimulates cAMP production, but had no significant effect on melanogenesis or proliferation in human melanocytes (Suzuki et al., 1996).
Five MC receptors (MC1R - MC5R) have been cloned and characterized from human tissues; most of them show tissue-specific expression patterns. MC1R serves as the key control of pigmentation as it has high affinity for α-MSH and signals through cAMP-dependent pathway ((Lalli & Sassone-Corsi, 1994). In vitro, the melanogenic effects of α-MSH can be mimicked by pharmacologic agents such as forskolin, a direct activator of adenylate cyclase (Englaro et al., 1995). Any mutation in MC1R or Gαs protein that controls cAMP level as well as a decrease of αMSH expression leads to inhibition of melanin synthesis (Valverde 1995; Krude et al., 1998; Schwindinger et al., 1992). cAMP increases melanogenesis mainly through the MITF-dependent stimulation of TYR expression. Patients with altered function of the cAMP/PKA pathway present with skin pigmentation defects, thereby demonstrating the importance of the cAMP/PKA pathway in the regulation of melanogenesis (Schwindinger et al., 1992).

Through activation of PKA and the transcription factor CREB, cAMP promotes an increase in the expression of microphthalmia-associated transcription factor (MITF) (Bertolotto 1998). MITF is a melanocyte-specific transcription factor of the basic-helix-loop-helix-leucine-zipper family (bHLH-LZ) (Hodgkinson et al., 1993; Steingrimsson et al., 1994). MITF binds as a dimer to the M-box (11bp motif AGTCATGTGCT), a highly conserved DNA sequence (Goding & Fisher, 1997; Sato et al., 1997; Yasumoto et al., 1997). Recent studies revealed that MITF is able to transactivate several other gene promoters in addition to pigmentation genes. For example, in cultured human osteoclasts MITF upregulates endogenous cathepsin K, which are required for the bone resorption. Several related bHLH-Zip transcription factors can heterodimerize with MITF and bind
to identical DNA sequences. These related factors, namely transcription factors EB (TFEB), TFE3 and TFEC, together with MITF are called the MIT family (Hemesath et al., 1998). MITF is expressed only in specific cell types, while other members of the MIT family are ubiquitously expressed. MITF mediates both the differentiation effect of α-MSH (Price et al., 1998; Bertolotto et al., 1998) and promotes the survival of pigment cells by up-regulating the expression of the major antiapoptotic protein BCL2 (McGill et al., 2002).

The MITF gene has a multi-promoter organization. At least nine distinct promoter-exon units direct the initiation of specific MITF isoforms that differ in their first one or two exons spliced to the common downstream exons (Hershey & Fisher, 2005). The promoter that is most proximal to the common downstream exons is known as the M promoter and is selectively expressed in melanocytes (Fuse et al., 1996). MITF-M promoter expression is cell-type specific despite its regulation by the ubiquitous cAMP–CREB pathway. Structural analysis of the MITF gene revealed that it encodes four isoforms of MITF proteins, termed MITF-A, -C, -H and –M (Shibahara et al., 2001; Amae et al., 1998). The longest isoform, MITF-A, is expressed in many cultured cells, including RPE cells (Amae et al., 1998). MITF-C is also expressed in many cultured cells, but is undetectable in melanocytes (Fuse et al., 1999), whereas MITF-H is enriched in the rodent heart tissue (Hodgkinson et al., 1993).

Regulation of MITF gene expression in melanocytes has been studied relatively well. In these cells, the activation of MC1R receptors by UV radiation or by binding to α-MSH peptides leads to activation of adenylyl cyclase and the subsequent increase in the
production of cAMP. Increase in cAMP leads to the phosphorylation of CREB transcription factors which activate MITF-M promoters (Price et al., 1998). In addition to CREB, other transcription factors, namely, paired box gene 3 (PAX3), SRY (sex determining region Y)-box 10 (SOX 10), lymphoid enhancer-binding factor 1 (LEF 1/TCF), one cut domain 2 (ONECUT-2) also bind to the MITF-M promoter (Widlund 2003, Yasumoto 2002). PAX3 is expressed in both melanocytes and melanoblasts cell lines (Galibert et al., 1999) and recognizes the MITF promoter for both in vitro and in vivo conditions (Watanabe et al., 1998). Mutations in PAX3 are known to impair its ability to regulate transcription by inhibiting its capacity to regulate MITF promoter activity. Sox10 proteins are also expressed in melanocytes and in melanoma cell lines (Kuhlbrodt et al., 1998; Southard-Smith et al., 1998) and are required in the development of the melanocyte lineage. Mutations in Sox 10 are responsible for Waardenburg-Hirschsprung syndrome (WS4) in humans (Pingault et al., 1998; Southard-Smith et al., 1998). This disease is characterized by a combination of deafness and pigmentary disorder.

Pigment cell formation is also promoted by ectopic expression of β-catenin in NCCs. Promoter regions of MITF (Takeda et al., 2002) contain binding sites for Tcf/Lef transcription factors (Dorsky et al., 1998), which form a complex with β-catenin and act as a transcriptional activator of Wnt target genes (Seidensticker & Behrens, 2000). Mutations in the Tcf/Lef binding sites of MITF-M regulated promoters inhibit the function of MITF-dependent genes in murine melanocytes (Takeda et al., 2002).
Exogenously applied WNT protein transactivates the MITF-M promoter and upregulates MITF-M RNA (Takeda et al., 2002). WNT signaling promotes this event possibly by up-activation of dishevelled protein, which is shown to inhibit GSK3β-phosphorylation and subsequent ubiquitination/degradation of β-catenin (Seidensticker & Behrens, 2000). Insulin-like growth factor has also been shown to enhance melanocyte growth (Edmonson SR, 1999). However, insulin itself downregulates tyrosinase and inhibits tyrosinase activity in melanoma cells (Ando et al., 1995; Fuller et al., 1987).

In addition to transcriptional regulation, MITF is also subjected to post-translational modifications, particularly phosphorylation by mitogen-activated protein kinase (MAPK), ribosomal S6 kinase (RSK), glycogen synthase kinase-3β (GSK3β) and p38 (Weilbaecher et al., 2001; Wu et al., 2000; Takeda et al., 2000). Particularly, activation of the Ras/ERK cascade inhibits melanogenesis by phosphorylating the MITF transcription factor that leads to its degradation (Hemesath et al., 1998; Wu 2000). This pathway is thought to be a feedback mechanism preventing an excessive production of melanin (Englaro et al., 1998). The transcriptional activity of MITF is also affected by sumoylation, a post-translational modification mediated by protein inhibitor of the activated STAT3 (PIAS3). On promoters with multiple MITF binding sites, sumoylated MITF seems to be less active than unsumoylated MITF. Therefore, post-translation modifications regulate MITF activity in a way that alters its target-gene specificity based on cellular contexts or signals (Miller et al., 2005; Murakami & Arnheiter, 2005).

NIH/3T3-L1 fibroblasts are known to differentiate into myoblasts or adipocytes in response to ectopic expression of specific lineage-determining transcription factor genes
Ectopic expression of MITF differentiates 3T3-L1 fibroblasts into cells with melanocyte characteristics (Tachibana et al., 1996).

UVR is the primary cause of melanoma and a potent inducer of ROS, including superoxide radical as well as hydrogen peroxide and hydroxyl radicals. ROS production results in inflammation and oxidative stress (Miller & Weinstock, 1994). Skin exposure to UVR induces a range of biologic responses such as erythema, inflammation, hyperpigmentation, hyperplasia, skin cancer (Norris et al., 1992; Gilchrest et al., 1996), DNA damage and mutations, induction of early and late gene responses (Sachsenmaier et al., 1994; Herrlich et al. 1994), systemic immunosuppression (Kripke, 1994), and activation of dormant viruses (Herrlich et al., 1994). UVR also induces pigmentation through the process that consists of three main steps, i.e., the proliferation of melanocytes (Rosdahl & Szabo, 1978), the synthesis and activation of tyrosinase (Imokawa & Mishima, 1982; Mishima & Imokawa, 1983), and the transfer of melanosomes to keratinocytes (Okazaki et al., 1976). The oxidative stress produced due to UVR, particularly UVB, selectively triggers the p38 signaling pathway and, to a lesser extent, ERK1/2 pathway (Peus et al., 1999; Rouse et al., 1994; Galibert et al., 2001), thus inducing the expression of POMC, MC1R, and TYR. This finding has been confirmed in experiments involving preincubation of skin explants with a specific inhibitor of p38 kinase (Galibert et al., 2001).

UVR also stimulates cleavage and turnover of cell membrane phospholipid constituents by phospholipases (DeLeo et al., 1988). This process generates a number of
signaling mediators, such as diacylglycerols, that activates the PKC-β pathway and upregulate tyrosinase expression (Nishizuka, 1988). Stimulation of PKC-β pathway has been shown to be associated with both differentiation and proliferation of melanocytes (Yada et al., 1991). UVR also stimulates epidermal cell proliferation and induces the secretion of several cytokines, including bFGF, α-MSH, nitric oxide, SCF and endothelin-1 (ET-1). Due to the paracrine linkage between keratinocytes and melanocytes within the epidermis, these cytokines regulate melanocytes growth and survival playing a critical role in cutaneous physiology and pathology (Grabbe et al., 1994).

ET-1 peptides that stimulate proliferation, chemotaxis, and pigment production in melanocytic cells are secreted by keratinocytes (Imokawa et al., 1996). Endothelins not only act to regulate melanocyte progenitor cell number, but also to induce pigment production in these cells. Endothelins also stimulate differentiation of the melanocyte progenitors into fully mature melanocytes. UVR induces a marked and sustained increase in ET-1 secretion by keratinocytes (Imokawa et al., 1992) and over expression of ET-B and its receptor EDNRB in human skin (Kadono et al., 2001). Studies in mice have established that Ednrb signaling is not necessary for the specification of pigment cells but is involved in the proliferation and migration of pigment cells (Pavan & Tilghman, 1994; Hosoda et al., 1994). Mutations of genes for EDNRB and its ligand endothelin 3 (EDN) lead to deficiencies in neural crest-derived melanocytes and neurons in mice (Hosoda et al., 1994; Baynash et al., 1994) and humans (Puffenberger et al., 1994). An EDNRB antagonist has also been shown to inhibit the growth and to induce cell death in the human melanoma cell lines (Lahav et al., 1994).
Besides ET-1, UVB upregulates secretion of the scatter factor (SCF) that regulates haematopoiesis, melanogenesis and gametogenesis (Besmer et al., 1993; Blume-Jensen et al., 2000; Fujio et al., 1994) and modulates MITF and, therefore, tyrosinase expression (Bertolotto et al., 1998, Hemesath et al., 1998). Basically, SCF/c-Kit and ET-1/(ETB_R) utilize a common signaling pathway, as they act synergistically to regulate melanin synthesis and proliferation (Imokawa et al., 1996). This synergistic cross-talk between SCF and ET-1 signaling is initiated by phosphorylation of c-kit in the signaling cascades. This promotes dimerization, autophosphorylation and transphosphorylation of several substrates at specific tyrosine residues, and results in activation of the Shc-Grb2 complex, which is, in turn, followed by the stimulation of mitogen-activated protein kinase signaling in human melanocytes (Imokawa et al., 2000). Activation of c-KIT in melanocytes results in phosphorylation of MITF at serine 73 by extracellular-signal regulated 2 (ERK2) and at Ser409 by p90 ribosomal S6 kinase (p90RSK). Phosphorylation at Ser73 increases the recruitment of the transcriptional coactivator p300 (CREB) (Price et al., 1998). Recent studies have shown that the inhibition of c-kit hinders communication between keratinocytes and other cell types, causing inappropriate tissue localization and disruption of integrin-mediated survival and proliferative signals (Bendall et al., 1998; Lahav, 2005) In addition, mutations of the EDNRB result in cutaneous and ocular hypopigmentation (i.e., Waardenburg Syndrome Type IV) (Reid et al., 1996). Piebaldism, a disorder presenting at birth, characterized by amelanotic patches on acral and/or ventral skin surfaces is also caused by mutations in
the genes encoding \textit{c-kit} (Bendall \textit{et al.}, 1998; Lahav, 2005) and \textit{ETBR} (Puffenberger \textit{et al.}, 1994; Amiel \textit{et al.}, 1996).

Fig 6: Melanogenesis signal transduction pathways.
Chapter III

Hypothesis and Research Objectives

Our specific hypothesis behind the proposed research is that the melanin synthesis pathway is functional in the adipocytes and that its activation may result in the formation of melanin.

The basis for our hypothesis is formed by following observations:

1) Our previous expression profiling experiments revealed statistically significant overexpression of melanogenesis-related genes in the visceral fat of obese individuals (Baranova et al., 2005) as compared to lean individuals.

2) α-MSH, a melanocortin peptide produced after the posttranslational cleavage of POMC, is required both for pigmentation control and energy homeostasis. Melanocortin peptides signal through G-coupled melanocortin receptors (MCR). Particularly in adipocytes, α-MSH signals through MC1R, MC2R, MC3R and MC4R receptors, whereas in the skin it exerts its action through MC1R and the protein kinase A (PKA) pathway. The expression of MC1R is more pronounced in the adipocytes of morbidly obese subjects as compared to normal subjects (Hoggard et al., 2004). Therefore, there is a possibility that under morbidly obese conditions,
melanogenesis-related genes may become activated ectopically in response to the signal propagated by MC1R activation.

3) Extracutaneous production of melanin has already been described in human central nervous system (Substantia nigra) of the Parkinson disease patients (Scherer, 1939 & Zecca et al., 1996) and in abnormal smooth muscle cells seen in the lungs of the patients with lymphangioleiomyomatosis (LAM) (Ferrans VJ et al., 2000). Therefore, this is not the first time that extracutaneous production of melanin has been documented.

4) Dermatological changes have been reported in patients with obesity. These changes include the hyperpigmentation disorder acanthosis nigricans that usually presents as periorbital darkening or darkening of the neck or knuckles (Garcia Hidalgo L. 2002).

It was hypothesized that hyperpigmentation in these obese subjects may be caused by increased levels of ROS characteristic of morbid obesity and compensatory increase in the expression of tyrosinase.

5) Sumo wrestlers, the grotesquely obese behemoths who measure more than 182 Kg are formally classified under morbidly obese category, but they are found to be devoid of any metabolic disorders as observed in other morbidly obese subjects. In these wrestlers, except for blood glucose and total cholesterol, mean serum levels of triglycerides, phospholipid, uric acid and total protein were significantly higher than in controls. Even though the basal metabolic index and serum composition of sumo wrestlers is similar to morbidly obese subjects, they don’t show any symptoms of
disorders like diabetes type II and arthrosclerosis. This phenomenon could be explained by the suppression of the production of ROS by intense exercise training sessions (Chinda D et al., 2003). Some percentage of morbidly obese subjects do not have sign of metabolic disorders (Jarrar et al., 2008), which suggests there should be something different about these patients, which is compensating for production of the ROS.

It was hypothesized that the ectopic melanin may serve as an agent capable of the buffering the ROS under morbidly obese conditions. The specific aims of this project are designed to support that melanogenesis may indeed take place in the adipose tissue of morbidly obese subjects.

**Specific aims**

1) **Quantification of the melanin pigment present in the adipose tissues obtained from morbidly obese subjects and from lean controls.**

   a) The melanin pigment present in the adipose tissue was analyzed by the Fontana Masson assay, which is specific for melanin, lipofuscin and argentaffin granules. Positive staining for melanin granules can be differentiated from lipofuscin staining by controlling the time period for staining.

   b) The eumelanin content present in the adipose tissue sections was measured by HPLC.

2) **Gene expression analysis of melanogenesis-related genes in adipose tissue sections of morbid obese and of lean subjects.**
Under normal circumstances, melanin is produced by specialized cells (melanocytes) in membrane-bound organelles termed melanosomes. Melanosomes gradually develop melanin pigment as they mature and are transported along the dendritic projections of melanocytes before being passed into adjacent keratinocytes of the skin and hair follicles. The list of the proteins required for melanin formation include tyrosinase (TYR), tyrosinase-related protein-1 (TYRP1), tyrosinase related protein-2 (TYRP2), microphthalmia transcription factor (MITF) and a structural protein gp 100 (PMEL17). The main substrate for the formation of the melanin is the tyrosine which is converted to L-3, 4-dihydroxy phenylalanine (L-dopa) which is ultimately converted to orange-colored dopachrome that is further processed to melanin. TYR enzyme performs one of the most critical steps required for melanin formation.

a) In this study expression analysis of the melanogenesis-related genes in the tissue samples obtained from obese subjects undergoing gastric bypass surgery and from lean subjects was performed. Gene expression study was performed for TYR, TYRP1, TYRP2 and MC1R that were analyzed by Real Time-PCR technique.

b) Localization of mRNA for TYR in morphologically preserved frozen tissue sections was established by in situ hybridization.

3) Enzyme activity assays of the adipose tissue extracts of morbidly obese and of lean subjects:
An assessment of the total melanogenic pathway activity was performed by radioactive C\textsuperscript{14} assay on adipose tissue samples from obese and from lean subjects as well as on liver and gastric tissue samples.

4) **Protein expression analysis in adipose tissue obtained from morbidly obese and from lean subjects.**

a) The melanogenesis-related enzymes TYR, TYRP1 and TYRP2 were localized by immunohistochemistry in adipose tissue sections of morbidly obese and of lean subjects.

b) The protein expression for TYR was assessed by western blotting in adipose tissue of morbidly obese and of lean subjects as well as on liver and gastric tissue samples.

5) **Gene expression analysis of melanogenesis-related genes in cultured human and murine adipocytes.**

Melanocortins interact with melanocytes through MC1R receptor. In these cells, stimulation by α-MSH enhances the expression, de novo synthesis, and activation of tyrosinase (Hunt *et al.*, 1994). In cultured human melanocytes synthetic α-MSH has been shown to induce a significant increase in the level of eumelanin but has lesser and varied effects on the level of pheomelanin (Hunt, 1995). Therefore, at least in melanocytes, the melanogenic pathway remains active when cells are cultured *in vitro*.

α-MSH plays a major role in energy homeostasis in adipose tissue and it is also required for activation in melanocytes of TYR, the rate limiting enzyme required for
pigmentation. Adipocytes constituting the major component of adipose tissue express MC1R, MC2R, MC3R, and MC4R receptors. There is a possibility that α-MSH is capable of inducing the expression of melanogenesis-related enzymes in the fully mature adipocytes. This hypothesis was attempted to confirm in cell culture based experiments. For this purpose, human and murine adipocytes and preadipocytes were incubated at different levels of differentiation with different concentrations of α-MSH for a number of time intervals. This helped to find the specific stage and concentration of α-MSH required to induce the expression of TYR.

4) **Enzyme activity assay analysis in human and murine adipocytes:**

Because mRNA expression does not necessarily reflect protein expression or function, the following assays were performed.

a) Tyrosinase enzyme activity was measured by L-dopa assay in cell lysates of murine adipocytes and preadipocytes incubated with α-MSH.

b) Tyrosinase enzyme activity was measured in human adipocytes at different stages of differentiation after incubation with α-MSH by specific C\(^{14}\) radioactive assay.

5) **Protein expression analysis in human adipocytes.**

Tyrosinase (TYR) is absolutely required for melanogenesis, but other melanosomal proteins, such as TYRP1, DCT, and gp100, also play important roles in regulating mammalian pigmentation. However, pigmentation does not always correlate with the expression of TYR mRNA or protein, as its function is also regulated at the post-
translational level. Even though TYR is ectopically expressed in a variety of tissues, it does not necessarily exist in these tissues in a catalytically active form, as its post-translational activation is required for melanin biosynthesis. After its synthesis, TYR is translocated into the ER and eventually into the Golgi, where its post-translational processing and glycosylation take place (Olivares et al., 2003; Francis et al., 2003). The fully glycosylated form of TYR is delivered to stage II melanosomes where it starts catalyzing the rate limiting step of melanogenesis. In order to catalyze the reactions, the enzyme should be properly glycosylated and have proper conformation.

a) The melanogenesis related enzymes TYR, TYRP1 and TYRP2 were localized in human adipocytes at different stages of differentiation after incubation with different concentrations of α-MSH by immunohistochemistry.

b) The protein expression for TYR, TYRP1, and TYRP2 was analyzed in human adipocytes at different stages of differentiation after incubation with different concentrations of α-MSH by western blotting.

c) The protein extracts obtained after incubation with different concentrations of α-MSH at different time periods at different stages of differentiation were checked for TYR enzyme activity by C\textsuperscript{14} assay.

d) After confirming the activity it was very interesting to see whether or not the TYR enzyme is properly glycosylated. Human TYR enzyme is a rate-limiting enzyme and has seven potential glycosylation sites and two copper binding sites. If the enzymes
are not properly glycosylated, they are considered to be inactive. In order to prove that the enzyme is properly glycosylated, glycosidase digestions were performed and obtained lysates were run the SDS gels along with the positive control to see the unglycosylated form of TYR.
Chapter IV
Materials and Methods

1) Patient tissue samples:
The visceral adipose (N=7) and gastric (N=2) tissue samples had been collected with informed consent during bariatric surgery as a part of ongoing Epidemiology of Nonalcoholic Fatty Liver Disease (EPI-NAFLD) study according to the protocol approved by Institutional Review Board of Inova Fairfax Hospital. Adipose samples from the non-obese kidney donors (N=2) and liver specimens from patients undergoing liver resections for liver mass with no underlying liver disease (N=2) were used as controls. Samples were snap-frozen in liquid nitrogen immediately after collection and stored at -80°C until the time of assay. Skin samples were obtained by shave biopsies, 4mm in diameter, as described previously (Tadokoro et al., 2003; Yamaguchi et al., 2004) and used as positive control.

2) Culture and maintenance of cell lines:
Experiments were performed on cultured adipocytes in parallel with a study of the tissue specimens. Four different cell lines were used:

   a) Mouse 3T3-L1 fibroblast cell line (ATCC, Manassas, VA)
a) Culture and differentiation of mouse 3T3-L1 fibroblast cell line:

The preadipocytes at passage 3 were purchased from ATCC (Manassas, VA). The media for maintenance and differentiation of cell line was purchased from Zen-Bio (Research Triangle Park, NC). Preadipocytes were cultured in 75cm² flasks (Corning, Corning, NJ) in preadipocyte medium (DMEM containing 10% bovine calf serum) and passaged three times without being allowed to reach confluence, then frozen in aliquots of 10⁷ cells/vial at -80°C for future use. For experimental purposes, the preadipocyte cells were grown in a 5% CO₂ atmosphere and preadipocyte medium was changed every other day until the cells reach 100% confluency. One day after confluence, the preadipocyte medium was replaced with 3T3-L1 adipocyte differentiation medium (DMEM containing 10% fetal bovine serum (FBS), 10µg/ml dexamethasone, and 0.5mM isobutylmethyl xanthine (IBMX) for 3 days. After removing the differentiation medium, cells were fed with 3T3-L1 adipocyte maintenance media (DMEM containing 10% FBS, 10µg/ml dexamethasone and 10µg/ml insulin) every other day. All cell-based assays and measurements were performed on preadipocytes, 2 days, 4 days and 6 days old differentiated cells after incubation with 100nM, 1µM, 5µM or 10µM concentrations of α-MSH peptides respectively. RNA and proteins were extracted after incubation at 0, 2hr, 6hr and 24 hr time periods for further analysis.

b) Culture and differentiation of primary adipocytes:
Human pre-adipocytes at passage 3 (Zen-Bio, Inc., Research Triangle Park, NC) were obtained from 5 female patients, age range from 29-43 yrs with BMI ranges of 38-52 after abdominal fat reduction surgeries. The preadipocytes were cultured in T25 cm² flasks with Preadipocyte Growth Medium (Zen-Bio, Inc.) at 37°C in 10% CO² atmosphere. Once the cells reached confluence, differentiation was induced by treating the cells with differentiation medium (DM) which contained insulin, dexamethasone, IBMX and a PPARγ agonist (Zen-Bio, Inc.). The medium was changed after seven days with adipocyte medium (DM without IBMX.) (Zen-Bio, Inc.), and subsequently the medium was changed every two days. All cell based assays and measurements were performed on preadipocytes, 7 days, and 14 days and 21 days old differentiated cells after incubation with 100nM, 1µM, 5µM or 10µM concentration of α-MSH peptides respectively. RNA and proteins were extracted after incubation at 0, 2hr, 6hr and 24 hr time periods for further analysis.

c) Culture and maintenance of MNT1 cell line:

The MNT-1 cells (Valencia et al., 2005) are highly pigmented human melanoma cells used as a positive control. Cells were grown at 37°C in an atmosphere of 95% air/5% CO₂ in 75-cm² culture dishes in MEM (all culture reagents from Life Technologies, Grand Island, NY). The medium is supplemented with heat-inactivated FBS to a final concentration of 20% (Atlanta Biologicals, Norcross, GA), 10% AIM-V medium, 20 mM Hepes, 1% antibiotic-antimycotic solution, 0.1 mM nonessential amino acids, 1 mM sodium pyruvate, 2 mM L-glutamine, 30 ng/ml gentamycin, and 3.7 µg/ml sodium bicarbonate.
d) Cultivation and maintenance of HeLa cell line:

The cells (human cervical carcinoma cells) were obtained from the American Type Culture Collection (Manassas, VA, USA) and used as a negative control. The cells were grown in Dulbecco’s modified Eagle’s medium supplemented with 10% bovine serum and antibiotics (30 mg/l penicillin, 50 mg/l streptomycin sulphate) at 37°C and 10% CO₂. RNA and protein were extracted from the cell line after incubation with α-MSH peptide in the same way as described from human adipocyte cell line and mouse 3T3-L1 fibroblast cell line.

3) Methods and procedures:

1) Real time PCR: Real time PCR is a technique that analyses the amplification of genomic DNA in Real Time. This means it is possible to evaluate the size of amplicons without having to run the finished product on a gel, though this is still recommended for confirmation of the result (Bustin, 2000). Real Time PCR is considered to be the most sensitive method for the detection of low-abundance mRNA (Bustin, 2000; Giuliani et al., 2001). The technique starts with extraction of RNA, cDNA synthesis, Primer designing which are explained below in detail.

A) RNA extraction

a) RNA extraction from cell lines:
The preparation of messenger RNA (mRNA) from eukaryotic cells and tissues is an important step which requires cell or tissue disruptions, denaturation of RNases and finally separation of mRNA from residual DNA contaminants. The important thing while performing the extraction of RNA is to maintain an RNase-free environment, because the ubiquitous RNase A is a highly stable and active ribonuclease that can contaminate extractions and degrade RNA.

The RNA was extracted at exponential phase, when cells were proliferating maximally. The cells were washed twice with PBS, trypsinized and centrifuged at 800g for 5mins. The cell pellet was washed twice with PBS and RNA extraction was performed using an RNeasy Mini Kit (Qiagen, Valencia, CA) according to manufacturer’s protocol with some modifications. The cells were disrupted by adding buffer RLT followed by homogenization. 1 volume of 70% ethanol was added to the homogenized cell lysate and mixed well by pipetting. Out of the total volume, 700µL of the sample was transferred to an RNeasy spin column and centrifuged for 30sec at 8000xg. Buffer RW1 (700µL) was added to the column and further it was centrifuged for 30sec at 8000xg, then 500µL of buffer RPE were added to the column twice and the column was centrifuged for 1min at 8000xg. RNA was eluted by adding 50µL of RNase-free water (Ambion, USA) and finally stored at -80°C.

b) RNA extraction from tissues: RNA extraction and analysis:

RNA was extracted from tissues obtained from morbidly obese and from lean subjects. Total RNA was extracted by following the procedure provided with the RNeasy
Lipid Tissue Midi Kit (75842, Qiagen, USA). For RNA extraction, 200mg of tissue was homogenized in 1ml of QIAzol lysis reagent by sonication until the tissue mass was converted into homogenous liquid. The contents after homogenization were transferred to 1.5ml test tube and incubated for 5min at room temperature to allow dissociation of nucleoprotein complexes. Chloroform (200µL) was added to the test tube and the tube was shaken vigorously for 15sec and incubated at room temperature for 5mins. The contents were centrifuged at 12,000xg for 15min at 4C. The upper, aqueous phase was transferred to a new tube and 600µL of 70% ethanol was added and mixed thoroughly by vortexing. The content was transferred to RNeasy Mini spin column placed in a 2ml collection tube and centrifuged for 15sec at 10000xg at room temperature. The flow through was discarded. This step was repeated again. Buffer RW1 (700µL) was added to the same column and centrifuged for 15 sec at 8000xg and the flow through was discarded. Buffer RPE (500µL) was added and centrifuged for 2 min at 10000rpm. The RNeasy spin column was placed in a 2ml collection tube and 50µL of water was added. The column was centrifuged for 1 min at 10000rpm to collect the RNA sample.

B) Assessment of the concentration and purity of RNA:

The concentration and purity of RNA was evaluated by using a Nanodrop ND-1000 Spectrophotometer (Nanodrop Technologies), RNA samples with the ratio of absorbance at 260nm and 280nm of 2.0 were accepted. The readings at 260nm are specific to nucleic acids while the readings at 280nm can detect proteins. If the ratio of 260:280 is between 1.8 and 2.0, there is little to no protein contamination within the sample. RNA integrity was checked by electrophoresis in a 2% agarose gel. Two strong
Ribosomal RNA bands were evident in each lane without smearing. This indicates a good quality sample without RNA degradation. DNase I treatment of RNA samples is a prime requirement in RT-PCR. The isolated RNA was treated with DNA-Free kit (Ambion Inc, Austin, TX) according to manufacturer’s protocol to get rid of chromosomal DNA to a degree that no background product is visible whereas in untreated samples false positives are produced due to cDNA synthesized from chromosomal DNA. Basically 0.1 volume of 10X DNase I reagent was added to the RNA sample, mixed gently and incubated at 37°C for 30 mins. After 30 mins 0.1 volume of DNase inactivation reagent was added and the samples were incubated at room temperature for 2 mins with occasional mixing. The samples were centrifuged at 8000xg and the supernatant was transferred to sterile test tubes.

C) Synthesis of the cDNA

Synthesis of the cDNA was performed using a first-strand cDNA synthesis kit (Invitrogen, Carlsbad, CA) according to manufacturer’s instruction with some modifications. RNA samples along with reaction components were thawed and vortexed. First strand reverse transcription cDNA synthesis was performed on 2μg RNA from each sample in 20μL reaction volume in a single reaction. Briefly, each RNase-free reaction tube contained 2μg of RNA, 250ng random hexamers (Invitrogen, Carlsbad, CA) and nuclease-free water (Ambion). The contents of the test tubes were mixed and centrifuged briefly. The tubes were heated at 70°C for 10 mins, and then cooled down to 4°C. After incubation, tubes were centrifuged to ensure that all reactants were located at the bottom
of the tube. Buffer (5x concentration, 4µL) (Invitrogen, Carlsbad, CA), 2µL of DTT and 1µL Superscript (Invitrogen, Carlsbad, CA) were added to the test tubes, the contents were briefly centrifuged and the test tubes were incubated at 42°C for 1 hour and then stored at 4°C.

D) Primers/Probe selection:

Primers were designed by using the DNA sequences published in GenBank and Primer3 (Frodo.wi.mit.edu), a free, web-based software (Fig. 7) with desired product size set to 100-150 bp size range for Real Time-PCR. To avoid degeneracy of primer sequence, sequences were compared to the sequence of the human or mouse genome by BLAST server of NIH. To avoid genomic DNA amplification the primers were either located in different exons or across an exon-exon boundaries. The program was set to pick forward and reverse primers 20 bp of length with Tm around 55°C. Desirable product size was automatically set to 100-120bp and default settings were used for rest of the parameters. The fragment length was confirmed by agarose gel electrophoresis of the product of the real-time PCR.
Primer3output

No mispriming library specified
Using 1-based sequence positions

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<tr>
<th>Primer</th>
<th>Start</th>
<th>Length</th>
<th>Tm</th>
<th>GC%</th>
<th>ANV</th>
<th>3' Seq</th>
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| LEFT PRIMER | 15    | 22     | 58.20  | 40.91| 4.00| 1.00
          |       |        |        |      |     | agcggaaagaagaatctgtga |
| RIGHT PRIMER | 120   | 20     | 58.52  | 50.00| 4.00| 2.00
          |       |        |        |      |     | saactccacagcggcagta |

SEQUENCE SIZE: 120
INCLUDED REGION SIZE: 120

PRODUCT SIZE: 106, PAIR ANY COMPL: 5.00, PAIR 3' COMPL: 0.00

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1 actctgtgtagtagtagctggaaagaaactctgtgacactccaacagccagtgcctgtgaga

61 ccttctgaggactagagggaaatgtcctgtgcctggttttgtactgcctgcgtgtggagttt
```

Fig 7: Primer design using Primer3 software. Part of the output produced by Primer3 shows the predicted Tm for the left (forward) primer and right (reverse) primer (58.20 C and 58.52 C resp.) and the size of PCR amplicon (106 bases). Output file highlights the positions of the primers on the template sequence provided.

In addition to the general criteria employed by Primer3, the free NetPrimer software (premierbiosoft.com) was used to estimate possible dimer formation, 3' hairpins, self-complementarity, palindromes, repeats, and G+C content. Special care was taken to match the melting temperatures ($T_m$s) of primer pairs (Fig. 8). Primers’ ∆G values and 3’end stability were also taken in account. The primers with 100.0 score ratings were selected for real-time PCR.
Fig 8: Net primer software used to check the compatibility of the designed primers.
Table 2: Primers designed for Real-Time PCR analysis of the murine 3T3-L1 cell line:

<table>
<thead>
<tr>
<th>Gene</th>
<th>Sense Primer</th>
<th>Antisense Primer</th>
<th>Temp</th>
<th>PCR fragment size</th>
</tr>
</thead>
<tbody>
<tr>
<td>18S</td>
<td>AGGAATTCCCAAGTAAGTGCG</td>
<td>GCCTCCTAAACCATCCAA</td>
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<td>140 bp</td>
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<tr>
<td>TYR</td>
<td>TCTGGCGCTTAGCAGTAGGC</td>
<td>CTGTTGGTAGTCGTCCTTTGTCC</td>
<td>56</td>
<td>107 bp</td>
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<tr>
<td>MITF</td>
<td>TCCCTATCCCAATCCACC</td>
<td>TGGATACCCAGATTGTGTCA</td>
<td>55</td>
<td>140 bp</td>
</tr>
<tr>
<td>POMC</td>
<td>GGAGACGCCGCTGTTTTC</td>
<td>ACTCGGCTCTGGAAGTCT</td>
<td>56</td>
<td>189 bp</td>
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<tr>
<td>TYRP1</td>
<td>TAAATCCAGAAGAGGAAGGGC</td>
<td>CATTTGTCGGTCGTGTCACC</td>
<td>54</td>
<td>140 bp</td>
</tr>
<tr>
<td>MC1R</td>
<td>CCTGTATGTCCTCCACCCAG</td>
<td>GATGATGATGTCCTCCAGAAGC</td>
<td>56</td>
<td>176 bp</td>
</tr>
<tr>
<td>MC2R</td>
<td>ACCATGACCGAGACCACCAGC</td>
<td>CGACGGACGAGGACAATC</td>
<td>56</td>
<td>103 bp</td>
</tr>
<tr>
<td>MC3R</td>
<td>TGTTTCATCGGCAATGGTCGC</td>
<td>TACGATGACCCAGAGGC</td>
<td>57</td>
<td>140 bp</td>
</tr>
<tr>
<td>MC4R</td>
<td>GGAACAGAAGGATGCTAAGGAGC</td>
<td>CCACGCCAGTGAGACTGTCACC</td>
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<td>101 bp</td>
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<tr>
<td>MC5R</td>
<td>AACTCCAGAAGAACCACCAGC</td>
<td>GGGAAGATTTCAATACACAGGC</td>
<td>56</td>
<td>183 bp</td>
</tr>
</tbody>
</table>

Table 3: Primers designed for human cell lines and tissues:

<table>
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<th>Sense Primer</th>
<th>Antisense Primer</th>
<th>Temp</th>
<th>PCR fragment size</th>
</tr>
</thead>
<tbody>
<tr>
<td>18S</td>
<td>AGGAATTCCCAAGTAAGTGCG</td>
<td>GCCTCCTAAACCATCCAA</td>
<td>55</td>
<td>140 bp</td>
</tr>
<tr>
<td>TYR</td>
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<td>GATGTCGATTGTCCATGGT</td>
<td>60</td>
<td>108 bp</td>
</tr>
<tr>
<td>MITF</td>
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<td>GACAGCTTGGAGATATAGTCCAC</td>
<td>60.2</td>
<td>118 bp</td>
</tr>
<tr>
<td>TYRP1</td>
<td>TGGTTTATTTCACGCTGCTTC</td>
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<td>59.9</td>
<td>106 bp</td>
</tr>
<tr>
<td>TYRP2</td>
<td>GGAAGCTGCTGAGTAGCCTT</td>
<td>GCCTGTTCTGCTCCATTTGATT</td>
<td>59.7</td>
<td>103 bp</td>
</tr>
<tr>
<td>MC1R</td>
<td>TGGTTTCCCGATGCTGCC</td>
<td>GCCACGATCAGATAGTTC</td>
<td>59.9</td>
<td>140 bp</td>
</tr>
</tbody>
</table>
E) Real Time PCR technique:

In this study, iCycler iQ real-time PCR (Bio-rad, Hercules, CA) was used for quantitative PCR analysis. The real-time PCR experiments were performed using the SYBR Green fluorescent probe, which binds non-specifically to any double-stranded DNA. The detection and quantification of fluorescence was directly proportional to the amount of PCR product in the reaction. The method quantifies the initial amount of the template specifically, sensitively and reproducibly, and is considered the preferred method over standard PCR. Real-time PCR monitors the fluorescence emitted during the reaction as an indicator of amplicon production during each PCR cycle. Primer dimerization needs to be limited by using minimal primer concentrations during the reaction. The optimized temperature of each PCR reaction was determined using the temperature gradient function of the iCycler iQ. Eight reaction tubes, using the same dilution of PCR product as template, were run at a range of annealing temperatures between 55°C and 62°C.

The real-time PCR reactions were set up in 96 well plates in a total volume of 15µL. Gene-specific master mixes were prepared by adding 7.5 µL of SYBR Green master mix (Bio-rad, Hercules, CA) to 4.5µL of nuclease-free water and 1µL of forward and reverse primers of 4nM concentration each. cDNA was added later to the reactions separately. In additions to reactions with cDNA, no template control (NTC) wells for every primer set were included, which contain all reaction components with 1µL of nuclease-free water (Ambion) instead of template cDNA. In addition to NTC, other controls were also used, for example, no primer control, to see if anything else in the
reaction mix will permit amplification of the cDNA target. Thermal cycling parameters for SYBR Green dye included first incubation at 94°C for 10 min, for 1 cycle, then 94°C for 20 seconds, 60°C for 20 seconds and 72°C for 30 seconds for 40 cycles. This set of cycles was followed by an additional extension step at 72°C for 5 min. For all RT-PCR reactions, melting curves were performed at the end of SYBR Green qPCRs to check for primer dimers and non-specific product formation. Only the reactions with a single melt curve were analyzed. The amplification of a single specific product was further confirmed by electrophoresis in 2% agarose gels. Real-time PCR results were calculated by using Pfaffl method (Pfaffl, 2001). All the reactions were performed in triplicates. Normalized gene expression values for each gene of interest and the housekeeping gene 18S rDNA were calculated. The expression values for each well were quantified as the “threshold cycle method” by the iCycler computer software (Bio-Rad, Hercules, California).

2) Western blotting:

A) Antibodies:

Rabbit polyclonal antibodies αPEP1h, αPEP7h and αPEP8h were specific to KLH-conjugated synthetic peptides that correspond to the carboxyl termini of human TYRP1, TYR and DCT, respectively. The peptides were as follows: PEP1h = CLLTDQYQCYAEERIWCOOH; PEP7h = CPLLMEKEDYHSLLYQSHL-COOH; and PEP8h = CMETHLSSKRYTEEA-COOH (Virador V, 2001).
B) Protein extraction from cell lines:

The cell pellets were obtained in the same way as described above for RNA extraction method. The cell pellet was solubilized in 500µL of M-PER mammalian protein extraction reagent (Pierce Biotechnology, Rockford, IL) and Protease Inhibitor mixture (Roche, Mannheim, Germany). Protein concentrations of extracts were quantified using the bicinchoninic acid protein (BCA) assay kit (Pierce, Rockford, IL). All steps were performed at 4°C to minimize the risk of proteolysis.

C) Protein extraction from adipose tissue:

Tissue (1gm) was homogenized in 1ml of RIPA buffer supplemented with protease inhibitors (Complete; Boehringer Mannheim, Indianapolis, IN), until all the tissue was converted into liquid. The liquid was snap frozen in liquid nitrogen and thawed at room temperature. The samples were centrifuged at 4000xg for 30mins at 4°C. All steps were performed at 4°C to minimize the risk of proteolysis. The infranatant was collected carefully and the protein concentration was measured by BCA protein assay (PIERCE, Rockford, IL) according to manufacturer’s instruction. Briefly, standards were made from BSA solution of 2000ug/ml, 1500ug/µL, 1000ug/µL, 750ug/µL, 500ug/µL, 250ug/µL and 125ug/µL with RIPA buffer, which was used for extraction of proteins. Each standard or unknown sample replicate (25µL) was added to a well of 96-well plates. Working reagent made by mixing 50 parts of BCA reagent A with 1 part of BCA reagent B. The working reagent (200µL) was then added to each well with standard or sample.
The plate was mixed thoroughly, covered and incubated at 37°C for 30 mins. Readings were taken with an ELISA reader (Bio-Tek Instruments Inc., USA).

D) Western blotting technique:

Protein extracts from adipocyte (15ug), Hela (15ug) and MNT-1 (15ug) cell lines were separated on 4-20% gradient denaturing SDS polyacrylamide gels (Invitrogen) at 160V for 1.5 hour. For tissue samples 15ug of protein from all tissues and 5ug of protein from MNT-1 cells were loaded into 4-20% gradient Tris-glycine gels (Invitrogen). Briefly, lysates were mixed in Tris-glycine SDS sample buffer (2X) (Invitrogen, Carlsbad, CA) containing 2-mercaptoethanol and were boiled for 5 mins at 95°C. After electrophoresis, proteins were transferred electrophoretically from the gel to Invitrolon PVDF transfer membranes (Invitrogen). Blots were blocked in 5% non-fat milk in 0.05% Tween-20 (Bio-Rad) in PBS for 1 hour at room temperature and were then incubated with antibody to Tyrosinase (rabbit, diluted 1:5000) and GAPDH (rabbit 1:10000; Santa Cruz Biotechnology) in 1% non-fat milk in 0.05% Tween-20 in PBS overnight at 4°C. After six washes (10 min each) with 0.05% Tween 20 (Bio-Rad) in PBS, the blots were incubated in horseradish peroxidase-linked goat anti-rabbit whole antibody (1:10000) (Amersham, Piscataway, NJ) in 1% non-fat milk with 0.05% Tween 20 in PBS for 1 hour at room temperature. After three washes (10 min each) with 0.05% Tween 20 in PBS, the immunoreactivity of the blots was detected using an ECL-plus Detection System (Amersham), according to the manufacturer’s instructions. Each experiment was performed at least in triplicate.

E) Glycosidase digestion performed on lysates of human adipocytes:
The methods used for predicting glycosylation sites analyzed both potential O-linked and N-linked sites. Endoglycosidase H (Endo H) and peptide: N-glycosidase F (PNGaseF) was purchased from New England Biolabs (Beverly, MA). MNT-1 cell lysates (1 µg), adipocyte cell lysates (1µL) and Hela cell lysates (1µL) were subjected to each glycosidase digestion according to the manufacturer's instructions. Protein extracts were denatured in 0.5% SDS, 1% 2-mercaptoethanol at 100°C for 10 min. One-tenth volumes of 0.5 M sodium phosphate (pH 7.5) and 10% NP-40 were added. N-Glycosidase F (2 µl of 500 U/µl) or Endoglycosidase H (2 µl of 5 mU/µl) was used for digestion at 37°C for overnight. Reactions were stopped by addition of an equal volume of 2x Lammeli loading buffer. After the digestion reaction, each cell lysate was subjected to SDS-PAGE, and immunoreactive bands were detected by western blotting using antibodies against TYR (dilution 1:5000).

3) Protein activity assays:

A) Protein activity assays performed on human tissues

a) Melanogenic assay: Materials: the radioactive substrates (L-[U-14C] tyrosine and L-[3,5-3H]tyrosine were obtained from both Amersham/Searle (Arlington Heights, IL) and Schwarz/Mann. BSA, chloramphenicol, cycloheximide and penicillin G were purchased from Sigma Chemical Co. (ST. Louis, MO, U.S.A.); Celite 545 and Norit A were purchased from Fisher Scientific Co. (Fair Lawn, NJ, U.S.A.).

a) 14C -tyrosinase enzyme activity: The method was a modification of the original technique described by Hearing and Ekel (1975). The tissue samples obtained from morbidly obese and from lean subjects were used as experimental samples whereas a
protein extract obtained from human melanocyte cell line was used as a positive control. Other controls used in this procedure were blanks from which substrate and/or enzyme were omitted. The assay (final volume 50µL) contained 30µL (50µg of protein) of sample in 10µL 14C isotope (25µCi/ml, specific radioactivity 100µCi/mmol) and 10µl of buffer solution containing made up of 1M potassium phosphate (pH 7.2), 1mg/ml chloramphenicol, 1mg/ml cycloheximide, 0.1mg/ml serum albumin, 1000U/ml penicillin B, and 0.25mM DOPA cofactor. The assay was set up on ice in triplicates in a 96-well round bottom microtiter plate and incubated for 16hr at 37°C. After incubation the plate was put on ice to stop the reaction and 40µL of sample were removed to Whatman 3MM filter-paper discs. The filters were dried and then washed as follows: one 15 min wash with 0.1N HCl (1l/50 filters) containing unlabelled tyrosine (1g/l), two 15 min washes with 0.1N HCl, two 5 min washes with 95% (v/v) ethanol (200ml/50filters) and one 5 wash with acetone (200ml/50liters). The filters were allowed to dry in air and the amount of radioactivity was determined using 0.4% (w/v) 2, 5-diphenyloxazole in toluene (with 70% efficiency).

**b) Melanogenic assay:** ³H-tyrosine assay for tyrosine hydroxylase: TH activity was determined by a modified method of Reinhard *et al.* (1986) . The assay was set up in the same way as described above and same controls were used except for the label, which was [³H] tyrosine (250µCi/ml, specific radioactivity 1µCi/mmol) and the cofactor DOPA was replaced by (6R)-R-L-erythro 5,6,7,8 tetrahydrobiopterin dihydrochloride (6BH4) at a concentration of 0.5mM. 6BH4 regulates tyrosinase through uncompetitive inhibition, whereas tyrosine hydroxylase requires 6BH4 as a cofactor for the hydroxylation reaction.
of L-tyrosine to L-DOPA. This method specifically measures the tritiated water produced during the hydroxylation of tyrosine to DOPA. Briefly, for each assay a 1 uCi aliquot of L-[3,5-3H]tyrosine was diluted 1:5 with a 500µM solution of cold L-tyrosine, HCL and used immediately. Each assay contained: 5.0 mM dithiothreitol (DTT) to ensure reducing conditions; 0.5M 6BH4; 9200units C-100catalase; 25uM L-tyrosine solution and 30µgm of protein sample. After incubation for 16hr at 37°C the activity was quenched by adding 40µL of sample from each well into plastic tubes containing 1ml of stirred charcoal slurry, 7.5% (w/v) in 1.0M HCl. The tubes containing the mixture were agitated vigorously every 10min for 1 hr and centrifuged at 1000g for 5 min. 100µL of supernatant was carefully removed from each tube into a scintillation vial containing 5ml suitable cocktail for aqueous samples. The radioactivity was measured in liquid scintillation counter and reported as, pmol product.

B) Protein activity assays performed on cells:

a) L-dopa assay:

Tyrosinase activity will be measured using L-DOPA as a substrate. The melanocyte cell line was used as the positive control, RPMI and HL60 cell lines were used as the negative control. Murine adipocyte cell line 3T3-L1 was used as the experimental samples and was incubated with the required concentrations of α-MSH as explained above for the respective time intervals. Since tyrosinase catalyzes the conversion of L-Dopa to dopachrome, this will permit measurement of the conversion of colorless DOPA to the dark orange dopachrome. α-MSH induces the expression of
tyrosinase enzyme, so we believe that after incubation tyrosinase enzyme will be present in the cell lysate. The standard reaction mixture contained 5mM L-DOPA, 0.1 M sodium phosphate buffer (pH 6.8) and the enzyme solution in a total volume of 3ml. Enzymatic activity was determined by the change of color of L-dopa to an orange color. The reaction was monitored continuously at room temperature using a spectrophotometer at 475nm. Phenoloxidase activity was assayed with L-DOPA as a substrate. It is based on the conversion of L-DOPA to L-DOPA-quinone by tyrosinase and results in the formation of dopachrome which is measured at 475nm.

\[
\text{L-DOPA} + \text{tyrosinase} \rightarrow \text{L-DOPA-tyrosinase complex} \rightarrow \text{L-DOPA quinone} + \text{tyrosinase}
\]

One unit of tyrosinase is defined as the amount of enzyme required to oxidize 1µM of L-DOPA per min under the above conditions.

6) Histochemical stainings

A) Staining performed on cellular model:

a) Immunocytochemistry:

Immunocytochemistry is a method of analyzing and identifying cell types based on the binding of antibodies to specific components of the cell. After three washes in PBS, the cells were fixed in 4% paraformaldehyde for 15 min at 4 °C. After three further washes in PBS, the cells were permeabilized with 100% methanol for 15 min at 4 °C and then blocked with 10% normal goat serum (Vector Laboratories, Burlingame, CA) for 1 hour at room temperature. The cells were incubated with primary antibodies against
TYR, TYRP1 and TYRP2 (gift from Dr Hearing, NIH) (at dilutions of 1:700, 1:700 and 1:7500, respectively) with 5% goat serum overnight at 4 °C in a humid chamber. After three washes with 0.05% Tween20 in PBS, the slides were incubated with Alexa Fluor® 594 goat anti-rabbit IgG (H+L) (Molecular Probes, Eugene, OR) at room temperature for 1 hour at 1:500 dilution with 5% goat serum and 0.05% Tween20, Nuclei were counterstained with DAPI (4,6-diamidino-2-phenylindole, blue fluorescence; Vector Laboratories). Alexa fluorescence was observed and analyzed using Leica DMR B/D MLD fluorescence microscope (Leica, Wetzlar, Germany) and ScionImage software (Scion, Frederick, MD).

B) Staining performed on tissues:

a) Preparation of frozen tissue sections:

The tissues were dissected and mounted in optimum cutting temperature compound (O.C.T) (Internal medical equipment, San Marcos, CA) then frozen at -80°C. The next day the frozen block of adipose tissue was sectioned into 10µm thickness in the cryostat chamber at a temperature of -18 to -20°C. Sections were stored at -80°C until further processing.

b) Preparation of frozen tissue sections:

The tissues were dissected and immersed in 70%, 90% and 100% ethanol sequentially for 30 min each at room temperature. Tissues were immersed in toluene for 20min at room temperature and embedded in paraffin (Paraplast, Fisher Scientific) two times for 60 min each at 58°C. The paraffin embedded tissues were cut into 10µm thick
tissue sections using a rotary microtome and floated in 56°C water bath, followed by mounting onto histological slides. The slides were dried overnight and stored at room temperature. Most of the work was performed on frozen tissue sections instead of paraffin embedded sections, because RNA in the frozen sections is more stable and it can be analyzed by in situ hybridization.

c) Fixation of frozen tissue sections:

Fixation of the tissue is the cornerstone for all types of staining. The main purpose of tissue fixation is to preserve tissue morphology and is essential for accurate histopathologic evaluation of the tissue. It arrests autolysis and putrefaction, coagulates soluble and structural proteins, fortifies tissues against the deleterious effects of subsequent processing, and facilitates staining. Preservation of tissue antigens and their availability for immunohistochemical staining and mRNA for in-situ hybridization depends on the type and concentration of fixative, on fixation time, and on the size of the tissue specimen to be fixed. Poor fixation or overfixation of the tissue sections can be detrimental for the histology technique.

Frozen tissue sections were fixed in 4% paraformaldehyde for 6 hr at 4°C. Fixation was followed by washing with PBS three times. The tissues were incubated with 70%, 90% and 100% methanol for 5 min each at 4°C for permeabilization and washed in PBS three times. The slides were dried with tissues and marked with a DAKO pen.

For paraffinized tissue sections, fixation was done after deparaffinization of tissue sections with xylene twice for 5 min each at room temperature. Deparaffinization was followed by dehydration step with 100%, 95%, 70% and 50% methanol for 3 min each at
room temperature. The slides were rinsed with PBS three times at room temperature followed by antigen retrieval step with 2% unmasking solution and heated in a microwave for 12 min and cooled down to room temperature for 20 min. The slides were dried with tissues and the tissue sections were marked with a DAKO pen.

d) Fontana-Masson staining:

Fontana-Masson staining (American Master Tech Scientific. INC, CA, USA) was performed according to the manufacturer’s recommendations with modifications. After fixation the tissue sections were rinsed with distilled water three times, followed by a quick rinse with running tap water followed by another quick rinse with distilled water. Fontana silver solution was prepared by adding ammonium hydroxide solution drop by drop to 10% silver nitrate solution followed by continuous stirring until a faint opalescence appears. The slides were placed in ammoniacal silver solution and incubated in a 60°C water bath for 35 min. After incubation the slides were rinsed with distilled water 3 times and placed in 0.1 % gold chloride for 1 min. The slides were rinsed with distilled water 2 times. The slides were placed in 5% sodium thiosulphate for 2 min. The slides were rinsed in running tap water for 3 min twice. Then the slides were placed in Nuclear Fast Red Stain for 5 min followed by rinsing in tap water twice and dehydrated with 3 changes of fresh absolute alcohol. The slides were cleared with 3 changes of fresh xylene. Finally the slides were covered with cover slips using the permanent mounting media. The pigment was observed with a light microscope. Commercially available skin samples were used a positive control.
e) Immunohistochemistry:

Immunohistochemistry involves the detection of specific chemical substances (infectious agents or cellular markers) within tissue by the use of derived antibodies to the substances. Antibodies (polyclonal or monoclonal) are applied to tissue sections and allowed to bind to their corresponding antigen. IHC was performed on adipose tissues slides obtained from obese and from lean subjects as explained above, skin sections were used as positive controls. After fixation the slides were washed with PBS twice for 3 min each. The tissue sections were marked with a DAKO pen. The slides were incubated in 20% goat serum (Vector Laboratories, Burlingame, California) for 1 hr at room temperature, then washed with PBS 3 times for 3 min each and incubated with the primary antibody overnight at 4°C in a humid chamber. The rabbit polyclonal antibodies against carboxyl termini of TYR (αPEP7h), TYRP1 (αPEP1h) and TYRP2 (αPEP8h) were diluted in PBS containing 2% goat serum up to 1:500, 1:500 and 1:1500, respectively. The next day the slides were given 3 washes with PBS containing 0.05% Tween-20 for 5 min each and incubated with Alexa Fluor(R) 594 goat anti-rabbit IgG (Molecular Probes, Eugene, OR) for 1 hr at room temperature at 1:200 dilutions with 2% goat serum and 0.05% Tween-20 in PBS. The slides were washed three times with PBS containing 0.05% Tween-20 for 5 min each in dark. After the final wash, the slides were mounted with DAPI plus Vector shield (Vector, CA) and glass cover slips were placed on the tissue sections and were fixed in place with nail polish. Red and blue florescence signals were observed and analyzed with Leica DMRB/DMLD laser microscope (Leica Microsystems, Bannockburn, IL) and ScionImage software (Scion, Frederick, MD).
IHC staining was also performed by using the Envision+ system-HRP (DAB) kit (Dakocytomation, Carpinteria, CA) for detection of the antibodies listed above. Briefly, after incubating the fixed slides overnight with primary antibody in the same way as for immunohistochemistry, the sections were washed three times with PBS for 5 min each. This step was followed with an application of enough hydrogen peroxide for peroxidase blocking and incubated for 5 min. The slides were gently rinsed with buffer and incubated with the secondary antibody: Labelled Polymer, HRP anti-rabbit (DAKO Corp., Carpinteria, CA) for 1 hr. Sections were rinsed 3 times with PBS containing 0.1% Tween-20 for five min each. For development the slides were incubated with 3,3'-diaminobenzidine tetrahydrochloride (DAB) substrate and chromogen for 10 min at room temperature. The color development was observed under the microscope to prevent background staining. The slides were rinsed again with PBS containing 0.1% Tween-20. After the washes, the slides were counterstained with hematoxylin (Sigma–Aldrich; St Louis, MO) for 3 min at room temperature, dehydrated and permanently mounted. For negative controls, primary antibodies were omitted.

f) In situ hybridization:

The purpose of the technique is to determine when and where the mRNA of a particular gene is localized, mainly by detection of complementary single-stranded nucleic acid strands (RNA or DNA), commonly known as "probes", which will hybridize to a specific mRNA. This technique is very time consuming and divided into cloning, PCR, labeling of riboprobe and hybridization with labeled riboprobe.
1) Cloning:

The human TYR and MITF clones in pSPORT6 vector were purchased from ATCC (Manassas, VA). The identity and orientation of the inserts were confirmed by sequence analysis. Re-cloning into the SP6 and T7 promoter containing vector pSPORT1 (Invitrogen, CA, USA) was performed for the tyrosine hydroxylase (TH) PCR4-TOPO clone (ATCC, Manassas, VA). RNA probes can be readily created utilizing SP6, T3, or T7 promoters in both sense and antisense orientations to provide non-specific (control) and specific probes.

a) Transformation:

The competent cells JM109 (Gene Choice) were transformed with pSPORT1 plasmid DNA and grown on agar plates overnight at 37°C for 16-18hr. Transformation was done by adding 50-100ng of plasmid DNA into 5µL of competent cells. The test tube containing both plasmid DNA and competent cells was placed on ice for 30min. After 30 min the test tube was immersed in 42°C water bath for 35sec, followed by immediate addition of SOC media (1ml) and the mixture was spread on agar plates containing the required amount of ampicillin. The next day single colonies were used to inoculate liquid broth containing ampicillin. Cultures were incubated overnight at 37°C at 225 rpm. Plasmid DNA was extracted the next day by using Plasmid mini kit (Qiagen) according to the instructions provided by manufacturer. Briefly, the bacterial cells were harvested by centrifugation at 5000 rpm for 15 min at 4°C. The bacterial pellet was resuspended completely in 300µL of Buffer P1 by vortexing or pipeting. Buffer P2 (300µL) was added to this mixture and mixed thoroughly by inverting the tube 4-6 times. The tube was
incubated at room temperature for 5 min. Chiled Buffer P3 (300µL) was added and mixed immediately by inverting the tube 4-6 times. The tube was incubated on ice for 5mins. The mixture was centrifuged at 14000 rpm for 10 min and supernatant which contains the plasmid was removed immediately. Qiagen-tip 20 was equilibrated with 1ml of Buffer QBT and allowed to empty by gravity flow. The supernatant containing plasmid was added to the equilibrated tip and allowed to empty by gravity flow. The tip was later washed twice with 2ml of Buffer QC and the DNA was eluted with 800µL of Buffer QF. The DNA solution obtained was precipitated by 560µL of isopropanal at room temperature. The mixture was centrifuged at 10000rpm for 30 min and supernatant was removed. The DNA pellet obtained was washed with 70% ethanol and air-dried. The DNA was redissolved in 50µL of pellet. The DNA concentration was calculated by using NanoDrop™ 1000 Spectrophotometer and the quality of the plasmid was determined by agarose gel electrophoresis.

Plasmid DNA of the PCR4-TOPO clone was extracted in the essentially the same way.

b) Restriction digestion:

To determine if the insert is of the correct size, the plasmids from both clones were restriction digested with EcoRI (New England Biolabs) in a total volume of 20µL. To each test tube 2µL of compatible buffer (New England Biolabs), BSA (10X), enzyme EcoRI each, 5ng of plasmid DNA and DEPC-treated water were added. The mixtures were incubated at 37°C for 2 hr and separated by electrophoresis on a 2% agarose gels.
To excise the TH cDNA insert from the PCR4-TOPO backbone, the plasmid was restriction digested with EcoRI as described above and the mixtures were separated on 2% agarose gels. The band on the gel corresponding to TH insert was extracted from the gel by Gel Extraction Kit (Qiagen, USA) according to manufacturer’s instructions. Briefly, the DNA fragment was extracted from the agarose gel with a clean and sharp scalpel. Three volumes of Buffer QG was added to the extracted gel in a sterile tube. The tube was incubated at 50°C for 10 min and 1 volume of isopropanol was added to the sample and mixed. The Qiaquick spin column was placed in a 2 ml collection tube and the sample was applied to the column and centrifuged for 1 min. Buffer QG (0.5µL) was added to Qiaquick column and centrifuged for another min. The column was washed with 750µL of Buffer PE and centrifuged for 1 min. The Qiaquick column was placed in a sterile test tube and the DNA was eluted with water. The quality and size of the insert was checked by agarose gel electrophoresis and the concentration of the insert was checked by NanoDrop™ 1000 Spectrophotometer.

c) Ligation:

Ligation was performed between the pSPORT1 plasmid (used as vector backbone) and the insert from PCR4-TOPO obtained after gel elution. The reactions were performed in duplicates. Prior to ligation the vector was restriction digested with EcoRI for 1 hr and 30 min and dephosphorylated with 1unit/µL of CIP for another 30 min. After two hours the test tubes were incubated at 65°C for 2 hr to inactivate the restriction enzyme and the CIP. For ligation the vector backbone and insert were used in the ratio 1:6 along with 2µL of ligase buffer and 1µL of ligase enzyme (New England biolabs).
The samples were incubated in 16°C waterbath overnight. Transformation was performed with these samples next day as explained above. The pSPORT1 vector expresses LacZ operon which makes the blue and white screening possible with IPTG and X-gal added to the agar medium. The transformed cells were allowed to grow on these agar plates for 16-18hr at 37°C. Next day only the big white colonies were selected, whereas the small colonies near the large colonies (satellite colonies) were disregarded. The correct orientation of the insert was confirmed by sequencing.

2) **PCR for in situ hybridization:**

The cDNA inserts for TYR, MITF and TH present in their respective vectors were amplified by PCR. Primers specific for amplification of human TYR, MITF and TH were designed. Target sites were selected based on the analysis of sequence matches and mismatches in BLAST (GenBank). The primers showed no evidence of crossreaction with sequences of other genes including other genes of the tyrosinase family. In case of tyrosinase, best results were obtained with the following probes: sense primer 5'-CATACGATTTAGGTGACACTATAG-aggcagcggaggcagagga-3'; antisense primer 5'-CGGCCGAATACGACTCACGATAGGG-gctgctcagctcgccgatgtcc-3'. The PCR reaction was performed in triplicates in PCR test tubes. Briefly, to each test tube 0.2 µg of plasmid DNA was added to, 1µL of dNTP (10µM final concentration), 2.5µL of EasyA transcription buffer (Roche) and EasyA polymerase (Roche), 1µL of reverse and forward primer each (10µM final concentration) and DEPC-treated water to make the final volume 25µL. The samples were incubated in the thermocycler that was programmed as follows: one cycle at 95°C for 2min, 28 cycles at 95°C for 30sec, 60°C for 30sec and
72°C for 1 min. This set of cycles was followed by an additional extension step at 72°C for 5 min and finally the products were cooled down to 4°C. The amplified product of the insert obtained after PCR was run on a 2% agarose gel electrophoresis, and extracted from the gel by using the gel extraction kit as explained above. The concentration of the DNA extracted from the gel was measured with NanoDrop™ 1000 Spectrophotometer.

c) **DIG labeling of the riboprobe:**

The amplified probe was labeled using a digoxigenine (DIG) RNA labeling kit (Roche) according to manufacturer’s protocol. Briefly, to each test tube 20 ng of template or amplified product was added to 2 μL of dNTP mix, 2 μL of 10X transcription buffer, 1 μL of RNase inhibitor, 2 μL of RNA polymerase (T7 or SP6) and water to make the final volume of 21 μL. The reaction mixture was incubated at 37°C for 2 hr. DNase I was added after 2 hr and was further incubated for 15 min. The riboprobe obtained was further purified using a RNeasy minikit (Qiagen) according to manufacturer’s instructions and stored at -80°C.

d) **Hybridization:**

The probes were 3’ tailed with digoxigenin-11-dUTP with a DIG RNA labeling kit (Roche, Basel, Switzerland), according to recommendations of the manufacturer. TISH was carried out as described previously with minor modifications. Briefly, after fixation of the tissues as described above, slides were then washed in glycine solution (2 mg/ml in PBS) for 10 min, then washed twice in PBS, and placed in 200 ml of acetylation buffer (0.1 M triethylamine, pH 8.0, containing 0.25% acetic anhydride) for 15 min. After washing in 4xSSC for 10 min, samples were incubated in prehybridization
solution (2xSSC, 50% deionized formamide) for 1 hr at 47°C. After overnight hybridization at 47°C, samples were placed in hybridization solution (3) containing 10 l of purified DIG-labeled antisense riboprobe. Samples then were incubated in 10 mM Tris-HCl, 0.5 M NaCl, and 0.25 mM EDTA (TNE) buffer, treated with RNaseA for 30 min, and returned to TNE buffer for 3 min, all at 37°C. After washing in 0.1xSSC for 15 min at 47°C, samples were blocked for 30 min and incubated with anti-DIG/HRP conjugate (DAKO, Carpinteria, CA) for 40 min at room temperature. For detection, we used the tyramide signal amplification system (GenePoint kit, DAKO) and VIP solution (Vector Laboratories) according to the manufacturers' instructions. Samples were observed and photographed in a Leica DMRB microscope.

7) LC-MS Analysis of Pyrrole-2,3,5-tricarboxylic acid (PTCA):

A) Extraction of melanin from hair:

The acid/base extraction of melanin from human hair was performed according to Bolt’s procedure (Bolt AG., 1967) with some modifications. Hairs were washed with detergent, water and acetone to remove dirt and lipid. The air dried hair was minced into 2-5 mm pieces and digested in 1N NaOH overnight. Concentrated HCl was added to the whole mixture to precipitate brown gum sediment. NaOH (1N) was used to dissolve the sediment, which was again precipitated by adding HCl. Base solubilization and acid precipitation was repeated 15 times until the supernatant of the acid wash was almost colorless. Melanoprotein concentrate obtained through the above procedure was stirred in 1M HCl for 4 hr and centrifuged. This step was repeated 10 times until the supernatant
was colorless. The resultant melanoprotein concentrate was washed with distilled water 6 times ethanol twice and finally washed with ether and lyophilized in a SpeedVac.

**B) Extraction of melanin from adipose tissues:**

One gm samples of adipose tissue were homogenized by sonication in 4 ml of RIPA buffer until the mass of tissue was converted to liquid. The samples were then were centrifuged at 4000 rpm. The fat was removed and the liquid phase was stored at -80°C. Pellets consisting of cell debris and black pigment were washed twice with PBS and centrifuged again at 4000 rpm. The pellets were dissolved in 1M NaOH to dissolve the pigment. HPLC was used to analyze the extracted pigments.

**C) Permanganate oxidation of synthetic and extracted melanin:**

Oxidation experiments were performed using 1mg of synthetic melanin, 1 mg of lyophilized hair melanin and melanin extracts obtained from 1 g of sonicated tissue samples as described above. To achieve complete solubilisation of the melanin, the mixtures were sonicated for 24 hr in 1ml of 1M NaOH. Each sample underwent permangate oxidation essentially as described by the original Ito and Fujita method (Ito and Fujita, 1985). Each sample was assayed in duplicate. Aqueous suspensions were prepared by sonicating 100µL of respective homogenates in separate test tubes with 100µL of BSA (20mg/ml) and 800µL of 1M H2SO4 at room temperature overnight. This step is required for the breakdown of melanin to PTCA which serves as an indicator of eumelanin presence that can be quantified by HPLC. To this, 3% KMnO4 was added in portions of 20µL while mixing until the purple color persists. At 10 min after the first
addition, the residual KMnO$_4$ and newly formed MnO$_2$ were decomposed by the addition of 100µL of 10% Na$_2$SO$_3$.

D) Sample preparation:

Samples obtained after permanganate oxidation were evaporated to dryness using a Centrivap sample concentrator (Labconco, Kansas City, MO). Samples were reconstituted in 1 mL of 18 M ultra-pure water and filtered into 12 x 32 mm deactivated amber autosampler vials (National Scientific Co., Rockwood, TN) using Millex hydrophilic PTFE syringe filters with 0.45 µm pores (Millipore Corp., Billerica, MA).

F) LC-UV-MS Analysis of Pyrrole-2,3,5-tricarboxylic acid (PTCA) protocol:

2.3.5-pyrroletetracarboxylic acid (PTCA) standards were obtained as a gift from Dr. S. Ito (Fujita Health University, Toyoake, Japan). Synthetic melanin (Sigma) and melanin extracted from hair were used as a positive control whereas a HeLa cell extract was used as negative control. The calibrating standards for the assay were prepared in Millipore-filtered water. A PTCA stock solution (1 mg/mL in water) was serially diluted in standard curve matrix to prepare a 10-point calibration curve ranging from 1 to 1000 ng/mL. A 10 µg/mL working solution of 3-nitro-tyrosine (internal standard) was prepared in 10mM ammonium acetate with 0.05% ammonium hydroxide. The applicability of this substitute matrix to accurately and precisely quantify PTCA in biological samples was established by using quality control (QC) samples prepared by pooling and spiking Caucasian skin sample extracts with known amounts of PTCA to prepare QC samples at five concentration points across the assay calibration range.
Samples were analyzed using a Waters 2695 Alliance Separations Module, a Waters Atlantis dC-18 column of 2.1 x 150 mm with 3 µm beads, a Waters 996 photodiode array detector (PDA) and a Waters ZQ2000 single-quadrupole mass spectrometer with an electrospray ionization (ESI) probe (Waters Corp., Milford, MA). Chromatographic separations were achieved with a binary gradient consisting of 0.1% v/v acetic acid in 18 M ultra-pure water as mobile phase A and 0.1% v/v acetic acid in acetonitrile as mobile phase B. Mobile phase B was increased linearly at a flow rate of 0.2 mL/min from 5% to 100% over 20 min and held at 100% for 10 min to purge the column. The gradient was returned to the initial conditions over 5 min and held for a 10-min column equilibration. Samples were kept in the refrigerated autosampler at 4 °C, and the column was heated to 35 °C.

The detectors were configured serially with the non-destructive PDA upstream to the MS. The PDA monitored the UV spectrum from 190 to 400 nm and recorded the absorbance at 270 nm for additional confirmation of PTCA. The MS ESI probe was operated in negative ionization mode with a capillary voltage of 3.5 kV and an extractor voltage of 5 V. The source temperature was 150 °C, and the desolvation temperature was 350 °C. The nitrogen desolvation gas flow was 250 L/hr and the cone flow was 50 L/hr. The mass spectrometer was operated in selected ion recording (SIR) mode monitoring ions 198, 154 and 110 m/z with a dwell time of 0.5 sec for each ion. The cone voltage was alternated between 14V and 25V during each 1.5 sec MS scan. The 198 m/z parent ion was produced at 14V. The higher cone voltage of 25V increased the kinetic energy
of the PTCA ion resulting in collision-induced fragmentation in the source to produce the 154 and 110 m/z secondary ions.

Quantitative analysis was accomplished using Waters Corp. MassLynx ver. 4.0 software. Five calibration standards were made in 18 M ultra-pure water at concentrations of 100, 250, 500, 750 and 1000 ng/mL. A peak from an unknown sample component partially co-eluted with the primary ion 198, therefore, the area response of secondary ion 154 m/z which was free from interference was used for external standard quantitation, and the ions 198 and 110 m/z were used for identification confirmation. A five-point linear calibration curve with forced 0, 0 axis generated a coefficient of determination value of 0.9997 for ion 154 m/z.
Initially adipose tissue was merely considered as a fat storage organ, but later with the discovery of leptin hormone less than a decade ago, in 1994, adipose tissue joined the list of endocrine organs. Besides being an endocrine organ, under morbidly obese conditions adipose tissue also acts as a source of inflammation by promoting the secretion and production of inflammatory molecules. Recently, a plethora of papers have been published on the adipocytic secretion of various adipokines and pro-inflammatory agents, uncovering additional regulatory roles of adipose tissue. Even though several novel signaling pathways have been identified in adipose tissue recently, additional studies are required for further characterization of the genes expressed in adipose tissue in order to evaluate the physiological role of this tissue. These studies may provide us with new approaches for the treatment of the metabolic consequences of excessive adipose accumulation and the lipodystrophy. In this study we describe a novel phenomenon of the expression of melanogenesis-related genes and biosynthesis of the melanin in the adipocytes obtained from morbidly obese subjects.
A STUDY OF THE MELANOGENESIS IN THE HUMAN ADIPOSE TISSUE

The expression of melanogenesis related genes TYR, TYRP1, TYRP2 and MC1R in human adipose tissue.

Aim of the study: To quantitative the difference in gene expression for TYR, TYRP1, TYRP2, MC1R and MITF in gastric tissue and adipose tissues from obese and from lean subjects.

A previous study performed in our lab revealed statistically significant overexpression of genes encoding tyrosinase-related protein 1 (TYRP1), dopachrome tautomerase (DCT/TYRP2), melanosome transport protein RAB27a and Melan-A (MLAN-A) in visceral adipose samples of morbidly obese patients (Baranova et al., 2005). These findings were made using cDNA-based microarray profiling of approximately 40000 different human mRNAs in human adipose tissue samples from morbidly obese subjects undergoing gastric bypass surgery and from lean kidney donors (Baranova et al., 2005). The TYR gene was not present on the cDNA microarray, so no conclusion concerning an expression of TYR was made at that time. To confirm these findings and to extend the list of the genes profiled, we performed real-time PCR to quantify mRNAs levels for DCT, TYRP1, MC1R and TYR in visceral adipose tissue of lean and of morbidly obese subjects and in gastric tissue controls.

In addition to the above mentioned melanogenesis-related genes we also quantified mRNAs of tyrosine hydroxylase (TH) gene. TH gene encodes for an enzyme present in the brain regions that are involved in the formation of neuromelanin. Similarly
to TYR, this enzyme carries out a rate-limiting step in the biosynthesis of catecholamines, by catalyzing the conversion of L-tyrosine to DOPA (Nagatsu et al., 1964); therefore, TH could theoretically substitute for TYR activity. Since much less is known about the biosynthesis of extracutaneous melanins, there is a possibility that TH is involved in melanin formation in adipose tissue.

The reliability of the results obtained with Real-Time PCR technique critically depends on the integrity and purity of mRNA preparation, so special consideration was paid to the RNA extraction process and further manipulations with these extracts. Each RNA preparation was characterized by three separate bands visible after electrophoresis of RNA on 1% agarose gel (Fig 9). The intactness, sharpness, and clearness of 28S, 18S and 5S ribosomal RNA bands indicated the integrity of the RNA samples. The size and brightness of 28S band was double than that of 18S band; the 2:1 ratio (28S:18S) provided an indication of RNA being intact. No evidence of smearing was found, which would otherwise have indicated degradation of RNA.
Fig 9: An example of the agarose gel used to confirm the integrity of the RNA samples. Three bands (28S, 18S and 5S) are visible at their respective positions on the gel.

Primers play a very important role in real-time PCR technique. It is also very crucial for successful real-time PCR that primers specific for the gene of interest amplify a single product. Primer design for TYR was evaluated by analyzing the amplicon viability and nonspecific product formation by standard PCR followed by electrophoresis in 1% agarose gels (Fig 10). As can be judged by the size of the obtained amplicon the primers amplified the correct product. No primer dimer or non specific product formation was recorded.
Fig 10: Agarose gel electrophoresis assuring the specificity of the PCR primers in the given condition: The figure shows the amplified fragment of DNA in adipose samples collected from morbidly obese (OA) and from non-obese (NOA) subjects by TYR primers.

Every Real-Time PCR reaction was monitored by the melting curves of the products obtained. Melting curve analysis allowed distinction of non-specific amplification products from true PCR amplicons. Melting curves were used as specific profiles that depicted the rate of change of fluorescence over time as a function of temperature for each sample. The products with larger target area (longer products) melt before smaller amplicons (http://DNA-9.int-med.uiowa.edu/realtime.htm.). SyberGreen dye changed the fluorophoric characteristics upon interacting with DNA, which offered a method of calculating the amount of DNA produced in a given PCR reaction by measuring the increased fluorescence (Moen et al., 2002). The cycle number at which the
level of fluorescence rose above a background threshold value was inversely proportional to the log of the number of initial template copies (Burgos et al., 2002; Mocelin et al., 2003a, 2003b). The use of melting curve analysis eliminated the necessity for a PCR sample to be analyzed via agarose gel electrophoresis because the melting temperature of the specific amplicon was analogous to the detection of an electrophoretic band (Giglio et al., 2003). Nevertheless, in our study we performed both types of the analysis, as described above.

![Melt curve diagram]

Fig. 11. The melt curve characteristics of the products for 18S and for the gene of interest.

In each reaction tube, only one product is formed (Fig 11). Absence of extra peaks (or humps) confirms the efficacy of the primer pair and reaction, as no primer-dimer or non-specific product formation was observed.
The quantification of expression for every gene of interest was based on the characteristic sigmoid curve produced during the real-time PCR reaction. Each curve depicted the increase in the intensity of the fluorescent signal that correlates with an increase in the amount of the product produced. Each curve reflects three phases of the polymerase chain reaction, the lag phase (little product accumulation), the exponential phase (rapid product accumulation), and the plateau phase (no further product is amplified) (Ririe et al., 1997; Bustin, 2000; von Ahsen et al., 2001; Wittwer et al., 2001; Bernard and Wittwer, 2002; Burgos et al., 2002; Mocelin et al., 2003a, 2003b). The point at which the lag phase crosses over into the exponential phase is known as the threshold cycle and was used for the quantification of the product.
Fig 12: An example of Real-Time PCR curves with the adjusted threshold.

There is always a possibility of errors in the quantification of mRNA transcripts by Real-Time PCR. These errors mostly arise due to variation in the amount of starting material between samples. Errors may result in the misinterpretation of the expression profiles of the target genes. In order to minimize these errors a cellular housekeeping gene/RNA (18S rRNA) was profiled as an internal reference against which other RNA values were normalized. This gene is assumed to be expressed at a constant level among different tissues, at all stages of development and not affected by the experimental treatments.
The figure shows the results of real-time PCR (RT-PCR) assays, which represents Figure 13. Relative abundance of MC1R, TYR, TYRP1 and TYRP2 transcripts in adipose samples collected from morbidly obese (OA) and from non-obese (NOA) subjects after the 18S RNA normalization procedure. Height of each bar corresponds to the level of the gene expression in the given sample. MCR1 expression values are shown in dark blue, TYR expression values in purple, TYRP1 – in yellow and TYRP2 – in light blue.

To determine the change in steady-state mRNA levels for each individual gene across multiple samples, we performed relative quantification of mRNA to the level of a reference gene 18S rRNA, a housekeeping gene co-amplified on the same plate. The relative abundance of mRNA obtained from each respective gene was normalized to the concentration of 18S rRNA in the same sample. The relative quantification of each
profiled gene was calculated by \( \Delta\Delta C_t \) model (Livak & Schmittgen, 2001). For each gene, \( \Delta C_t \) was calculated by subtracting the threshold cycle (Ct) value of target sample from that of reference gene sample. The concentration of template for a particular species of cDNA relative to the housekeeping genes was calculated using the formula \( 2^{\Delta C_t} \) for gene of interest.

**Figure 14:** PCR quantification curve for 18S and TH gene in brain tissue and adipose tissue.

Real-Time PCR assays revealed that the TYR mRNA was indeed expressed in adipose tissue, and its relative levels were higher in adipose tissue obtained from morbidly obese subjects than the adipose tissue from lean subjects (Fig 13). The attempt to detect mRNA for TYR in the sample from gastric lining yielded negative results (Fig 13). TYRP1, TYRP2 genes were also found to be expressed in the adipose tissue samples.
from morbidly obese subjects at higher levels than in adipose tissue samples from lean subjects. However, the differences in mRNA expression for TYRP1 and TYRP2 were somewhat less pronounced as compared to TYR. MC1R gene was also expressed in the obese adipose tissue at higher levels that in the samples form lean subjects, thus, confirming the findings of Hotche et al., 2007 (Fig 13). Tyrosine hydroxylase encodes an enzyme involved in formation of neuromelanin pigment in the brain. Gene expression analysis did not detect TH mRNA in adipose tissue of either obese or lean subjects. cDNA from brain samples showed amplification with crossing threshold cycle 26, whereas amplification curve obtained when amplifying TH on cDNA template from adipose samples crossed the threshold at cycle 44 or higher (Fig 14). The findings confirm the absence or extremely low levels of TH expression in adipose tissue.

Hybridization of a human TYR probe to the sections of the visceral adipose

Once the gene expression analysis confirmed the presence of the mRNAs of melanogenesis related genes in adipose tissue, we attempted to localize the expression of these genes within adipose tissue. The adipose tissue is a heterogenous organ that consists of various types of cells, including macrophages, fibroblasts, endothelial cells and adipocytes. Fully mature adipocytes mainly are comprised of central fat vacuole, with nucleus and cytoplasm pushed to the periphery of the cells. Hybridization of the DIG-labeled cRNA complementary to the tyrosinase mRNA on frozen adipose tissue (in situ) revealed an intense staining in the periphery of the adipocytes. Sense (control) RNA probes for tyrosinase revealed no signal. TYR mRNA expression levels were
significantly higher in adipose samples obtained from obese subjects as compared to those from lean subjects.

**Figure 16.** In situ hybridization of a human TYR RNA probe on the sections of the visceral adipose from morbidly obese and from non-obese subjects demonstrated cytoplasmic and membrane-associated staining pattern (20x magnification). A and B. T7 (cTYR) probe, visceral adipose from morbidly obese subject. C. T7 (cTYR) probe, visceral adipose from lean subject. D. SP6 (control) probe, visceral adipose from morbidly obese subject.
Protein expression for TYR, TYRP1 and TYRP2 in visceral adipose tissue samples from obese and from lean individuals revealed by immunohistochemistry.

Aim of the study: To confirm that the above mentioned melanogenesis related enzymes are expressed in the adipocytes and not in other cells of adipose tissue. The second aim was to compare the difference in protein expression of enzymes between adipocytes of obese and of lean adipose tissue.

Gene expression does not always correlate with protein expression (Nie et al., 2007). Particularly, high expression of mRNA may not lead to similarly prominent levels of the encoded cellular protein due to the changes in its posttranslational processing that may enhance the degradation of the protein. Therefore, after confirming the expression of the genes encoding melanogenesis related enzymes we evaluated expression of their corresponding proteins and localized them within adipose tissue. Particularly, the expression of enzymes participating in the melanin biosynthesis was localized by immunohistochemical staining of cryosliced adipose samples with αPEP7h, αPEP1h, and αPEP8h antibodies specific for TYR, TYRP1 and TYRP2, respectively.

Standardizing the Immunohistochemistry technique:

Immunohistochemistry is now accepted as the most useful ancillary method in diagnostic histopathology, apart from simple histochemical stains. The specificity of IHC is affected by a variety of factors, including the quality of the primary antibodies, fixation of the tissues, unmasking of epitopes and sensitivity of the detection system. Before performing the experiment, we standardized the IHC technique by taking into account all the above factors. Particularly, the processing of the adipose tissue represented the major problem. Visceral fat is mainly composed of lipids that melt at room temperature. Moreover, this type of tissue cannot be fixed by paraformaldehyde. In order to maintain
the histomorphology of adipose tissue, most of the work on the tissues specimens was performed at 4°C.

Prolonged fixation of tissue in paraformaldehyde and interruption of the formaldehyde fixation process before it is completed may result in irreversible damage to some protein epitopes and the cross-linking reactions only at the periphery of the tissue block, respectively. Prolonged fixation may also result in complete absence or weak staining that depends largely on the susceptibility of individual epitopes. Short fixation time results in more intense staining of the center or of the periphery of the tissue. Therefore, we performed a number of protocol modification trials that helped us to establish the correct conditions for tissue fixation. As melanogenesis related genes were expected to be expressed in adipose tissue in smaller quantities as compared to the skin, we adjusted antibody dilutions to the concentration allowing minimization of the background staining of tissue.

At appropriate antibody dilutions of 1:500, 1:500 and 1:5000 for TYR, TYRP1 and TYRP2 respectively, background staining was low, and the actual signal was high. Staining for the melanogenesis-related enzymes was evident in adipose tissues from obese and from lean subjects. The staining patterns were the same as seen for in-situ hybridization, revealing the expression of melanogenesis-related enzymes was limited to adipocytes only (Fig. 17 & 18). The cytoplasmic distribution of TYR, TYRP1 and TYRP2 was mainly restricted to the periphery of the cells. Expression of all three melanogenesis-related proteins was higher in the visceral adipose tissue from morbidly
obese as compared to non-obese subjects. TYR expression was higher than that of TYRP1 and TYRP2.
Figure 17. Immunohistochemical staining of the visceral adipose tissue sections from morbidly obese and non-obese subjects for human TYR, TYRP1 and TYRP2 proteins (20x magnification). Red: TYR, TYRP1 or TYRP2 stainings. Blue: DAPI (nuclei). A, C, E, G: Visceral adipose from morbidly obese subject. B, D, F, H: Visceral adipose from non-obese subject. A and B: Tyrosinase; C and D: Tyrosinase-related protein Tyrp1 (dopachrome tautomerase); E and F: Tyrosinase-related protein Tyrp1; G and H: Negative control (secondary antibodies) and DAPI.
**Figure 18.** DAB staining of the visceral adipose tissue sections from morbidly obese and non-obese subjects for human TYR, TYRP1 and TYRP2 proteins (20x magnification). Red: TYR, TYRP1 or TYRP2 stainings. Blue: DAPI (nuclei). A, C, E, G: Visceral adipose from morbidly obese subject. B, D, F, H: Visceral adipose from non-obese subject. A and B: Tyrosinase; C and D: Tyrosinase-related protein Tyrp1 (dopachrome tautomerase); E and F: Tyrosinase-related protein Tyrp1; G and H: Negative control (secondary antibodies) and DAPI.

**Optimizing the western blotting technique:**

*Aim of the study:* To validate the results of immunohistochemistry by western blotting on the gastric tissues and adipose tissues from obese and from lean subjects.

The separation of the proteins extracted from the adipose tissue in the gel and their transfer to a membrane was a challenging task that required optimization. Adipose tissue from morbidly obese subjects contains a substantial amount of lipids, and produces a very prominent albumin band that is the most visible band on the gel. Albumin molecules tend to bind to other proteins and interfere with their migration through the gel. Attempts to perform chloroform/phenol extraction of proteins from adipose tissues uniformly failed. For the purpose of this study, the proteins were extracted from adipose tissue after flash freezing the specimen in the liquid nitrogen, followed by homogenization of the tissues, another flash freezing of the homogenate and thawing it at the room temperature. The infranatants of the tissue homogenates were collected while carefully separating the fat. Samples were centrifuged before adding to the gel.

Expression of tyrosinase protein in visceral adipose samples from morbidly obese subjects was confirmed by western blotting with αPEP7h antibodies (Fig. 19). The 140KDa band corresponds to the dimer of the normal fully glycosylated tyrosinase.
was the only specific band observed after Western blot staining. Tyrosinase specific bands were not detected in the gastric lining sample nor in the visceral adipose from non-obese subjects.

**Figure 19.** Western blot reveals presence of tyrosinase in the protein extracts from visceral adipose tissue. OA1-4: adipose samples from morbidly obese individuals. NOA1-2: adipose sample from non-obese subjects; Ga: gastric sample; MNT1: highly pigmented human melanoma MNT1 cells; M: weight marker.
Measurement of the tyrosinase enzyme activity in tissue extracts by $^{14}\text{C}$ assay:

Aim of the study: To find whether the melanogenic pathway is fully functional in the adipose tissue and to compare the activity of the tyrosinase in the extracts of different tissues.

Tyrosinase is the rate-limiting enzyme defining the efficiency of the melanogenesis. Human tyrosinase protein possesses seven putative N-linked glycosylation sites and 17 lumenal Cys residues that participate in disulfide bond formation. Fully glycosylated Tyrosinase enzyme catalyzes ortho-hydroxylation of monophenols (cresolase activity), i.e. hydroxylation of tyrosinase to DOPA. Mutations resulting in loss of glycosylation disrupt the melanogenesis pathway by inactivating the tyrosinase enzyme.

In order to confirm whether the TYR expression in adipocytes of adipose tissue obtained from obese and from lean individuals is active and fully glycosylated, $^{14}\text{C}$ assay was performed. The total output of the melanogenic pathway initially carried out by tyrosinase was quantitatively evaluated by incorporation of the labeled L-$^{14}\text{C}$ tyrosine into its final product. Protein extracts of human gastric and liver tissues were used as negative controls, while an extract of the highly pigmented human melanoma MNT1 cells served as positive control. Tyrosinase activity in the liver and the gastric side controls was similar to blank negative controls with no protein extract added, while activities in the adipose samples of obese subjects were much higher (Fig 20) and were characterized by marked heterogeneity. Activity in the adipose sample of non-obese individual was approximately 2-fold lower than the mean activity value in adipose samples of obese subjects (Fig. 20).
Figure 20. Results of the L-[U-14C] tyrosine assay. OA1-7: adipose samples from morbidly obese individuals. NOA: adipose sample from non-obese subject; Ga: gastric sample; Liv: liver sample; MNT1: highly pigmented human melanoma MNT1 cells. Recorded values of melanogenic pathway activity for blanks (no protein extract added) and for gastric and liver side controls were similar (Blanks: 234.3 +/- 36.50 cpm; Ga: 237.0 +/- 15.2 cpm; Liv: 237.7 +/- 23.2 cpm). Recorded values for obese adipose samples were different from that of non-adipose samples (OA1-7: 845.29 +/- 99.38; Ga&Liv: 237.4 +/- 0.5). Recorded value for non-obese sample was 427.7 +/- 25.0. For representation purposes recorded cpm value were adjusted by subtraction of blank values and transformed to pmol product/µg/hr values. Activity of melanogenic pathway per µg of protein extract of MNT1 cells was approximately 20 times higher that that of the adipose samples, and was plotted to the different scale on the right side of the bar graph.
Melanin staining was confirmed by Fontana Masson staining:

_Aim of the study:_ to visualize staining for melanin in the adipose tissue from obese and from lean subjects and particularly in the adipocytes.

The histological examination of adipose tissue revealed an accumulation of melanin pigment in the periphery of adipocytes of adipose tissue from morbidly obese patients (Fig 21 A&B). The staining pattern revealed the specificity of the staining to adipocytes. This cytoplasmic staining was localized to the periphery of the cells with no staining observed in the lipid-filled central vacuoles of adipocytes, and in other cells of adipose tissue, including microvessels and fibroblasts (Fig 21E). Little or no melanin pigment was detected in the adipocytes from adipose tissue of lean subjects (Fig 21 C&D).
Figure 21. Masson-Fontana stain of human adipose tissue demonstrates melanin pigment (black staining) mainly in the periphery of the adipocytes. A and B. Multiple conglomerates of melanin granules are present at the periphery of the adipocytes of adipose tissue from morbidly obese subjects (20x magnification). C and D. Melanin granules are scarce in the adipocytes of adipose tissue from non-obese subjects (20x magnification). E. No melanin granules observed in the microvessels located in the adipose tissue (20x magnification). D. Melanin staining in skin tissue used as a positive control (10x magnification).

The presence of the melanin was revealed by LC-UV-MS measurement of PTCA in adipose tissue extracts:

Human body contains a variety of pigments which are similar in terms of color to the melanin, but differ in their chemical composition. Chemical composition of these pigments cannot be determined by histological methods used for melanin determination, such as the Fontana-Masson staining method, or immunohistochemical staining for melanocytes. Moreover, staining-based methods provide only subjective quantification due to the difficulty in discriminating the small differences in amounts of the pigments. In order to differentiate melanin from other pigments, for example, lipofuscin, an LC-UV-MS assay was developed.

Previous attempts to quantify melanin in human skin by direct analysis yielded little success owing to the relative insolubility of melanin over a broad pH range, lack of well-defined physio-chemical properties, and susceptibility to structural alteration by oxidative reactions during isolation procedures (A. Napolitano et al, 1995; S. Ito and K. Jimbow 1983; K. Wakamatsu and S. Ito, 2202; S. Alaluf et al, 2002). Quantitative
isolation of melanin from natural sources is hampered further by its strong affinity to protein components of the tissues (L. Novellino et al., 2000). The LC-UV-MS assay performed as part of this study was based on the HPLC analysis coupled with analysis of UV absorption of its 2,3,5-pyrroletricarboxylic acid (PTCA) derivative. The assay involved chemical degradation of the pigmented tissues followed by high performance liquid chromatography (HPLC) identification and quantification of a specific structural component of eumelanins, PTCA.

Initial evidence for the previous notion of melanin in adipose tissues was provided at the stage of homogenization of the tissue samples. A higher amount of black pigment was observed in adipose tissue from mobidly obese subjects, whereas no or little melanin was present in adipose tissues from lean individuals (Fig 24).

Before performing the actual assay, attempts were made to reveal possible spectrophotometric differences between synthetic melanin and melanin extracted from hair. The spectrophotometric analysis performed at 470 nm on the tow types of melanin yielded similar results Fig 22.
Fig 22: Spectrophotometric measurement for synthetic and melanin extracted from hair at 470nm

Both synthetic melanin and the melanin extracted from the human hair were used as positive controls. Confluent HL-60 cells collected from 75 cm$^2$ flasks were used as negative controls. Custom-synthesized stable PTCA was used as a standard. The degradation of the melanin that resulted in the production of the pyrrole-2,3,5-tricarboxylic acid (PTCA) was performed with potassium permanganate (Ito & Fujita 1985; Ito & Wakamatsu 1994). Samples of adipose tissue (1g) were liquefied by sonication, the resulting pellets were dissolved in NaOH, subjected to permanganate oxidation and used for PTCA quantification by LC-UV-MS.
A linear calibration curve ranging from 0.01 to 1.0 ng concentration was established for PTCA (Fig 23). Standard PTCA samples showed an excellent correlation with the weight of the sample with a linearity coefficient ($r^2$) of 0.999.

![Calibration Graph](image)

Fig 23: Standard calibration graph generated by HPLC analysis of PTCA.

LC-UV-MS analysis of PTCA positively identified and quantified this compound and its secondary ions in visceral adipose samples (Fig. 24 C&D). The collisionally activated dissociation of the PTCA precursor ion at mass-to-charge ratio (m/z) 198 produced abundant product ions at m/z 154 and 110 (Szekely-Klepser G et al., 2005) (Fig 24B). These product fragments were formed by loss of either one or two carboxylic acid functional groups from the PTCA molecule in the mass spectrometer collision cell. In
adipose samples PTCA ion 193 m/z co-eluted with an interfering component of the tissue extract, whereas ion 154 m/z did not co-elute with a tissue component ion. Hence, ion 154 m/z was used as the quantitative ion for all samples.

Quantification of the 154 m/z PTCA ion in profiled visceral adipose tissue sediments of three morbidly obese subjects revealed the presence of the ion in concentrations ranging from 0.19 to 0.12 ng/µL, while its concentrations in oxidized adipose tissue sediment from two non-obese subjects were 0.05 ng/µL (abdominal visceral adipose) and 0.0009 ng/ µL (perirenal fat). PTCA ion 154 m/z was not detected in the sediments of the gastric tissue sample and of the cultured HL-60 cells (appr. 10^7 cells).

The absorbance spectra at 270nm for PTCA product obtained from synthetic, hair and melanin extracted from obese adipose tissues were at almost at the same retention time 5.90 min (Fig 24 E). No peaks were observed for lean adipose tissues or from gastric tissues.
Figure 24. The presence of the melanin in adipose extracts as revealed by LC-UV-MS. A. Homogenized samples of adipose tissue separate in three phases: supernatant (fat), aqueous and sediment. Cell debris sediments from adipose samples of morbidly obese subjects contain visible amounts of black pigment. a and b: adipose sediments of non-obese subjects; c and d: adipose sediments of obese subjects. B: The CAD of the PTCA precursor ion at mass-to-charge ratio (m/z) 198 produces abundant product ions at m/z 154 and 110 peaks. C. LC-MS Multi-ion SIM chromatogram of PTCA peak at retention time of 6 min. D. Negative ESI mass spectra of PTCA peak at 6 min. E. HPLC – UV/VIS Chromatograms at 270 nm.
A summary of attempts to develop an in vitro model of adipocytic melanogenesis.

Expression of melanogenesis-related genes was demonstrated in the experiments with the human adipose tissue samples both at mRNA and protein levels as described above. Furthermore, by analyzing the chemical composition of the adipose pigment by LC/UV/MS proved that the pigment produced in these adipocytes is the melanin. To pave the way for the study of the regulation of the ectopic melanogenesis in adipocytes, attempts were made to reproduce the findings in the primary human samples in adipocytic cellular model on various stages of differentiation cultivated in vitro. Two types of cells were used: murine fibroblastic cell line (3T3-L1) differentiated into adipocytes and primary human pre-adipocytes purchased from Zen-Bio. At the different stages of differentiation, these adipocytes were exposed to different concentrations of α-MSH 100nM, 1µM, 5µM and 10µM at different time periods. The physiological concentrations of α-MSH tested in the experiment were chosen according to the existing literature. Preadipocytic cells and differentiated cells at different stages were incubated with 100nM, 1uM, 5uM and 10uM concentrations of α-MSH for 2hr, 6hr and 24hr time periods. The main objective was to determine at which stage and which concentration of α-MSH the biosynthetic pathway will be induced.

After incubation with α-MSH the expression levels of tyrosinase mRNA and protein expression were recorded in both cell lines. Both in 3T3-L1 preadipocytes and 2 day differentiated 3T3-L1 adipocytes gene expression profiles of TYR were similar same as its mRNA levels were increased in the first two hours of incubation with α-MSH, then
decreased at 6 hours and finally rose again at 24 hours of incubation (Fig 25 A&B), thus, demonstrating biphasic response. The 4 days differentiated cells showed a drastic decrease in the expression of TYR in response to α-MSH (Fig 25 C). The 6 days differentiated cells responded similarly to preadipocytic and 2 day differentiated cells (Fig 25 D). The overall expression levels of TYR mRNA were lower in 6 days differentiated adipocytes as compared to preadipocytes.

**Figure 25.** Relative abundance of TYR transcripts in 3T3-L1 cell line (A) Expression level of TYR in preadipocytes after incubation with 1μM α-MSH at baseline, 2hr, 6hr and 24hr. (B) Expression level of TYR in 2 days differentiated cells after incubation with
1uM α-MSH at baseline, 2hr, 6hr and 24hr. (C) Expression level of TYR in 4 days differentiated cells after incubation with 1uM α-MSH at baseline, 2hr, 6hr and 24hr. (D) Expression level of in 6 days differentiated cells after incubation with 1uM α-MSH at baseline, 2hr, 6hr and 24hr.

In human preadipocytes, expression levels of TYR mRNAs decreased after 2, 6 or 24 hour of exposure to α-MSH as compared to the baseline (Fig. 26 A). In cells differentiated for 7 days, expression levels of TYR mRNAs decreased after 2 hours of exposure to α-MSH as compared to the baseline. Six and 24 hours incubation of these cells to α-MSH resulted in less pronounced decrease of TYR mRNA levels (Fig 26 B). Cells differentiated for 14 days reacted to 2 and 6 hours of exposure to α-MSH by a decrease of TYR mRNA levels, while 24 hours exposure resulted in an increase in these levels (Fig 23 C). Cells differentiated for 21 days (full differentiation) reacted to any length of exposure to α-MSH by an increase in TYR mRNA levels. This increase was most pronounced after 2 hours exposure as compared to 6 and 24 hours exposure (Fig 23 D).
As seen from the results described above, the melanogenesis-related genes display inconsistent expression profiles in real-time PCR. One explanation for this inconsistency
is the possible differential regulation of the expression of the tyrosinase gene in cells at the different stages of differentiation grown in media with different compositions. There is also a possibility that differentiating cells do not attain maturity at the same time, which may result in inconsistency of the results. Also, adipocytes embedded in the adipose tissue are accompanied by a wide variety of supporting cells, particularly fibroblasts, macrophages and stromal vascular cells that are not present \textit{in vitro}. The murine 3T3-L1 model adipocytes might differ from the primary adipocytes by their transcriptional programs as they has been shown to express only MC2R and MC5R (Boston & Cone, 1996).

Tyrosinase activity in the 3T3-L1 cell line was analyzed by direct L-DOPA assays. In 3T3-L1 preadipocytic cells TYR activity was lower after 2, 6 or 24 hour of exposure to \( \alpha \)-MSH as compared to the baseline (Fig. 27 A). In 2 days differentiated cells, TYR activity remained the same as the baseline after 2 hours of exposure to \( \alpha \)-MSH. Six and 24 hours incubation of these cells to \( \alpha \)-MSH resulted in a slight decrease of TYR activity (Fig 27 B). Cells differentiated for 4 days and for 6 days showed a decrease of TYR activity after 2, 6 or 24 hour of exposure to \( \alpha \)-MSH as compared to the baseline (Fig. 27 C). Six and 24 hours incubation of the 6 days differentiated cells with \( \alpha \)-MSH resulted in less pronounced decrease of TYR activity (Fig 27 D).
Figure 27. Assay of tyrosinase activity in 3T3-L1 cell line through the biocatalyzed oxidation of L-DOPA into orange colored product dopachrome, measured at 475nm wavelength. (A) Absorbance spectra of dopachrome in preadipocytes after incubation with 1μM α-MSH at baseline, 2hr, 6hr and 24hr. (B) Absorbance spectra of dopachrome in 2 days differentiated cells after incubation with 1μM α-MSH at baseline, 2hr, 6hr and 24hr. (C) Absorbance spectra of dopachrome in 4 days differentiated cells after incubation with 1μM α-MSH at baseline, 2hr, 6hr and 24hr. (D) Absorbance spectra of dopachrome in 6 days differentiated cells after incubation with 1μM α-MSH at baseline, 2hr, 6hr and 24hr.
Immunohistochemical staining of the human adipocyte cell line for human TYR proteins (20x magnification). Red: TYR staining. Blue: DAPI (nuclei). A, B: TYR staining performed on preadipocytes at baseline and 24hr exposure to 1µM α-MSH, C, D: TYR staining performed on 7 days differentiated adipocytes at baseline and 24hr exposure to 1µM α-MSH. E, F: TYR staining performed on 14 days old adipocytes at baseline and 24hr exposure to 1µM α-MSH. G and H: Negative control (secondary antibodies) and DAPI.

In the experiments described below the expression of tyrosinase was also analyzed at the protein level by immunohistochemistry. The presence of the expression of TYR protein was demonstrated in primary human adipocytes (Fig 28). The expression of tyrosinase was easily detectable in adipocytes before exposure to α-MSH, while after prolonged exposure to α-MSH the amount of this enzyme decreased. Interestingly, in fully differentiated adipocytes tyrosinase demonstrated both cytoplasmic and nuclear staining, while in the less differentiated cells the staining pattern remained only in cytoplasm. At all stages of differentiation, TYR expression was found to be more pronounced than TYRP1 and TYRP2.

Since TYR expression was recorded in cultured adipocytes, C\textsuperscript{14} assay was performed: a radioactive assay to analyze the net output of melanogenic pathway. C\textsuperscript{14} assay of the protein extracts of human adipocytes at different stages differentiation after exposure to α-MSH at different time periods did not reveal significant differences. Although the values obtained were slightly higher than the background values, no difference was registered when values were compared. The observation supports the presence of the residual tyrosinase activity in cultured cells, but fails to reveal any signs of its differential regulation in the adipocyte development.
Table 4: C_{14} assay readouts (CPM values) of the total outputs of the melanogenic pathway in the extracts of the primary human adipocytes differentiated in culture 14 days and 21 days as compared to the extracts of the adipose samples from morbidly obese patients

<table>
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<th>Replicate 2</th>
<th>Replicate 3</th>
<th>Average</th>
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<td>387</td>
<td>400</td>
<td>392.00 +/- 7</td>
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<td>14 Days baseline</td>
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<td>668</td>
<td>650.33 +/- 22</td>
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<td>628</td>
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Tyrosinase activity was registered by C_{14} assay (Table 4). The enzyme activity of TYR was very low as compared to the obese adipose tissue, but it remained the same during the course interval of incubation with α-MSH. In contrast, western blotting performed on the protein extracts obtained after incubation with α-MSH at different time points showed decrease in the TYR protein expression (Fig 27 A).
Chapter VI
DISCUSSION

White adipose tissue is a heterogeneous organ with major part consisting of loose association of lipid-filled cells called adipocytes, which are held in a framework of collagen fibers. In addition to adipocytes, adipose tissue contains stromal-vascular cells including fibroblastic connective tissue cells, leukocytes, macrophages, and pre-adipocytes (not yet filled with lipid), which contribute to the structural integrity. During the past 10 years, the understanding of the physiologic and pathophysiologic role of the adipose tissue has completely changed. Traditionally, adipose tissue was considered to be a passive type of connective tissue involved only in storing excess energy as triglycerides. In 1987, adipose tissue was for the first time described as a source of bioactive molecules carrying out various functions in the human body. These proteins are collectively known as adipokines and act at both the local (autocrine/paracrine) and systemic (endocrine) level (Siiteri, 1987).

Nowadays, adipose tissue has been established as a true endocrine organ coupling endocrine and metabolic signaling (Kershaw and Flier), with adipocytes secreting a majority of the hormones. This critical change in our perspectives on white adipose tissue came about due to the discovery of the cytokine-like factor, leptin (Zhang et al., 1997). Secretory products of both mature adipocytes and stromal adipose cells are referred to as adipocytokines (2), which regulate energy homeostasis, appetite/satiety, reproduction, insulin sensitivity and influence neuroendocrine, endothelial, immunological,
hematological, angiogenic and vascular functions in endocrine, paracrine and autocrine manners. These adipokines consist of polypeptides but also nonprotein factors that are known to affect immune function (complement factors and haptoglobin), endocrine function (leptin, sex steroids, and various growth factors), metabolic function (fatty acids, adiponectin and resistin), angiogenesis (Vascular endothelial growth factor) and cardiovascular function (angiotensinogen and plasminogen activator inhibitor-1) (Trayhurn & Beattie, 2001; Trayhurn & Wood, 2004). With the exception of adiponectin, most circulating factors are elevated. Even though a large group of peptides have been discovered to be expressed and secreted by the components of adipose tissue, there are undoubtedly more of these molecules yet to be discovered.

In previous studies expression of melanogenesis-related genes was observed in the primary adipose tissue samples. The results were obtained by studying transcriptome of the visceral adipose of morbidly obese individuals (Baranova et al., 2005). Moreover, when visceral fat of morbidly obese individuals was compared to that of lean subjects, these genes were found to be expressed at the higher levels. In the current study, the expression of melanogenesis-related mRNAs and proteins, namely TYR (Tyrosinase), DCT (Tyrp1), TYRP2 and MITF (microphthalmia transcription factor) in human adipose tissue was confirmed by real-time PCR and by immunohistochemical (IHC) staining. Additionally, TYR mRNA signal was registered by in situ hybridization in the visceral adipocytes, and TYR protein in the protein extracts of the visceral adipose tissue of morbidly obese subjects. The staining of the western blot with antibodies to human TYR
revealed the 140 kDa band which corresponds to mature tyrosinase homodimer representing properly folded enzyme (Francis Parag 2003).

The presence of the melanin pigment in human adipose tissue was revealed by Masson–Fontana staining (Fig. 21) and was further characterized by permanganate degradation of melanin coupled with Liquid Chromatography/ Mass Spectrometry (LC/UV/MS) determination of pyrrole-2,3,5-tricarboxylic acid (PTCA) derivative of the eumelanin.

To exclude an obvious possibility that a certain level of melanogenic activity is a common characteristic for many human tissues, total outputs of the melanogenic pathway in adipose tissue and other human tissues was compared by incorporation of the labeled L-[U-14C] tyrosine into its final product, an acid insoluble melanin. Activities of the pathway in the adipose samples of obese subjects were much higher than that in the non-obese adipose tissue (Fig. 20), while activities in the protein extracts of human gastric and liver tissues were similar to that of blank controls. Marked heterogeneity of the melanogenic activities of individual adipose tissue extracts was noted. The connections between extracutaneous accumulation of the melanin and morbid obesity were never described before, although a hyperpigmentation of the skin known as acanthosis nigricans has been associated with either clinical or subclinical insulin resistance, a component of the metabolic syndrome.

What would be a plausible explanation for the fact that adipose tissue of morbidly obese patients produces higher levels of melanin compared to lean subjects? Ectopic synthesis of melanin may serve as a compensatory mechanism that utilizes anti-
inflammatory (Mohaghheghpour et al., 2001) and oxidative damage absorbing (Rózanowska et al., 1999; Seagle et al., 2005) capacities of this compound.

The maintenance of the BMI within the healthy standards is recommended for the prevention of conditions associated with oxidative injury and increased inflammation. Weight gain generally results in obesity, which is mainly related to an increase in number and size of adipocytes, infiltration of adipose deposits by mononuclear cells, and the relative rarefaction of blood vessels and neural structures (Wellen & Hotamisligil, 2003). Obesity is associated with a state of chronic, low-grade inflammation characterized by abnormal cytokine production, increased acute-phase reactants, and activation of inflammatory signaling pathways (Hotamisligil, 2003). Through the activation of ‘stress-sensing pathways’, metabolic and endocrine alterations are produced. These pathways probably contribute to the pathogenesis of co-morbidities associated with obesity. The cluster of diseases associated with obesity includes development of insulin resistance and subsequent diabetes type 2, atherosclerosis and other cardiovascular diseases, dyslipidemia and hypertension that feature a number of common abnormalities of both metabolic and inflammatory pathways.

TNFα was the first inflammatory cytokine recognized in white adipose tissue, where it acts as powerful local regulator within adipose tissue and influences a wide range of processes (Prins et al., 1997; Coppack, 2001) including production of cytokines and adipokines. TNFα is markedly increased in morbidly obese subjects (Hotamisligil et al., 1993). Plasminogen activator inhibitor-1 (PAI-1), another important bioactive substance produced in adipose tissue by adipocytes, has been shown to play an important regulatory
role in the fibrinolytic process and in thrombus formation (Shimomura et al., 1996). The increase in the level of expression and secretion of these and other inflammatory agents are linked to development of the manifestations of the metabolic syndrome, particularly in diabetes type II and atherosclerosis. In addition to inflammatory agents, adipose tissue also secretes anti-inflammatory agents, for example, adiponectin; however, the decline in the concentration of these agents has been documented in the state of obesity (Ouchi et al., 2000; Arita et al., 1999). The levels of adiponectin could be downregulated by an increase in expression and activation of TNFα and peroxisome proliferators-activated receptors α (PPARα) (Moore et al., 2001). Overproduction of several circulating markers, particularly, IL-18, MIF, SAA, IL-6, TNFα, C-reactive proteins and haptoglobin, has already been reported in morbidly obese subjects (Yudkin et al., 1999; Das, 2001; Festa et al., 2001).

Adipocytes themselves are considered as the immediate source of these inflammatory markers, as an increase of their production parallels an increase in white adipose mass. In addition to that, some new models propose that as a consequence of the production of proinflammatory molecules inflammation is enhanced through infiltration of adipose tissue by macrophages and by interactions between the adipocytes and macrophages (Xu et al., 2003; Weisberg et al., 2003). The recent series of publications by Fain and co-authors indicated that the net production of pro-inflammatory molecules by adipose tissue is supported mostly by non-fat cellular components of this tissue, including macrophages (Fain et al., 2007; Fain et al., 2006; Fain et al., 2004).
One possible mechanism for the increased inflammation within white adipose tissue is the hypoxic conditions. White adipose tissue is not highly vascularized as compared to brown adipose tissue and moreover the cross-sectional area of blood vessels per unit weight of adipose tissue is reduced in obesity (Summers, 1996, Summers, 1999). Due to less oxygen provided to the growing adipocytes, the clusters of adipocytes become relatively hypoxic, which leads to an induction of leptin and VEGF expression in adipocytes through hypoxia-inducible factor-1α (HIF-1α). Immunoreactive HIF-1α has been reported in 3T3-L1 adipocytes (Lolmede et al., 2003). The initiation of hypoxia may also leads to the stimulation of the release of inflammatory cytokines, chemokines and angiogenic factors, the function of which is to increase blood flow and stimulate vascularization (Trayhurn & Wood, 2004). Since production of ROS is regulated by oxygen tension, ROS have been hypothesized to be a signaling mechanism in hypoxia-induced pulmonary vasoconstriction and vascular remodeling. Increased production of ROS selectively in adipose tissue of obese mice was found to be accompanied by augmented expression of NADPH oxidase and decreased expression of antioxidative enzymes. Furthermore, an increased ROS production is also implicated in arteriosclerosis, which is also indirectly linked to an increase of inflammatory agent secretion by adipocytes.

Another source for an increase of inflammatory processes in the adipose tissue is through the enhancement of the lipid peroxidation. Subcutaneous and visceral adipose tissues are thought to be composed of 60 to 85% of lipids with 90-99% being triglycerides, while the rest is composed of small amounts of free fatty acids, diglyceride,
cholesterol and phospholipids. These mixes of lipids are under continuous pressure for lipid peroxidation. Lipid peroxidation is a continuous process which proceeds by a free radical chain reaction mechanism and results in the formation of ROS. Excessive production of these molecules creates "oxidative stress," which can damage cellular structures and trigger an inflammatory response (Hotamisligil, 2006). The primary cellular defense against the toxicity of free radicals of oxygen is the antioxidant activity of several enzymatic systems such as superoxide dismutase, catalase, and glutathione peroxidase.

Recently, the melanin as been proposed as the non-enzymatic antioxidant compound (Sichel et al., 1987) capable of the reduction of the oxidative stress in the skin. Melanin could be also involved in the prevention of oxidative stress generated by lipid peroxidation in other pigmented tissues. In one study, a comparison of liver tissue from albino rats and melanin-pigmented frog liver demonstrated superior resistance to lipid peroxidation related oxidative stress in amphibian tissue (Scalia et al., 1990).

As has been mentioned before, an increase in the inflammatory and oxidative stress is the common feature of obesity. In most cases, these conditions lead to the development of the obesity-associated metabolic syndrome. Nevertheless, some morbidly obese patients with BMIs > 45 remains relatively disease free, having a normal degree of resistance to insulin (Jarrar et al., 2007). These findings speculate that the existence of some differential factors capable of counteracting the proinflammatory oxidative pressure within adipose tissue and actively suppressing the secretion of harmful adipocytokines.
Speaking generally, these, yet unknown, factors, can be regarded as endogenous protectors, allowing some morbidly obese patients to remain relatively healthy. So it can be proposed that the endogenous adipocytic melanin as one of these hypothetical factors.

Below is the summary of the hypothesis that will serve as the basis for the future studies aimed at the testing of this hypothesis. With the progression of obesity and an increase of the cellular fat depot, adipocytes become more exposed to endogenous apoptotic signals, especially ROS. To counteract pro-apoptotic ROS effects, the adipocytes, in turn, may ectopically activate the genetic program of melanogenesis, thus, neutralizing excessive ROS. Adipocytic melanin also suppresses secretion of pro-inflammatory molecules (Mohaghheghpour et al., 2001), therefore, decreasing the pro-inflammatory background in the obese body and alleviating the metabolic syndrome.

Very high levels of polymorphism in human genes regulating melanin biosynthesis may account for the highly individual melanogenic response of adipocytes that may account for the differences in an individual’s propensity to develop secondary complications of obesity. Interestingly, limited clinical information available for 5 out of 9 patients sampled for visceral adipose allowed us to reveal a positive correlation between fasting glucose levels and total outputs of the melanogenic pathway in adipose tissue \((R = 0.9685, \ p <= 0.007)\). This observation might indicate a connection of adipocytic melanogenesis to insulin resistance.

Alternatively, an explanation for an activation of the melanogenic pathway in adipose tissue may be related to the significantly elevated levels of an endogenous melanogenic peptide \(\alpha\)-MSH in the serum of obese subjects (Hoggard et al., 2004). \(\alpha\)-
MSH exerts its effects through the melanocortin receptors which have been shown to relay the signal resulting in the execution of the wide range of relevant effects both in the nucleus and in cytoplasm of the human cells. For example, in melanocytes MC1-R triggers \(\alpha\)-MSH-induced melanogenesis in melanocytes, whereas in other tissues this receptor propagates an anti-inflammatory signal. Recently, MC1-R has been shown to be expressed by adipocytes of morbidly obese subjects (Hoch et al., 2007). Other MC receptors, for example, MC2-R, are also found on the surface of the adipocytes, where they participate in lipolysis, whereas adipocytic MC3R and MC4R are involved in energy homeostasis (Cone, 2005). Etopic melanogenesis in human adipose tissue may result from excessive exposure of human adipocytes to melanogenic signals, including \(\alpha\)-MSH. The described phenomenon is possibly more complex than that and being connected to other, not yet uncovered facets of human obesity.

Interestingly, the G-coupled melanocortin receptors signal through the PKA pathway resulting in the induction of cAMP. Recently, the cAMP-activated protein kinase (AMPK) cascade (Heymans & Bouckaert, 1930) has come to prominence as a sensor of metabolic stress that appears to be ubiquitous throughout eukaryotes (Hardie, 2003). AMPK is activated by many types of the metabolic disorders, including heat shock and metabolic poisons in hepatocytes (Corton et al., 1994), exercise in skeletal muscle and ischemia and hypoxia in the heart (Marsin et al., 2000). An explanation of whether increased levels of \(\alpha\)-MSH in the serum of morbidly obese subjects are capable to activate AMPK.
What are the factors inducing the program of ectopic melanogenesis in adipose tissue? A variety of genes has been shown to be differentially expressed in adipose tissue under morbidly obese conditions (Baranova et al., 2005; MacLaren et al., 2008; Marrades et al., 2006; Lee et al., 2005; Nair et al., 2005). Other factors change their states on post-translationally. For example, HIF1-α is rapidly degraded under normoxic conditions of lean adipose tissue and is stabilized in obesity. Hypoxia is known to result in disturbances in adipokine secretion and increased macrophage infiltration in adipose tissue, events that are frequently observed in obesity (Goossens, 2007). On top of that, hypoxic conditions prompt human tissue to paradoxically release ROS (Guzy & Schumarker et al., 2006). A novel mechanism of tyrosinase activation by H₂O₂ was proposed recently, providing a connection between melanogenesis and pro-inflammatory background in the ROS-overproducing obese adipose tissue (Schallreuter et al., 2008).

Taking into account all written above, the plausible explanation for an increase in the melanin biosynthesis seen in the adipose tissue of the morbidly obese patients is the protective properties of the melanin. The above study performed on adipose tissues raised more questions than it reduces answers. Both the causes and the consequences of the melanin production in adipose tissue remain unknown; one obvious candidate for regulatory molecule is α-MSH that is intimately connected to both melanogenesis and energy homeostasis. The immediate attempts will be aimed at the development of the appropriate cellular system allowing study of the adipocytic melanogenesis in vitro and at exploring the connection between melanogenesis and metabolic syndrome.
Interestingly, attempts to register melanogenesis in human and mouse adipocytes differentiated from primary cultured preadipocytes repeatedly failed. Despite substantial expression levels of *TYR, TYRP1, DCT* and *MITF* mRNAs neither an enzymatic activity of tyrosinase, nor the production of the enzyme was detected *in vitro*. Cultured adipocytes are probably unable to support adequate post-translational modifications or the folding of tyrosinase and, therefore, are deficient in this sense. This notion supports previous evidence that the molecular networks activated during adipocyte development *in vivo* and *in vitro*, although overlapping, are in many respects quite different (Soukas *et al.*, 2001). Next plan is to continue efforts aimed at the development of the suitable *in vitro* system for the studies of the regulation of the production of ectopic melanin. Particularly, next attempt will be to induce melanogenesis in the cultured human adipocytes by the stimulation with melanogenic peptide hormone α-MSH and/or compounds inducing differentiation of adipocytes, e.g. by the insulin and the agonists of *PPAR-γ*.

In conclusion, the study performed demonstrates for the first time that the biosynthesis of the melanin is indeed taking place in visceral adipose tissue of morbidly obese subjects. Thus, an entirely new phenomenon never reported before is uncovered in this study. Further research into this area is warranted.
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