Transcriptional Regulation of Metabolism

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Desvergne, Béatrice, Liliane Michalik, and Walter Wahli. Transcriptional Regulation of Metabolism. Physiol Rev 86: 465–514, 2006; doi:10.1152/physrev.00025.2005.—Our understanding of metabolism is undergoing a dramatic shift. Indeed, the efforts made towards elucidating the mechanisms controlling the major regulatory pathways are now being rewarded. At the molecular level, the crucial role of transcription factors is particularly well-illustrated by the link between alterations of their functions and the occurrence of major metabolic diseases. In addition, the possibility of manipulating the ligand-dependent activity of some of these transcription factors makes them attractive as therapeutic targets. The aim of this review is to summarize recent knowledge on the transcriptional control of metabolic homeostasis. We first review data on the transcriptional regulation of the intermediary metabolism, i.e., glucose, amino acid, lipid, and cholesterol metabolism. Then, we analyze how transcription factors integrate signals from various pathways to ensure homeostasis. One example of this coordination is the daily adaptation to the circadian fasting and feeding rhythm. This section also discusses the dysregulations causing the metabolic syndrome, which reveals the intricate nature of glucose and lipid metabolism and the role of the transcription factor PPARγ in orchestrating this association. Finally, we discuss the molecular mechanisms underlying metabolic regulations, which provide new opportunities for treating complex metabolic disorders.
I. INTRODUCTION

Our knowledge of metabolism mainly results from the long-standing and very extensive work of a myriad of biochemists. They first achieved the identification of enzymatic steps, their functional characterization, and the discovery of regulatory loops with which they are associated. For decades, allosteric controls linked to substrate availability constituted the best of our knowledge of metabolic control systems. A crucial step was then accomplished with the deciphering of the galactose operon in bacteria, which represented a major discovery for the study of metabolism. It revealed how organisms can adapt their metabolic activity to environmental nutritional changes by modifying the level of expression of specific enzymes and linked modulation of enzymatic activity to the transcriptional control of gene expression for the first time.

It is now commonly accepted that metabolic regulation in complex organisms relies on three main types of control. The first corresponds to the classic allosteric control of the activity of a key enzyme along a metabolic pathway triggered by the binding of an activator, which often is the enzyme substrate itself. The second mechanism involves various posttranslational modifications such as proteolytic cleavage, phosphorylation, glycosylation, sumoylation, and acetylation, which may shift the equilibrium between an inactive and active enzyme within seconds and/or affect protein stability. In these two types of control, subsequent changes in protein-protein interaction may participate in producing the active/nonactive enzymatic complex. The third mechanism is transcriptional regulation, which affects the level of expression of key enzymes and is effective on a longer time scale. It clearly appears that most metabolic regulations benefit from a coordination of these various mechanisms. The purpose of this review is to highlight the recent progress in understanding when and how transcriptional regulation participates in the control of metabolic homeostasis.

Transcriptional control requires specific signals to be transduced to the cell nucleus where defined sets of genes are targeted. Thus understanding the transcriptional control of metabolism relies on three complementary pieces of information: 1) events upstream of transcriptional activity, which define the signals involved and their route to the nucleus; 2) the molecular mechanisms by which transcription factors operate; and 3) events downstream of transcriptional activity, which depend on the groups of genes that are targeted and how further signals are generated to reach the dynamic equilibrium of homeostasis. Virtually all transcription factor families are in one way or another involved in metabolic regulation. However, a few of them have a clear predominant role and seem mainly dedicated to metabolic regulation. For the sake of clarity, the transcription factors most often cited herein are briefly presented. Appendix A presents the nuclear receptor family and is accompanied by Figure 1A, which shows the main characteristics of the transcription factors belonging to this family. The notion of “metabolic sensor” receptors was more particularly developed with respect to these nuclear receptors, as also explained in Appendix A with the accompanying Figure 1B. Appendices B–F describe the main features of some of these “sensors,” which belong to the nuclear receptor family, with the peroxisome proliferator activated receptor (PPAR) in Appendix B, the liver X receptor (LXR) in Appendix C, the farnesol X receptor (FXR) in Appendix D, the hepatocyte nuclear factor 4 (HNF4) in Appendix E, and the retinoid X receptor (RXR) in Appendix F. Appendix G details the amazing characteristics of the sterol response element binding proteins (SREBPs), which play a major role in lipid and cholesterol metabolism. Finally, the heterogeneous family of proteins initially grouped under the name of liver-enriched transcription factors and which comprises the CAAT enhancer binding proteins (C/EBP) is discussed in Appendix H.

In recent years, interest has increased in cofactors that bridge proteins and allow the DNA-bound transcription factors to transmit their activation or repression properties to the transcriptional machinery. These cofactors are characterized by 1) their ability to interact with a wide variety of transcription factors and 2) their ability to assemble a protein complex that will be the transcriptional effector. Importantly, these cofactors are direct targets of certain signaling pathways, as seen with the insulin-dependent phosphorylation of the CREB-binding protein (CBP). With respect to the role of cofactors in transcriptional regulation of metabolism, for example, a clear picture has emerged from study of the PPAR gamma coactivator 1 (PGC1), which is implicated in thermogenesis and in associated metabolic responses, and of CBP in contributing to neoglucogenesis (see appropriate sections). Specific metabolic roles for steroid receptor coactivator 1 (SRC-1), transcriptional intermediary factor 2 (TIF2), and the receptor interacting protein 140 (RIP140) are also emerging, and further work should guarantee several important new developments in this field. However, it is beyond the scope of this review to discuss transcriptional cofactors specifically, and we refer to reviews that have recently been devoted to the subject (e.g., Refs. 85, 207, 250).

The aim of this review is to summarize recent knowledge concerning the transcriptional control of metabolic homeostasis. Analyses of the main transcriptional controls occurring in the regulation of intermediary metabolism is followed by an integrative approach which illustrates how these regulations can take place during the alternation between fasting and feeding to achieve energy homeostasis. In a pathological context, the disruption of energy homeostasis reflected by the metabolic syndrome highlights the intricate link between glucose and lipid.
metabolism. The last section discusses transcription factors as targets for treating complex metabolic disorders.

II. TRANSCRIPTIONAL CONTROL BY GLUCOSE

A. Introduction

Glycemia is a parameter over which the organism establishes tight control. In humans, blood glucose levels are kept constant in a narrow range from 4 to 7 mM, despite discontinued supply due to the alternation between feeding and fasting. One main danger of prolonged hypoglycemia is acute brain damage. At the other end of the scale, acute hyperglycemia is a serious complication of decompensated diabetes mellitus. The associated ketoacidosis and hyperosmolar hyperglycemic state might be fatal due to dehydration and electrolyte imbalance. Chronic hyperglycemia is a major cause of neuropathy and vasculopathy, as seen in diabetes.

Glucose homeostasis is maintained by a hormonal network in which insulin and glucagon are the main agents. Synthesis and secretion of insulin are stimulated by increased glucose levels, particularly after feeding. Insulin release allows the quick removal of glucose from circulation by stimulating the entry of glucose into peripheral tissues, mainly in muscle and adipose tissue cells. In parallel, insulin increases energy storage by inducing glycogen synthesis in liver and muscle, and fatty acid synthesis in liver and adipose tissue. When insulin levels are low, between meals or upon fasting, the hormone glucagon increases the hepatic production and release of glucose by increasing glycogenolysis and stimulating gluconeogenesis. The pancreas is the chief organ of these dual regulations, as it senses glucose levels and produces insulin and glucagon accordingly. The liver functions as the main “buffer,” providing glucose when nutrients are scarce and storing glucose as glycogen when food is abundant. Once the liver glycogen store is full, the adipose tissue converts glucose into triacylglycerol for longer term storage as fat. Muscles mainly consume rather than store energy, although they efficiently accumulate glycogen for their own use. The brain is a particular target organ that can use glucose and/or ketone...
bodies as an energy source. However, the fact that glucose represents the sole source of energy for some of its cells imposes a tight control over glycemia. In this organ, the entry of glucose in cells is mediated by the Glut3 transporter, which maintains a constant supply of glucose to brain cells until glycemia drops to very low levels close to its $K_m$ value, i.e., when approaching 2.2 mM.

The aim of this section, which cannot be exhaustive, is to discuss the main threads of the transcriptional network which result in glucose homeostasis. As is the case for many pathways regulated by nutrients, glucose is both an end product and the nutrient substrate that triggers regulation. Therefore, two opposing situations are considered for simplicity: that of high and that of low glucose levels. For each situation, we will describe the signals that are triggered and their action in transcription.

B. Transcriptional Regulation of Metabolism by High Glucose Levels

High glucose levels influence gene expression either directly or through the stimulation of insulin production by the $\beta$-cells of the pancreas. We first analyze the transcriptional regulation of insulin$^1$ and the regulation of insulin secretion. We then review the mechanisms by which glucose and insulin, independently or together, modulate gene transcription.

1. Transcriptional control of insulin expression and secretion

Pro-insulin is synthesized in the $\beta$-cells of the pancreatic Langerhans islets and is then cleaved by proconvertases in insulin and peptide C. Insulin is stored in secretory vesicles, and its secretion is directly linked to a mechanism sensing glucose availability via an increase in the intracellular ATP/ADP ratio that correlates with the entry and metabolism of glucose in the $\beta$-cells (Fig. 2, Ref. 230). The entry of glucose into the $\beta$-cells requires a glucose transporter, Glut2 in rodents but Glut1 rather than Glut2 in humans (56), whose expression and membrane localization are independent of glucose or insulin signaling. The posttranscriptional control of insulin expression and processing, as well as the control over the secretory mechanism, which is dependent on glucose sensing, are key features of the regulation of insulin signaling. However, the pathologies exhibited by patients in whom the regulation of insulin gene expression is altered emphasize the importance of the transcriptional level of control.

While insulin mRNA and protein expression have been found in various tissues (151) in different diabetic mouse and rat models, insulin is normally produced in highly specialized $\beta$-cells in the pancreatic islets. The tissue-specific expression of insulin is tightly regulated at the transcriptional level, and the major regulatory elements are located in the 5’-flanking region of the insulin gene. Among the set of transcription factors involved, PDX1 (pancreatic duodenum homeobox) is a key component (214) (Fig. 2). PDX1 is the main determinant in the cell lineage of the developing endocrine pancreas and in combination with other transcription factors confers tissue-specific expression of insulin (reviewed in Ref. 215). PDX1 is also the glucose-sensitive transcription factor of the insulin gene transcription machinery. Indeed, glucose triggers the phosphorylation of PDX1, via the phosphatidylinositol 3-kinase (PI3K) pathway, which induces the nuclear translocation of PDX1 and increases insulin expression (185, 237). Other transcription factors and/or coactivators activated by glucose likely contribute to the PDX1-mediated glucose response of insulin (189).

In addition to HNF3$\beta$/FOXA2, which positively regulates PDX1 expression (83), other members of the HNF family, HNF1$\alpha$, HNF1$\beta$, and HNF4$\alpha$ are expressed in the pancreatic $\beta$-cells. The maturity-onset diabetes of youth (MODY) has highlighted the importance of this network of transcription factors, acting directly or via a cascade of transcriptional regulation on insulin gene expression, and possibly on insulin secretion (310, 319) (see also Fig. 2 and Appendix H). MODY is characterized by the appearance in children or young adults of a non-insulin-dependent form of diabetes mellitus, inherited as an autosomal dominant trait. Except for MODY2, which is caused by a mutation in the enzyme glucokinase, MODY is due to mutations in genes encoding transcription factors involved in insulin gene expression (see Fig. 2). Alteration of HNF1$\alpha$, which causes MODY3, is the most frequent transcription factor defect leading to MODY, whereas MODY1 is rare and due to mutations in HNF4$\alpha$. MODY4 is characterized by a primary defect in insulin synthesis and secretion due to mutations in PDX1. Finally, two Japanese families with mutations in HNF1$\beta$ responsible for MODY5 have been reported (reviewed in Ref. 319). The fact that mutations in any of these genes result in altered insulin secretion reveals that each of these transcription factors is crucial for the control of cell specificity and metabolic adjustment of insulin expression.

2. Insulin-regulated transcription of genes involved in glucose metabolism

In the insulin-targeted cells, transduction of the insulin signal from the cell surface to key regulatory factors in the cell nucleus occurs in a very short time frame, allowing the immediate adaptive response of the cells (Fig. 3). In short, circulating insulin interacts with its membrane insulin tyrosine kinase receptor, ex-
pressed in most cells in vertebrates. This interaction drives the activation of the Ras/mitogen-activated protein kinase (MAPK) pathway. The cascade of phosphorylation events starts with the phosphorylation of insulin receptor substrate 1 and/or 2 (IRS1, IRS2). The successive activation of son-of-sevenless (SOS), Ras, and Raf-1 subsequently activates MEK (mitogen-activated, ERK-activating kinase), which in turn phosphorylates MAPK. Activation of this pathway seems to mostly target cellular growth and proliferation, rather than direct metabolic actions, and will not be discussed further.

An important relay of the metabolic action of insulin is the activation of PI3K leading to that of PDK1, which in turn phosphorylates the serine/threonine kinase Akt (also called PKB). Activated Akt/PKB phosphorylates several factors, including GSK3β and FOXOs, which directly or indirectly mediate the effects of insulin on the transcription of genes involved in glucose metabolism. There are three highly homologous Akt/PKB members, Akt1/PKBα, Akt2/PKBβ, and Akt3/PKBγ. Specific gene deletion or gene silencing of Akt1 and Akt2 demonstrates the primary role of Akt2 in insulin signaling and glucose metabolism, while significant redundancy of Akt1 and Akt2 still exists (80, 129). In addition to activating Akt, insulin triggers the tyrosine phosphorylation of the protooncogene Cbl in a PI3K-independent manner. Phosphorylated Cbl stimulates the translocation of the glucose transporter 4

Fig. 2. Transcriptional regulation of the insulin gene in the presence of high glucose. The roles of the HNF family of proteins and the central role of the transcription factor PDX1 in the regulation of the expression of the insulin gene are highlighted. Gene alterations responsible for the maturity onset diabetes of the young (MODY) are shown in red. For further details, see section II.B. Insulin secretion is triggered by an increase in cytosolic ATP/ADP, which closes K+ channels in the plasma membrane, thereby causing membrane depolarization and opening of voltage-gated Ca2+ channels. The resulting rise in the cytosolic Ca2+ concentration activates exocytosis of insulin-containing granules (230). Factor X, hypothetical factor acting on insulin secretion; F6P, fructose-6-phosphate; F1,6-P2, fructose 1,6-diphosphate; Glut, glucose transporter; G6P, glucose-6-phosphate; HNF, hepatocyte nuclear factor; PDX1, pancreatic duodenum homeobox; PI3K, phosphatidylinositol 3-kinase; TFs, transcription factors.
(Glut4) to the cell surface membrane, independently of transcriptional regulation (reviewed in Refs. 176, 265). Together, the biological effects of the transduction cascades are Glut4 translocation to increase glucose uptake in adipose tissue and muscle, the activation of glycogen synthase and thus of glycogen synthesis, and the activation of the S6-kinase that will increase protein synthesis.

In addition to or as a result of these signaling cascades in target cells, the expression of more than 150 genes expressed in various tissues is transcriptionally regulated by insulin. The diversity of mechanisms of the insulin-mediated transcriptional regulation is extremely wide, as indicated by the variety of promoter sequences that are responsible for insulin-mediated action. A classification of the insulin response sequences or insulin response elements (IRS/IRE) into seven groups has been proposed (212). Whereas there is still much to learn about the factors that are modified by insulin and bind to these elements, recent results have identified sterol response element binding protein 1c (SREBP-1c) as a major contributor. SREBPs are transcription factors of the helix-loop-helix family highly expressed in the liver. Their three forms, SREBP-1a, SREBP-1c, and SREBP-2, were first explored for their role in cholesterol and lipid homeostasis (see Appendix G). Interestingly, SREBP-1c expression in the liver is upregulated by insulin independently of glucose levels.

**Fig. 3.** Insulin and glucose action on the regulation of gene expression. Four main outcomes of the interaction of insulin with the membrane insulin receptor (IR) receptor and insulin receptor substrate 1 and 2 (IRS1/IRS2) are represented. The important reduction of intracellular levels of cAMP counteracts the action of glucagon (see also Fig. 5). The RAS/MAPK pathway leads to the activation of genes, which are mainly involved in cell growth (not reviewed herein). The activation of the phosphatidylinositol 3-kinase (PI3K) mediates most of the action of insulin on intermediary metabolism, via activation of Akt and activation of the SREBP-1c gene expression. PI3K-dependent activation of SREBP-1c is presently considered to be a major event for insulin-mediated gene induction. In contrast, Akt/PKB activation inhibits the activity of the transcription factors FOXO. This insulin-mediated inhibition of FOXO mainly results in target gene repression. Other consequences of Akt activation are also indicated but are not discussed in the present review, due to the fact that little is known about their action at the transcriptional level. The mechanism of insulin action on Glut4 expression remains unknown (dotted line), while Cbl is involved in the insulin-mediated increase of Glut4 translocation in muscle and adipose tissue. The intricate roles of glucose and insulin as regulators of gene expression are shown on the right of the scheme and comprise three aspects: the insulin-mediated translocation of the glucose transporter Glut4 as it occurs in the adipose tissue and muscle cells but not in the liver, while expression of Glut4 in the liver is dependent on the transcription factor FOXA3 (HNF3γ); the increased glucokinase (GK) expression in the liver; the independent activation of common target genes by both insulin via SREBP-1c and by glucose via carbohydrate response element binding protein/carbohydrate response factor (ChREBP/ChoRF). Xylose-5-phosphate (xylose 5P), glucose-6-phosphate (G6P), and hexosamine are glucose metabolites indicated as possible signal molecules directly responsible for the transcriptional response of the cell to glucose. For more details, see section iiB.
This regulation occurs at the transcriptional level, and detailed functional analyses of the SREBP-1c promoter revealed a complex interplay of transcription factors (24, 33). Whereas SREBP itself and nuclear factor Y act to maintain the basal level of SREBP-1c expression, the LXR (see Appendix C), a nuclear receptor activated by oxidized derivatives of cholesterol, plays a crucial role in its insulin-mediated increased expression. The proposed mechanism by which insulin-increased LXR activity would involve the insulin-dependent production of a ligand for LXR (33) is yet to be demonstrated. In addition to this transcriptional up-regulation of SREBP-1c expression, insulin triggers the proteolytic cleavage of SREBP-1c in a PI3K-dependent manner to produce the mature active form of this transcription factor (104).

In turn, SREBP-1c acts as an important mediator of insulin action, at least with respect to the transcriptional regulation of glycolytic and lipogenic genes in the liver. Overexpression of a dominant negative form of SREBP-1c counteracts insulin-mediated induction of the expression of liver pyruvate kinase (L-PK), spot 14 (S14), and fatty acid synthase (FAS), three canonical genes with respect to glucose*insulin2 responsiveness (see below and Ref. 73). The expression of glucokinase (GK) in the liver is also regulated by insulin, independently of extracellular glucose levels. GK phosphorylates glucose in glucose-6-phosphate (G6P), a first reaction required for any further intracellular metabolism of glucose. This regulation requires an intact PI3K pathway (124) and might also occur via SREBP-1c (73, 145). Less is known of insulin-mediated activation of SREBP-1c in the adipose tissue. The overlap in the gene expression profile of 3T3-L1 adipocytes subjected to insulin treatment with that of cells overexpressing the mature form or a dominant negative form of SREBP-1c strengthened the notion of a correlation between insulin-induced gene expression and SREBP-1c activity. It also revealed that, in these cells, the transcription factor CAAT/enhancer binding protein β (C/EBPβ) is responsive to insulin stimulation of SREBP-1c (169a).

Insulin also negatively regulates transcription, particularly that of genes involved in hepatic glucose production, such as those encoding IRS2, phosphoenolpyruvate carboxykinase (PEPCK), insulin growth factor binding protein-1 (IGFBP-1), and glucose-6-phosphatase (G6Pase) (for review, see Ref. 213). A particular sequence element, often contained in a broader insulin response unit, was identified in the promoter region of these genes as a mediator of negative regulation by insulin. Several transcription factors, such as members of the C/EBP, HNF-3/FOXA, and FOXO families, can bind to this element. Of particular interest are the FOXOs, represented by three members, FOXO1, FOXO3a, and FOXO4 (previously called FKHR, FKHLR, and AFX, respectively), which are phosphorylated by Akt-1 upon insulin-mediated activation of the PI3K pathway (reviewed in Ref. 299).Phosphorylated FOXO has a high affinity for protein 14-3-3, which relocates FOXO from the nucleus to the cytosol. In addition, insulin enhances ubiquitination of phosphorylated FOXO and its further degradation (193). Thus, in a simple mechanistic model, insulin would mediate repression via removal from the nucleus and accelerated degradation of the positive transcriptional regulator FOXO. Whereas the correlation between the activity levels of FOXO, HNF3, and C/EBP in mammals and gene repression by insulin remains unclear, their complex intertwined functions are illustrated in insulin-mediated PDX1 regulation. In the pancreatic β-cells, nuclear FOXO1 acts as a repressor of the positive activity of HNF3β on the PDX1 promoter, while insulin signaling relieves this repression by excluding FOXO1 from the nucleus (149).

It is interesting to note that the insulin signaling pathway going through PI3K and Akt activation is also present in Drosophila and in Caenorhabditis elegans. Indeed, the identification of FOXO (as DAF-16) and its involvement in insulin signaling was first described in C. elegans, triggering its characterization in mammals. A unique homolog of FOXO, dFOXO, has now been reported in Drosophila (134, 235) where insulin plays a crucial role in cellular growth. As seen in mammals and C. elegans, Drosophila Akt (dAkt) sequesters dFOXO in the cytoplasm when the insulin pathway is active. This results in an inhibition of dFOXO target gene expression, including that of the insulin receptor gene itself.

In summary, the general pathways followed by insulin to trigger many changes in gene expression are beginning to be understood. However, a lot more needs to be done to decipher the molecular mechanisms of this transcriptional regulation. The initial signal is at the cell membrane, and all subsequent events occur via phosphorylation cascades that mainly go through the PI3K pathway, but also possibly via the phosphorylation of the Cbl protooncogene. Thus it is possible that most of the insulin action on gene expression results from posttranslational modifications of various transcription factors, a process that would account for the pleiotropic effects of this hormone.

3. Glucose*insulin regulated transcription of genes involved in glucose metabolism

As mentioned above, glucose, independently of insulin, can regulate the expression of genes involved in carbohydrate metabolism. Upon entry into the cells, glucose is phosphorylated to G6P by GK in hepatocytes and by

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2 The experimental difficulty in discriminating between insulin and glucose effects in some of the gene transcriptional responses is reflected in the present review by the usage of the associated words glucose*insulin.
hexokinase in all other cells. This step is required for glucose to either undergo glycolysis, be used in the glycogen synthesis pathway, or enter the pentose phosphate pathway. This first metabolic transformation of glucose is also required for generating the signal that acts in transcriptional regulation. Some reports suggest that G6P itself might be the signaling molecule. Alternatively, other metabolites such as xylitol produced by the pentose phosphate pathway or intermediates of the hexosamine biosynthetic pathway might also act in tissue-specific regulations (reviewed in Ref. 305).

The analysis of glucose signaling is, however, often difficult to dissociate from insulin signaling (see Fig. 3). First, the entry of glucose into muscle and adipose tissue cells, which are two main insulin target organs, operates through the translocation of the Glut4 transporter via an insulin-mediated transduction signal. In contrast, the expression of the glucose transporter Glut2 in the liver and pancreas, Glut3 in the brain, and the widely distributed Glut1 is insulin independent, and their translocation is constitutive. Second, in the liver and to a lesser extent in the pancreas, the initial metabolic modification of glucose into G6P by GK is required for transcriptional regulation by glucose and is strongly dependent on insulin. Thus the actions of glucose and insulin are often interdependent and, in this review, we refer to this ambiguity by the use of the associated words glucose*insulin when relevant.

Three genes have been important tools for the analysis of the ability of glucose to direct transcriptional regulation; they encode the L-PK (acting on the glycolytic pathway from glucose to pyruvate), S14 (associated to lipogenesis but with an unclear function), and FAS (a key enzyme in lipogenesis). Analyses of the promoter region of these genes have identified response elements called carbohydrate response elements (ChoRE) or glucose response elements (GIRE), which have similarities. The main common feature is the presence of at least one E-box. The GIRE/ChoRE of L-PK and S14 comprises two E-boxes in tandem, in addition to a binding site for an ancillary factor which is HNF4 in the case of L-PK (57, 175, 276). The glucose*insulin responsiveness of FAS also requires a complex array of promoter elements, a complexity that has generated some controversy. Three glucose*insulin response sites are now proposed. A region between −150/+50 centered around an E-box was the first proposed IRE/GIRE/ChoRE. This sequence efficiently binds SREBP-1c and mainly represents an insulin responsive element. A second element is located at −332, but its role in vivo is unclear. Finally, a third far-upstream element located around −7 kb closely resembles the GIRE/ChoRE found in L-PK and S14 (152, 201, 261).

E-boxes are binding motifs for helix-loop-helix transcription factors and can bind the abundant and ubiquitous upstream stimulatory factor (USP), whose two forms USP1 and USP2 can heterodimerize. However, various studies in vivo (with knock-out animals) and in vitro (electromobility shift assays) aimed at elucidating the link between the ability of USFs to bind to the E-box and glucose responsiveness have failed to prove the concept accurate. An alternate hypothesis involves the negative transcription factor COUP-TFI, which is also able to bind to the L-PK GIRE/ChoRE. The equilibrium resulting from the competition between USF1/USF2 and COUP-TFI would then create the dynamic modulation and glucose-dependent regulation (305). A new factor, initially cloned as WBSCR14 (Williams-Beuren syndrome deleted DNA region, Ref. 51), exhibits a GIRE/ChoRE binding activity that could account for glucose responsiveness. Based on its interaction with the L-PK ChoRE, this helix-loop-helix factor has been renamed the ChoRE binding protein (ChREBP) (329). ChREBP is mainly expressed in liver, kidney, and adipose tissue. A mouse line null mutant for ChREBP provided evidence for a direct and dominant role of ChREBP in the glucose-mediated upregulation of LPK, ACC, and FAS gene transcription, coordinating synthesis of fatty acids and triglycerides in vivo in response to high levels of glucose (118, 123). Under basal conditions, ChREBP is phosphorylated by protein kinase A (PKA) and remains cytosolic. The glucose-dependent activation of ChREBP is a two-step process, with a first dephosphorylation at serine-196 which triggers its nuclear translocation, and a second dephosphorylation in the nucleus at serine-568 and threonine-666, which allows it to bind to DNA. This activation requires GK expression, as demonstrated in GK knock-out mice (53), and the glucose metabolite xylulose 5-phosphate from the pentose phosphate pathway is the proposed functional link between high glucose and ChREBP activation (136). Indeed, xylulose 5-phosphate can activate protein phosphatase 2, triggering ChREBP dephosphorylation in the cytosol as well as in the nucleus (reviewed in Ref. 52). Finally, the transcriptional activity of ChREBP requires its heterodimerization with the bHLH/LZ factor Max-like protein X (Mix) (183), which would allow the complex to specifically target E-box binding sites in glucose-responsive gene promoters.

Thus we have gained extremely interesting new understanding of glucose*insulin regulation of gene expression in the last few years. Present works are now aimed in part at understanding when and how these factors respond to an altered metabolic environment, such as in obesity, insulin resistance, or type 2 diabetes.

C. Transcriptional Regulation of Metabolism by Low Glucose Levels

The prevalence of diabetes, i.e., a deregulation characterized by high glucose levels due to impaired insulin signaling, demonstrates the preeminent role of insulin over the action of all counteracting hormones. This fact possibly explains why less is known about the hormonal
regulation of genes in situations of low glucose availability. In periods of starvation, even between regularly spaced meals, the liver and to a lesser extent the kidney are responsible for the glucose production required for a sufficient supply to the brain. The small intestine also provides glucose upon prolonged starvation. Hormonal controls of this adaptation associate increased glucocorticoids, decreased insulin levels, and, importantly, glucagon secretion by the pancreas in response to low glucose.

Glucagon is processed from proglucagon in the α-cells of the pancreatic islets and is secreted in response to low blood glucose levels. In the liver, glucagon interacts with a membrane receptor coupled to GTP-binding proteins, inducing a rise in intracellular cAMP (Fig. 4), which in turn activates PKA. By this mechanism, glucagon counteracts some of the glucose-insulin-mediated responses. For example, increasing cAMP levels in primary hepatocytes decrease SREBP-1c expression via a mechanism requiring de novo protein synthesis (75, 279). Also, the PKA-dependent phosphorylation of ChREBP sequesters it in the cytosol and inhibits its lipogenic activity (140). Modulation of cAMP levels is the major mechanism by which the liver adjusts glycogenolysis and gluconeogenesis, which produce and release hepatic glucose in the blood. Several transcription factors such as the cAMP response element binding protein (CREB), the cAMP response element modulator (CREM), and the activation transcription factor-1 (ATF-1) are positively activated upon phosphorylation. All three belong to the bZIP family of transcription factors and share a conserved phosphorylation box and glutamine-rich transactivation domain (reviewed in Ref. 48). Their PKA-dependent phosphorylation allows the recruitment of the CREB-binding protein (CBP) coactivator, which contributes to the transcriptional activity of the DNA-bound complexes. CREB is a ubiquitously expressed transcription factor that induces the expression of key genes involved in the gluconeogenesis pathway, such as those encoding PEPCK, G6Pase, and pyruvate carboxylase. Accordingly, CREB binding sites have been identified in the PEPCK and G6Pase promoters (81, 112, 223), but not in that of pyruvate carboxylase. Additional mechanisms for cAMP-mediated transcriptional response are required to explain the full range of responsive genes and the specificity of the response in gluconeogenic tissues, i.e., liver, kidney, and small intestine. For example, the gene for the cofactor PGC1 is strongly activated by CREB in the liver (106). As a cofactor, PGC1 was shown to increase the transcriptional activity mediated by both HNF4α and the glucocorticoid receptor bound to the PEPCK promoter (334) (Fig. 4). The occurrence of such an indirect mechanism via PGC1 for the positive glucagon-dependent induction of pyruvate carboxylase is yet to be examined. HNF4α together with C/EBPα and C/EBPβ also constitutively binds to the G6Pase promoter. In this context, glucagon further induces gene transcription via CREB binding to its cognate site and further recruitment of CBP (81). CBP might be of prime importance for cessation of gluconeogenesis upon feeding, as it is also a target of insulin-dependent phosphorylation at Ser-436 (339). This phosphorylation impairs CBP recruitment to CREB, thereby inhibiting CREB target genes, as demonstrated for PGC1 (341).

**Fig. 4.** Transcriptional adaptation of the metabolism in liver cells upon low levels of glucose. Most of the responses to low glucose are mediated by the lack of insulin, associated with increased levels of glucagon and, to some extent, by the stimulation of adrenergic receptors. The subsequent increase in cAMP levels triggers the phosphorylation of the transcription factor cAMP response element binding protein (CREB), responsible for increased expression of gluconeogenic enzyme genes. In addition, C/EBPβ participates in increasing cAMP levels, whereas C/EBPα independently activates genes involved in gluconeogenesis. The inset shows the combined interaction of transcription factors and the coactivator PGC-1 involved in the transcriptional regulation of the phosphoenolpyruvate carboxykinase (PEPCK) gene, which plays a crucial role in gluconeogenesis. For more details, see section II C.
The importance of C/EBPα (see Appendix H and Fig. 4) in the transcriptional control of gluconeogenesis has been revealed by the phenotype of C/EBPα null mice (312). The major metabolic disturbance seen in these mice is a lethal neonatal hypoglycemia. This hypoglycemia is due to the combination of two deficiencies; first, reduced glucose homeostasis via the transcription of key genes. The Aminotransferase deficiency in these mice results in a decreased expression of SREBP-1c (72), reduced transcriptional activation of ChREBP, in parallel with a decreased expression of AMPK in the liver decreased the refeeding-increased. In the liver, carbon skeletons of the gluconeogenic AA (e.g., alanine and serine in the liver and glutamine in the kidney and the gut) are catabolized into pyruvate or into one of the metabolites of the citric cycle, which can then be converted into glucose. In contrast, ketogenic AA (e.g., leucine, isoleucine, phenylalanine) degraded into acetyl-CoA or acetoacetate are precursors of the ketone bodies that can be used as an alternate source of energy, in particular by brain cells. The AA carbon skeleton used as the energy source results from an AA deamination process, which leads to the cytosolic accumulation of toxic free ammonia (NH₄⁺). For further processing, NH₄⁺ is transported as glutamate and/or glutamine to the liver and kidneys (see Fig. 5), where ammonia is freed and processed in the urea cycle. Urea is a diffusible molecule, which is then excreted in the urine.

In conclusion, this short presentation highlights some of the best-characterized features of the regulation of glucose homeostasis via the transcription of key genes. However, this summary cannot take into account the specificity of these regulations within each tissue, which is essential for the homeostasis at the level of the whole organism. An effort to integrate some of the pathways described above into the global balance of metabolic regulations will therefore be presented in section VI.

III. TRANSCRIPTIONAL CONTROL OF AMINO ACID AND PROTEIN METABOLISM

A. Introduction

In addition to being substrates for the synthesis of specialized products such as neurotransmitters, hemes, nucleotides, and polyamines, amino acids (AA) also play an important role in energy supply. Among the 20 AA involved in protein synthesis, around half can be synthesized de novo and 9 essential AA must come from the diet. However, during growth or other situations of high-energy expenditure, some of the nonessential AA such as arginine, whose synthesis is energy demanding, should rather be provided by food (reviewed in Ref. 241).

In western countries, under normal nutritional and physiological conditions, proteins/AA are often ingested in excess and are neither stored nor excreted but catabolized and used as an important source of energy that fuels the production of glucose and fatty acids. With a western diet, degradation of AA provides 10–15% of the total energy requirement of the organism. During starvation, the use of AA degradation for energy supply is increased. In the liver, carbon skeletons of the gluconeogenic AA (e.g., alanine and serine in the liver and glutamine in the kidney and the gut) are catabolized into pyruvate or into one of the metabolites of the citric cycle, which can then be converted into glucose. In contrast, ketogenic AA (e.g., leucine, isoleucine, phenylalanine) degraded into acetyl-CoA or acetoacetate are precursors of the ketone bodies that can be used as an alternate source of energy, in particular by brain cells. The AA carbon skeleton used as the energy source results from an AA deamination process, which leads to the cytosolic accumulation of toxic free ammonia (NH₄⁺). For further processing, NH₄⁺ is transported as glutamate and/or glutamine to the liver and kidneys (see Fig. 5), where ammonia is freed and processed in the urea cycle. Urea is a diffusible molecule, which is then excreted in the urine.

Regulation of these pathways, i.e., regulation of AA synthesis and degradation, depends heavily on substrate availability and allosteric mechanisms, which are mainly used in the short-term regulation of energy homeostasis. However, transcriptional regulation of genes for key enzymes is important for long-term adaptation to specific diets. Here, we will emphasize three aspects of AA metabolism for which transcriptional regulation has been shown to play a major role. We will first describe the regulation of glutamine homeostasis and of the urea cycle, two key determinants of the maintenance of the nitrogen balance under different physiological conditions. Then the specific regulatory pathways triggered by AA deprivation, which might be encountered in cases of general malnutrition or deficiency in any of the essential AA, will be discussed.
organism. In addition to being an important energetic and metabolic substrate, it provides sufficient amino groups for AA and nucleotide synthesis. It is also the main transporter of ammonia in the blood towards the liver, and glutamine generation in the brain is crucial for avoiding highly neurotoxic hyperammonemia. Thus it has a dual importance: during growth for anabolism and under catabolic conditions for limiting ammonia levels in peripheral tissues and blood. Glutamine is required for normal growth and proliferation of cells, particularly of enterocytes. Additionally, glutamine requirements are particularly high at times of severe sepsis, when proliferation of the immune cells is necessary (reviewed in Ref. 208). Conversely, depletion of glutamine due to a high cellular metabolic rate, often associated with high catabolism of AA, also occurs in cancer patients.

Glutamine formation via the ATP-dependent glutamine synthase (GS) occurs in most tissues (Fig. 5), with, in adults, the greatest activity in skeletal muscle, lung, brain, and adipose tissue. Whereas the GS turnover is increased by a high concentration of the end-product glutamine, regulation of GS activity also occurs at the transcriptional level, as best characterized by the responsiveness of GS to glucocorticoids in the lungs and muscles. Two broad regions mediating this response have been identified in the upstream promoter and in the first intron of GS. Detailed studies of GS expression in chicken brains showed that glucocorticoids act by relieving the repression mediated by a silencer element located upstream of the glucocorticoid receptor binding site (5). This glucocorticoid-mediated induction of GS occurs particularly in conditions of trauma or high catabolic rate and results in increased glutamine synthesis at the expense of the AA that purvey the amino groups. It is thus believed that the action of glucocorticoids on glutamine metabolism is responsible for some of the deleterious effects of corticotherapy on muscle physiology, such as muscle atrophy. Transcriptional regulation of GS has also been studied in adipocyte differentiation, where the high expression of GS is controlled through a C/EBP responsive element in the distal 5′-flanking promoter region (96). However, little is known about the physiological significance of GS activity in the adipose tissue.

Conversely, glutamine homeostasis for the whole organism is also largely controlled in the liver, and to a lower extent in the kidneys, by the glutaminase activity which participates in the reverse pathway, allowing the disposal of $\text{NH}_4^+$ (see Fig. 5) and providing gluconeogenic substrate. The hepatic-type glutaminase expression is increased during starvation, diabetes, and protein-rich diets, when AA degradation is increased. At the molecular level, the transcription of hepatic-type glutaminase during fasting (e.g., in conditions of low insulin/high glucagon) is highly activated by cAMP, as well as by glucocorticoids via a promoter element that has been identified (42). In contrast, the kidney-type glutaminase is mainly responsive to metabolic acidosis, triggered by prolonged starvation or uncontrolled diabetes. The catabolism of glutamine in metabolic acidosis has a dual role, both facets contributing to the restoration of metabolic homeostasis. First, the generation of $\text{NH}_4^+$ from the conversion of glutamine to glutamate and $\alpha$-ketoglutarate facilitates the excretion of acids; second, further catabolism of

![Figure 5](http://physrev.physiology.org)
\( \alpha \)-ketoglutarate is linked to increased gluconeogenesis, most notably via the increased activity of PEPCK. Whereas the increased expression of glutaminase is mainly the result of mRNA stabilization via an element located in the 3'-UTR region of its mRNA (160, reviewed in Ref. 156), the expression and response to acidosis of PEPCK in the kidney depends on an HNF1 binding site present in its promoter (28).

Thus, whereas glutamine homeostasis seems to be one physiologically important knot of AA metabolism, most of the molecular mechanisms governing the regulation of these enzymatic activities remain to be analyzed. As glutamine synthesis is required, particularly in conditions of cell proliferation, decreasing glutamine intake has been proposed to control cell growth in cancer. It now appears that depletion of glutamine in cancer patients contributes to a global degradation of their health status. Consequently, a supplementation in glutamine is now on trial in these patients, as well as during severe sepsis for reinforcing the immune system (195).

**C. Transcriptional Regulation of Urea and Ammonia Homeostasis**

Urea production in the liver, via the activity of the five enzymes of the urea cycle, carbamoyl phosphate synthetase 1 (CPS-1), ornithine transcarbamoylase (OTC), argininosuccinate synthase (ASS), argininosuccinate lyase (ASL), and arginase, is the main pathway for ammonia detoxification (Fig. 6). There are two main sources of ammonia production: the diet with its protein content, and endogenous protein degradation which occurs when there is a relatively low energy supply. The regulations occurring on a short time scale are mainly allosteric reactions, such as CPS-1 activation in the presence of N-acetylglutamate. However, there is a coordinated regulation of the expression of these five enzymes in response to dietary changes or to metabolic challenges during development and in adulthood. Hints regarding the molecular mechanism of this transcriptional regulation and the nature of the coordinating factor are starting to emerge.

Some important transcription factors for the regulation of ureagenesis have been revealed by the phenotype of several mutant mice. Liver-specific deletion of the nuclear receptor HNF4\( \alpha \) results in hyperammonemia and hypoureemia due to a dramatic reduction of the expression of OTC, consistent with the presence of two functional HNF4 response elements in the promoter of this gene. In contrast, the expression of the four other genes of the urea cycle remains unchanged or is only slightly increased (122). Thus the loss of the HNF4\( \alpha \) function in ammonia detoxification might be the cause of the premature death of the mice carrying a liver-specific disruption of HNF4\( \alpha \).

PPAR\( \alpha \), on the other hand, is a negative regulator of the urea cycle, consistent with the PPAR\( \alpha \)-mediated down-regulation of CPS-1, OTC, ASS, and ASL in mice treated with a fibrate (141). While the mechanism by which PPAR\( \alpha \) exerts this coordinated down-regulation of urea enzyme gene expression is not yet known, it has been proposed that HNF4\( \alpha \) positively regulates PPAR\( \alpha \) expression via a binding site for HNF4\( \alpha \) in the promoter of this gene (231). Accordingly, the relatively low levels of PPAR\( \alpha \) in HNF4\( \alpha \) null mice could thus explain the normal or higher levels of CPS-1, OTC, and ASL enzymes in these mice.

Severe hyperammonemia has been observed in mice carrying a liver-specific deletion of C/EBP\( \alpha \). This phenotype correlates with a dramatic fall in CPS-1 and arginase expression, together with a disturbed intrahepatic lobular distribution of OTC expression (147). Glucocorticoids,
which have a strong impact on protein degradation in muscle, are efficient inducers of the urea cycle in the liver, allowing excretion in the form of urea of the excess ammonia produced by protein degradation. Intriguingly, the response of arginase to glucocorticoids via binding of the glucocorticoid receptor to its specific response element requires C/EBPβ expression and the integrity of the C/EBP binding sites in the 5′-flanking region of the gene (89). The same C/EBPβ dependency of the glucocorticoid response is also true of CPS-1 (146). The CPS-1 promoter contains a complex regulatory module composed of multiple sites for glucocorticoid receptor, HNF3 and C/EBP family members, as well as for unknown factors that control its specific pattern of expression and regulation. The glucocorticoid response unit itself combines a glucocorticoid response element with sites for HNF3β and C/EBP (41). Whereas these in vitro studies performed in primary hepatocytes in culture attributed a preferential role to C/EBP (41), Whereas these in vitro studies performed in primary hepatocytes in culture attributed a preferential role to C/EBPβ, C/EBPβ null mice did not present any perturbation of ureagenesis (40, 147), and it seems reasonable to propose that in vivo C/EBPα is the functional partner of the glucocorticoid receptor. With respect to HNF3β, an embryonic lethality of HNF3β null mutation preempts the analysis of the role of this factor in vivo, whereas HNF3α and HNF3γ null mice do not exhibit any perturbation of ammonia metabolism (40, 147), and it seems reasonable to propose that in vivo C/EBPα is the functional partner of the glucocorticoid receptor. With respect to HNF3β, an embryonic lethality of HNF3β null mutation preempts the analysis of the role of this factor in vivo, whereas HNF3α and HNF3γ null mice do not exhibit any perturbation of ammonia metabolism.

The above discussion on the search for a coordinator of the ureagenesis pathway underscores the complex interplay established by transcription factors and pinpoints a general mode of homeostasis regulation, in which equilibrium is obtained via the simultaneous control of opposite pathways. One example here is the glucocorticoids that activate the urea cycle via C/EBPs, but at the same time increase the levels of PPARα, which is an inhibitor of the same cycle. Another example is the regulation of PPARα expression by HNF4α, which contributes to keeping a balance between activation and inhibition of urea cycle activity. A third example is that of GS and glutaminase, which have opposite activities but are both up-regulated in the liver by glucocorticoids (see sect. mB). While somewhat counterintuitive, such a mode of regulation should lead to a fine-tuning that limits the oscillations of feed-back and feed-forward regulations.

D. Amino Acid Deprivation and Induction of Gene Expression

1. The search for an amino acid response element

Two genes have been extensively explored for their ability to respond to amino acid deprivation. C/EBP homologous protein (CHOP) is induced by various stresses. One of these is the unfolded protein response pathway triggered by the accumulation of unfolded proteins in the endoplasmic reticulum (ER), which activates chaperones resident in the ER. CHOP is related to the C/EBP family of nuclear factors with which it forms heterodimers. Interestingly, global AA deprivation or starvation in individual AA induces CHOP expression independently of the ER stress pathway. A short promoter sequence or AA response element (AARE) conveys the AA sensitivity of CHOP. This sequence combines the features of a C/EBP consensus element and a cAMP response element (22). The second gene that is used as a model to explore the transcriptional response to amino acid deprivation is the gene encoding asparagine synthetase (AS). AS catalyzes the glutamine- and ATP-dependent conversion of aspartate to asparagine. AS mRNA accumulates in mammalian cell cultures in response to asparagine starvation. Amazingly, deprivation of a wide range of individual AA also induces this accumulation, therefore suggesting that AS responds to a signal reflecting AA deprivation more broadly. Two discrete response elements, called nutrient sensing response elements NSRE-1 and NSRE-2, were found in a short promoter region of human AS. Both of them are required for AS activation and form a nutrient-sensing response unit that mediates not only the response to AA deprivation, but also the response to glucose deprivation (284 and references therein).

The identity of the factors that bind to these elements and are thus responsible for the response to nutrient deprivation remains disputed. They belong to the activating transcription factor (ATF)/CREB family, which includes members sharing a basic leucine zipper motif and a consensus ATF/CRE DNA binding site “TGACGTCA” (reviewed in Ref. 97). The CRE-binding protein 1 (CREBP1 or ATF2) binds to the C/EBP-ATF composite site forming the AARE of CHOP, either as a homodimer or as a heterodimer with an unknown dimerization partner (22). ATF4, but not ATF2, binds as a complex with C/EBP to NSRE-1 of CHOP and is required for the response of AS to nutrient deprivation (283). ATF4 itself is transcriptionally and posttranscriptionally regulated by both AA and glucose deprivation (100, 283), which suggests that ATF4 is an important transcriptional mediator of the nutrient-sensing response. However, the response of AS to glucose deprivation is mediated via the unfolded protein response pathway and may be independent of the response to AA deprivation.

Taken together, the differences between and similarities in the regulation of these two genes underline the existence of at least two related but independent pathways, via ATF2 and ATF4, respectively, which control gene expression in response to AA deprivation (21).

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2. TOR as a master switch for catabolism versus anabolism

Studies carried out to decipher the yeast response to AA deprivation have identified the target of rapamycin (TOR) proteins as master switches for protein/AA catabolism versus anabolism (125). TOR belongs to the PI3K-related kinase family and appears to function as a nutrient-sensing check-point by controlling many aspects of mRNA translation. Inhibition of TOR proteins by rapamycin in yeast mimics nutrient deprivation. In fact, TOR modulates the transcription of genes involved in AA biosynthesis and the activity of permeases that allow AA transport into the cells. It also inhibits autophagy in yeast and in mammalian cells, a process that degrades cytoplasmic proteins and organelles for scavenging AA when nutrient levels are low. When a sufficient amount of nutrients is sensed, TOR proteins act as a permissive signal for growth and protein synthesis. A single mammalian homologous TOR protein, alternatively called mTOR, FRAP, or RAFT1, has been cloned in various species with a remarkable level of AA identity, suggesting well-conserved functions of the TOR-dependent regulatory pathways. In mammalian cells, as well as in Drosophila, TOR is not only sensitive to the presence of sufficient levels of AA, but also integrates energy and growth signals through the AMPK and PI3K pathways, respectively. Increased Akt activity, via insulin signaling for example, would trigger the phosphorylation of the tuberous sclerosis complex (TSC) relieving the constitutive inhibition that TSC exerts on mTOR activity (reviewed in Ref. 125). In addition to this regulatory interaction, recent evidence has demonstrated that rapamycin-sensitive mTOR kinase activity requires the direct interaction of the small GTPase RheB-GTP with the TOR-containing complex TORC1 (177).

The best-known molecular mechanism of TOR action is a posttranscriptional action on the phosphorylation status of the initiation and elongation factors involved in translational control (reviewed in Ref. 240). However, it also regulates the abundance of the components of the translation machinery both at the transcriptional and translational levels. This results, for example, in controlling the translational events that regulate mammalian cell size (71). At the transcriptional level, TOR modulates the expression of numerous enzymes involved in multiple metabolic pathways. In yeast, this transcriptional control is mainly exerted by the cytoplasmic sequestration of transcription factors. TOR controls ribosomal protein (RP) gene transcription by maintaining the corepressor CRF1 in the cytoplasm, thereby allowing the forkhead-like transcription factor FH1 and its coactivator IFH1 to efficiently activate RP gene transcription. Upon TOR inhibition, phosphorylated CRF1 rapidly translocates to the nucleus inhibiting RP transcription (131, 190).

IV. TRANSCRIPTIONAL CONTROL OF LIPID METABOLISM

Lipids occur in three major classes of molecules in multicellular organisms: 1) fatty acids, as a source of energy, mainly stored in form of triglycerides in fat tissues; 2) phospholipids and cholesterol, which are major structural components of the cellular membranes; and 3) lipid-derived small bioactive molecules, such as the steroid hormones, the arachidonic acid derivatives (prostaglandins and leukotrienes), and other intracellular messengers that are present in small quantities and have crucial roles in signaling. In this section, we concentrate on the transcriptional mechanisms that govern the synthesis, storage, release, uptake, and oxidation of fatty acids as the main participants in energy homeostasis and on the intricate control of cholesterol metabolism.

A. Transcriptional Control of Fatty Acid Metabolism

Transcriptional regulation of the genes involved in fatty acid metabolism is presently considered as the major long-term regulatory mechanism controlling lipid homeostasis. It is executed by a variety of transcription factors among which the SREBPs, the C/EBPs, and members of the nuclear receptor family are particularly active agents.

1. Fatty acid synthesis and storage: control by an intricate array of transcription factors

Adipogenesis is a developmental process by which cells become determined and differentiate to fulfill adipose tissue specific functions, among which are fat stor-
age and production of adipose hormones (also called adipokines). Lipogenesis results in cellular lipid accumulation, via the uptake of lipogenic substrate from the diet, endogenous fatty acid synthesis, and fatty acid storage as triglycerides. It should be noted that lipolysis, which releases fatty acids into the blood, is the reciprocal important function of the white adipose tissue (WAT). However, little is known on the transcriptional regulation of lipolysis that will be discussed in the context of adaptation to fasting (see sect. VII).

A) SREBP-1, A MAJOR TRANSCRIPTION FACTOR INVOLVED IN FATTY ACID SYNTHESIS. A brief overview of the SREBPs is shown in Appendix G. The maturation process of SREBPs via a cholesterol sensing mechanism initially emphasized their role in cholesterol homeostasis (see Fig. 7). However, whereas the main role of SREBP-2 is indeed geared toward cholesterol metabolism, SREBP-1a and -1c have been clearly associated with both cholesterol and fatty acid homeostasis. SREBP-1a appears to be constitutively expressed at quite low levels and mainly in the liver (116, 280). Overexpression of the nuclear form of SREBP-1a in transgenic mice leads to a massive engorgement of the liver with triglycerides and to a lesser extent with cholesterol (277). SREBP-1c, whose lower transcriptional activity has been related to its shorter NH₂-terminal tail, acts more specifically on genes involved in fatty acid synthesis. It was initially identified in the WAT and named adipocyte determination and differentiation factor 1 (ADD-1) (297). It is highly expressed in the liver and the WAT and is sensitive to multiple regulatory signals. Overexpression of its nuclear form strongly increases the triglyceride content of the liver, with no parallel accumulation of cholesterol. While a null-mutation of SREBP-1 (deleting both SREBP-1a and SREBP-1c) provokes an embryonic lethality that is partially penetrant, analyses of surviving SREBP-1 −/− mice show reduced synthesis of fatty acids, whereas the cholesterol synthesis pathway is increased due to a compensatory expression of SREBP-2 (see Ref. 114 for review).

Although the posttranscriptional maturation of SREBPs is an important regulatory event, the transcriptional regulation of SREBP-1c expression parallels the activity of the transcription factor. As discussed in section II, one major metabolic signal that upregulates SREBP-1c is insulin, whereas glucagon represses it. Insulin, whose release is stimulated by high blood glucose levels, induces the production of fatty acids from glucose-derived pyruvate in the liver and the adipose tissue. Most of the lipogenic effects of insulin are dependent on the induced expression of SREBP-1c and the subsequent stimulation of the fatty acid synthesis pathway. SREBP-1c also positively regulates its own production in a feed-forward loop that ensures higher SREBP-1c production when the levels of the mature nuclear form are elevated (1). SREBP-1c expression is also stimulated by the LXR, via two LXR binding sites (LXRE) present in the SREBP-1c promoter. The primary role of LXR was thought to be in the control and protection of cells from cholesterol overload. Thus the reason why LXR would trigger a parallel increase in fatty acid synthesis via transcriptional regulation of SREBP-1c remains unclear. One hypothesis is that the blood transport and cell storage of excess cholesterol requires the formation of cholesteryl ester. This increases the demand for oleate production,
which is indeed stimulated upon overexpression of SREBP-1c (243, 272).

Conversely to these positive signals, high levels of polyunsaturated fatty acids (PUFA) repress SREBP-1c expression. Several mechanisms have been proposed for this downregulation. A functional analysis of the SREBP-1c promoter demonstrated a PUFA-dependent inhibition of LXR binding to the promoter (219, 336). PUFA may also accelerate the decay of SREBP-1c transcripts as demonstrated in rat hepatocytes (326) or may act on the process of SREBP-1c protein maturation, thereby decreasing SREBP-1c activity. Indeed, whereas the post-translational maturation of SREBP-2 depends tightly on the abundance of membrane cholesterol, the inhibition of the SREBP-1 cleavage and maturation seems to depend on the presence of PUFAs (reviewed in Ref. 63). However, little is known about the relative importance of this regulation. Further understanding the role of the PUFA-mediated repression of SREBP-1c is of importance as it may be the main mechanism by which dietary PUFAs inhibit lipogenesis.

As shown in Figure 8, the action of SREBP-1c in fatty acid synthesis is mediated by the activation of a set of target genes. The DNA elements in the promoter of SREBP-responsive genes are surprisingly diverse, and defining a consensus SRE is thus difficult. The intricate localization of an SRE with an E-box motif as seen in the FAS promoter does not seem to be the rule. In addition, SREBP-1 can bind to both an E-box motif or to a non-E-Box SRE (143). Nonetheless, the transcriptional activity of SREBPs often requires cooperation with other DNA binding transcription factors such as SP1, NF-Y, and CREB as well as with coactivators (reviewed in Ref. 63). Whereas further work is on-going to elucidate the molecular interactions involved in SREBP-mediated transcriptional activation, the evidence discussed above clearly emphasizes the role of SREBP-1c in increasing fatty acid synthesis in liver and in adipose tissue, particularly in response to insulin.

**B) PPARγ: A MAJOR REGULATOR OF FATTY ACID STORAGE AND ADIPOGENESIS.** Mature adipocytes are cells that sustain efficient lipogenesis and triglyceride synthesis and storage, but also release fatty acids and glycerol into the blood via lipolysis. The mature adipose tissue also secretes many endocrine signals whose importance and mechanism of action are presently intensively scrutinized in the context of whole body energy metabolism.

It is believed that the developing and mature adult adipose tissues constitute a life-long reservoir of preadipocytes that can be triggered to differentiate in adipocytes if the need for fat storage is signaled. The peroxisome proliferator activated receptor γ (PPARγ) has been clearly linked to the adipocyte differentiation program (reviewed in Ref. 258). At the protein level, there are two PPARγ isoforms, PPARγ1 and PPARγ2. PPARγ2 has 30 and 28 additional NH2-terminal amino acids in mice and humans, respectively. Both isoforms are produced from the same gene by alternative promoter usage and mRNA splicing. PPARγ1 is mainly expressed in adipose tissues but is also detected in the colon, spleen, retina, hematopoietic cells, liver, and skeletal muscle, whereas PPARγ2 expression is restricted to the adipose tissue (18, 66 and reviewed in Ref. 54). However, the two isoforms have identical ligand-binding properties and share the same target genes (257), for instance, those coding for the adipocyte fatty acid binding protein (aP2), lipoprotein lipase (LPL), acyl-CoA synthase (ACS), and fatty acid transport protein (FAT/CD36). PPARγ is a late marker of

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**FIG. 8.** Sterol response element binding protein (SREBP)-2 and SREBP-1c at the branching point of cholesterol and fatty acid metabolism. SREBP-1c is a transcription factor activating genes encoding enzymes involved in fatty acid synthesis and driving the formation of triglycerides and phospholipids (right). In contrast, SREBP-2 stimulates the transcription of genes encoding enzymes involved in cholesterol synthesis (left). These two pathways require the generation of NADPH, provided by the reactions shown in the middle of the figure. Both SREBP-1c and SREBP-2 positively regulate the three enzymes participating in these reactions, i.e., malic enzyme (ME), glucose-6-phosphate dehydrogenase (G6PDH), and 6-phosphogluconate deshydrogenase (PGDH). For more details, see section IV.
adipocyte differentiation, and its artificial expression is sufficient to force fibroblasts into the adipogenic program. Whereas PPARγ null mice are not viable, due to defects in placenta formation (9), the lack of adipocytes carrying the genotype PPARγ−/− in chimeric PPARγ+/−:PPARγ−/− mice has demonstrated the importance, in vivo, of PPARγ for adipogenesis (255).

The role of PPARγ in the transcriptional control of metabolism in mature adipose tissue is less clear. The overexpression of a dominant negative mutant of PPARγ in fully differentiated mature 3T3-L1 adipocytes results in increased lipolysis and decreased fatty acid uptake, concomitant with a decreased expression of known PPARγ target genes, including aP2, LPL, ACS, and CD36 (203). In vivo, synthetic PPARγ ligands, the glitazones, increase the number of adipocytes rather than modulate the mature adipocyte functions. The availability of full PPARγ antagonists should be an interesting tool for deciphering, in vivo, the role of PPARγ in adult tissues. However, the antagonists described so far either have a residual agonist activity or are hardly soluble and/or toxic (204, 211, 251). Recently, a new PPARγ antagonist has been tested in vivo, demonstrating that inhibition of PPARγ activity results in a decrease in white and brown adipose tissue fat deposits to a similar extent as that observed in PPARγ heterozygous mice. This reduction reflects a decrease in the average adipocyte volume (247), which could either be due to a lesser fat accumulation or to an increased lipolytic activity. The important role of PPARγ in the mature tissue is now confirmed by the phenotype of mice with an adipose tissue-specific deletion of PPARγ. As the expression of the CRE enzyme responsible for the gene deletion is under the activity of the aP2 promoter, the deletion occurs after adipogenesis has taken place. This deletion results in a severe reduction of the number of mature adipocyte, both in white and brown adipose tissues, whereas small and likely nascent adipocytes are appearing. Thus PPARγ seems to be essential for the survival of mature adipocytes (103, 121). The implication of PPARγ in the stimulation or inhibition of lipolysis has not yet been addressed.

Adipogenesis via PPARγ activation may occur in other tissues than the adipose tissue. A forced expression of PPARγ1 in hepatocytes induces the classic pattern of PPARγ-mediated gene activation and results in steatosis (338). These observations recapitulate the liver phenotype of murine models of diabetes and obesity that are marked by elevated levels of PPARγ in the liver. However, the normal expression of PPARγ in the liver is very low and, consistently, a liver specific deletion of PPARγ gives no phenotype unless this deletion is performed in a murine model of obesity such as ob/ob mice (192). Specific deletion of PPARγ in the skeletal muscle also results in a lack of phenotype with respect to growth and lipid distribution, whereas there is increased adiposity and development of insulin resistance upon exposure to a high-fat diet (107, 209) (see also sect. vi). This emphasizes the link between muscle metabolism and adipose tissue response, the nature of which remains to be elucidated. Accumulation of lipids via PPARγ might also occur in smooth muscle cells and/or in activated macrophages. At the same time, PPARγ has also been implicated in cholesterol export from macrophages, thereby counterbalancing the detrimental effects of lipid loading (reviewed in Ref. 162). However, the complex role of PPARγ in foam cell formation and atherosclerosis goes beyond the scope of the present review.

A puzzling observation made a decade ago was that glitazones, which were developed for treatment of insulin resistance, are PPARγ-selective ligands. The link between the promotion of adipocyte differentiation and lipid storage by PPARγ and the antidiabetic effects of these compounds is not fully understood and several hypotheses, based on a wealth of experiments, have been proposed. These are discussed in section vi.

C) THE PART PLAYED BY C/EBP IN ADIPOGENESIS AND LIPOGENESIS.

The involvement of C/EBPs in adipogenesis can be summarized as follows. C/EBPβ and -δ are expressed transiently in the earliest steps of adipocyte differentiation; they activate the expression of PPARγ and C/EBPα, both of which are involved in the terminal differentiation of the adipocyte. Overexpression of C/EBPβ triggers the differentiation of preadipocytes into adipocytes in the absence of adipogenic hormones, whereas an inhibition of C/EBPα results in a blockade of late adipocyte differentiation (173, 267).

Compared with these cell culture studies, the phenotype of mice carrying null allele mutations for the various C/EBPs is more difficult to interpret. C/EBPα null mice exhibit a severe reduction of brown and white fat mass (312), which may result from impaired lipogenesis, and thus lipid accumulation, rather than from altered adipocyte differentiation. This distinction is also suggested by the phenotype of either C/EBPβ or C/EBPδ null mice. Whereas little alteration is seen in the adipose tissue of these animals, mice carrying the double null allele mutation exhibit an important hypoplasia of WAT reflecting an impaired adipocyte differentiation program, and a moderate reduction of the brown adipose tissue mass attributed to reduced lipogenesis (295). Interestingly, these observations point to distinct regulations that may occur through the same transcription factors in brown and white adipose tissues (reviewed in Ref. 253).

The target genes of the C/EBPs that mediate their effects in fatty acid metabolism remain largely unknown. The transcription factor cascade described for adipogenesis suggests that C/EBPs coordinate the differentiation program by controlling the expression of transcription factors such as C/EBPα itself and PPARγ rather than by acting on direct effectors (274). Furthermore, because...
C/EBPs are considered as constitutive regulators, their activity has to be regulated by increased or decreased expression. The two main hormonal signals known so far to trigger C/EBP expression in the liver are glucocorticoids and glucagon, hormones that are secreted in response to stress, fasting, or hypoglycemia. Glucocorticoids increase C/EBPα and C/EBPβ levels in the liver (45, 89), which in turn stimulates gluconeogenesis (see sect. 2). Glucagon, and more particularly increased cAMP levels, are also an important metabolic signal that upregulates C/EBPβ expression in hepatocytes (44, 45). The regulation in fat tissue is somewhat different, since glucocorticoids decrease C/EBPα expression in WAT (184). In addition, and as mentioned above, insulin induces C/EBPβ in 3T3-L1 cells, via an increased SREBP-1c expression. Finally, C/EBPs are highly phosphorylated; however, the many studies aimed at deciphering how the phosphorylation pattern of C/EBPs might alter their metabolic-directed activities have not delivered a clear answer so far (238).

In summary, C/EBPs are linked to fatty acid metabolism via their implications in adipogenesis. However, their molecular mode of action remains elusive.

D) PPARβ: A ROLE IN ADIPOGENESIS? Reasons for the scarce knowledge gained so far of PPARβ functions are its ubiquitous expression and the lack of specific synthetic and natural ligands until recently. Carbaprostacyclin (cPGI), a stable analog of prostacyclin (PGl₂), acts as an agonist of PPARβ, supporting the notion that the cyclooxygenase-2 arachidonate metabolite PGI₂ might itself act as a bona fide natural ligand for PPARβ (273). Like PPARα and PPARγ, PPARβ binds fatty acids and, therefore, is also most likely a sensor of dietary lipids and lipid derivatives (reviewed in Ref. 196). In preadipocytes, PPARβ mediates long-chain fatty acid effects on the expression of adipose-related genes (2). Together with two additional transcription factors, C/EBPβ and C/EBPδ, PPARβ appears to be implicated in the induction of PPARγ expression (12, 110). In turn, high expression of PPARγ and C/EBPs in adipocytes establishes and maintains the terminal differentiation program. According to this scheme, activation of PPARβ by dietary lipids in preadipocytes would contribute to the expansion of the adipose tissue, a role consistent with the decreased amount of brown and white adipose tissues reported in the PPARβ null mice (8, 228). However, these observations contrast somewhat with other evidence reported by the same investigators. Adipose tissue specific deletion of PPARβ does not alter fat mass (8), whereas PPARβ overexpression and overactivation in the WAT trigger fatty acid mobilization and oxidation (313). Overexpression of PPARβ in C2C12 myoblasts participates in their transdifferentiation in adipocytes (110), while a PPARβ agonist induces fatty acid oxidation in differentiated muscle cells (111). Indeed, there is now compelling evidence which supports a role for PPARβ in fatty acid oxidation in muscle, as is discussed below. However, this activity in mature muscle cells does not preclude a specific role of PPARβ in the very early stages of adipocyte differentiation.

E) DIVERSE ARRAYS OF SIGNALS PARTICIPATE IN THE TUNING OF THE ADIPOGENESIS PROCESS. We have seen that adipogenesis is a highly regulated process in which C/EBPs and PPARγ play a major part. However, there is a diverse array of factors that completes the scheme. For example, HNF3β/FOXA2 is expressed in preadipocytes as well as in adipocytes and inhibits adipocyte differentiation (321). FOXC2 is a winged helix/forkhead transcription factor that induces the expression of the type Ia regulatory subunit of cAMP-dependent PKA (29). Consequently, FOXC2 increases the sensitivity to PKA activation (29) and was shown to inhibit white adipocyte differentiation (49). Other transducing signals acting on adipocyte differentiation involve SMAD3, GATA2, and GATA3 (reviewed in Ref. 101), but further work is needed to integrate these factors in a global representation of the transcriptional regulation network controlling adipogenesis.

2. Fatty acid oxidation: regulation by PPARα and PPARβ

In contrast with the numerous factors involved in fatty acid synthesis and storage, our knowledge of the transcriptional regulation of fatty acid oxidation has long been dominated by the major role of PPARα. However, in addition to recent work suggesting the potential role of PPARβ in fatty acid oxidation, other transcription factors certainly participate in the coordinated induction of the fatty acid oxidation enzymes. To gain some insight as to the nature of these factors, the regulation of the expression of the carnitine palmitoyltransferase I (CPT-1) gene, encoding a major rate-limiting enzyme in fatty acid oxidation, is used as a model and will be discussed in this section.

A) PPARα: A MAJOR TRANSCRIPTIONAL REGULATOR OF FATTY ACID OXIDATION. PPARα expression levels are the highest in brown adipose tissue and the liver, then in the heart, kidneys, enterocytes, and muscles. PPARα target genes constitute a comprehensive set of genes that participate in many if not all aspects of lipid catabolism. This includes fatty acid transport across the cell membrane (fatty acid transporter protein genes), intracellular binding (liver fatty acid binding protein gene), activation via β-oxidation in peroxisomes and mitochondria, and catabolism by α-oxidation in microsomes (acyl-CoA oxidase gene, CYP4A1 and CYP4A6 genes, medium-chain acyl-CoA dehydrogenase, and 3-hydroxy 3-methylglutaryl-CoA synthase genes) (reviewed in Ref. 54). The role of PPARα in fatty acid oxidation is particularly highlighted during fasting that results in an
enhanced load of fatty acids in the liver, to be used as energy source. Food deprivation provokes an increased expression and activity of PPARα, which stimulates β-oxidation. PPARα null mice, which are viable and exhibit only subtle abnormalities in lipid metabolism when kept under normal laboratory confinement and diet (163, 224), cannot sustain fasting. Their inability to enhance fatty acid oxidation results in hypoketonemia, associated with severe hypothermia and hypoglycemia (142, 170). Thus PPARα is crucial for the organism to adapt to increased demand in fatty acid oxidation, while it seems to play a marginal role in the basal situation with normal diets.

Most of the initial studies of PPARα and its role in fatty acid oxidation have been performed in the liver. However, the major site of fatty acid oxidation at rest and during exercise is the skeletal muscle, which expresses PPARα. Interestingly, a clinical study in lean women demonstrated that endurance training increases PPARα expression in muscle, together with increased expression of medium-chain acyl-CoA dehydrogenase (MCAD) and very-long-chain acyl-CoA dehydrogenase (VLCAD) (113), thereby increasing the oxidative capacity of the muscle. Surprisingly, PPARα null mice have a relatively mild phenotype in this respect, characterized by a precocious exhaustion when exercising. In normally fed animals, the fatty acid oxidative capacity in muscle is similar in wild-type and PPARα null mice. Upon starvation, the fatty acid oxidative capacity in muscle of null mice is reduced by only 30%, suggesting that other factors are important for the regulation of this pathway (206). This is in agreement with the coordinated induction of many genes involved in fatty acid oxidation in the muscles of streptozotocin-induced diabetic mice, which is paradoxically associated with a decreased expression of PPARα (333).

In cardiac muscle, PPARα is increased at the transition from fetal to adult cardiomyocytes. This step is characterized by a switch in the source of energy from glucose and lactate to fatty acids and by high expression levels of the PPARα target genes involved in their oxidation. PGC-1, which increases in parallel, may cooperate with PPARα for the induction of the genes encoding enzymes of the fatty acid oxidation pathway (11). In pathological cardiac hypertrophy, the expression of PPARα is down-regulated, the utilization of fatty acids as energy substrate is decreased, and the genes implicated in the utilization of glucose as the main energy source are reinduced (10). It is presently unclear whether the decline in PPARα activity and fatty acid oxidation is a cause or a consequence of cardiac hypertrophy. It is therefore also unclear if PPARα ligands would be beneficial or detrimental to the heart in this pathological context (337; reviewed in Ref. 167). In this context, it is interesting to note that PGC-1α is a crucial regulator of mitochondrial biogenesis (236). It interacts with the nuclear respiratory factor NRF1, which positively regulates the nuclear genes that are involved in the biogenesis of the respiratory chain (68, 323). The expression of PGC-1α also dramatically increases at birth when the heart has an increased requirement for mitochondrial respiration (166). Overall, in the heart, PGC-1 could coordinate the increased capacity of the respiratory chain via its interaction with NRF1 and the increased fatty acid oxidation which provides the fuel for the respiratory chain via a functional interaction with PPARα.

B) PPARβ AND FATTY ACID OXIDATION: OVERLAPS AND SPECIFICITIES WITH RESPECT TO PPARα FUNCTIONS. The first studies linking PPARβ to fatty acid oxidation in muscle were performed in cultures of primary muscle cells or muscle cell lines, where activation of PPARβ induced UCP2 and UCP3 gene expression (38, 206). In C2C12 cells, a PPARβ-dependent upregulation of H-FABP and FAT, and to a lesser extent of LPL, ACS, and CPT1 was observed (58, 110). In vivo, transgenic mice that overexpress PPARβ or a constitutively active PPARβ-VP16 fusion protein in muscle exhibit an enrichment of the muscle in red oxidative fibers, with an increased oxidative capacity assessed both at the gene expression and functional levels (182, 314). Similar results were obtained in wild-type mice treated with the PPARβ agonist (GW501516). Such a treatment results in a dose-dependent activation of fatty acid β-oxidation in the quadriceps muscles, sustained by the higher expression of genes encoding enzymes involved in mitochondrial fatty acid catabolism, such as fatty acid transport proteins (FAT and LCAD) as well as UCP2 and UCP3 (294). Overlapping activities of PPARβ with PPARα with respect to fatty acid oxidation are also described in the heart, where a mutation of PPARα is partially compensated by PPARβ (36, 50). In this organ, PPARβ may have a dominant role, as a cardiac specific disruption of PPARβ results in a cardiomyopathy that develops as early as 4 mo of age and is associated with a general decrease in the cardiac expression of all genes involved in β-oxidation (35).

In the adipose tissue, and in contrast to the proposed role of PPARβ in adipogenesis (see previous section), PPARβ may also be involved in fatty acid oxidation. Overexpression of the PPARβ-VP16 fusion protein in brown and white adipose tissues produces lean mice and increases the mobilization and oxidation of fatty acids (313). This is due, at least in part, to the concomitant increased expression of genes involved in fatty acid β-oxidation (long-chain acyl-CoA dehydrogenase, LCAD; VLCAD), lipolysis (hormone-sensitive lipase, HSL), and energy uncoupling in mitochondria (uncoupling proteins UCPI and UCP3). The increased catabolism of fatty acids in these transgenic mice is further illustrated by the protective effects of adipose PPARβ-VP16 expression against obesity, hyperlipidemia, and liver steatosis upon high-fat diet feeding.

An interesting aspect here is the consistent upregulation of the uncoupling proteins in various tissues. This suggests that in the adipose tissue and the skeletal muscle, PPARβ expression and activity are also linked to
energy dissipation, in addition to facilitating the use of fatty acid as an energy source (reviewed in Ref. 13).

C. Transcriptional Regulation of CPT-I: A Complex Array of Transcription Factors. CPT-I is a major rate-limiting enzyme of fatty acid oxidation, and functional analyses of CPT-I promoter have revealed the contribution of other transcription factors to the regulation of fatty acid oxidation. The CPT system controls the transfer of long-chain fatty acids inside the mitochondria. CPT-I, which is localized in the outer mitochondrial membrane, is considered as a rate-limiting enzyme for the oxidation of long-chain fatty acids. The major regulator of CPT-I activity consists of an allosteric control via malonyl-CoA, a substrate for fatty acid synthesis whose accumulation inhibits CPT-I activity.

CPT-I is expressed as two distinct isoforms, L-CPT-I (also called CTP-\(\alpha\) in the liver) and M-CPT-I (also called CPT-\(\beta\) in muscles and adipose tissue). Transcription of L-CPT-I is under the control of the thyroid hormone receptor and PPAR\(\alpha\), through appropriate response elements found in the promoter of the gene (128). A sequence element in the first intron is also proposed to mediate a PPAR-independent transcriptional regulation by long-chain fatty acids (126, 178, 180). In addition, glucagon can induce L-CPT-I expression through increased cAMP levels. Intriguingly, the cAMP response unit is a composite of the DR1 sequence, previously shown to strongly activated by CREB in liver in response to cAMP induction, and of a CREB binding element. As already mentioned (see sect. iiC), PGC1 is strongly activated by CREB in liver in response to cAMP and potentiates HNF4\(\alpha\) and glucocorticoid/glucocorticoid receptor action onto the promoter of the gluconeogenic enzyme PEPCK (106). A similar mechanism is now proposed to activate L-CPT-I via the formation of a multiprotein complex containing PGC1, CREB, and HNF4\(\alpha\) (179). Chromatin immunoprecipitation might now be of help to analyze the composition of the transcriptional complex that binds to this composite element.

Much less is known about the regulation of M-CPTI in muscle. An interaction of PGC1 with the muscle specific myocyte enhancer factor 2 (MEF2) has been proposed to increase the expression of M-CPT. USF proteins would in contrast repress M-CPT-I gene expression, possibly by interacting with the cofactor PGC1, and therefore reduce MEF2 activity (202). Thus the functional network of transcription factors must be broadened to include that of the coactivators. While not unexpected, this leads the way for further understanding of the complexity of metabolic regulations.

Overall, our broad understanding of the mechanisms controlling lipogenesis and adipogenesis differs strikingly from our relatively poor understanding of the transcriptional control of fatty acid oxidation. Does this reflect the respective diversity and paucity of the mechanisms involved in these two processes? Recent progress towards understanding the transcriptional control of fatty acid oxidation will help to build a more comprehensive model of the necessary balance between storage and consumption.

B. Transcriptional Control of Cholesterol Homeostasis

1. An outline of cholesterol metabolism and its main regulatory factors

Cholesterol is an essential component of cell membranes and is therefore critical for cell viability and growth. It is also the precursor from which steroids and bile acids are synthesized.

The main source of cholesterol is the diet. Its inclusion in micelles, formed by bile acids in the intestine, facilitates its uptake as well as that of fatty acids by the enterocytes. In these cells, lipids are incorporated into chylophilomicros, then secreted into the lymph for delivery into the bloodstream. Chylomicros deliver triglycerides to peripheral tissues via LPL-mediated hydrolysis, and their cholesterol-enriched remnants are taken up by the hepatocytes. In the liver, cholesterol is channeled to bile acid synthesis or is reincorporated into the very-low-density lipoprotein (VLDL) particles that are redelivered into systemic circulation. Lipoproteins are complex particles that allow the transport and circulation of lipids such as phospholipids, free cholesterol, esterified cholesterol, and triglycerides, in the blood. The proteins associated with the particles, or apoproteins, serve as lipid acceptors, enzyme cofactors, or ligands for receptor-mediated cellular uptake. Each lipoprotein particle has distinct properties and distinct destinations. Among them, VLDL particles are assembled in the liver and redistribute triglycerides and cholesterol to peripheral cells. They become LDL after a partial depletion in triglycerides has occurred, due to peripheral lipoprotein lipase activity. LDL are considered as the “bad” cholesterol compartment, due to their poor clearance from the blood, in contrast to HDL, which contain apoAI and apoAll proteins that serve as cholesterol acceptor and which are efficiently cleared from the blood by the liver. VLDL form the primary source of cholesterol for peripheral tissues, mainly via the low-density lipoprotein (LDL) receptor expression which is a main regulatory step for adjusting the import of cholesterol (reviewed in Ref. 340). Its export from cells requires the expression of transporters of the ATP binding cassette (ABC) superfamily and the extracellular presence of specific apolipoproteins as free cholesterol acceptors. These proteins are major components of the reverse cholesterol pathway and are therefore crucial for the efflux of excess cellular cholesterol. At the level of the whole organism, if cholesterol is in excess, its
conversion into bile acids is increased, allowing its elimination in the feces, which is the sole way of excreting cholesterol. This excretion is further regulated by the enterohepatic cycle of bile acids [i.e., secretion in the gut lumen, followed by reabsorption in the ileum and reentry to the liver (see Fig. 9)], which is in part aimed at saving complex and energy-costly molecules such as cholesterol and its bile acid derivatives. In contrast, if the cholesterol supply is low, de novo synthesis takes place mainly in the liver and the intestine.

The factors acting on the transcriptional regulation of cholesterol homeostasis can be presented schematically as follows. When there are sufficient amounts of cholesterol, membrane-bound SREBP-2 is not processed to its mature form, and cholesterol synthesis is not stimulated. In addition, if the cholesterol supply is high, LXR is activated by the high levels of oxysterols that are the intermediate compounds in the process of cholesterol degradation. In peripheral cells, LXR activity enhances cholesterol efflux, while in the liver it increases the formation of bile acids to avoid an accumulation of cholesterol in excess. In turn, FXR that senses high levels of bile acids inhibits their synthesis, thereby limiting the elevation of bile acid levels and their deleterious effects on liver cell functions. Conversely, low cholesterol levels are also sensed by the cells, which respond by increasing the processing and activity of SREBP-2, thereby stimulating cholesterol de novo synthesis.

2. **SREBP-2 is required for the transcriptional activation of the cholesterol synthesis pathway**

SREBP-2 is present in most tissues at rather low levels but is more particularly expressed in the liver. As mentioned previously, high levels of cholesterol in the ER membrane inhibit the activity of the SREBP-cleavage activating protein (SCAP). This occurs through Insig-1 and Insig-2, which are ER proteins that bind SCAP in the presence of high levels of cholesterol. Insig interaction retains the SCAP/SREBP complex in the ER (331) (see Appendix G and Fig. 7). When cholesterol is low, this interaction is released, and the activated SCAP triggers the transport of the SCAP/SREBP complex from the ER to the Golgi membrane where proteases free the cytosolic domain of SREBP-2. This domain then translocates to the nucleus and affects the expression of key enzymes in cholesterol biosynthesis and uptake (19). In addition to this control, SREBP-2 positively regulates its own gene transcription via an SRE present in its promoter region (268). In contrast, upregulation of SREBP-2 in LXR null mice suggests that LXR has a negative action on SREBP-2 expression and, consequently, on the expression of its target genes (226). However, the mechanism of this downregulation and its relevance in humans has not yet been addressed. The tight posttranscriptional control of SREBP-2 by cholesterol also raises the question of whether transcriptional regulation of SREBP-2 participates in the regulation of its activity.

![Fig. 9. The coordinated action of FXR, LXR, and SREBP-2 in the cholesterol metabolism pathway.](http://physrev.physiology.org/)

For more details, see section IV B.
In transgenic mice, overexpression of the active nuclear form of SREBP-2 dramatically increases the expression of the genes involved in cholesterol synthesis, particularly that encoding HMG-CoA reductase. In addition to cholesterol synthesis, SREBP-2 enhances the expression of the LDL receptor gene. A subsequent massive cholesterol overload of the liver is induced, with only a comparatively mild increase in fatty acid accumulation (reviewed in Ref. 114). Whereas little knowledge has been gained from SREBP-2 null mice which die in utero for nonelucidated reasons, a liver-specific mutation of SCAP almost completely blunts cholesterol synthesis (191).

In short, the synthesis of cholesterol by the liver and peripheral cells is strictly dependent on the presence of SREBP-2, whose activity is tightly regulated by the levels of membrane cholesterol sensed by SCAP.

3. LXR: a player in the reverse cholesterol pathway

LXR (see Appendix C) was initially characterized by its role in the positive regulation of the gene encoding cholesterol 7α-hydroxylase (CYP7A), the rate-limiting enzyme in the neutral bile acid biosynthetic pathway. The nature of LXR endogenous ligands, the cholesterol metabolite oxysterols, further emphasized the importance of this receptor in cholesterol metabolism (127, 168). This function has been confirmed by the phenotype of LXRα null mice, which display a severely impaired cholesterol and bile acid metabolism. Indeed, they fail to induce CYP7A and consequently suffer from a dramatic accumulation of cholesteryl ester in the liver with no increase in bile acid production (226). LXR in mice was thus credited for a major role in the disposal of excess cholesterol by increased elimination via bile acid synthesis (see Fig. 9). However, it is now clear that the human CYP7A is not responsive to LXR and might even be repressed by LXRα activation (34, 87). This difference between mouse and human is of interest as it might explain, at least in part, both the higher capability of mice to face a high-cholesterol diet and their increased resistance to developing atherosclerosis.

As a consequence, research now focuses more on other important LXR-mediated regulations that converge on the reverse cholesterol transport pathway. This pathway limits the exposure of peripheral cells to cholesterol excess, and its modulation by LXR has been reported in both humans and mice, at least at three levels (see Fig. 9). First, LXR upregulates the expression of several genes coding for members of the ABC transporter family. ABCG5 and ABCG8, which are expressed almost exclusively in the liver and small intestine, favor the secretion of sterols from the liver epithelial cells to the bile duct and from the gut epithelial cells to the intestinal lumen (14). Activation of these two genes by LXR is considered to be the main mechanism by which an LXR agonist in mice causes a total blockade of cholesterol absorption (232, 242). Another important target is the widespread ABCA1 transporter, which promotes the efflux of intracellular and plasma membrane cholesterol to the nascent high-density lipoprotein (HDL) particles via interaction with ApoA1, thereby increasing HDL levels (245). Second, LXR increases ApoE expression. Effluxed cholesterol from cell membranes can also be charged on HDL particles by ApoE, increasing their total capacity for accepting cholesterol. In addition, ApoE-containing particles can interact with the scavenger receptor that increases uptake of these particles in the liver. Third, LXR increases the expression of the cholesteryl ester transfer protein (CETP), which promotes cholesteryl ester transfer from VLDL to HDL and from HDL to LDL, a lipoprotein which is also efficiently taken up by the liver. By these means, LXR increases cholesterol clearance from the blood (reviewed in Ref. 292). Some of these regulations are shared by LXRα and LXRβ. If LXRβ null mice do not have the dramatic phenotype described for LXRα, the double mutant mice are more strongly affected than the LXRα null mice (244). However, no functional compensation by LXRβ is seen in LXRα null mutant mice, and a specific role for LXRβ has not yet been clearly defined.

In short, LXR is the major transcription factor that acts as a sensor of cholesterol levels via its interaction with oxysterols and, in turn, drives the disposal of excess cholesterol. It also acts at the level of individual cells by increasing the ABC transporter molecules responsible for cholesterol efflux, and at the level of the organism by decreasing the cholesterol uptake from the diet. In mice livers, it also increases the conversion of cholesterol into bile acid.

4. FXR and the inhibition of bile acid synthesis

The farnesol X receptor (FXR) senses bile acids and responds by inhibiting bile acid synthesis (see Appendix D), as illustrated in FXR null mice. These mice, which have no overt phenotype except increased bile acid levels in the blood, cannot sustain a cholic acid-enriched diet. They suffer a severe wasting syndrome with hypothermia, and ~30% of them die by day 7 on such a diet (282), which is probably mainly due to the known toxicity provoked by accumulation of bile acids in hepatocytes (252).

As discussed above, CYP7A expression controls the neutral pathway of bile acid synthesis (Fig. 9). CYP7A, which is positively regulated by LXR in mice livers, is negatively regulated by FXR via an indirect mechanism. The constitutive expression of CYP7A in the liver depends on the nuclear factor LRH-1 (NR5A2), which binds to a response element located in the promoter of the CYP7A gene. Activated FXR does not bind to the promoter of this
gene but upregulates the expression of the short heterodimerization partner (SHP-1, NR0B2), which binds to LRH-1 to form an inactive heterodimer. This results in downregulation of CYP7A expression, even in the presence of activated LXR (86, 181). Increased expression of SHP-1 may also be responsible for the FXR-mediated decreased expression of the sodium taurocholate cotransporting polypeptide (NTCP). In hepatocytes, this transmembrane protein mediates the uptake of bile acids that are returning from the gut to the liver in the enterohepatic cycle. Finally, a direct action of FXR:RXR positively regulates the gene encoding the bile salt export pump (BSEP). This protein, only expressed in the liver, belongs to the ABC transporter superfamily and allows the extrusion of bile acids from hepatocytes into the biliary canaliculus. These coordinated actions of FXR on CYP7A, NTCP, and BSEP result in lowering the potentially deleterious high levels of bile acids to which hepatocytes are exposed (reviewed in Ref. 77). In contrast, the FXR-mediated induction of the ileal bile acid binding protein (IBABP), an intracellular carrier of bile acids expressed in the ileal epithelial cells, favors the reuptake of bile acids from the gut lumen (91). One proposed hypothesis is that reabsorption of bile acids in the ileum would diminish the micelle formation and solubilization of cholesterol and thus limit its absorption. Because bile acids are more abundant in times of cholesterol excess, this would limit further cholesterol uptake by the intestine. This hypothesis might seem incomplete at first glance, as bile acid reabsorption mainly occurs in the ileum, whereas that of cholesterol mainly happens in the jejunum. However, ABCG5 and ABCG8, which are important regulators of cholesterol absorption, are also expressed in the ileum (59), and clinical studies in subjects with ileal pouch-anal anastomosis suggest that ileal dysfunction diminishes cholesterol absorption (153).

FXR might also act directly on circulating lipoproteins, by inducing the expression and secretion of hepatic apoCII (138), and by increasing the levels of the secreted enzyme phospholipid transfer protein (PLTP), which facilitates the transfer of cholesterol and phospholipids, respectively, from triglyceride-rich lipoproteins to HDL (303). However, the prominent effects of FXR on bile acid metabolism in part mask the specific effects of the increased apoCII and PLTP expression.

In short, FXR is the transcription factor that senses the intracellular levels of bile acids and is required for limiting liver bile acid accumulation. It inhibits bile acid synthesis via the downregulation of CYP7A and increases bile acid efflux in the bile via increased BSEP expression. In the ileal enterocytes, the reabsorbed bile is taken in charge by the cytosolic IBABP whose expression is also increased by FXR.

In conclusion, because cholesterol synthesis is quite energy-demanding, an efficient transport system for picking up cholesterol from the diet is probably advantageous for any organism. This efficiency is reflected in the increased cellular cholesterol levels that correlate with a cholesterol-rich diet. Once again, the mammalian organism is quite well-adapted for sparing the important and high-cost cholesterol molecules, but is ill-prepared for coping with an excess of cholesterol intake, which adds to cardiovascular risks.

C. Intricate Regulation of Cholesterol and Fatty Acid Metabolism

Recent progress in understanding the regulatory pathways of fatty acid metabolism and cholesterol homeostasis are progressively unraveling more and more intricate regulatory events, shrinking the gap between these two main branches of lipid metabolism. In this section, we present these intersecting pathways, although some have already been mentioned in the appropriate sections above.

This meshing can be seen from two points of view. First, lipoprotein metabolism itself participates in both cholesterol and fatty acid metabolism and controls the circulation and tissue delivery of these molecules in a very fine-tuned and orchestrated manner. Second, transcription factors might themselves be knitting points in the intricate pattern of regulation. This is the case for the two SREBP isotypes, but also for LXR, which was initially credited with a major role in cholesterol homeostasis but is also now integrated in many aspects of fatty acid metabolism. RXR, by being a promiscuous partner for each of the nuclear receptors that participate in the control of lipid metabolism, could also act as a converging regulatory node. However, this is only partially true as discussed below.

1. Regulation of the lipoprotein system and the particular role of PPARs

As discussed above, activation and repression of various components of the reverse cholesterol pathways are regulated by LXR and FXR, thereby directly affecting the circulating lipoproteins. We will discuss below how PPARs, which mainly target fatty acid metabolism, also affect cholesterol metabolism via their action on lipoproteins.

The blood lipoprotein profile is first modulated by PPARs via PPAR-mediated increased expression of LPL (reviewed in Ref. 15). Both PPARα and PPARγ can upregulate LPL expression. This is associated with a decreased expression of apoCIII, an inhibitor of LPL activity. These joint effects increase the free fatty acid delivery to tissues such as muscle and adipose tissue. PPARα also participates in an increase in the reverse cholesterol transport by upregulating the expression of the genes encoding the cholesterol acceptor apolipoprotein apoAI.
and apoAI. These two proteins participate in the formation of HDL particles, which carry the cholesterol transferred from chylomicron and VLDL remnants to the liver. Finally, PPARα increases the hepatic expression of the scavenger receptor BI (SR-BI)/CLA-1, thereby increasing the selective uptake of HDL cholesteryl esters from the blood (reviewed in Ref. 54). These actions of PPARα on the lipoprotein profile, associated with its stimulation of fatty acid oxidation, counteract the highly atherogenic situation characterized by high levels of triglycerides and low levels of HDL in the blood, often associated with clinical dyslipidemia. Indeed, these observations unveil the molecular mechanisms by which fibrates have hypolipidemic properties that are very useful for the treatment of human lipid disorders.

The role of PPARβ in regulating the lipoprotein transport system has been best demonstrated in obese rhesus monkeys, used as a relevant animal model for human obesity and the associated metabolic disorders. Treatment of these animals with a selective PPARβ agonist caused an increase in the level of serum HDL cholesterol, while lowering the level of small-dense LDL, fasting triglycerides, and fasting insulin (216). In mice, the administration of a PPARβ-selective synthetic agonist to obese and diabetic db/db mice raised total plasma cholesterol levels primarily associated with HDL particles and decreased the expression of the lipoprotein lipase in WAT (169). However, under nonchallenging conditions, PPARβ null mice have normal plasma lipid composition with no changes in the levels of total cholesterol, HDL-cholesterol, triglycerides, and free fatty acids compared with wild-type littermates. Therefore, the contribution of PPARβ to lipoprotein metabolism in healthy mice remains to be investigated, especially in the context of different diets and levels of physical exercise (reviewed in Ref. 13).

2. SREBPs and LXR at the branching point between fatty acid and cholesterol metabolism

The combined role of SREBP-1c and SREBP-2 in fatty acid and in cholesterol metabolisms can be effectively highlighted using a scheme representing the two pathways emerging from the single initial precursor acetyl-CoA (63, 114) (see Fig. 8). Acetyl-CoA is either condensed in acetocetyl CoA via acetacetyl CoA thiolase for cholesterol synthesis, or is transformed to malonyl-CoA by acetyl-CoA carboxylase for fatty acid synthesis. Alternatively, acetyl-CoA is oxidized in the citric acid cycle for ATP production. The relative importance of each of these three pathways is determined by the levels of activity of their respective enzymes. These are, at least in part, dependent on the transcriptional expression of the corresponding genes controlled by SREBP-1 and SREBP-2. In addition, both cholesterol and fatty acid synthesis require NADPH. NADPH is produced either by the malic enzyme (ME), which transforms cytosolic malate into pyruvate, or via the pentose phosphate pathway, which produces ribulose and xylulose from G6P by activation of the glucose-6-phosphate dehydrogenase (G6PDH) and 6-phosphogluconate dehydrogenase (PGDH). Interestingly, the genes coding for these three enzymes (ME, G6PDH, PGDH) are targets of both SREBP-2 and SREBP-1c (281).

Links between fatty acid and cholesterol metabolisms also exist via the mechanisms regulating VLDL assembly. SREBP-1 and SREBP-2 are both capable of inducing the expression of the liver microsomal triglyceride transfer protein (MTP), which is essential in the process of VLDL assembly. VLDL particles are composed of the apoB protein, phospholipids and free cholesterol at the surface, and triglycerides and cholesterol esters in the core. The more that cholesterol is synthesized or present in the liver, the more fatty acids are associated with the forming particles. In addition, cholesterol esterification also requires fatty acids, and more particularly oleate. The unusually high level of oleate in mice overexpressing SREBP-1c might reflect a role for SREBP-1c in promoting the supply of the appropriate fatty acid. SREBP-1 and SREBP-2 also increase the expression of the LDL receptor in hepatic cells (115). Thus, by increasing the turnover of the VLDL particles, SREBPs contribute towards enhancing the supply of fatty acids and cholesterol to peripheral cells and limiting hyperlipoproteinemia.

The elegant design of a microarray experiment has given a recapitulation of most of these observations and casts light on new features with respect to SREBP target genes (115). The set of upregulated genes in transgenic mice overexpressing either SREBP-1a or SREBP-2 were compared with that of downregulated genes in mice specifically deleted of SCAP in the liver. The small subset of genes found in both is likely composed of genes directly regulated by SREBP activation. All others, in contrast, likely respond to secondary events. Two major LXR target genes involved in cholesterol metabolism, ABCG5 and ABCG8, but not LXR itself, are found in the latter class. Indeed, SREBP activation leads to the production of endogenous LXR ligand and increases its activity. Reciprocally, activated LXR also acts on fatty acid synthesis via stimulation of SREBP-1c. In addition, experiments performed in HepG2 cells demonstrate that LXR plays a direct role in enhancing the expression of FAS (132). Together these observations highlight the intertwined regulation of cholesterol and fatty acid metabolisms and might explain the steatosis and the massive increase in VLDL and triglyceride blood levels observed in mice treated with pharmacological doses of an LXR ligand (90). In contrast to the results discussed above and with the general pattern of LXR activities, a study performed in 3T3-L1 aimed at understanding the role of LXRα in the adipocytes reports an increased release of nonesterified
fatty acids. In the same report, treatment of mice with an LXRα agonist increased blood levels in glycerol and non-esterified fatty acids, which suggests an LXR-mediated increased lipolysis in the WAT (259). These discrepancies remain to be elucidated.

3. RXR: a pivotal element of sensor-regulated pathways

RXR occupies a particular position, as it participates in most of the fatty acid and cholesterol metabolic regulations via its interaction with several of the nuclear receptors, a simple means of creating regulatory cross-talk. The role of RXR in metabolic regulation must be analyzed from two different angles.

RXR is an obligatory partner for many nuclear receptors (186). Thus its deletion putatively affects a very diverse array of developmental and physiological pathways. This has been clearly underlined by the thorough phenotypic analyses of the developmental defects occurring in mice carrying various RXR gene deletions or mutations. Whereas RXRβ and RXRγ null mutations give rise to quite minor developmental defects, RXRα null mice die at early embryonic stages (188). This lethality might be due to defects in the PPARβ and PPARγ signaling pathways. Indeed, invalidation of either PPARβ or PPARγ provokes an embryonic lethality due to placental defects, appearing at the same early point as in RXRα null placenta. Other defects analyzed in the various RXR mutants and seen in tissues such as the skin, the eye, the heart, and the testis, reflect alterations in the pathways of many other receptors, such as the thyroid hormone, vitamin D, and retinoic acid receptors (188, 316). From the metabolic point of view, analyzing the importance of RXRα in adult tissues required the generation of tissue-specific knockouts. Specific invalidation of RXRα has been generated in the liver, and metabolic studies were performed to identify which pathways were most affected. As expected, many PPARα-mediated functions in fatty acid oxidation were altered by the lack of RXRα. However, other pathways that include LXR and FXR pathways were also compromised, at least partially, by the absence of RXRα. These effects could not be compensated for by RXRβ and RXRγ (309). RXRα was also invalidated in the adipose tissue. The deletion of the two RXRα alleles in the adipocytes of adult animals resulted in an alteration of preadipocyte differentiation as well as in a resistance to induced obesity. These results are reminiscent of the observations obtained with PPARγ heterozygous mice, suggesting that most of the effects due to the lack of RXRα expression in adipocytes reflect altered PPARγ functions (120). Finally, as mentioned above, the impaired lipolysis observed in these mice might be related to an alteration to LXR:RXR heterodimer signaling.

RXR might also be important not only as a partner for other receptors but also as a bona fide receptor. As already stated, the nature of the endogenous ligand of RXR remains elusive, and all observations made so far were in a pharmacological context, treating mice with 9-cis-retinoic acid (RA) or synthetic RXR agonists. Upon such treatments, RXR may regulate transcription as homodimers (RXR:RXR), binding to DR-1-like response elements. While the formation of these homodimers is difficult to assert in vivo and has been neglected, new approaches by chromatin immunoprecipitation demonstrated that RXR homodimers do form in vivo and efficiently trigger the activation of PPRE-containing genes, such as ME in the liver (119). In contrast, there is major interest in exploiting the ability of RXR ligands to activate the so-called permissive heterodimers. These RXR-containing heterodimers allow RXR to bind its ligand and subsequently turn on the transcriptional activation of the gene to which the heterodimer binds. Thus, in many experimental reports, 9-cis-RA was used in combination with a PPAR ligand to amplify the PPAR biological response. In in vivo studies, potentiation of PPARα action by RXR ligands is mainly observed in the liver (220). In another example, in vivo administration of synthetic RXR ligands mimics, and increases when given in combination with TZD, the metabolic effects of PPARγ ligands, by decreasing hyperglycemia, and improving insulin sensitiv-
adaptive mechanisms to cope with the alternation between feeding and fasting, with a main objective of keeping glucose levels high enough for brain cells and other glucose-dependent tissues while maintaining an energy supply sufficient for all organs. These regulations mainly focus on the production, distribution, use, and saving of energy-rich molecules, and not on constituting optimal energy stores. Here, transcription factors serve two main purposes: regulating specific metabolic adjustments within each peripheral organ and coordinating intertissue regulation for robust homeostasis. This section is aimed at presenting the role of transcription factors involved in the functional pathways detailed in the previous sections, in the context of this daily challenge constituted by the alternating status of fasting and feeding.

Fasting represents a situation that is reflected in the blood by decreased levels of glucose, and accordingly decreased levels of insulin and increased levels of glucagon. To maintain energy supply during fasting, the organism must 1) have efficient lipolysis in the WAT for the systemic release of fatty acids and glycerol, 2) promote the use of fat over glucose in all tissues where possible, and 3) provide glucose in sufficient amounts for glucose-dependent tissues such as the brain and erythrocytes. Upon refeeding, a large glucose load from nutrients taken up by the intestine triggers the hormonal response with high insulin and low glucagon, which coordinates metabolic adjustments in peripheral tissues. This situation is often used to analyze insulin-mediated metabolic action. During this period, energy stores are replenished, in the liver (glycogen and triglycerides), in muscle (glycogen), and in WAT (triglycerides).

A. Fasting-Feeding: Metabolic Adjustment in the WAT

In the WAT, free fatty acid release (i.e., lipolysis), in response to fasting, is a critical step aimed at maintaining whole body energy homeostasis in the absence of external energy supply.

Catecholamines are the most studied signals that trigger lipolysis, but other hormonal signals in the adipose tissue include thyroid hormone, leptin, and growth hormone (reviewed in Ref. 79). HSL is a key enzyme for the mobilization of triglycerides deposited in adipose tissue, and lipolysis is triggered by cAMP/PKA-dependant phosphorylation of HSL, as provoked by adrenergic stimuli (reviewed in Ref. 109). Interestingly, FOXC2, which increases the sensitivity to cAMP/PKA-dependant signals, induces an upregulation of HSL mRNA levels in WAT. Free fatty acids can then be funneled into the mitochondria whose biogenesis is also increased by FOXC2 (29). In this context, the associated increased expression of Ucp1 redirects the energy in thermogenesis (99). It now remains to be determined whether FOXC2 is involved in the non-adrenergic events controlling lipolysis especially in response to fasting or starvation.

A role for nuclear tissue receptors is suggested from the results of the adipose tissue-specific deletion of RXRa in adult mice, which results in impaired lipolysis. This translates into an absence of free fatty acid increase in the blood upon starvation, a lack of adipose tissue reduction, and profound hypothermia, all effects which cause premature death if starvation is prolonged (120). RXR is thought to act in partnership with many other nuclear receptors, and the responsible heterodimer complex(es) that fail to form in the adipose tissue of these mutant mice is not yet known. It is unlikely to be PPARγ, since PPARγ1 and PPARγ2 are strongly downregulated during fasting (306). Based on microarray analyses, LXR has been proposed to be one of these potential RXR partners (259). However, this is difficult to reconcile with the strong LXR-mediated lipogenesis found in both liver and WAT of mice treated with a LXR agonist. Whereas the expression of LXR in the WAT of fasted mice has not been reported, the downregulation of both Glut4 and SREBP-1c, which are considered as LXR target genes (157), suggest that LXR activity itself is also downregulated upon fasting. The question of the RXR partner involved in fasting-induced lipolysis is still open. It might be worth considering PPARβ as a candidate, since its overexpression in the adipose tissue results in an increased fatty acid consumption (313), although PPARβ null mice have a reduced adiposity (8). Leptin also drives lipolysis leading to increased plasma levels of glycerol that are not accompanied by increased free fatty acids. The fact that leptin-mediated lipolysis is inefficient in PPARα null mice (164) suggests that in wild-type mice the increased expression of PPARα caused by leptin triggers an immediate oxidation of the released fatty acids, and the subsequent release of glycerol in the blood.

Refeeding is first characterized by the immediate action of insulin on Glut4 translocation and the coordinated control of enzymatic activities along the glycolytic and fatty acid synthesis pathways. At the transcriptional level, insulin action translates into higher expression of PPARγ in the WAT, with reciprocal positive interaction of SREBP-1c and PPARγ on their respective gene expression, both leading to increased fatty acid synthesis and storage as triglycerides. Fat accumulation is also permitted by the insulin-mediated upregulation of the expression of the transporter Glut4. Although the corresponding transcriptional mechanism involved is not fully elucidated (reviewed in Ref. 31), the insulin-stimulated expression of Glut4 in mouse epididyimal WAT is dependent on LXRα expression, consistent with the induced Glut4 expression by LXR agonists (46). In addition, insulin is strongly antilipolytic via at least two mechanisms. First, insulin lowers cAMP levels through activation of phosphodiesterase...
3B. Second, insulin also provokes the immediate reesterification of the triglycerides which constitutively undergo hydrolysis in WAT cells. These few understood general features mask our ignorance of many facts concerning the transcriptional regulation of the metabolic adjustment to refeeding. The time courses of the patterns of expression in the WAT of the main transcription factors, such as C/EBPα, SREBP, and PPARγ, remain largely unexplored. In contrast, there is a lot of interest in the identification of adipokines, which are molecules released by the WAT, depending on its level of replenishment. These cytokines are thought to play an important role as paracrine and/or endocrine signaling molecules (78, 79). Leptin, which has major effects in reducing food consumption, is the most studied. Other adipokines include adiponectin, resistin, and fasting-induced adipose factor (FIAF), which reach the systemic circulation, but also other cytokines such as tumor necrosis factor-α, interleukin-6, and plasminogen activator inhibitor-1 which may rather act as paracrine factors. While their roles and molecular mechanisms of their actions are beyond the scope of this review, there are high expectations concerning our understanding of energy homeostasis from analyses of the functions of these signaling molecules.

B. Fasting-Feeding: Metabolic Adjustment in Muscles

In muscles, adaptation to fasting is aimed at sparing glucose, first by switching from glucose to fatty acid oxidation. In that respect, analysis of the regulation of the pyruvate dehydrogenase complex (PDC) during starvation is of principal importance. Pyruvate dehydrogenase activity irreversibly converts pyruvate to acetyl-CoA in the mitochondrion, leading to a complete glucose oxidation into CO₂ and H₂O, concomitant with ATP production (see Fig. 10A). This reaction links glycolysis to the citric cycle in organs of high energy demand and provides the carbon for fatty acid and ketone body synthesis in adipose tissue and in the liver. In the opposite situation, i.e., during starvation, the inhibition of PDC in tissues that can meet their energy needs independently of the complete oxidation of glucose is essential to spare pyruvate that can be channeled to the liver. This inhibition is particularly efficient in skeletal muscle via activation of pyruvate dehydrogenase kinase 4 (PDK4), which phosphorylates and thus inactivates PDC. This results in the conversion of pyruvate into lactate, which is used in the liver for gluconeogenesis. PPARα and glucocorticoids increase the expression of PDK4, in both muscles and the liver (102, 117). Thus the activation of PPARα during stress or starvation spares pyruvate for gluconeogenesis via increased levels of PDK4, while it concomitantly stimulates the use of fatty acids as an energy source via increased β-oxidation. Simultaneously, lipolysis of the triglyceride store is also activated in the muscle, but remains a minor source of free fatty acids compared with the supply coming from the adipose tissue and transported to the liver as albumin-bound molecules. However, as mentioned above, the fatty

![Metabolic adjustment of glucose metabolism upon fasting](http://physrev.physiology.org/)

**FIG. 10.** Metabolic adjustment of glucose metabolism upon fasting. **A:** in the liver, pyruvate carboxylase directs pyruvate toward gluconeogenesis, producing glucose for other demanding tissues. In contrast, the pyruvate dehydrogenase complex favors the complete oxidation of pyruvate to provide energy to the cells. The pyruvate dehydrogenase kinase 4 (PDK4) can repress this pathway by inhibiting the activity of the pyruvate dehydrogenase complex. PDK4 is under the positive transcriptional control of PPARα. During fasting in muscle, the activation of PDK4 spares the pyruvate for its redirection to the liver, depicted in **B.** Upon starvation, the glucose-alanine cycle in the muscle redirects the pyruvate formed by glycolysis to the liver in the form of the amino acid alanine that will be used as a gluconeogenic substrate for hepatic glucose production.
acid oxidative capacity of muscle is reduced by only 30% upon starvation in PPARα null mice, suggesting that PPARβ may compensate for the lack of PPARα in these mice (206). This hypothesis is further supported by the demonstration that PPARβ is a transcriptional regulator of fatty acid oxidation in muscle (see above), although the role of PPARβ during fasting in wild-type mice remains to be specifically explored.

Amino acid metabolism in the muscle is also affected by starvation. It must be kept in mind that, while fatty acid oxidation participates in glucose formation by providing the required energy, fatty acids cannot form glucose. In contrast, glycerol from triglyceride lipolysis normally contributes up to ~10% of hepatic gluconeogenesis. Thus, once glycogen is depleted, amino acids, which mainly come from contractile proteins of skeletal muscles, remain the major source of substrate to be converted into glucose. In the glucose-alanine cycle (see Fig. 10B), oxidation of the branched chain amino acids is the privileged source of alanine that is transferred to α-ketoglutarate to form glutamate. Glutamate in turn either donates its amino group to pyruvate forming alanine for the hepatic gluconeogenesis pathway, or is further aminated into glutamine that is used as an energy source by the intestine. In muscles, the transcriptional regulation of contractile protein degradation and amino acid metabolic adaptations are not yet clarified.

Refeeding completely reverses PDK4 induction in both heart and skeletal muscle. The insulin-mediated increase of Glut4 expression and membrane translocation facilitates glucose uptake and glycogen synthesis. Depending on the nature of the fiber, glycolytic or oxidative, preferential consumption of glucose or fatty acids can resume.

C. Fasting-Feeding and Gluconeogenesis: Metabolic Adjustment in the Liver, Kidney, and Small Intestine

While hepatic gluconeogenesis is essential upon fasting, the kidney and small intestine also contribute to glucose output, more particularly upon prolonged fasting (64, 200). In these tissues, the main gluconeogenic substrate is glutamine which undergoes a partial decarboxylation, releasing a three-carbon compound that enters the mitochondrial tricarboxylic acid cycle. In these tissues the expression of G6Pase, which hydrolyzes G6P in glucose and P1, is essential for glucose secretion. G6Pase expression is highly dependent on an insulin-deprived status and is markedly induced during fasting and diabetes.

However, in the normal postabsorptive state, the liver is the site of high glucose production, via glycogenolysis and gluconeogenesis. As we have seen, this requires the use of substrates such as glycerol, which comes from lipolysis, and gluconeogenic amino acids among which are alanine coming from the muscles, and lactate provided by muscle glycology. Glucocorticoids and glucagon are increased upon fasting, and both hormones increase C/EBPβ as well as C/EBPα (reviewed in Ref. 253), which participate in the enzymatic induction responsible for this increased hepatic glucose production. A main conveyer of this adaptive response is CREB, whose PKA-dependent phosphorylation, triggered by glucagon, turns on its activity as a key regulator of gluconeogenic gene transcription. Gluconeogenesis is energy-demanding and requires that fatty acid oxidation provides the necessary amounts of acetyl-CoA, NADH, and ATP as an energy source and a proper directional flux of the enzymatic reactions from pyruvate to glucose. In this respect, PPARα plays a major role in allowing an adaptive response to fasting. Its stimulation by glucocorticoids and sustained increased activity upon fasting correlates with the enzymatic induction of the fatty acid oxidation pathway and with the concomitant production of ketone bodies. At the same time, the lipogenic transcription factor gene SREBP-1c is repressed (172), and its activity further inhibited by increased Insig levels, which retain unprocessed SREBP in the ER membrane. In fact, Insig-2 is expressed via two mRNA species Insig-2a and Insig-2b, the former being upregulated upon fasting (328). With respect to amino acid metabolism, even though there is an increased degradation of muscle proteins, the consequence of fasting is an overall decrease in urea formation, reflecting the lack of dietary proteins that degrade. It is thus interesting to note that both C/EBPα, which increases urea cycle enzyme activities, and PPARα, which is an inhibitor of the urea cycle, are induced upon fasting. These concomitant activities are likely to permit a fine tuning of urea production, according to the needs rather than unilaterally imposing a directional flux in one direction or the other.

The phenotype of PPARα null mice is extremely revealing regarding the intricate links between the regulatory pathways of fatty acid, glucose, and amino acid metabolisms during fasting. Unchallenged PPARα null mutant mice do not present any overt phenotype. However, they cannot sustain fasting due to a severe hypoglycemia and lack of ketone body production, while levels of urea production remain high (141). The severe hypoglycemia of fasted PPARα null mice has been initially attributed to two facts: the low levels of the hepatic glycogen store and an impaired gluconeogenesis, probably due in part to the inability of fatty acids to provide the necessary energy. If we take the example of the PGC1 interaction with HNF4 and the glucocorticoid receptor for induction of the gluconeogenic PEPCK gene, it might be possible that PGC1, which also efficiently interacts with PPARα, also serves PPARα target genes that would belong to the gluconeogenic pathway. However, the hypothesis of an impaired gluconeogenesis in PPARα null mice is challenged by the
observation that PPARα null mice have an increased hepatic glucose production (327). This production might rely on glycerol coming from the adipose tissue as a result of starvation-induced lipolysis of triglycerides. However, the upregulation by fasting of several genes involved in the hepatic metabolism of glycerol (glycerol transporters aquaporin-3 and -9, glycerol kinase, and cytosolic and mitochondrial glycerol-3-phosphate dehydrogenase) is abolished in PPARα null mice (225). Alternatively, it is possible that amino acids are increasingly used for gluconeogenesis in PPARα null mice, which would be consistent with the maintenance of high urea levels in the blood of fasted mice (141). The association between a higher production of glucose and severe hypoglycemia in PPARα null animals indicates that a high disposal of glucose is maintained in the peripheral tissue, due to the lack of an alternative energy source such as ketone bodies and to the impaired ability to directly use fatty acids. As glucose sparing, via PDK4 activation, may not work in fasted PPARα null mice, the muscle could be a glucose sink, participating in the death-frightening hypoglycemia. Such inhibition of the fasting-induced PDK4 expression has indeed been observed in the liver of PPARα null mice (290). The analyses of stable isotopomer flux demonstrated a diminished response of these mice to meal-induced regulation, resulting in a decreased substrate/futile cycling of glucose, despite increased gluconeogenesis (325). However, other mechanisms might be involved. A recent study demonstrates a lack of Glut4 downregulation in the adipose tissue of fasted PPARα null mice. This is associated with an increased transport of glucose in the WAT, which participates in the synthesis of new fat (R. Burcelin and B. Thorens, personal communication).

Refeeding is marked by a delay in the metabolic adjustment of the liver, which is likely due to a lingering action of the gluconeogenic and lipolytic signals. This characterizes an early fed state during which fatty acid β-oxidation remains high and the liver continues in its gluconeogenic mode using lactate and dietary amino acids as gluconeogenic precursors. After this initial stage, the major transcriptional coordination comes on one hand from the downregulation of PPARα and on the other from the upregulation of SREBP-1c, which mediates many of the insulin-dependent actions on lipid but also glucose metabolism (88). Intriguingly, while Insig-2a is strongly repressed by insulin, Insig-1 is itself a target gene of SREBP-1c, thus increasing in parallel during refeeding (331). This stimulation results in controlling SREBP processing and limiting the insulin-mediated high lipogenic activity at the time of refeeding (4, 65). Finally, the feeding-induced flooding of the liver with lipoprotein particles triggers the main pathways of lipid and cholesterol redistribution, as described previously.

D. Metabolic Adjustment and Circadian Rhythm

Circadian rhythms correspond to rhythms established over the day, observed at the level of gene expression, physiology, and behavior. These rhythms are essential for our daily well-being, allowing for anticipatory regulations, concerning for example heartbeat, blood pressure, hormone secretion, to prepare the upcoming activity or rest. The suprachiasmatic nucleus (SCN) is the main controller of these rhythms and is itself regularly reset by the mean of light intensity sensors, located in the retina (246). However, each tissue, if not each cell, contains molecular oscillators, as demonstrated by the brief exposure of fibroblasts to high concentrations of serum that provokes circadian gene expression which persisted for several days (7, 269). The nature of the mammalian circadian clock proteins that compose the molecular oscillator system is now being unraveled and will not be presented here (for a recent review, see Ref. 246). However, one major question is how all these lessors can become synchronized in a complex organism. Indeed, reciprocal interactions between feeding, metabolism, and circadian rhythm correspond to an important part of this systemic cycling.

The metabolism of the liver and kidneys is in part under the control of circadian regulation. In these tissues, genes whose expression follows a circadian rhythm mainly encode enzymes or regulatory proteins involved in food processing or energy homeostasis, such as the cholesterol 7α-hydroxylase, PEPCK, glycogen synthase, and PPARα. As shown in the liver, many clusters of circadian-regulated genes are rate-limiting steps in their respective pathways (222). In the intact organism, the SCN seems to be the main synchronizer of the circadian rhythm seen in these peripheral tissues. However, the role of the daily feeding/fasting alternation in setting or altering the circadian rhythm, and the nature of the cues that reset the SCN according to environment sensors other than light sensors raise interesting questions. In addition, the SCN might use cues to override peripheral clocks, to keep control over genes linked to circadian rhythm, thereby allowing the organism to proceed via anticipatory rather than reactive metabolic adjustments. The role of circadian rhythm as imposed by the SCN in energy metabolism is demonstrated by the obesity and metabolic syndrome developed by mice mutant for the transcription factor Clock, a key component of the molecular clock in the SCN (300).

In this respect, recent observations have enlightened our understanding of these regulations. One of these stems from an observation made in nocturnal rodents, such as mice. Restricting their feeding to a short and unusual period of the day (daytime instead of night) results in a progressive phase shift of the circadian rhythm of liver gene expression. Interestingly, this resetting does not alter the activity of the SCN, which continues in its own phase (47, 288). The observed phase shifting is rather
slow, requiring several days for adjustment. In contrast, the reverse shift, i.e., by rescheduling feeding at night, results in a quick resetting. What is the signal that leads to anticipation of food intake with changes in gut motility or in liver gene expression, such as increased expression of Cyp7a hydroxylase and bile acid synthesis? One hypothesis is that the small circadian oscillations in body temperature might be one of these signals (20). Similarly, the redox status that can be sensed by some clock proteins reflects energy metabolism and might participate in the rhythm imposed by fasting-feeding (262, 270). Other blood-borne signals were looked at in light of experiments showing that numerous signaling factors are able to reset circadian rhythm in the ex vivo system represented by cells in culture. Particularly, glucocorticoids for which we have discussed the impact on many aspects of metabolism were good candidates. Their levels are oscillating during the day, influenced by the circadian rhythm of adrenocorticotropic hormone (ACTH) that drives glucocorticoid release. Interestingly, glucocorticoid administration is indeed able to induce a phase shift of the general circadian rhythm, demonstrating that glucocorticoids might participate in the physiological control of the circadian rhythm synchronized by the SCN. However, they are not responsible for, and rather inhibit, the liver phase shift provoked by the challenge of restricted feeding time, since mice without glucocorticoid receptors shift their liver gene expression rhythm faster than wild-type mice.

This short section leads to two main conclusions with respect to metabolism and its regulation. The first is that any in vivo experimentation must take into account feeding behavior as well as light/dark alternation. The second is that while we are used to thinking in terms of metabolic reaction and adjustment, the anticipation conditioned by the circadian rhythm and the ability of the latter to readjust to feeding time introduces a level of complexity that we are not yet able to fully grasp.

VI. TRANSCRIPTION FACTORS AND INSULIN RESISTANCE: A MAJOR FOCUS ON PEROXISOME PROLIFERATOR ACTIVATED RECEPTORS

A. Introduction

Our westernized societies are witnessing a frightening increase in metabolic disorders, linked in good part to inappropriate nutritional and life-style habits and favored by genetic predisposition. The current definitions of various terms used for characterizing these metabolic disorders are presented in Appendix I. In many cases, but not all, the pathophysiology progression has the following pattern. Obesity and little physical activity are often the initial triggers of the disorders, characterized by high levels of triglycerides and low levels of HDL in the blood. Glycemia levels in the postprandial period or upon fasting are normal at this first stage, reflecting the ability of β-cell insulin secretion to manage high glucose and compensate for nascent insulin resistance. Impaired glucose tolerance, characterized by an increased postprandial glycemia associated with increased insulinemia, signals the second stage with a progression of the disease and an increased risk of macro- and microangiopathy complications (92). The third stage is that of an established type 2 diabetes mellitus (T2DM), which requires pharmacological treatment aimed at controlling the high levels of glycemia, insulin resistance, dyslipidemia, and the vascular complications.

The metabolic syndrome, or syndrome X, corresponds to an aggregation of metabolic disorders clustered around insulin resistance, which does not necessarily evolve towards T2DM (see Appendix I; Ref. 6). The overall effect adds to the insulin resistance and the impaired glucose regulation, a combination of components that comprise altered blood lipid and lipoprotein profiles, hyperpertension, central obesity, and microalbuminuria. These perturbations lead to pathologies, mainly in the cardiovascular area, such as atherosclerosis, which have an extremely high prevalence in westernized societies where they represent the major cause of death.

For both pathologies, T2DM and the metabolic syndrome, insulin resistance is the core initial defect and is the focus of the following discussion.

B. Insulin Resistance: A Mixed Defect in Glucose and Lipid Metabolism

Glucose homeostasis relies on the concerted production of insulin by the pancreatic β-cells, the uptake of glucose by muscle and liver, and the production of glucose by the liver. Insulin resistance is the inability of the peripheral tissues to respond properly to insulin. Fat cells are the most sensitive to insulin levels, responding by inhibition of lipolysis and stimulation of glucose uptake, followed by liver cells in which hepatic glucose production is inhibited (reviewed in Ref. 194). However, quantitatively speaking, the major intracellular metabolic defect in insulin resistance is an impaired muscle glucose uptake and glycogen synthesis (reviewed in Ref. 229). In insulin resistance, hepatic glucose production and lipolysis from fat cells are unduly maintained, while glucose disposal by muscle is less efficient, provoking an increased secretion of insulin by the pancreatic cells to maintain euglycemia (reviewed in Ref. 264). In parallel, chronic high levels of fatty acids in the β-cell and in blood impair insulin secretion by the pancreatic β-cells and participate in pathogenesis. Obesity, which is responsible for high circulating fatty acid levels, is one major cause of insulin resistance. Accordingly, weight reduction and physical activity of
patients clearly improve insulin sensitivity. However, the nature and the mechanism of the initial perturbation that provokes insulin resistance remain unclear.

Although it remains difficult to identify a single phenomenon at the origin of insulin resistance, the possible intricate causes and diverse consequences of insulin resistance are discussed below. Either as a cause and/or consequence, elevated fatty acid levels appear to be a crucial parameter in the development of insulin resistance. Obesity, even in the absence of other metabolic abnormalities, is accompanied by increased free fatty acid release in the blood, in proportion to the fat mass. These elevated levels of circulating free fatty acids divert triglyceride clearance from the adipose tissue towards nonadipose tissues. Visceral fat (i.e., in the splanchnic area) in particular becomes less sensitive to the antilipolytic action and fatty acid reesterification effects of insulin (308). Once insulin resistance is established, insulin signaling becomes even more important, leading to an even greater flux of free fatty acids towards the liver. In the liver, the increased efflux of free fatty acids augments VLDL production, reinforcing the pool of blood triglycerides. Furthermore, the liver maintains a high glucose production, aggravating hyperglycemia. In muscle, increased free fatty acids also impair insulin-stimulated glucose transport. The mechanism proposed is that free fatty acids negatively interfere with the insulin signaling cascade, at levels which are not clearly defined but may encompass the phosphorylation of IRS-1, the level of P3K activity, leading to that of glycolgen synthase activity, and finally Glut 4 gene expression and/or protein translocation (reviewed in Ref. 264).

The diversion of free fatty acids towards nonadipose tissues not only affects liver and muscle metabolism, but may also alter pancreatic β-cell functions. In the Randle hypothesis, a substrate competition occurs in the oxidation pathway between free fatty acids and glucose, to satisfy the energy need of a given tissue. This leads to an impaired utilization of glucose when a high free fatty acid flux leads to increased acetyl CoA in mitochondria (239). Alternately, it is now suggested that free fatty acids might directly impair insulin signaling.

C. Insulin Resistance: On the Molecular Nature of the Causal Mechanism

The failure of the pancreas to adjust its insulin secretion according to the needs of the organism might lead to a long-term modification of the insulin-sensitive tissue responses and is thought to be one causal mechanism. In particular, the MODY subset of diabetes is connected to this concept. All but one of the factors involved in MODY are transcription factors acting on insulin synthesis and secretion pathways, as has been described in section 1B and Figure 3. However, these monogenic determinants of diabetes do not concern the vast majority of T2DM associated with obesity. Failure of the pancreas may also arise from exposure to high levels of free fatty acids. The actions of an excess of free fatty acids on pancreas function are complex, ranging from regulatory events to lipo toxicity. An acute increase in free fatty acid levels provokes an elevated glucose-stimulated insulin secretion. In obese T2DM patients, insulin secretion is initially high and parallels the increased demand of tissues for an insulin response. Pancreas failure could occur at a later stage, due to exhaustion of the gland. Alternatively, it was shown that chronic high levels of free fatty acids in obese patients inhibit insulin secretion and participate in accumulation in the β-cells of long-chain acyl CoA in the presence of glucose (249, 301). In the model of lipo toxicity, the accumulation of acyl CoA not only alters insulin secretion but also increases β-cell apoptosis, leading to pancreas failure (reviewed in Refs. 289, 302). However, results in experimental models seem to differ between in vitro and in vivo experiments and appear to depend on the species analyzed (rat, mouse, or human islets). The effects also vary depending on whether basal insulin secretion or high glucose-stimulated insulin secretion is being analyzed (reviewed in Ref. 171). Not surprisingly, the role of transcription factors such as SREBPs and PPARs in the accumulation and metabolism of lipids in the pancreatic β-cells and their effects on insulin secretion are presently being analyzed (67, 150, 254, 311), but no working hypothesis can yet be clearly defined. In any case, the pancreas, whose alteration is a critical feature of the progressing disease, does not seem to be the master player in the onset of the T2DM.

At the level of the peripheral cells, mutations in the insulin receptor (IR) gene or in the genes encoding key insulin signaling molecules, such as the transducers IR substrate-1 (IRS1) and IR substrate-2 (IRS2), were examined as a straightforward explanation for insulin resistance. However, polymorphisms in IRS1 or IRS2 are not clearly associated with insulin resistance but may impair β-cell function and insulin secretion (234).

Thus a simple and single causal explanation has to be set aside, and research is now geared to understanding what, in the functioning of the main insulin-sensitive tissues and their cross-talk, initiates and worsens T2DM. Particular attention has been given to muscle, with the hypothesis that the primary defect could be in glucose uptake due to decreased levels of glucose transporters. This hypothesis could not be confirmed so far, even though a decreased expression of Glut4 in skeletal muscle cells may participate in the disorder (229). Along the same lines, PKC-α can be activated by fat metabolites in skeletal muscle where it impairs insulin signaling and glucose transport via decreased tyrosine phosphorylation of IRS-1. Genetic deletion of PKC-α prevents fat-induced defects in insulin signaling and glucose transport in skel-
et al muscle, pointing to the importance of this pathway in initiating insulin resistance (144).

There is also much interest in the hormone-like substances that might play an important role in the cross-talk between tissues and that have at one point or another been suggested as a possible starting point for the disease. Interestingly, among the molecules described, more than 14 hormone-like substances are peptides secreted by the adipose tissue, identifying it as a major sensor and potential regulator of metabolic status (79). As changing the diet and life habits that primarily affect the adipose mass is a successful method of preventing and improving insulin resistance, some of the adipokines probably participate in the disturbed intertissular communication seen in insulin resistance. The particular physiopathological context of lipodystrophic diabetes highlights the importance of adipokines. Lipoatrophy is characterized by a reduced body-fat mass responsible for a disorder in which insulin resistance and hyperglycemia are associated. In mouse models for lipodystrophy, transplantation of adipose tissue reverses the metabolic abnormalities associated with lipoatrophy (82). A similar benefit is obtained with leptin (217) as well as adiponectin (330). This clearly emphasizes the role of adipokines in regulating insulin sensitivity of peripheral tissues, independent of the ability of the organism to store fat. The nature and mechanisms of the regulatory events taking place in the responsive tissues have yet to be deciphered.

D. The Thiazolidinediones as a Tool for Understanding the Physiopathogeny of the Metabolic Syndrome

The discovery that the thiazolidinediones (TZDs), specific activators of PPARγ, modify the insulin-resistance status in humans represents one major step in the search for the molecular mechanisms of the metabolic syndrome. TZDs first decrease plasma-free fatty acid concentrations and fasting hyperglycemia via an insulin-sensitizing effect (137, 266). This class of compounds also has other beneficial effects on T2DM by lowering hypertension if present and improving the overall lipoprotein profile, thereby reducing the risk of atherosclerosis. These benefits, extended to reduced occurrences of heart attack, stroke, and premature death, are now clearly highlighted by the results of the major clinical trial PROActive (www.proactive-results.com). Not surprisingly, however, TZDs enhance adipocyte differentiation, likely explaining the increased weight gain often observed upon its use (84, 98). Other side effects include edema and an idiosyncratic acute hepatotoxicity that seems to be a specific effect of troglitazone, one of the first TZD compounds initially used in clinics. Although it is clear that TZDs are potent and high-affinity ligands for PPARγ, it should be kept in mind that not all TZD effects are mediated by PPAR activation.

1. Hypothesis for the mechanism involved in PPARγ-mediated improvement of insulin sensitivity

A reasonable hypothesis is that by increasing the ability of the adipose tissue to store the excess of circulating fat, PPARγ diverts the fat that has unduly accumulated in muscle, liver, and pancreas towards fat tissues. It may be particularly efficient as it refurbishes more particularly subcutaneous fat, which is itself more sensitive to insulin than visceral fat (95, 308). This is consistent with the effects of TZD treatment in patients, which does not further increase the deleterious adipogenesis in muscle or the liver, but rather diminishes it. Thus potentially deleterious effects of an increase of the general amount of fat are counterbalanced by a redistribution of these lipids, which prevents excessive fat accumulation in the peripheral organs, including visceral fat, improving their insulin sensitivity (reviewed in Ref. 159).

In addition to fat redistribution, TZDs may also trigger other mechanisms. A direct action of PPARγ activation on the production of adipokines has been investigated, and some data support the hypothesis that adiponectin might be a crucial component connecting PPARγ activation in the adipose tissue and the metabolic response of the peripheral organs (95). Leptin has also been often cited as a potential effector of TZDs. However, consistent clinical data are difficult to assemble, and rodent models have generated some controversies. PPARγ might also directly regulate tumor necrosis factor (TNF)-α expression. A global analysis by DNA microarrays of PPARγ target genes has suggested an interesting mechanism by which PPARγ activation inhibits NF-κB, thereby blocking the TNF-α-mediated inhibition of adipogenesis (260).

One difficulty is revealed by the fact that rodent models of lipodystrophy with associated insulin resistance also benefit from TZD treatment, suggesting the presence of an adipose tissue-independent mechanism. A direct action of TZD acting via PPARγ on the skeletal muscle appears difficult to reconcile with the rather low expression of PPARγ in this tissue. Two reports describe opposing results when TZDs are used in mice selectively lacking PPARγ in muscle, either with persistence of a TZD beneficial response (209) or a loss of TZD response (107) in mutant mice. While these discrepancies are likely due to small but significant differences in the experimental design and methodology, they underline the urgent need for further mechanistic explorations.

Indeed, to better understand these observations and their therapeutic outcome, there is strong hope in learning from selective deletions of PPARγ in various tissues (reviewed in Ref. 148). Muscle specific deletion of PPARγ is reported to increase adiposity and insulin resistance, suggesting an important intertissular cross-talk, for which the molecules involved remain to be identified (107, 209). Mice in which PPARγ has been specifically deleted in the
liver do not exhibit any phenotype unless they are challenged by an additional mutation with deleterious metabolic effects (ob/ob mice and lipodystrophic A-ZIP/F-1 mice; Refs. 23, 192). This situation likely reflects the very low expression of PPARγ in the normal liver, while it increases and plays a role upon steatosis, as it may occur in the metabolic syndrome (61, 187). Selective deletion of PPARγ in adipose tissue demonstrates its importance for mature adipocyte survival (121). With respect to the metabolic consequences of PPARγ-selective deletion in the adipose tissue, a first report showed that it causes insulin resistance in the adipose tissue and in the liver but not in muscle (103). With the use of a very similar mouse model, a second independent report suggested opposing results, with such mice being protected against high-fat diet-induced obesity and insulin resistance. While the muscle exhibits a diminished glucose uptake, the authors proposed that the protective effect may come from increased glucose utilization by the liver (130). Altogether, these results are difficult to reconcile, but they contribute towards a wealth of knowledge which with more time, experiments, and mouse models, will help in defining the molecular network that controls metabolic homeostasis.

2. The paradox of PPARγ and insulin resistance

One more level of complexity was revealed by the striking observation that PPARγ +/− heterozygous mice, rather than being prone to insulin resistance, are partially protected from high-fat diet-induced or monosodium glutamate-induced weight gain and insulin resistance, compared with their wild-type littermates (154, 199). Lower activation of PPARγ can also be obtained via the use of PPARγ antagonists. Few of these molecules have been identified: GW0072 is a partial agonist in transactivation assays, which inhibits adipocyte differentiation in cell culture (211); BADGE inhibits adipocyte differentiation in cell culture (322) but is highly cytotoxic; PD068235 also blocks adipocyte differentiation but does not revert the phenotype of terminally differentiated adipocytes (26); and LG100641 blocks adipocyte differentiation and stimulates glucose uptake in 3T3-L1 adipocytes (204). However, these inhibitors have not been tested in vivo to verify whether they may prevent obesity and reduce insulin resistance. An in vivo study in mice was performed using SR-202 [dimethyl α-(dimethoxyphosphinyl)-p-chlorobenzyl phosphate], a novel PPARγ-specific antagonist that has no PPARγ agonist activity and partially inhibits adipocyte differentiation induced either by thiazolidinediones or by the combination of dexamethasone, insulin, and IBMX. In vivo, this compound partially blocks PPARγ activity, decreases fat deposits, and increases insulin sensitivity, setting most of the metabolic parameters at the levels of those seen in PPARγ +/− mice (247).

An additional surprise is that when PPARγ activity is decreased via monoallelic disruption of PPARγ, exposure to the agonist TZD does not further enhance insulin sensitivity of the PPARγ +/− mice but has almost the opposite effect, restoring a “wild-type” like phenotype to these mice (198). In this action, TZD might appear as a corrector of insulin sensitivity: it does not affect insulin sensitivity in normal humans and animals, it restores normal insulin sensitivity in human and rodent models with insulin resistance, but it diminishes insulin sensitivity in abnormally sensitive rodents. We have seen above that a TZD treatment, via PPARγ activation, might act by redirecting the triglyceride load from skeletal muscle to adipose tissue, restoring muscle sensitivity to insulin, and thereby improving glucose disposal. PPARγ +/− mice have smaller adipocytes and a smaller fat deposit. Treating these mice with TZD would simply help to restore a “normal” fat mass and would thus place them equal to wild-type mice in the face of a metabolic challenge such as high-fat diet. In contrast, a PPARγ antagonist would inhibit triglyceride accumulation in fat tissue and, as in PPARγ +/+ animals, would not allow fat accumulation to occur in ectopic tissues such as muscle, thus preserving insulin sensitivity.

Altogether, these results and pending questions suggest that potent PPARγ agonists might not be the most appropriate treatment of obesity and T2DM. Modulators, which may share antagonist and/or partial agonist properties, might represent a better perspective than thiazolidinediones. However, whereas experiments in mice are promising, the demonstration that these PPARγ modulators can act similarly in adult and obese humans is still lacking.

3. A possible role of PPARα and PPARβ in the metabolic syndrome

Thanks to its consistent role in stimulating fatty acid oxidation, PPARα could also be an interesting target for controlling the insulin resistance that accompanies dyslipidemia. Whereas fibrates act on lipid profiles mainly through their interaction with PPARα, little evidence for a fibrate action on insulin sensitivity was initially provided. Studies aimed at exploring this possibility demonstrated that the synthetic PPARα ligand Wy-14643, along with fenofibrate and ciprofibrate, are able to improve insulin sensitivity in rodent models of insulin resistance (93, 332). A better consumption of fatty acids by the skeletal muscle, thereby decreasing the overall fat accumulation, is proposed as the main mechanism of PPARα-induced insulin sensitivity. However, in contradiction with the latter observation, PPARα null mice subjected to a high-fat diet or more simply to aging are protected from developing insulin resistance (94).

Notwithstanding these apparent contradictions, a large effort is presently being made by numerous phar-
maceutical companies to test dual agonist compounds, which associate PPARα and PPARγ ligand properties, as a means of controlling some undesired side effects of TZD, which include hypertriglyceridemia, weight gain, and edema. Ragaglitazar, one of the first of these compounds to be tested clinically, seems to be promising, although its development is hampered by the persistence of side effects already mentioned for TZD, e.g., peripheral edema and anemia (285).

A role played by PPARβ in lipid metabolism has been envisioned in the context of obesity, diabetes, and dyslipidemia, based on both the aforementioned effect of a PPARβ ligand on HDL-cholesterol levels in obese monkeys and the decreased adiposity of PPARβ −/− mice (8, 228). However, the tissue-specific gene deletion of PPARβ in the WAT does not alter the fat mass, suggesting a cell-nonautonomous action. It is now reported that PPARβ −/− mice challenged with a high-fat diet are prone to obesity. In addition, an artificially overexpressed and overactive PPARβ in mouse WAT increased fatty acid oxidation and energy dissipation, features that in turn reduce adiposity and improve the lipid profiles (313). This latter observation, while difficult to interpret with respect to physiology, is consistent with a proposed role of the latter observation, while difficult to interpret with respect to physiology, is consistent with a proposed role of PPARβ in muscle fatty acid oxidation (206; reviewed in Ref. 13). Altogether, these observations emphasize the difficulties that lie in addressing the complexity of physiopathology mechanisms that take place in metabolic diseases and the general caution that should be applied in interpreting in vivo observations. It also demonstrates that much work needs to be performed to clarify the benefit that we may gain from targeting PPARβ for diabetes or syndrome X therapeutic strategies.

VII. TRANSCRIPTION FACTORS AS TARGETS
FOR THERAPEUTIC APPROACHES
OF METABOLIC DISORDERS

In this section, we briefly present how the modulation of transcription factor activities might be used for the treatment of metabolic disorders. The example of nuclear receptors, already discussed above for PPARs, and which are a major focus of pharmaceutical companies at the present time, will be extended to some other members of this family. Other drugs that will be analyzed are those aimed at treating unrelated specific pathologies, but which have serious side effects in metabolic regulation. The glucocorticoid analogs and the protease inhibitors are among the compounds in this category.

A. How to Modulate Transcription Factor Activity

The activity of transcription factors is the result of a number of events on which therapeutic strategies may act. First, the level of expression of a factor might be modulated, often via a known regulator of that gene. For example, glucocorticoids can increase PPARα expression. Understanding the network created by nuclear receptors (see Fig. 11) and their reciprocal interactions on their own gene expression is clearly of importance when establishing the molecular mechanism of action of a ligand, with the prospect of predicting clinical effects. A second general strategy that is commonly used by cells to regulate their metabolic pattern is by acting on the nuclear translocation or exclusion of a transcription factor. This may occur either via dissociating or associating cytoplasmic complexes, or via operating posttranscriptional modifications, such as phosphorylation, which precludes or favors entry of the transcription factor into the nucleus. For example, cytoplasm retention of FOXOs via phosphorylation is suggested to be a mechanism of insulin-mediated gene repression. With respect to drug design strategy, it is somehow difficult to specifically modify the status of a factor of interest, using the physiological means by which protein modification is achieved, since the effect of one single signaling molecule is most often distributed to a full range of factors that affect various pathways. In a third mode of action, the activation of transcription factors in the nucleus relies either on interaction with a ligand, such as for the nuclear receptors, or upon posttranslational modification (phosphorylation, acetylation, or other modifications). Activation implies the formation of specific transcriptional complexes that may vary according to the nature and extent of the modification that occurs. This is particularly explored in the context of nuclear receptors, where the ultimate objective...
is to generate specific ligands that might trigger a conformation change specific enough to provide selectivity upon the transcriptional complex formed, allowing a specific pattern of gene activation.

B. Nuclear Receptors as Targets for New Therapeutic Approaches

The general interest of pharmaceutical companies in the nuclear receptor family stems from the “druggability” of nuclear receptors. They naturally bind and respond to small lipophilic molecules; the search is for both agonists and antagonists. The notion that modulators differentially affect transcriptionally active complexes, together with the pattern of expression of these receptors and of their coactivator and corepressors, make it possible to develop tissue-specific responses. In the next paragraph, we give a brief description of the present achievements and hopes with respect to nuclear receptor targeting in metabolic diseases, without mentioning PPAR ligands since this has been thoroughly discussed above.

RXR ligands have been considered as a therapeutic approach for metabolic perturbations and have been tested in mice. A heterodimer composed of RXR and a permissive partner such as PPAR, LXR, or FXR can be activated by a RXR ligand, independently of the presence or absence of a ligand for the partner receptor. Rexinoids are particularly attractive for their potential ability to trigger PPAR:RXR activation (220). Indeed, benefits such as potent antidiabetic effects have been reported (203). However, these are accompanied by a massive increase in circulating triglycerides (248), possibly due to the parallel activation of the LXR:RXR pathway. In addition, RXR agonists also interfere with thyroid hormone receptor signaling, provoking a profound hypothyroidism due to a decrease in TSH levels (174). As rexinoids also trigger the LXR:RXR and FXR:RXR pathways, their action on cholesterol metabolism has been evaluated. Oral treatment with rexinoids results in a very efficient inhibition of cholesterol uptake by the gut. This effect was reported to involve at least two mechanisms: 1) the activation of LXR: RXR in the intestine, causing an increased cholesterol efflux in the lumen, and 2) the activation of FXR:RXR in the liver, producing an inhibition of bile acid synthesis, thus reducing micelle formation and cholesterol uptake in the gut (245). In a different context in humans, oral bezafibrate (Targrettin capsules corresponding to the well-characterized rexinoid LG1069) has been used successfully since the 1980s for the treatment of refractory or persistent early-stage cutaneous T-cell lymphoma. It acts by inhibiting mitogen-induced interleukin-4 production by the peripheral blood cells and inducing apoptosis of malignant T cells. However, studies of phase 2 and 3 clinical trials of cutaneous T-cell lymphoma by bezafibrate reported high triglyceridemia, hypothyroidism, but also hypercholesterolemia (60). As these observations helped to illustrate, rexinoid effects might not be as specific as wished for, and therefore, the potential therapeutic use of rexinoids requires a careful evaluation. Alternatively, the search for and development of heterodimer-specific ligands for RXR might allow a better targeting of the pathways of interest (70, 197).

LXR is a metabolic sensor for cellular cholesterol and a major regulator of cholesterol homeostasis, designating it as a pertinent target for drug development (76). More particularly, and as discussed in previous sections, LXR agonists should raise HDL levels, increase reverse cholesterol transport, increase cholesterol elimination via bile acid synthesis, and decrease intestinal cholesterol uptake. Accordingly, the LXR ligand T0901317 increases HDL levels. However, consistent with the role of LXR in lipogenesis, this is accompanied by an increased hepatic production of triglycerides and elevated blood VLDL, together with an accumulation of liver triacylglycerol. The direct effect on glucose tolerance seems to be modest (39). Thus the dual action of LXR on cholesterol disposal and fatty acid synthesis makes the therapeutic potential of LXR agonists rather difficult to predict. The description of the anti-atherosclerotic activities of a new LXR ligand GW3965 suggests that specific modulators might be at hand (133). This clearly needs further evaluation.

FXR is a metabolic sensor for bile acids that mediates their effects on gene expression, particularly by creating a negative-feedback loop controlling their synthesis. Treatments aimed at decreasing the body content of cholesterol should thus antagonize FXR, allowing for an increased production of bile acids from cholesterol in excess. Amazingly, such molecules seem to have been used for ages in Indian traditional medicine. While the gum resin from the Commiphora mukul tree was shown to have hypolipidemic activities, it has now been established that its active phytosterol, the guggulsterone, is an effective FXR antagonist (for review, see Ref. 304). Clinical studies have reported a 10–20% reduction in triglyceride levels and a 20–30% reduction in cholesterol levels. Side effects seem to be mild and do not affect basic metabolic pathways. However, 20–30% of patients are nonresponsive to the drug. Finally, it is also important to note that gugulipid interacts with the pregnane X receptor (PXR), raising the concern of drug-drug interference via activation of PXR-mediated increase of the expression of drug metabolizing enzymes. Ironically, this rediscovery underlines the continuing importance of empirical approaches for therapeutic strategies.

In contrast, orphan receptors may represent interesting alternative therapeutic targets. Understanding of their main function is still lacking, often because of the lack of adequate tools for inhibiting or activating them. However, many of them share a ligand-binding domain structure

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that suggests the existence of natural and possibly of synthetic ligands that so far remain unidentified. With this in mind, orphan receptors can be considered as an almost untouched reservoir of innovation.

C. Adverse Effects of Drugs on Energy Metabolism

A number of drugs not initially aimed at treating any metabolic disorder cause severe problems because of important metabolic deregulations, some of which are mediated by transcription factor alterations.

Perhaps the best example is glucocorticoid analogs, mainly used as anti-inflammatory drugs. As already seen, glucocorticoids affect metabolic pathways, particularly those of insulin sensitivity, amino acid degradation, and muscle physiology. Not all mechanisms that cause such disturbances are known, although a direct action of the nuclear glucocorticoid receptor on some target genes is likely to be the main one. Glucocorticoid receptors may also act indirectly by increasing the expression of PPARs. Here again, identifying specific modulators that would provide a better focus for the therapeutic activity in a given tissue or on a given set of target genes represents a major challenge for the future.

An intriguing question concerns the metabolic consequences of the use of protease inhibitors to treat human immunodeficiency virus (HIV) infection. Dyslipidemia occurs in up to 70–80% of the treated patients, often with associated hypertriglyceridemia and hypercholesterolemia. A severe lipodystrophy is also found in many but not all occurrences. The mechanism of this deregulation remains hypothetical thus far. A first hint is the identity, ~60%, of the catalytic domain of the HIV protease with that of the cytoplasmic retinoic acid binding protein type 1. The hypothesis is that the protease inhibitor might erroneously bind to this protease-related domain, affecting the retinoic acid signaling pathway. It also might similarly affect the low-density lipoprotein receptor related protein (LRP), which also encompasses a domain sensitive to protease, resulting in disturbed lipoprotein metabolism. Such perturbations could explain both the alteration in lipid metabolism seen in nontreated HIV patients and the aggravation upon treatment with protease inhibitors (reviewed in Ref. 25). Alternatively, it has been suggested that protease inhibitors might affect the SREBP maturation process, resulting in impaired nuclear accumulation of SREBP-1 (27, 317). A better understanding of the molecular mechanisms at work would clearly be extremely helpful for designing the best therapeutic strategy.

VIII. CONCLUSIONS

A conclusion for such a long story must be short. Thus let us highlight only one important feature of the transcriptional regulation of metabolism. Most of the observations gathered for one pathway and its regulation via a given transcription factor show that this regulation not only targets the expression of rate-limiting enzymes but also coordinately regulates the expression of the enzymes working at many intermediary steps of the pathway. The initial concept of regulation operating at the level of the rate-limiting step is losing importance in favor of coordinated regulation. This feature must now be put in parallel with what should also strike us from the present review, i.e., the intense networking between transcription factors by which homeostasis can be achieved (see Fig. 11). Thus, in addition to classical approaches, metabolic explorations must be included in global approaches, such as analyses of the transcriptome, proteome, and metabolome. However, even more than for classical approaches, such experiments must be designed to answer well-defined and physiologically relevant questions. While the first reports on complex metabolic networks drawn from bacteria models are beginning to appear, the realization of such a vision of mammalian metabolic regulation still seems to be some years away.

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APPENDIX

Appendix A: The Nuclear Receptor Family

Nuclear receptors are transcription factors that are characterized by two important properties: 1) they are activated upon binding of specific ligands, and 2) they bind to specific response elements located in the vicinity of the promoter of their target genes. Thus, in a simplified view, the effector function of nuclear receptors in a cell is to adapt the gene expression program according to signals received as specific ligands. An official nomenclature of these receptors across species is now used, organized according to their phylogeny (210; reviewed in Ref. 3).
1. Nuclear receptors share a common structural and functional organization (Fig. 1A)

The poorly structured NH₂-terminal domain encompasses a ligand-independent transactivation domain in some of the receptors. The DNA binding domain (DBD) with its two zinc fingers is the hallmark of the nuclear receptor family. The hinge region links the DNA binding domain to the ligand binding domain (LBD). The general fold of the ligand binding domain is structured by 12 α-helices and 3 β-sheets defining the ligand binding pocket (see Fig. 1A, top left).

In general, the nuclear receptors bind to DNA in the form of dimers, either homodimers, such as HNF4α or the estrogen receptor, but more often as heterodimers with the receptor for 9-cis-retinoic acid known as RXR (i.e., NR2B according to the unified nomenclature of nuclear hormone receptors; Ref. 210). The DNA response element of nuclear receptors is comprised of two sequence motifs corresponding or closely related to the hexamer AGGTCA. The organization of these two motifs in direct repeats (DR) or palindromic arrays and the length of the spacing between the two hexamers determine the specificity of these response elements towards each receptor dimer (Fig. 1A, top right).

The general scheme for transactivation via nuclear receptors is thought to occur in at least two steps. In the absence of ligands, nuclear receptor dimers may bind a corepressor protein that inhibits their transactivation properties. In the presence of ligands, or due to an alternative pathway of activation such as phosphorylation, the corepressor is released and a coactivator is recruited, allowing further interactions with the transcription initiation complex (TIC), eventually resulting in transcription enhancement. Corepressors and coactivators work at least in part by modulating the chromatin status via histone deacetylation and acetylation, respectively. Other modifications such as histone methylation have also been shown (Fig. 1A, bottom).

2. The three functional classes in the nuclear receptor family (Fig. 1B)

Analyses of the human genome have identified 48 nuclear receptor genes, most of them generating more than one receptor isoform. Based on their ligand binding properties, nuclear receptors can be ordered into the three following classes.

The classic hormone receptors bind molecules such as glucocorticoids, thyroid hormone, retinoic acid, and estrogen, with a high affinity. As key factors of endocrine homeostasis, their activation is associated with many metabolic adjustments (Fig. 1B, left).

The orphan receptors possess the structural characteristics of nuclear receptors including a sequence consistent with the presence of a ligand binding domain. However, no ligand has thus far been identified for these receptors. In addition, a tight structure of the 12 helices has been shown in the Nurr1 receptor to preclude the formation of a ligand binding pocket (315). The functions of many orphan receptors remain elusive (Fig. 1B, right).

The receptors of the third class are metabolic sensors. This class comprises receptors that bind to a broad range of molecules with, as a corollary, a relatively poor affinity. Rather than responding to hormones secreted by endocrine glands with tight feedback controls, these receptors can bind to molecules that are components of metabolic pathways as substrates, intermediates, or end-products, such as fatty acids, eicosanoids, and oxysterols. These receptors are sensors of the metabolic status, respond to both incoming dietary signals and metabolites generated in the organism, and are responsible for the metabolic adaptation at the cell, organ, and whole organism level. In this class are PPARs and HNF4α, which play a major role in energy metabolism, as well as LXR and FXR, which are closely involved in cholesterol metabolism, together with their common partner RXR. RXR itself is difficult to assign to a distinct class, as it behaves as a classic receptor with respect to 9-cis-retinoic acid. However, the nature of its endogenous ligand is still unclear, and the functions of RXR rather plead for its belonging to the sensor receptor class (see also Appendix F). HNF4 is accompanied by * since its mode of interaction with ligands is disputed (see also Appendix E) (Fig. 1B, middle).

Appendix B: Peroxisome Proliferator Activated Receptors

PPARs were the first nuclear receptors identified as “sensors” rather than classic hormone receptors. They are nuclear, lipid-activatable molecules that control a variety of genes in several pathways of lipid metabolism (reviewed in Ref. 56).

Three isotypes of PPAR, PPARα (NR1C1), PPARβ (NR1C2) (also called PPARγ, NU1, and FAAR), and PPARγ (NR1C3), have been cloned in Xenopus, rodents, and humans. Two PPARγ isomers, PPARγ1 and PPARγ2, are splice variants in their NH₂-terminal domain.

PPARα is highly expressed in tissue with high activity levels of lipid catabolism, e.g., liver, brown adipose tissue, and skeletal and heart muscle. PPARβ is ubiquitously expressed. PPARγ1 is mainly expressed in adipose tissues but is also detected in the colon, spleen, retina, hematopoietic cells, and skeletal muscle. PPARγ2 has been found mainly in the brown and white adipose tissue.

Their modular structure is that of all nuclear receptors. The less conserved NH₂-terminal region bears a ligand-independent activation domain, at least in PPARα and PPARγ. The DNA binding domain is extremely well conserved. The ligand binding pocket of PPARs is much larger than that of the other nuclear receptors and relatively easily accessible (324 and references therein).

PPARs bind to DNA as heterodimers with RXR, on PPAR response elements (PPRE) comprising direct repeats of two hexamers closely related to the sequence AGGTCA and separated by one nucleotide (DR-1 sequence). The five nucleotides that flank the 5'-end of this core sequence are also important for the efficiency of PPARα:RXR binding.

The first molecules to be recognized as PPARα activators, and later characterized as ligands, belong to a group of molecules that induce peroxisome proliferation in rodents, thus explaining the name of peroxisome proliferator activated receptor given to this receptor. This diverse group of substances includes, for example, some plasticizers and herbicides. More interestingly, various fatty acids, more particularly unsaturated fatty acids, and some eicosanoids mainly derived from arachi-

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Acid storage in the adipose tissue, via PPAR/HR9253 rodents and human.
The ligands. The most potent of these are 22-nucleotides.
related to the sequence AGTTCA, separated by four response element, formed from a direct repeat of two hex-
Expressed ubiquitously.
The heterodimer can also be triggered by RXR ligands.
Role of PPAR

Liver X Receptors

The liver X receptors (LXR; NR1H3) are members of the nuclear receptor family. Their endogenous activators are oxysterols and other derivatives of cholesterol metabolism. As such, they participate in the cholesterol sensing processes and regulate important aspects of cholesterol and fatty acid metabolism.

Two isotypes, LXRα and LXRβ, share 77% amino acid identity in their DBD and LBD and are highly conserved between rodents and human.

LXRα is highly expressed in the liver but is also found in kidney, intestine, adipose tissue, and macrophages. LXRβ is expressed ubiquitously.

LXRs heterodimerize with RXR to bind to their DNA response element, formed from a direct repeat of two hexamers related to the sequence AGTTCA, separated by four nucleotides.

Mono-oxidized derivatives of cholesterol are potent LXR ligands. The most potent of these are 22(R)-hydroxycholesterol, 24(S)-hydroxycholesterol, and 24(S),25-epoxycholesterol, which activate both LXRα and LXRβ. Little is known about the steroid hydroxylases that produce these metabolites, but it is assumed that oxysterol concentrations parallel those of cholesterol. Importantly, oxysterols are found at micromolar concentrations in tissues that express high levels of LXRα or LXRβ. Activation of the heterodimer can also be triggered by RXR ligands.

LXRα and LXRβ null mutant mice have been generated and confirm the important role of these receptors, and more particularly that of LXRs in cholesterol homeostasis. These models also pinpoint the role of LXR in fatty acid metabolism.

Farnesol X Receptor

The farnesol X receptor (FXR, NR1H4) is a member of the nuclear receptor family, which acts as a bile acid sensor and is involved in a negative-feedback regulation to control and moderate excess of bile acid production (reviewed in Ref. 62).

FXR was initially cloned as a RXR interacting protein called RIP14.

FXR expression is restricted to adrenal cortex, intestine, colon, kidney, and liver.

FXR forms heterodimers with RXR to bind to DNA response elements consisting of two hexamers (GGGTCA), spaced by one nucleotide, organized in a palindromic configuration (also called inverted repeat-1).

Initially proposed as a receptor of farnesol metabolites, and thus renamed FXR from RIP14, it was shown to bind and be activated by physiological concentrations of free and conjugated bile acid products: chenodeoxycholic acid, lithocholic acid, and deoxycholic acid. These bile acids are the end products of the neutral and acidic bile acid biosynthetic pathway. The heterodimer FXR:RXR can also be activated by RXR ligands.

FXR acts as a bile acid sensor that regulates bile acid metabolism. Because bile acids are an important means for disposal of cholesterol in excess, FXR thereby affects cholesterol metabolism.

Hepatic Nuclear Factor 4

Hepatocyte nuclear factor 4α (HNF-4α, NR2A1) is a highly conserved member of the nuclear receptor family.

HNF-4α was first identified as a member of a heterogeneous family of transcription factors named the liver-enriched nuclear receptor factor family. At least nine splice variants have been described.

HNF4α is mainly expressed in the liver, intestine, and pancreatic β-cells.

HNF4α mainly binds as homodimer to response elements closely related to DR1 elements, thus resembling PPREs (see Appendix B).

The question of the status of HNF4α as orphan or liganded receptor is still unsolved (286). While a report indicated that fatty acyl CoA thioesters could bind to HNF4α and modulate its activity (105), these compounds were not able to alter HNF4α interaction with cofactors (16). Its transactivation properties, readily present when assessed in transfection assays, also support the proposition that it acts as a constitutively active orphan receptor. These observations may be reconciled by the crystallographic structure of the ligand binding domain, which showed that saturated or monounsaturated C14-C18 fatty acids not only fill the ligand binding pocket but appear to be locked in the protein, suggesting that it plays a structural role rather than acts as an activating signal (320).

Possibilities other than ligand binding for modulating HNF4 activity have thus been proposed. Alternative splicing of the mRNA, phosphorylation of the protein (13 phosphorylation sites exist), and/or inhibitory interaction with phosphorylated FOXOs (108) are among the numerous possible ways of regulating HNF4α activity.

HNF4α affects the expression of genes involved in glucose, fatty acid, and cholesterol metabolisms.
Appendix F: Retinoid X Receptor

Except for HNF4α, the nuclear receptors of the metabolic sensor class are active as heterodimers with RXR. The important part that RXR may play is further emphasized by the fact that RXR is itself a nuclear receptor that can be activated by specific ligands. PPAR-RXR, LXR-RXR, and FXR-RXR are permissive heterodimers within which RXR can bind its own ligand, in the absence of a ligand for its partner, and can thereby activate the transcription of the heterodimer target genes.

There are three isotypes of RXR, α, β, and γ, and several isoforms for each of them (reviewed in Ref. 30).

Each isotype and isoform has its specific expression pattern. However, any single tissue contains one or several forms of RXR.

In addition to the various heterodimers for which RXR is an obligatory partner, RXR can form homodimers that bind to DB1 elements (see also Appendix B). The in vivo relevance of these homodimers is still under study.

RXR can be activated by 9-cis-retinoic acid, an isomer of all-trans-retinoic acid. While the occurrence of this molecule in vivo has been questioned, the identification of two enzymes that participate in the isomerization of all-trans-retinoic acid to form 9-cis-retinoic acid lends support for its relevance in the whole organism. However, some doubts linger about the nature of the major natural RXR ligand. An oleic acid molecule was found in the RXR ligand binding pocket in a crystal of RAR:RXR ligand binding domains (17). Whether this indicates that fatty acids might be RXR ligands remains to be confirmed. In support of this notion, RXR is activated by the long-chain polyunsaturated fatty acid docohexaenoic acid in the adult mouse brain (51).

RXRa null embryos die in utero, whereas the RXRγ and RXRβ null animals do not display any severe phenotype (188, 316). To overcome RXRa embryonic lethality, a mouse allowing tissue-specific invalidation of the RXRa gene has been generated. Deletion of the RXRa gene in the liver allowed the identification of the most affected pathways (309). As expected, many PPARα-mediated functions were altered, and the activity of LXR and FXR were also compromised, suggesting that the absence of RXRa cannot be compensated by RXRβ and RXRγ in the liver.

Appendix G: Sterol Regulatory Element Binding Proteins

The sterol regulatory element binding proteins (SREBPs) are transcription factors of the helix-loop-helix family. Starting out wrapped in the ER membrane, their activation as transcription factors requires a maturation process tightly controlled by the levels of cholesterol present in the membrane (see Fig. 2). They are thus important regulators of cholesterol metabolism, most notably with respect to cholesterol biosynthesis, but also have an important role in fatty acid metabolism (reviewed in Refs. 63, 114, 218).

SREBPs are encoded by two genes, SREBP-1 and SREBP-2. Alternative promoter usage and alternative splicing of SREBP-1 drive the production of two isoforms, SREBP-1a and SREBP-1c. The 29 additional amino acids present in the SREBP-1a NH2 terminus are enriched in acidic residues and might be responsible for the higher transcriptional activity of SREBP-1a compared with that of SREBP-1c. SREBP-1c was initially cloned in rats and called adipocyte determination and differentiation factor-1 (ADD1) (reviewed in Ref. 256).

SREBP-1a has been mainly studied in cell lines, where it is the major form produced, while its expression in animal tissues is relatively weak. In contrast, SREBP-1c is highly expressed in liver and WAT. SREBP-2, also expressed in cell lines, is also predominantly present in the liver and adipose tissue but has a rather weak expression in animal tissues overall.

The three proteins SREBP-1a, SREBP-1c, and SREBP-2 are synthesized as precursors of ~125 kDa, entangled in the ER membrane. Maturation of these proteins requires the activation of the SCAP, a chaperone protein for SREBPs. SCAP acts as a sensor of the content of cholesterol in the ER membrane. In the presence of high levels of cholesterol, SCAP remains anchored in the ER membrane due to its interaction with the polytopic INSIG proteins.

Low membrane cholesterol levels lead to the transport of SCAP/SREBP to the Golgi membrane where activation of the site 1 serine protease results in a first cleavage. A second enzyme, the site 2 metalloproteinase, completes the maturation of SREBPs and releases the 68-kDa NH2-terminal domain of SREBP from the membrane (reviewed in Ref. 63). This fragment contains a basic helix loop helix (HLH) leucine zipper domain, which functions as a transcription factor upon translocation into the nucleus (see Fig. 7).

The mature forms of SREBPs bind to elements initially characterized as featuring an enhancer sequence called E-box that is recognized by members of the HLH transcription factor family. SREBPs also bind to sites related to the direct repeat TCANCCAC. However, it remains difficult to define a consensus sterol response element (SRE). SREBP-1 and SREBP-2 have preferential target genes, which may be determined, at least in part, by interaction of the mature forms with different cofactors. However, this remains to be demonstrated.

Germline deletion of SREBP-1 (eliminating both SREBP-1a and SREBP-1c) or SREBP-2 leads to partial or fully penetrant embryonic lethality, respectively. In contrast, specific deletion of the SREBP-1c transcript is not lethal, suggesting an important role of SREBP-1a and SREBP-2 in embryonic development.

The SREBP maturation process via membrane cholesterol sensing is consistent with their important role in cholesterol homeostasis. However, and in a schematic view, SREBP-2 is indeed mainly involved in cholesterol metabolism, whereas SREBP-1c has an important implication in fatty acid synthesis (see also Fig. 8). The range of target genes activated by SREBP-1a is wider, encompassing many key genes for both cholesterol and fatty acid synthesis.

Appendix H: The Liver-Enriched Transcription Factors

The class of liver-enriched transcription factors regroups very heterogeneous factors initially described for their role in liver development and hepatic cellular functions. However, their expression and subsequent actions are often not restricted to the liver (reviewed in Ref. 271). Among these factors, two main subgroups are involved in metabolic regulations, which are briefly presented below.
1. Hepatocyte nuclear factors

Hepatocyte nuclear factors form a heterogeneous subgroup of transcription factors, each of them being evolutionarily conserved.

HNF4α belongs to the family of the hormone nuclear receptors and is described in Appendix E.

HNF1α (or LF-B1) and the related vHNF1 (HNF1β or LF-B3) contain a homeodomain motif through which they bind to DNA. HNF1α is crucial for the liver-specific expression of albumin and has strong implications in normal pancreatic β-cell functions (233).

HNF3α, -β, and -γ are members of the forkhead family of transcription factors (263), therefore also called FOXA1, -2, and -3, respectively. They are identified by a DNA binding domain structured as a winged helix motif (i.e., an helix-turn-helix motif with an adjacent additional DNA interaction region). Whereas their roles have mainly been explored in the context of development, these factors are also involved in signaling cascades in pancreatic β cells, particularly affecting the regulation of expression of the insulin gene.

2. CCAAT/enhancer-binding proteins

C/EBPs are members of the basic leucine zipper family (bZIP) and were predicted to play a major role in energy metabolism (reviewed in Refs. 238, 253).

Six genes coding for the various isotypes of C/EBP have been identified (renamed α, β, γ, δ, ε, and ζ, according to a systematic nomenclature), some of them coding for several isoforms. C/EBPα, -β, and -δ are the isotypes involved in the transcriptional control of metabolic pathways.

C/EBPα, -β, and -δ are highly expressed in liver, adipose tissues, lung, and intestine.

C/EBPs bind DNA via a basic amino acid-rich domain, adjacent to a leucine zipper motif responsible for interaction with the homologous domain of the dimerization partner. The high conservation of this bZIP domain among C/EBPs results in an apparent lack of discrimination between the different isoforms for interaction with C/EBP DNA binding sites and dimerization between the different isoforms.

The transactivation function of C/EBPs, which are considered as positive constitutive transcriptional regulators, is localized in their NH₂ terminus. C/EBPγ, which lacks the transactivation domain, and C/EBPζ, which has impaired DNA binding properties, can form inactive heterodimers with other isoforms and thus have a dominant negative activity with respect to the α-, β-, δ-, and ε-isoforms. However, the assessment of the physiological importance of these two negative forms needs further investigation.

C/EBPs and more particularly C/EBPα affect the three main branches of the intermediary metabolism, i.e., glucose, amino acid, and lipid metabolisms.

Appendix I: Insulin Resistance: Definition and Characteristics

Diabetes is considered as a major world-wide health problem. Type 1 diabetes is due to the absolute lack of insulin, most often linked to a specific destruction of the β-cells of the pancreatic islets, specialized in insulin production and secretion. They represent 5–10% of all cases. Type 2 diabetes is far more common, and it is estimated that ~25% of the adult citizens in western countries are affected by this syndrome. Several terms are currently used to describe this pathology, and efforts have been made to unify the definition and characterization of the various aspects of the disease. The World Health Organization (WHO) has proposed a series of criteria, which is not easy to evaluate in clinical practice. The National Cholesterol Education Program Expert Panel (NCEP) proposed a more practical approach but does not define the insulin resistance status. Below are some of the definitions proposed by the European group for the study of insulin resistance (EGIR) and the WHO. However, these definitions are still being tested for their relevance in epidemiological studies (6, 155).

Impaired glucose tolerance is assessed by the glucose tolerance test, during which a charge of glucose (75 g) is given orally to a patient and glycemia is followed up to 2 h after the initial charge. A persistent high level of blood glucose (≥7.8 mM) 2 h after glucose intake is a criterion for impaired glucose tolerance.

The diabetes status corresponds to chronic hyperglycemia, with high fasting plasma glucose levels (≥6.1 mM) and impaired glucose tolerance with a glycemia 2 h after glucose intake that is ≥10 mM.

Insulin resistance syndrome (according to EGIR) is defined as hyperinsulinemia associated with two or more of the other components: hyperglycemia, hypertension, dyslipidemia, and central obesity. Insulin resistance is best assessed by euglycemic hyperinsulinemic clamp, which consists of a constant infusion of insulin with a parallel infusion of glucose to ensure euglycemia. The amount of glucose needed to maintain euglycemia under the controlled hyperinsulinemia is used to evaluate the status of insulin sensitivity of the patient. Alternatively, for large epidemiological studies and for practical reasons, EGIR proposes to use hyperinsulinemia, defined as fasting insulin levels in the upper 25%, as a surrogate marker of insulin resistance.

The metabolic syndrome (according to WHO) associates at least insulin resistance and/or impaired glucose tolerance with two or more of the other components: insulin resistance, impaired glucose regulation, hypertension, dyslipidemia (high triglycerides and low HDL cholesterol), central obesity, and microalbuminuria.

REFERENCES

TRANSCRIPTIONAL REGULATION OF METABOLISM


32. Christoffers VM, Grange T, Kaestner KH, Cole TJ, Darlington CJ, Groniger CM, and Lamers WH. Glucocorticoid receptor, C/EBP, HNF3, and protein kinase A coordinately activate the glu-


Horowitz JF, Leone TC, Feng W, Kelly DP, and Klein S. Effect of endurance training on lipid metabolism in women: a potential...


