

# Trabajo Fin de Máster

Función de los microARNs del tejido adiposo subcutáneo en la regulación de la esteatosis hepática. Role of microRNAs from subcutaneous adipose tissue in the regulation of hepatic steatosis.

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## Abstract 📃

Background: Hepatic steatosis arises from excessive triglyceride buildup in hepatocytes and correlates with the progression of metabolic dysfunction-associated steatotic liver disease (MASLD). Molecules secreted by subcutaneous white adipose tissue (scWAT), such as microRNAs (miRNAs), appear to influence MASLD by regulating key molecular pathways. This study aimed to identify miRNA biomarkers linked to steatosis in the scWAT of individuals with obesity and varying degrees of hepatic steatosis.

*Methods*: Subcutaneous white adipose tissue (scWAT) samples were collected from 78 individuals with obesity undergoing bariatric surgery. The patients were categorized into four steatosis groups based on liver fat content: "<5%", "5–33%", "33–66%", and ">66%". Small RNA sequencing data from scWAT were processed with the *nf-core/smrnaseq pipeline* (v2.4.0), and gene expression analysis utilized the *DESeq2* and *isomiRs* R packages.

Results: A total of 374 miRNAs were found to be differentially expressed between obese patients with varying degrees of hepatic steatosis. Among these, two miRNAs (hsa-miR-372-3p and hsa-miR-144-3p) remained significant after multiple test correction. Bioinformatic analysis of predicted target genes revealed their involvement in pathways related to hepatocellular carcinoma and hepatitis B.

Conclusions: The differential expression of hsa-miR-372-3p and hsa-miR-144-3p in scWAT suggests that these miRNAs may play a role in the regulation of hepatic steatosis and the progression of MASLD. Their association with hepatocellular carcinoma and hepatitis B pathways underscores their potential relevance to liver pathology. These findings open avenues for research on miRNA profiles as biomarkers or therapeutic targets for MASLD, with a focus on distinguishing simple steatosis from advanced disease.

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## Chapter 1

### **Introduction and Antecedents**

# 1.1 Overview of Hepatic Steatosis and Metabolic dysfunction- associated steatotic liver disease

Hepatic steatosis, defined by the excessive accumulation of triglycerides within hepatocytes, arises from a range of factors<sup>[1]</sup>. These include drug-induced effects, such as those caused by steroids and chemotherapy, infections like hepatitis C virus, and both nutritional and metabolic causes<sup>[2]</sup>. This condition is closely linked to steatotic liver disease (SLD), which encompasses two primary metabolic categories: alcoholic-related liver disease (ALD) and metabolic dysfunction-associated steatotic liver disease (MASLD)<sup>[3]</sup>.

MASLD and its more advanced form, metabolic dysfunction-associated steatohepatitis (MASH), were previously known as nonalcoholic fatty liver disease (NAFLD) and nonalcoholic steatohepatitis (NASH), respectively<sup>[4]</sup>. MASLD is diagnosed in adults who exhibit hepatic steatosis, identified through imaging techniques, blood biomarkers, or liver histology, in conjunction with being overweight or obese<sup>[5]</sup>, or in the presence of type 2 diabetes mellitus (T2DM) or at least two metabolic risk abnormalities (hypertension, hyperlipidaemia or insulin resistance)<sup>[6,7]</sup>. Although in many cases it may be asymptomatic in its early stages, this condition can progress to more severe stage pending on its cause<sup>[8]</sup>. It encompasses a spectrum of liver pathologies ranging from simple hepatic steatosis to metabolic dysfunction-associated steatohepatitis (MASH), which is marked by lobular inflammation and hepatocellular ballooning, potentially progressing to fibrosis and cirrhosis, ultimately leading to liver failure<sup>[9]</sup>.

The prevalence of MASLD is affecting more than a third of the adult population worldwide, making it the most common chronic liver disease globally<sup>[10]</sup>. Among adults, the prevalence (MASLD) is approximately 3 [22] [11]. MASLD is particularly prevalent in overweight or obese individuals, with a global prevalence of approximately 50%, rising to nearly 60% is a dividuals with type 2 diabetes (T2D), a condition that affects up to 10% of the world's adult population<sup>[12]</sup>.

Over the last 30 years, the global prevalence of MASLD has experienced significant growth, rising from 17.6% in 1990 to 23.4% in 2019, reflecting an average annual increase of approximately 1.0%. As of 2019, it was estimated that there were around 1.66 billion prevalent cases of MASLD worldwide<sup>[13]</sup>. This condition is widespread, with the highest rates reported in South America (44.4%), Middle East East and North Africa (36.5%) and followed by South Asia (33.8%), South-East Asia (33.1%), North America and Australia (31.2%), East Asia (29.7%), Asia Pacific regions (28.0%)

and Western Europe  $(25.1\%)^{[11,14]}$ . Approximately one-quarter of the European population is affected by this liver disease<sup>[7]</sup>. In Europe, the prevalence of NAFLD variates between countries ranging from 5% to  $44\%^{[15]}$ . Data from Spain reflect similar rates, indicating a NAFLD prevalence of 25.8% in the adult population<sup>[16]</sup>.

This growing prevalence of MASLD is closely intertwined with the global rise in obesity and metabolic syndrome, conditions largely influenced by the complex interplay of adipose tissue function and dysfunction. Understanding the role of adipose tissue in metabolic regulation is therefore critical for elucidating the pathophysiology of MASLD.

#### 1.2 Role of Adipose Tissue in Metabolic Regulation

#### 1.2.1 Metabolic and Endocrine Functions of Adipose Tissue

Adipose tissue (AT) is a complex and dynamic organ with both metabolic and endocrine functions<sup>[17]</sup>. It plays a pivotal role in energy balance, insulin sensitivity, immune responses and overall health<sup>[18]</sup>. Its role extends beyond simple fat storage, influencing whole-body physiology and contributing to various pathologies, notably obesity and its associated complications like MASLD<sup>[19]</sup>.

#### 1.2.2 Types and Location of Adipose Tissue

Adipose tissue in mammals exists in three primary forms: white adipose tissue (WAT), brown adipose tissue (BAT), and beige or brite (brown-in-white) adipose tissue. Each type is distinguished by its unique functions and cellular composition<sup>[20]</sup>. WAT primarily stores energy in the form of lipids, while BAT specializes in heat production through the uncoupling of oxidative phosphorylation, a process critical for thermogenesis<sup>[21]</sup>. Beige AT, a metabolically flexible tissue, can transition between energy storage and thermogenesis depending on physiological needs, highlighting its role in adaptive responses<sup>[22]</sup>.

The anatomical distribution of WAT further under cores its functional diversity. WAT is primarily located in subcutaneous depots, beneath the skir, and visceral depots, surrounding internal organs such as the liver and intestines<sup>[23]</sup>. Additionally, smaller WAT depots are present in areas like bone marrow and muscle tissue, contributing to localized metabolic regulation<sup>[21]</sup>.

White adipose tissue (WAT) is classified into two major subtypes based on Tocation and function: subcutaneous white adipose tissue (scWAT), which constitutes over 80% of total body fat and serves as the primary site for long-term energy storage, with its distribution and metabolic activity linked to protective effects on overall metabolic health; and visceral white adipose tissue (visWAT), which accounts for 10–20% of total body fat in men and 5–10% in women, is more metabolically active, and is strongly associated with adverse health outcomes such as insulin resistance and inflammation<sup>[24,25]</sup>.

#### 1.3 Subcutaneous White Adipose Tissue (scWAT)

Subcutaneous White Adipose Tissue (scWAT) is the most abundant type of adipose tissue, found in various locations such as under the skin and in clustered regions of the body, including the upper (deep and shallow abdomen) and lower (gluteofemoral) body areas<sup>[26]</sup>.

Under physiological conditions, scWAT serves as a metabolically inert, long-term triglyceride storage site<sup>[27]</sup>. Acting as a buffer for excess energy, scWAT protects other organs from ectopic lipid deposition and contributes to specific metabolic benefits<sup>[20]</sup>. This buffering role allows scWAT to mitigate the impact of excess dietary lipid consumption while also compensating for energy deficits during fasting, starvation, or strenuous exercise<sup>[27]</sup>.

Structurally, Sc WAT is composed predominantly of adipocytes, which constitute approximately 50% of its cellular content<sup>[28]</sup>. In addition to adipocytes, scWAT includes vascular cells, fibroblasts, adipocyte precursors, multipotent mesenchymal stem-like cells, nerve processes, and immune cells such as macrophages, lymphocytes, eosinophils, and mast cells. These components are embedded within an extracellular matrix that provides structural support<sup>[28]</sup>. Collectively, these cells secrete a wide array of signaling molecules, including adipokines such as leptin, adiponectin, and resistin, which play essential roles in maintaining metabolic homeostasis<sup>[22]</sup>

#### 1.4 Metabolic Functions of WAT

The main role of white adipose tissue (WAT) is to manage energy balance by storing and releasing fatty acids (FAs) according to variations in energy availability<sup>[22]</sup>. Furthermore, WAT produces a range of hormones and cytokines, collectively referred to as adipokines, which are crucial for regulating numerous physiological functions<sup>[29]</sup>.

#### 1.4.1 Lipids Storage and Mobilization

WAT maintains energy balance by storing and releasing fatty acids (FAs), a process controlled by the interplay between lipogenesis and lipolysis. This equilibrium is essential for sustaining energy homeostasis during periods of fasting or exercise<sup>[22]</sup>.

Lipogenesis is the process of synthesizing new lipids from excess glucose or dietary fatty acids. This process occurs in the cytoplasm of adipocytes and is tightly regulated by various hormones and enzymes<sup>[30]</sup>. Insulin, the key hormone involved in this process, facilitates the uptake of glucose and fatty acids into adipocytes and activates essential enzymes for lipid synthesis, including acetyl CoA-carboxylase (ACC) and fatty acid synthase (FAS)<sup>[31]</sup>.

Lipolysis, on the other hand, is responsible for breaking down stored lipids in adipose tissue to release energy for peripheral organs. This process is particularly important during fasting or exercise when glucose levels are low, prompting the body to utilize stored fat for energy<sup>[32]</sup>. Lipolysis is regulated by lipases, which are activated by signals from the sympathetic nervous system, primarily mediated by norepinephrine, with some influence from epinephrine. The primary lipases involved include adipose triglyceride lipase (ATGL), hormone-sensitive lipase (HSL), and monoacylglycerol lipase (MGL)<sup>[33]</sup>.

#### 1.4.2 Endocrin Inction

Additionally, WAT secretes various hormones and cytokines, collectively known as adipokines, which play essential roles in regulating multiple physiological processes such metabolic, inflammatory, and immune processes throughout the body<sup>[29]</sup>. The endocrine function of WAT is influenced by nutritional status, physical activity, hormonal levels, and environmental signals, and is closely related to its metabolic and storage functions<sup>[34]</sup>.

Current understanding of adipose tissue (AT)-derived adipokines encompasses over 100 proteins that interact with various cells and tissues<sup>[22]</sup>. Leptin and adiponectin are the most prevalent and well-studied adipokines. Leptin, the first adipokine to be discovered, has been shown to influence appetite and energy expenditure, serving as an important feedback mechanism to the brain regarding the size and condition of adipose tissue<sup>[35]</sup>. In contrast, adiponectin promotes insulin sensitivity and fatty acid oxidation in skeletal muscle and the liver, contributing to the maintenance of glucose and lipid homeostasis<sup>[36]</sup>.

Additionally, WAT releases extracellular vesicles (EVs), small membrane-bound particles containing various bioactive molecules such as microRNAs (miRNAs), proteins, and lipids. These EVs play a role in maintaining energy homeostasis both locally and systemically<sup>[37,38]</sup>.

The endocrine activity of WAT is influenced by factors such as nutritional status, physical activity, hormonal levels, and environmental signals. Dysregulation of WAT function, as observed in obesity and metabolic disorders, can result in insulin resistance (IR), chronic inflammation, and various metabolic and cardiovascular complications<sup>[39,40]</sup>.

#### 1.5 Dysregulation of Adipose Tissue and Its Implications

Under prolonged positive energy balance, a failure to expand AT mass via adipocyte hyperplasia (the formation of new adipocytes through precursor differentiation) results in adipocyte hypertrophy (an increase in cell size)<sup>[41]</sup>. This is a hallmark of AT dysfunction, as enlarged adipocytes have a diminished capacity to store excess dietary energy due to being already saturated with lipids<sup>[42,43]</sup>.

As adipocytes grow in size, they experience increased mechanical stress due to enhanced contact with neighboring cells and components of the extracellular matrix<sup>[22]</sup>. Additionally, adipocytes face hypoxia, as the hypertrophic expansion of adipose tissue outpaces angiogenesis<sup>[44]</sup>. Furthermore, hypertrophic adipocytes and damaged cells release pro-inflammatory cytokines, which recruit and activate immune cells. These factors collectively contribute to a chronic low-grade inflammatory state in adipose tissue, significantly impairing its functionality<sup>[45]</sup>.

In obese individuals, enlarged adipocytes exhibit partial resistance to the antilipolytic effects of insulin<sup>[22]</sup>. This increased lipolysis results in elevated levels of free fatty acids (FFAs) in the bloodstream, leading to ectopic lipid deposition and subsequent lipotoxicity. These disruptions can contribute to the onset of systemic insulin resistance, oxidative stress, and the progression of obesity-related comorbidities<sup>[46]</sup>.

The functionality of scWAT is closely linked to body fat distribution<sup>[22]</sup>. Under pathological conditions, its lipid-storage capacity becomes overwhelmed, leading to hypoxia, infiltration of pro-inflammatory macrophages, and ectopic fat accumulation in other tissues, such as the liver<sup>[47]</sup>. These alterations disrupt the balance of adipokine secretion, fostering a pro-inflammatory state and systemic insulin resistance, hallmark features in patients with MASLD and related disorders<sup>[47]</sup>.

Understanding the mechanisms underlying scWAT dysfunction highlights the critical role of adipose tissue expandability in metabolic health. The ability of adipose tissue to appropriately expand in response to excess energy intake is crucial for maintaining metabolic homeostasis. When the expansion capacity is compromised, it contributes to the pathogenesis of various metabolic diseases, including MASLD.

#### 1.6 Adipose Tissue Expandability and Liver Fat Deposition

The adipose tissue expandability hypothesis is crucial for understanding MASLD. This hypothesis posits that the body's capacity to store excess calories in subcutaneous adipose tissue (scWAT) is limited and varies between individuals<sup>[48]</sup>. Once scWAT reaches its maximum storage capacity, adipose tissue can no longer effectively store lipids, leading to the redistribution of lipids to other organs<sup>[49]</sup>. This results in ectopic fat accumulation, primarily in visceral adipose tissue (visWAT) and the liver, which contributes to insulin resistance and associated metabolic complications through mechanisms driven by lipotoxicity and inflammation<sup>[50]</sup>.

The disparity between energy intake and the storage capacity of adipose tissue is a critical factor in the onset of MASLD. When excessive caloric intake occurs, particularly in conjunction with the limited expandability of subcutaneous adipose tissue (scWAT), it results in fat accumulation in the liver. This hepatic fat buildup, known as steatosis, is a defining characteristic of MASLD and paves the way for additional liver damage<sup>[22]</sup>.

#### 1.6.1 Hepatic Response to Ectopic Fat Accumulation

In MASLD, hepatocytes are the primary cells impacted by ectopic fat deposition. The buildup of triglycerides in hepatocytes, initially serving as a protective mechanism against excess circulating free fatty acids, ultimately results in cellular stress and damage. This stress manifests in various ways:

- *Inflammatory Response*: The accumulation of ectopic fat in the liver triggers an inflammatory response, attracting immune cells and producing pro-inflammatory cytokines. Chronic inflammation induces notable histological changes, that are critical for the progression of the disease from steatosis to MASH and fibrosis, and may also contribute to hepatocellular carcinoma (HCC)<sup>[51]</sup>.
- Endoplasmic Reticulum (ER) Stress: The accumulation of lipids disrupts ER function in hepatocytes, triggering an unfolded protein response that further contributes to cellular stress and apoptosis<sup>[52]</sup>.
- Oxidative Stress: When lipid overflow exceeds the capacities of mitochondria and peroxisomes, respiratory oxidation becomes impaired, resulting in the generation of harmful metabolites and excess reactive oxygen species (ROS)<sup>[53]</sup>. These reactive molecules cause oxidative stress, worsening necro-inflammatory processes in the liver and damaging mitochondria. ROS and oxidized low-density lipoproteins (LDL) activate Kupffer and hepatic stellate cells, leading to collagen deposition and secondary liver fibrosis<sup>[54,55]</sup>.
- *Lipotoxicity*: Increased lipid levels, particularly saturated fatty acids and harmful lipid species such as diacylglycerol (DAG) and ceramide, lead to lipotoxic stress in hepatocytes, causing dysfunction and cell death<sup>[56]</sup>.
- *Altered Metabolism*: In a fatty liver, hepatocytes exhibit changes in carbohydrate and lipid metabolism, often linked to insulin resistance. This exacerbates lipid accumulation and impairs liver function<sup>[57]</sup>.

#### 1.7 miRNAs and Adipose Tissue Regulation

#### 1.7.1 miRNA biology and function

MicroRNAs (miRNAs) are a highly conserved family of small (21–25 nucleotides) noncoding RNAs<sup>[58,59]</sup>, first identified in *Caenorhabditis elegans* by Victor Ambros and Gary Ruvkun, who were awarded the 2024 Nobel Prize in Physiology or Medicine for their groundbreaking work on miRNAs<sup>[60–62]</sup>. miRNAs are considered negative regulators of gene expression, functioning by either suppressing translation or promoting mRNA degradation through base pairing with complementary sequences in the 3'-untranslated region (3'-UTR) of protein-coding mRNA transcripts<sup>[63]</sup>. Notably, mRNA degradation constitutes the primary mechanism of miRNA activity<sup>[64]</sup>.

miRNAs have the ability to regulate hundreds of mRNA targets, and a single mRNA can be influenced by multiple miRNAs<sup>[65]</sup>. In the human genome, approximately 2,500 miRNAs have been identified<sup>[66]</sup>, with about 60% of protein-coding genes being regulated by miRNAs at the post-transcriptional level<sup>[67]</sup>. Due to this regulation ability, miRNAs are key regulators within complex genetic pathways, providing a post-transcriptional layer of control over homeostatic and developmental processes<sup>[@6]</sup>. They play critical roles in a wide range of biological functions, including the regulation of the cell cycle, proliferation, lipid metabolism, inflammation, and fibrosis, among others<sup>[69]</sup>. However, the dysregulation of miRNA has been implicated in the pathogenesis of several many diseases and metabolic disorders, including MASLD<sup>[70]</sup> pecific miRNAs are known to be differentially expressed in metabolic tissues, including adipose tissue, in response to pathological states<sup>[68]</sup>. Their potential as biomarkers for disease progression and therapeutic targets has garnered significant interest in recent years<sup>[70]</sup>.

#### 1.7.2 miRNA biogenesis

The maturation of miRNAs occurs through a multi-step process that begins with transcription by RNA polymerase II<sup>[71]</sup>. This enzyme synthesizes single-stranded, non-protein-coding RNAs, which can be transcribed either as independent transcripts (intergenic miRNAs), often containing multiple miRNAs, or derived from the introns of protein-coding genes (intragenic or intronic miRNAs)<sup>[68]</sup>. The transcription of intergenic miRNAs produces primary miRNAs (pri-miRNAs) with a distinct hairpin or stem-loop structure and extends 1 kb or much longer,<sup>[72]</sup>, resulting in a long primary transcript<sup>[73]</sup>.

Then the pri-miRNA is cleaved by ribonulease III-type protein Drosha<sup>[74]</sup> and RNA binding protein DiGeorge Syndrome Critical Region 8 (DGCR8)<sup>[75]</sup> in the nucleus to produce a ~70 nucleotide long hairpin structure named precursor miRNA (pre-miRNA). Pre-miRNAs are transported to the cytoplasm by exportin-5, where they are cleaved by the ribonuclease III enzyme Dicer in complex with TRBP. This cleavage removes the terminal hairpin structure and generates a ~22-nucleotide miRNA:miRNA\* duplex<sup>[76]</sup>.

The selection of one strand of the miRNA:miRNA\* duplex is influenced by the thermodynamic stability at the duplex ends<sup>[71]</sup>. This duplex associates with an Argonaute (AGO) protein to form the RNA-induced silencing complex (RISC). The mature miRNA is generated when the miRNA\* strand is removed from the duplex. Generally, the strand with lower thermodynamic stability at its 5' end is preferentially loaded into an Argonaute protein (AGO). Notably, miRNA\* is not merely a byproduct of miRNA biogenesis; in many instances, it functions as a regulatory miRNA<sup>[77]</sup>.

The loaded strand, referred to as the guide strand (miRNA), directs the miRNA-induced silencing

complex (miRISC) to the mRNA target. In contrast, the unloaded strand, known as the passenger strand (miRNA\*), is unwound from the guide strand and subsequently degraded by cellular machinery<sup>[78,79]</sup>.

In addition, the diversity of miRNAs can lead to variations known as isomiRs, which arise from imprecise cleavage by Drosha and Dicer, 3' addition events, RNA editing, or single nucleotide polymorphisms (SNPs)<sup>[79,80]</sup>. These isomiRs contribute to the functional complexity of the miRNA landscape, further expanding the regulatory potential of miRNAs and highlighting their importance in the intricate mechanisms of gene regulation.

#### 1.7.3 Mechanisms of miRNA-mediated gene regulation

The interaction between miRNAs and their target mRNAs forms a complex miRNA-mRNA interactome, where miRNAs bind to various regions of mRNAs. In the 3' untranslated region (3' UTR), miRNA binding typically induces transcript degradation or translation inhibition. Interactions in the 5' UTR and coding regions can also repress gene expression<sup>[81]</sup>. Furthermore, miRNA binding to promoter regions may induce transcription, although this mechanism remains poorly understood<sup>[78]</sup>.

The classic mechanism of miRNA action involves inhibiting gene expression by binding to complementary sequences, typically 6–8 nucleotides long, within the 3' untranslated region (3' UTR) of target mRNAs<sup>[68]</sup>. The miRNA strand loaded into Argonaute (AGO) proteins guides the miRNA-induced silencing complex (miRISC) to miRNA response elements (MREs) on the target mRNA. The outcome of this interaction depends on the degree of complementarity: perfect matches result in transcript degradation, whereas partial complementarity inhibits gene expression either through mRNA degradation or translation repression. Interestingly, some miRNAs can act independently of the miRISC complex. However, the mechanisms underlying these miRISC-free actions are not yet fully understood and warrant further investigation<sup>[78,81]</sup>.

Due to the requirement for only a 6–8 nucleotide complementarity, a single miRNA can potentially target hundreds of mRNAs. In fact, most mammalian mRNAs are conserved targets of miRNAs<sup>[82]</sup>. Bioinformatic tools have become essential for predicting and validating miRNA-target interactions, facilitating the study of miRNA regulatory networks<sup>[83,84]</sup>.

#### 1.7.4 Association of miRNA with Hepatic Steatosis

miRNAs play a critical role in regulating key metabolic pathways in various liver cell types during the transition from steatosis to fibrosis, influencing processes such as lipid metabolism, oxidative stress, inflammation, and fibrosis<sup>[85]</sup>. Throughout the progression of MASLD, miRNAs have been shown to be involved at every stage of the disease spectrum through diverse regulatory mechanisms<sup>[86]</sup>.

Acting as negative posttranscriptional regulators of the expression, miRNAs also influence the expression of genes involved in lipid metabolism. Typically, one miRNA targets multiple mRNAs that are interconnected within the same metabolic pathway. This targeting capability allows miRNAs to regulate the expression of numerous other miRNAs, thereby contributing to a complex network of regulatory interactions<sup>[87,88]</sup>.

Given the critic ple of miRNAs in regulating metabolic pathways in the liver, it is essential to explore which specific miRNAs are associated with hepatic steatosis. These miRNAs can be

categorized according to their primary effects, allowing for a more detailed understanding of their involvement in this condition

- miRNAs Involved in Lipid Accumulation: *miR-20b* inhibits *PPARα* reducing free fatty acid oxidation and mitochondrial biogenesis, which increases hepatic lipid accumulation<sup>[89]</sup>. *miR-32* activates sterol regulatory element binding protein-mediated adipogenesis, promoting de novo lipogenesis and liver lipid accumulation<sup>[90]</sup>. *miR-30a-3p* upregulates lipid metabolism-related proteins and downresplates mitochondrial proteins leading to liver fat accumulation<sup>[91]</sup>. *miR-130b-5p* upregulated in MASLD murine models, increasing hepatic lipid accumulation via the IGFBP2-dependent AKT pathway<sup>[92]</sup>. *miR-451a* regulates THRSP, reducing triglyceride accumulation in the liver<sup>[93]</sup>.
- miRNAs Implicated in Insulin Resistance: *miR-188* targets *ATG12*, negatively regulating hepatic glucose and lipid metabolism, which worsens steatosis and insulin resistance<sup>[94]</sup>. *miR-27* disrupts insulin/AKT signaling by targeting *PDPK1* and *PIK3R1*, leading to insulin resistance<sup>[95]</sup>. *miR-30b* induces ER stress by targeting *SERCA2b*, contributing to insulin resistance<sup>[96]</sup>.
- miRNAs Regulating Fatty Acid Oxidation: *miR-100* regulates directly *CD36*, reducing fatty acid uptake in hepatocytes and downstream lipid metabolic mediators<sup>[97]</sup>.*miR-376b-3p* targets *FGFR1*, increasing fatty acid oxidation and improving hepatic lipid accumulation<sup>[98]</sup>.
- miRNAs Associated with Endoplasmic Reticulum (ER) Stress: miR-26a mitigates high-fat diet-induced ER stress and steatosis by targeting  $eIF2\alpha^{[99]}$ .
- miRNAs Modulating Inflammatory and Fibrotic Pathways: miR-222 regulates mitophagy in hepatic stellate cells (HSCs) by targeting PINK1, contributing to fibrosis in MASH<sup>[100,101]</sup>. miR-146a: Suppresses fibrosis and improves hepatic lipid/glucose metabolism by targeting WNT1 and  $WNT5a^{[102,103]}$ .
- miRNAs Affecting Lipogenesis and Lipolysis: *miR-199a-5p* reduces *MST1*, modulating hepatic lipogenesis and lipolysis<sup>[104]</sup>. *miR-103-3p* targets *ACOX1*, promoting hepatic steatosis and worsening MASLD<sup>[105]</sup>. Furthermore, *miR-802* inhibits *AMPK* expression, promoting NAFL/MASH<sup>[106]</sup>.
- Regulatory Insights: Part of the miR-17-92 cluster, exacerbates steatotic changes by regulating CYP7A1 signaling [107]. Aditionally, miR-132 is involved in lipid homeostasis by suppressing multiple regulatory transcripts [108].

Table 1 miRNAs and Their Molecular Targets in Hepatic Steatosis

miRNA	Molecular Target	Effect	Reference
miR-20b	$PPAR\alpha$	↑ Hepatic lipid accumulation	[89]
miR-188	ATG12	† Steatosis and insulin resistance	[94]
miR-27	PDPK1, PIK3R1	↑ Insulin resistance	[95]
miR-32	INSIG1	† Lipogenesis, hepatic lipid accumulation	[90]
miR-100	CD36	↓ Fatty acid uptake	[97]
miR-451a	THRSP	↓ Triglyceride accumulation	[93]
miR- = 199a-5p	MST1	Modulates hepatic lipogenesis and lipolysis	[104]

miRNA	<b>Molecular Target</b>	Effect	Reference
miR- 376b-3p	FGFR1	↑ Fatty acid oxidation, improves hepatic lipid accumulation	[98]
miR-103- 3p	ACOX1	Promotes steatosis, worsens MASLD	[105,109]
miR-222	ACOX1, PINK1	↑ Triglycerides in hepatocytes, regulates mitophagy	[100,101]
miR-802	AMPK	Promotes MASH by inhibiting AMPK	[106]
miR-30a- 3p	ACC, p-GSK-3β, FASN, CPT1, p-AMPK, UCP2	↑ Hepatic fat accumulation	[91]
miR-17	CYP7A1	↑ Steatotic changes	[107]
miR- 130b-5p	IGFBP2	↑ Hepatic lipid accumulation, insulin resistance	[92]
miR-146a	MED1, WNT1, WNT5a	Improves lipid and glucose metabolism, ↓ Fibrosis	[102,103]
miR-30b	SERCA2b	↑ Endoplasmic reticulum (ER) stress, insulin resistance	[96]
miR-26a	$e$ IF $2\alpha$	↓ ER stress and high-fat diet-induced steatosis	[99]
miR-132	SIRT1, P300, PTEN, CYP2E1, FOXO3	† Lipid homeostasis by suppressing regulatory transcripts	[108]



### Chapter 2

## Hypothesis and Objectives

#### 2.1 Hypothesis

The main hypothesis of this study is that microRNAs (miRNAs) derived from subcutaneous adipose tissue plate key role in regulating metabolic processes involved in the development and progression of hepatic steatosis. These miRNAs can alter fundamental metabolic pathways by modulating genes associated with lipid metabolism in human hepatocytes.

More specifically, it is proposed that:

- miRNAs secreted or expressed in the subcutaneous adipose tissue of obese patients are involved in intercellular communication that affects lipid accumulation in the liver, contributing to the development of hepatic steatosis.
- Bioinformatic analysis of RNAseq data will identify differential miRNAs in the subcutaneous adipose tissue of obese patients, associated with key metabolic pathways related to lipogenesis, lipolysis, and  $\beta$ -oxidation.
- The experimental modulation of specific miRNAs in *in vitro* cellular models of human hepatocytes, through overexpression or inhibition strategies, can reveal their functional role in processes related to hepatic steatosis.
- The functional characterization of these miRNAs may propose them as potential biomarkers or therapeutic targets for the prevention and treatment of obesity-associated hepatic steatosis.

### 2.2 Main objective

To study the function of microRNAs (miRNAs) expressed in subcutaneous adipose tissue in the regulation of metabolic processes involved in the development of hepatic steatosis, with the aim of identifying potential biomarkers and therapeutic targets for its management.

# 2.3 Specific Objectives

- Analyze RNAseq data data from subcutaneous adipose tissue of obese patients obtained from the FATe cohort (ADIPOFAT)
- Identify a set of differential miRNAs associated with metabolic pathways involved in lipogenesis,  $\beta$ -oxidation, and lipolysis related to hepatic steatosis.
- Establish in vitro cell models of human hepatocytes to study the function of candidate miRNAs.
- Modify the expression of specific miRNAs through transfection with synthetic analogs (overexpression and inhibition).
- Evaluate the expression of key genes related to lipogenesis, lipolysis, and  $\beta$ -oxidation in tran ted hepatocytes.
- Explore the potential of identified miRNAs as biomarkers or therapeutic targets for the treatment of obesity and its associated disorders.

## Chapter 3

## Methodological Framework

#### 3.1 Study Population

The analyzed data originates from the FATe cohort<sup>[110]</sup>, a longitudinal study of obese patients undergoing bariatric surgery at Miguel Servet University Hospital (HUMS, Zaragoza, Spain). For this research, 78 patients with varying degrees of adiposity were selected based on subcutaneous adipose tissue (scWAT) samples, which are registered at the regional Biobank (Biobanco Aragón) and approved by the CEICA ethics committee. Patients with alcohol or drug abuse, autoimmune diseases, chronic inflammatory conditions, or infectious diseases (HIV, HBV, HCV) were excluded during the screening process. The FATe cohort is characterized by a range of clinical and demographic variables, including sex, age, body mass index (BMI), steatosis, metabolic dysfunction-associated steatohepatitis (MASH), hepatocytic ballooning, lobular inflammation, diabetes, hyperlipidemia, and metabolic dysfunction-associated steatotic liver disease (MASLD).

# 3.2 Collection and characterization of subcutaneous white adipose tissue

Subcutaneous white adipose tissue Adipose tissue (scWAT) biopsies (~3 cm³) from the subcutaneous depot were obtained during laparoscopic bariatric surgery using a bipolar/ultrasonic device (Thunderbeat). The samples were extracted through a 12 mm trocar (Applied Medical) inserted into the left hypochondrium. An experienced pathologist evaluated pathological features such as steatosis, lobular inflammation, hepatocellular ballooning, and fibrosis according to the criteria established by the Nonalcoholic Steatohepatitis Clinical Research Network ([111]).

#### 3.3 RNA isolation

Total RNA was extracted from frozen biopsies of subcutaneous white adipose tissue (scWAT) and cell cultures using TRIzol (#T9424, Sigma Aldrich) following the manufacturer's instructions. For lysing adipose tissue samples, 1 ml of TRIzol was added per sample in a homogenizer, while for cell cultures, 1 ml of TRIzol was used per 10 cm<sup>2</sup> of the culture plate, along with a scraper. The resulting cell lysates or tissue disaggregates were transferred to a vial and incubated for 5 minutes at room temperature to dissociate nuclear components. Subsequently, 0.2 ml of 100% chloroform

per ml of TRIzol was added. The mixture was shaken vigorously, incubated for 15 minutes at room temperature, and then centrifuged for 15 minutes at 12,000 g and 4 °C.

Following centrifugation, three distinct phases were formed. The aqueous phase, which contained the ribonucleic acids, was collected, and 0.5 ml of 100% isopropanol was added to precipitate the RNA. This mixture was mixed, incubated on ice for 10 minutes, and then centrifuged at 12,000 g for 15 minutes at 4 °C. Afterward, the supernatant was carefully decanted, and the RNA pellet was resuspended in 1 ml of 75% ethanol for washing. The pellet was homogenized and centrifuged at 7,500 g for 5 minutes at 4 °C. The supernatant was discarded, and the pellet was allowed to dry for 10 minutes at room temperature. Finally, the RNA was resuspended in DEPC-treated water. To eliminate any genomic DNA contamination, all RNA samples were treated with RNase-Free DNase (Life Technologies). The concentration and purity of the RNA were assessed by measuring absorbance at 260/280 nm and 260/230 nm using a Nanodrop 2000 (Thermo Fisher). The quality of the extracted RNA was visualized using an agarose gel.

#### 3.4 RNA Sequencing

RNA integrity was evaluated using the RNA Integrity Number (RIN) on the Agilent 2200 TapeStation with the RNA ScreenTape assay. Stranded mRNA libraries were prepared with the Novogene NGS RNA Library Prep Set (PT042), which included mRNA isolation using poly-T oligo-attached magnetic beads, cDNA synthesis, adapter ligation, and PCR amplification. Libraries that passed quality control checks were sequenced (2x150 bp) on the Illumina Novaseq X Plus platform.

#### 3.5 Analysis of sRNA-seq Data with nf-core/smrnaseq

For the analysis of small RNA sequencing (sRNA-seq) data, version 2.4.0 of the *nf-core/smrnaseq* pipeline<sup>[112]</sup> was used, which is specifically designed for the automated processing of miRNAs data.

#### 3.5.1 Execution of the *nf-core/smrnaseq* Pipeline

The installation of nf-core/smrnaseq was carried out following the instructions provided by the authors in nf-core<sup>[113]</sup>, available at https://nf-co.re/smrnaseq/2.4.0.

To ensure the proper installation and execution of the pipeline, the following key components were installed beforehand:

- 1. *Nextflow*: Version 24.04.4 of Nextflow was used, following the detailed instructions at https: //nf-co.re/usage/installation.
- 2. *Java Runtime Environment (JRE)*: Version 11.0.25 of the Java Runtime Environment was installed, as it is required for compatibility with Nextflow and the *nf-core/smrnaseq* pipeline.

To ensure reproducibility and streamline the pipeline execution, one of the available Docker containers was utilized. These containers provide the necessary instructions and configurations required to run the pipeline. The configuration is specified at runtime using the profile argument. For this analysis, the Docker image *nf-core/smrnaseq*, available at Docker Hub, was employed.

The pipeline was executed on a server with 8 CPUs, 16 GB of RAM, and a Linux operating system. The following command was used in the terminal, which configures the main options, including

the reference genome, input data, and output file location:

```
nextflow run nf-core/smrnaseq -r 2.4.0
-profile docker,ci
--genome GRCh38
--input '/home/joshoacr13/Documentos/TFM/nfcore-smrnaseq/input/samples.csv'
--fasta 'https://github.com/nf-core/test-datasets/raw/smrnaseq/reference/genome.fa'
--mirtrace_species 'hsa'
--outdir /home/joshoacr13/Documentos/TFM/nfcore-smrnaseq/workdir
-resume -c /home/joshoacr13/Documentos/TFM/nfcore-smrnaseq/nextflow_memory.config
--filter_contamination
```

The pipeline was executed three times to accommodate the large number of samples, processing 26 samples per run.

#### 3.5.2 Description of the Parameters Used

- -profile docker, ci: Runs the pipeline inside a Docker container to ensure reproducibility and sets up a continuous integration (CI) profile.
- --genome GRCh38: Specifies the human genome (version GRCh38) as the reference for sequence mapping.
- --input: Provides the path to the CSV file containing metadata and the paths to the FASTQ files.
- --fasta: URL to the FASTA file of the reference genome.
- --mirtrace\_species hsa: Defines the species as Homo sapiens (hsa) for miRNAs analysis with miRTrace.
- --outdir: Sets the working directory for the processed results.
- -resume: Allows continuation of a previous analysis without restarting from the beginning.
- -c: Specifies a custom configuration file (nextflow\_memory.config) to adjust resource usage.
- --filter\_contamination: Enables the contamination filtering.

#### 3.5.3 Analysis Workflow and Tools Used

The *nf-core/smrnaseq* pipeline perform the following steps:

- 1. **Quality Control** An initial quality assessment of the raw reads was conducted using FastQC (version 0.12.1)<sup>[114]</sup>. Additionally, 3' adapter trimming was performed using fastp (version 0.23.4)<sup>[115]</sup>, followed by quality and length filtering. A second quality assessment of the trimmed reads was conducted with FastQC.
- 2. **miRNA Quality Control**: A more specific quality control process for miRNA sequencing was implemented, whereby samples that failed to meet the minimum quality thresholds established by miRTrace (version 1.0.1)<sup>[116]</sup> were excluded from further analysis. This tool incorporates the following steps to ensure the integrity and reliability of the data:
- **Verify Read Length Distribution**: The majority of reads fell within the expected range of 18–24 nucleotides, indicative of high-quality small RNA data.

- **Identify Contaminants**: Potential contaminants such as tRNA, rRNA, and other non-target molecules were flagged.
- **Taxonomic Classification**: Reads were classified taxonomically to ensure that most sequences originated from the organism of interest (*Homo sapiens*).

#### 3. miRNAs Quantification:

- **Alignment**: The filtered reads were aligned against mature miRNA sequences in the miRBase database using *Bowtie1* (version 1.3.1)<sup>[117]</sup>. Unmapped reads were aligned against "hairpin" sequences to identify miRNA precursors.
- **Post-Alignment Processing**: *SAMtools* (version 1.16.1)<sup>[118]</sup> was used to process the mapping results.
- 4. **IsomiR Annotation**: The collapsed reads were processed with *mirtop* (version 0.4.28)<sup>[119]</sup> to identify miRNA variants (isomiRs).
- 5. Analysis and Visualization of Results: The overall pipeline metrics, encompassing quality assessments, mapping statistics, and expression analysis results, were consolidated and summarized using MultiQC (version 1.25.1)<sup>[120]</sup>. Visualization of the results was performed using the ggplot2 package in  $R^{[121]}$ , which facilitated the creation of clear and informative graphical representations.

#### 3.6 Differential expression analysis according to steatosis using DEseq2

For the differential expression analysis, the R statistical software<sup>[122]</sup>, version 4.4.1 (2024-06-14) (https://cran.r-project.org/), was used. This analysis was performed using the RStudio integrated development environment (IDE)<sup>[123]</sup>, version 2023.12.0+369, designed for Ubuntu Jammy (https://www.rstudio.com/).

The script used to perform the differential expression analysis is available in the file "miRNA\_steatosis.qmd" which can be accessed at the following link: https://github.com/joshoandres13/miRNAs.

The analysis began by loading essential R packages: tidyverse (version  $2.0.0)^{[124]}$  for data manipulation,  $isomiRs^{[125]}$  (version 1.32.1) for analyze isomirs and miRNAs from small RNA-seq, DESeq2 (version  $1.44.0)^{[126]}$  for differential expresion analysis, org.Hs.eg.db (version  $3.19.1)^{[127]}$  for gene annotation.

#### 3.6.1 Data Preparation

Metadata were imported, and sample identifiers were stablished as row names. The variable *Steatosis* is categorized in four distinct groups. Isomirs count data were subsequently obtained where rows represent the identified isomiRs and columns correspond to the experimental samples.

Using isomirs count data and metadata, an object of class IsomirDataSeq was created. This object enables efficient management of information derived from small RNA sequencing studies, streamlining differential expression analyses and facilitating the interpretation of biological findings.

#### 3.6.2 Filtering and Processing of isomiRs

The filtering process enables the grouping of isomiRs into distinct categories, associating them with a single variant of a miRNA. This grouping is essential for ensuring consistency and accuracy in differential expression analyses. To reduce technical noise and highlight biologically meaningful signals, a stringent filtering criterion was applied: only isomiRs with a minimum of 20 counts in at least 40 samples were retained.

#### 3.6.3 Differential expression in scWAT

In this study, the *DESeq2* package was used to perform differential expression analysis on scWAT samples with varying degrees of liver steatosis. The *DESeq2* object was configured using the Likelihood Ratio Test (LRT), enabling the analysis to account for all four steatosis categories: <5% (no steatosis), 5–33% (mild steatosis), 33–66% (moderate steatosis), and >66% (severe steatosis). This approach allowed for a comprehensive assessment of gene expression changes across the entire spectrum of steatosis progression.

#### 3.6.3.1 Testing for differential expression

The criteria for identifying significant differentially expressed miRNAs across the steatosis categories involved applying a false discovery rate (FDR) cutoff of less than 0.05. Following the identification of significant miRNAs, the expression patterns of the selected miRNAs were further analyzed by representing their normalized counts in boxplots across the four steatosis groups. This approach enabled the observation of expression variations of these miRNAs across different steatosis degrees, offering insights into their potential roles in steatosis progression. Differences between steatosis groups were evaluated using the Kruskal-Wallis test

#### 3.7 Target mRNA Selection and Validation Using multiMiR

The selected miRNAs were used for the identification of mRNA targets through the *multiMiR* bioinformatics package<sup>[128,129]</sup>, version 2.4.0 in R. *multiMiR* facilitates a systematic search and annotation of miRNA targets, providing functional analysis to elucidate biological mechanisms. For this analysis, only validated interaction data were utilized.

#### 3.7.1 Filtering Parameters

The validated target table provided by *multiMiR* was used during the selection process. Key columns included:

- 1. **database**: Source database of validated interactions, such as *miRTarBase*, *TarBase*, or *miRecords*.
- 2. mature\_mirna\_id: Standard format identifier for the miRNA.
- 3. target\_symbol: Target gene symbol.
- 4. **experiment**: Experimental methods used for validation, including luciferase assays, Western blot, or qRT-PCR.
- 5. **support\_type**: Level of experimental support, such as "Functional MTI" (miRNA-mRNA functional interaction).
- 6. **pubmed\_id**: References to PubMed articles reporting the interaction.
- 7. **type**: Specifies whether the interaction is "validated" or "predicted."

#### 3.7.2 Selection Criteria

To ensure reliable results, databases were filtered according to update criteria and the following selection parameters:

- Databases up-to-date at the time of analysis were prioritized (*miRTarBase* and *TarBase*).
- Only interactions classified as "validated" were included.
- Interactions backed by robust experimental methods, such as luciferase assays or Western blot, were prioritized.
- Interactions with functional support ("Functional MTI") and verifiable references in PubMed were selected.

This approach ensured the identification of mRNA targets with high reliability and experimental backing, facilitating the analysis of potential regulatory functions of the studied miRNAs.

#### 3.7.3 Functional Analysis

To explore the biological functions associated with the validated target genes, an enrichment analysis was performed using the *KEGGREST* package<sup>[130]</sup> in R. Gene symbols for validated target genes associated with selected miRNAs were extracted using *multiMiR* previously, with duplicates removed to ensure a unique gene list. The Entrez IDs of these genes were mapped to specific metabolic pathways in KEGG, using parameters that controlled the false discovery rate (FDR) with the Benjamini-Hochberg method and significance cutoff values for qvalue and pvalue set at 0.05. The results included a bar plot displaying the top 5 significantly enriched KEGG pathways, highlighting their statistical significance and the number of genes associated with each pathway. This analysis provided insights into key metabolic pathways and biological processes involving miRNA-regulated target genes.

#### 3.8 Functional validation of miRNAs

#### 3.8.1 Cell Culture and Transfections with miRNA Mimics

The human hepatoma HepG2 cell line (American Type Culture Collection, ATCC® HB-8065™; Manassas, VA, USA) was cultured in an incubator at 37 °C and 5% CO₂ using high-glucose Dulbecco's Modified Eagle Medium L-GlutaMAX (DMEM) (Gibco, Thermo Fisher Scientific Inc., Waltham, MA, USA) supplemented with 10% fetal bovine serum (FBS; Gibco, Thermo Fisher Scientific Inc., Brazil). Experiments were carried out when the cells reached 70-80% confluence.

For the experiments, HepG2 cells were plated at a density of 100,000 cells per well in 12 well-plates for gene expression assays, in DMEM high-glucose (1 g/L) L-GlutaMAX supplemented with 10% FBS.

HepG2 cells were reverse-transfected with Lipofectamine RNAiMA; Reagent (Thermo Fisher Scientific Inc.) and 50 nM of the following mirVana<sup>TM</sup> miRNA mimics (Thermo Fisher Scientific Inc.): a scramble sequence as a negative control (mirVana<sup>TM</sup> miRNA Mimic, Negative Control #1), negative control (mirVana<sup>TM</sup> miRNA Inhibitor, Negative Control #1), hsa-miR-144-3p (5' – UACAGUAUAGAUGAUGUACU – 3'; assay ID H11051), and hsa-miR-372-3p (5' – AAAGUGCUGCGACAUUUGAGCGU – 3'; assay ID MC10165). The miRNA mimics were diluted in Opti-MEM I Reduced Serum Medium (Gibco, Thermo Fisher Scientific Inc.) and added to the

wells according to the manufacturer's instructions. Following this, Lipofectamine was added to the wells containing the diluted miRNA mimics and incubated for 15 minutes at room temperature to form miRNA mimic–lipofectamine complexes. HepG2 cells diluted in DMEM with 10% FBS were then plated into the wells. The cells were incubated for 24 hours to assess transfection efficiency. To induce lipid accumulation and simulate steatosis<sup>[131]</sup>, HepG2 cells were treated with oleic acid 24 hours after miRNA mimic transfections, mixed in DMEM GlutaMAX-I supplemented with 10% FBS, at a final concentration of 0.5 mM, which reflects the physiological range of fatty discussed to mimic hepatic steatosis<sup>[132]</sup>.

#### 3.8.2 RNA Isolation and Gene Expression Analyses

HepG2 cells were frozen on dry ice and stored at -80 °C until RNA extraction. Cells used for miRNA expression analyses were washed previously with phosphate-buffered saline (PBS) to completely remove potential unabsorbed miRNA mimics. Total RNA was extracted from frozen cell cultures using TRIzol (#T9424, Sigma Aldrich) following the manufacturer's instructions. For lysing cell cultures, 1 ml of TRIzol was used per 10 cm² of the culture plate, along with a scraper. The resulting cell lysates were transferred to a vial and incubated for 5 minutes at room temperature to dissociate nuclear components. Subsequently, 0.2 ml of 100% chloroform per ml of TRIzol was added. The mixture was shaken vigorously, incubated for 15 minutes at room temperature, and then centrifuged for 15 minutes at 12,000 g and 4 °C.

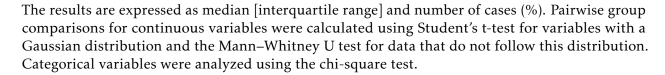
Following centrifugation, three distinct phases were formed. The aqueous phase, which contained the ribonucleic acids, was collected, and 0.5 ml of 100% isopropanol was added to precipitate the RNA. This mixture was mixed, incubated on ice for 10 minutes, and then centrifuged at 12,000 g for 15 minutes at 4 °C. Afterward, the supernatant was carefully decanted, and the RNA pellet was resuspended in 1 ml of 75% ethanol for washing. The pellet was homogenized and centrifuged at 7,500 g for 5 minutes at 4 °C. The supernatant was discarded, and the pellet was allowed to dry for 10 minutes at room temperature. Finally, the RNA was resuspended in DEPC-treated water. To eliminate any genomic DNA contamination, all RNA samples were treated with RNase-Free DNase (Life Technologies). RNA concentration were determined using Qubit 4 Fluorometer (Thermo Fisher Scientific Inc.).

mRNA expression was evaluated in HepG2 cells transfected with miRNA mimics for 48 h. RNA was reverse transcribed using PrimeScript Reverse Transcriptase (Takara Bio), with 100 ng of RNA utilized for each reaction in a total volume of 10  $\mu$ l. The process was carried out using an Applied Biosystems 2720 Thermal Cycler, for ying this protocol: 10 minutes at 25 °C, 2 hours at 37 °C, and finally, 5 minutes at 85 °C. Ten nanograms of the cDNA product were amplified using Quantitative-real time PCR (qPCR) in a total reaction volume of 15  $\mu$ l with SYBR Select Master Mix (Applied Biosystems), to which 0.5  $\mu$ l of gene-specific prime can a concentration of 10  $\mu$ M was added. The primers utilized are detailed in Table 3.1 cDNA amplification was performed on a StepOnePlus system (Applied Biosystems) with the following protocol: an initial step at 95 °C for 10 minutes, followed by 40 cycles of 15 seconds at 95 °C and 1 minute at 60 °C, concluding with 15 seconds at 95 °C. The gene  $\beta$ -actin was used as a housekeeping control to normalize gene expression levels ( $\Delta Cq$ ). Comparisons between gene expression levels after miRNA mimic transfections vs. scramble-sequence-transfected cells (negative control) were established using the the  $2-\Delta\Delta CT$  method, determining relative gene expression  $\alpha$ -actin were expression.

Table 3.1: Prime sesigned for qPCR mRNA gene expression analysis. Abbreviations: *ACTB-2* (Actin), *ACACA* (Acetyl-CoA Carboxylase Alpha), *DGAT2* (Diacylglycerol O-Acyltransferase 2), *FAS* (Fatty Acid Synthase), *PNPLA2* (Patatin-like phospholipase domain-containing protein 2), *PPARG* (Peroxisome Proliferator Activated Receptor Gamma)

Gene Na	Forward Primer $(5' \rightarrow 3')$	Reverse Primer (5'→ 3')
ACTB-2	ACCGAGCGCGGCTACAG	CTTAATGTCACGCACGATTTCC
PPARG	AGATGACAGCGACTTGGCAAT	ACTCAGGGTGGTTCAGCTTC
ATGL (PNPLA2)	TGGAGACTGAGGAGAACAAG	ATCCCTGCTTGCACATCTCT
FAS	AAGGACCTGTCTAGGTTTGATGC	TGGCTTCATAGGTGACTTCCA
DGAT2	AGTGGCAATGCTATCATCATCGT	TCTTCTGGACCCATCGGCCCCAGGA
ACCA	AGGTGCCTAGAGGGTTGAAGA	TCGGCCCTGCTTTACTAGGT

#### 3.9 Statistical Analysis



### Chapter 4

### **Results**

#### 4.1 Phenotypic Characterization

Table 4.1 describes the phenotypical features of FATe patients selected for this study. This cohort encomprisses 78 subjects (24% males; 76% females), aged between 22 to 61 with obesity as regards of BMI levels of  $46.11 \pm 6.13$  kg/m². These patients had an overall presence of general variables/descriptors of metabolic syndrome features of particular interest those related to liver/hepatic status (steatosis, lobular inflammation and the prevalence of associated metabolic diseases). According to the hepatic steatosis scale, 35.9% of patients had less than 5% liver fat, indicating a normal or minimal steatosis state. A total of 32.1% exhibited mild steatosis (5–33%), while 25.6% showed moderate fat accumulation (33–66%). Only 6.4% of patients presented severe steatosis (>66%), with no significant differences between men and women (p= 0.818).

Table 4.1. clinical characteristics of the FATe cohort. Data are presented as number of cases (%) or median [interquartile range]. Differences between groups were tested with the Mann–Whitney U test and chi-square test; *BMI*: Body Mass Index (kg/m²); *MASH*: Metabolic dysfunction-associated steatohepatitis.

Characteristic	Overall (n=78)	Female (n=59)	Male (n=19)	p
Age (years) median	47.03 [13.75]	46.53 [13]	48.58 [14]	0.408
Body Mass Index (kg/m²) median	46.11 [8.70]	45.68 [8.76]	47.42 [8.70]	0.284
MASLD Activity Score Category (%):				0.510
- 0	22 (28.2)	16 (27.1)	6 (31.6)	
- 1	16 (20.5)	13 (22.0)	3 (15.8)	
- 2	17 (21.8)	14 (23.7)	3 (15.8)	
- 3	9 (11.5)	7 (11.9)	2 (10.5)	
- 4	9 (11.5)	7 (11.9)	2 (10.5)	
- >= 5	5 (6.4)	2 (3.4)	3 (15.8)	
Hepatic Steatosis Scale (%):				0.818
- < 5%	28 (35.9)	21 (35.6)	7 (36.8)	
- 5-33%	25 (32.1)	20 (33.9)	5 (26.3)	
- > 33-66%	20 (25.6)	15 (25.4)	5 (26.3)	
- > 66%	5 (6.4)	3 (5.1)	2 (10.5)	
Hepatocytic ballooning Category (%):				0.489

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- None	58 (74.4)	45 (76.3)	13 (68.4)	
- Few cells	13 (16.7)	10 (16.9)	3 (15.8)	
- Many cells	7 (9.0)	4 (6.8)	3 (15.8)	
Lobular Inflammation Category (%):				0.596
- No foci	52 (66.7)	41 (69.5)	11 (57.9)	
- < 2 foci/200x	19 (24.4)	14 (23.7)	5 (26.3)	
- 2-4 foci/200x	4 (5.1)	2 (3.4)	2 (10.5)	
- > 4  foci/200x	3 (3.8)	2 (3.4)	1 (5.3)	
Diabetes:				1.000
- Yes (%)	21 (26.9)	16 (27.1)	5 (26.3)	
- No (%)	57 (73.1)	43 (72.9)	14 (73.7)	
Hyperlipidemia:				0.006
- Yes (%)	27 (34.6)	15 (25.4)	12 (63.2)	
- No (%)	51 (65.4)	44 (74.6)	7 (36.8)	
Metabolic dysfunction-associated:				0.249
- MASH (%)	12 (15.4)	7 (11.9)	5 (26.3)	
- Non-MASH (%)	66 (84.6)	52 (88.1)	14 (73.7)	

Regarding lobular inflammation, most patients (66.7%) had no inflammatory foci, while 24.4% exhibited fewer than two foci per microscopic field, and 3.8% showed severe inflammation (>4 foci). Hepatocyte ballooning analysis revealed that 74.4% of patients did not display significant damage, although 9.0% showed severe ballooning.

Among most prevalent comorbidities, 26.9% of patients were diagnosed with diabetes, and 34.6% presented hyperlipidemia, with the latter being significantly more prevalent in men (63.2%) compared to women (25.4%, p=0.006). Additionally, 15.4% of patients were classified with metabolic dysfunction-associated steatohepatitis (MASH), although this proportion showed no statistically significant differences between genders (p=0.249). Collectively, these data highlight heterogeneity in the clinical characteristics of the cohort, emphasizing the complexity of the relationship between obesity and liver disease in this group of patients.

#### 4.2 Quality Control (QC) and reads preprocessing

All QC results from the various steps in the *nf-core/smrnaseq* pipeline are summarized in the following figures and tables. The evaluation of samples, both before and after processing, was conducted using *fastp*, a rapid tool designed for preprocessing RNA sequencing data. This tool includes features for adapter trimming, quality filtering, and report generation, making it essential for assessing sample quality. Overall, Table 4.2 presents a descriptive analysis that includes key metrics related to the sequencing data.

Table 4.2: Descriptive statistics of the analyzed metrics with fastp. % *Duplication*: Duplication rate before filtering; *Reads After Filtering*: Total reads after filtering in millions; % *GC* content: GC content after filtering; % *PF*: Percent reads passing filter; % *Adapter*: Percentage adapter-trimmed reads

	Mean	sd	Median	Minimum	Maximum	Range
% Duplication	98.30513	0.68	98.45	94.31	99.09	4.78

Reads After Filtering (M)	25.23152	5.67	25.82	1.51	35.47	33.96
% GC content	46.71982	1.68	46.43	43.26	50.89	7.63
% PF	99.02358	1.21	99.36	90.13	99.76	9.63
% Adapter	99.38240	0.35	99.45	96.96	99.62	2.66

The metrics obtained from the fastp analysis provide a detailed overview of the preprocessing performance across the samples. The percentage of duplicated reads before filtering was notably high, with a mean of 98.31% ( $\pm 0.68$ ), a median of 98.45%, and a range between 94.31% and 99.09%. After filtering, the number of reads retained per sample averaged 25.23 million ( $\pm 5.67M$ ) suggesting a reasonable amount of reads obtained, with a median of 25.82M and a range from 1.51M to 35.47M.

The GC content showed consistency across the samples, with an average of 46.72% ( $\pm 1.68$ ), a median of 46.43%, and a range between 43.26% and 50.89%. The percentage of pass-filtered (PF%) reads was high, with an average of 99.02% ( $\pm 1.21$ ), reaching a maximum of 99.76%, as shown in Figure 4.1.

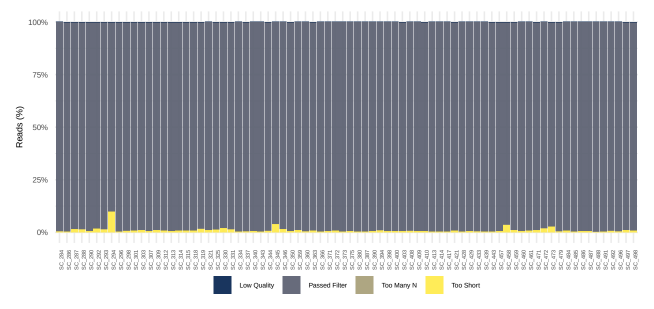


Figure 4.1: Fastp: Filtered Reads

The adapters used in library construction, specifically the *Illumina Universal Adapter* type, are optimized for the amplification and sequencing of diverse sample types. The adapter trimming was highly effective, achieving a mean success rate of 99.38% ( $\pm 0.35$ ) indicating a low level of adapter contamination in most samples, with a range from 96.96% to 99.62% (Table 4.2). In line with these findings, Figure 4.2 illustrates the adapter content before and after processing with *fastp*, where the mean trimming percentage of  $0.05 \pm 0.06\%$  for scWAT samples indicates that adapters were effectively removed from all samples. These results confirm the efficiency of the preprocessing steps, ensuring that high-quality reads were retained while effectively eliminating low-quality sequences and adapters.

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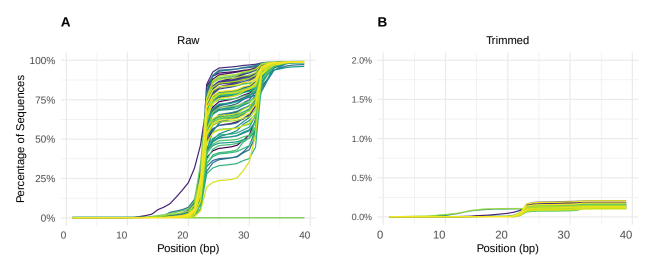


Figure 4.2: Adapters Content (%) across all bases before (A) and after (B) using *fastp* (version 0.23.4)

Figure 4.3 shows the analysis of the mean quality value of sequences across all bases, both before (A) and after (B) processing with fastp is evident from that the sequences from all samples consistently fall within an acceptable quality range, with mean Phred scores exceeding 30. This indicates a high base-calling accuracy, with an error probability of less than 0.1%, ensuring robust data integrity throughout the process. However, there is a noticeable drop in quality towards the extremes of the readings, particularly at position 31 bp. This decline can be attributed to the effects of adapter trimming in small RNA sequencing (smRNA-seq). Even though adapters have been removed, if they were present at the end of the sequence, their trimming may result in reduced quality for the subsequent bases. Bases near the adapter might have lower quality, leading to a set of bases that, while technically valid, originate from a lower-quality region. Additionally, it is common in smRNA-seq to encounter contaminants from various sources, such as rRNA or tRNA, which introduce background noise and can further diminish the quality of the readings obtained.

The Figure 4.3 illustrates the metric of the average quality value of each sequence across all samples. Before processing with fastp (C), the Phred scores consistently exceed 30, with most samples achieving scores around 35. This indicates that the average sequence quality is optimal, ensuring high reliability and low error rates. However, after trimming (D) a slight decrease in both Phred scores and read counts is observed. This reduction in the number of reads is expected after strict preprocessing with fastp, especially if contaminated or low-quality reads were present.

In Figure 4.3 (*E*), prior to trimming, the GC content distribution does not follow a normal pattern. Multiple peaks are observed, likely due to the presence of adapters, which typically have a fixed GC content that generates specific peaks. Additionally, since this is a *small RNA-seq* experiment, contamination from rRNA or tRNA is common, as these molecules often have a different GC content compared to other small RNAs. After processing with *fastp* (*F*), the GC content decreases, indicating that most of these contaminant sequences have been removed, leaving of reads with anomalies. This remaining may correspond to sequences that are difficult to classify or contaminants that are not easily eliminated by *fastp*.

Additionally, no issues with the presence of ambiguous bases (Ns) are detected in any of the samples before and after trimming. Figure 4.3 (G and H).

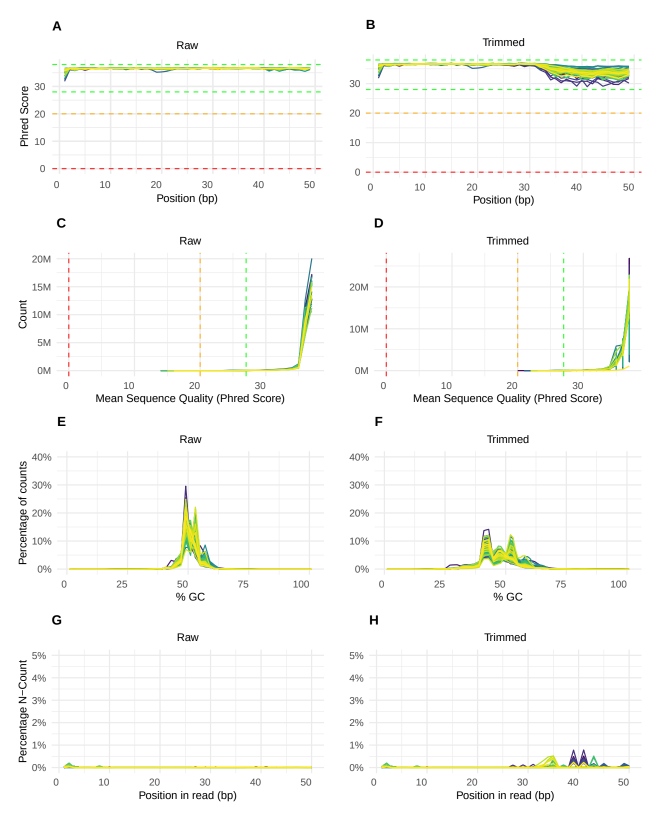


Figure 4.3: Quality Control Analysis. A and B: Mean quality values of sequences across all bases after and before using fastp (v0.23.4); C and D: Per Sequence Quality Scores across all bases after and before using fastp; E and F: Per Sequence GC Content Raw after and before using fastp; G and H: Read N content after and before using fastp.

However, the distribution of sequence lengths is irregular across all samples Figure 4.4. After trimming, the majority of sequences cluster around a length of 20-25 nucleotides, although a smaller subset of sequences with lengths between 29-32 nucleotides is also observed. This indicates the presence of different types of small RNAs. Generally, miRNAs are found in the range of 20-24 bp, with a peak at 22 bp potentially indicating an abundant population of miRNAs. In contrast, tRNAs can vary in length but are commonly found at lengths that may include higher peaks, such as at 31 bp. This may reflect the presence of tRNAs derived from the degradation of double-stranded RNA or transposons.

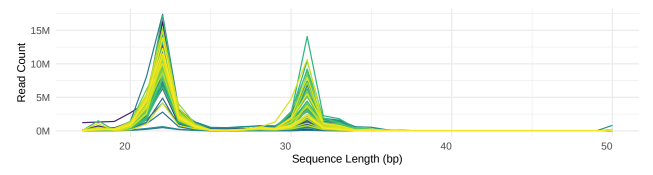


Figure 4.4: FastQC: Sequence Length Distribution

In the Figure 4.5 shows the total number of reads per sample is around 25 million, both in Raw and Trimmed, with means of 25.47 and 25.23 million, respectively. The interquartile ranges (IQR) of these measurements are similar, approximately 4.2 million, reflecting a high consistency among the samples.

Regarding duplicate reads, they dominate the data, with means of 25.10 million in Raw and 24.95 million in Trimmed, and an IQR of about 4.2 million in both conditions. This indicates that more than 90% of the sequences are duplicated, a high value but expected in smRNAseq samples, given the nature of the short reads of 20 to 24 nucleotides.

In contrast, unique reads are significantly less frequent, with means of 0.38 million in Raw and 0.28 million in Trimmed. The interquartile ranges for these measurements are also low, around 0.13 million in Raw and 0.10 million in Trim.

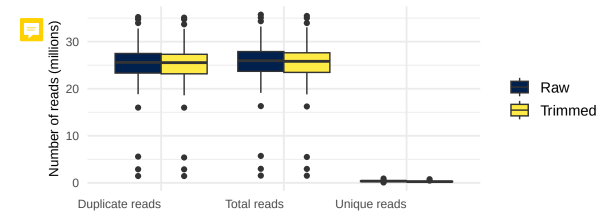


Figure 4.5: Number of reads from small RNA-seq. Total reads before and after trimming of adapters

## 4.3 miRNA Quality Control

The *nf-core/smrnaseq* pipeline performs a quality analysis specific to smkNAseq data using *miR-Trace*. This analysis assesses sequencing quality, identifies the presence of miRNA and undesired sequences from tRNA, rRNA, or Illumina artifact sequences, and identifies clade-specific miRNA profiles based on a comprehensive catalog of previously identified miRNA families.

In the annotation step Figure 4.6, the mapped reads against reference databases revealed that a mean of 65.06% of analyzed sequences per sample corresponded to miRNA precursors, with a range of 22.51% to 91.77%. Other categories included 21.16% tRNA sequences, 9.16% unknown sequences, 4.55% rRNA sequences, and 0.05% artifacts.

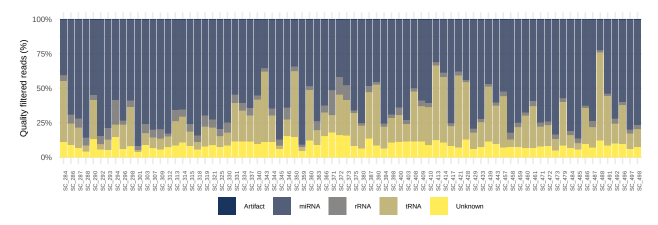


Figure 4.6: miRTrace (v1.0.1) Analysis: RNA Categories

During the contamination assessment step Figure 4.7, mapping miRNA precursor sequences against the clade-specific miRNA catalog showed that a mean of 94.14% of analyzed sequences belonged to the human category, with a range of 67.21% to 99.94%. Minor contributions from other clades, such as Rodentia (2.07%), Dicots (2.73%), Insects (0.58%), and Monocots (0.38%), were detected at low proportions. These identifications could result from contamination, such as incorrect index assignment during sample demultiplexing, or may have a biological origin.

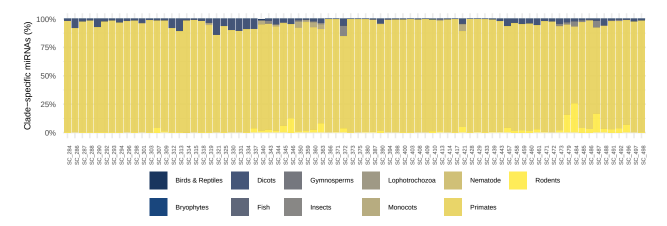


Figure 4.7: miRTrace (v1.0.1) Analysis: Contamination Check

### 4.4 miRNA Quantification

In this step of the *nf-core/smrnaseq pipeline*, the alignment of reads is conducted sequentially against databases of mature miRNAs, precursor miRNAs, and a combined database of both. These alignments enable the identification and quantification of miRNAs. The statistics obtained for mature miRNAs, precursor miRNAs, and the combination of both against the reference genome are presented in Table 4.3. On average, 52.07% of the mature miRNAs were mapped, with a range between 17.72% and 79.75%.

For precursor miRNAs, an average of 32.33% of sequences was identified, ranging from 6.31% to 68.83%. Subsequently, the reads of mature and precursor miRNAs were aligned against a reference genome, not for miRNA identification and quantification but as a quality control for the sequences. In this regard, the mean percentage of aligned reads was around 43% of the total reads. Across the sample set, minimum values of 10.28% and maximum values of 73.31% were observed. For more details, the metrics for each sample can be found in https://github.com/joshoandres13/miRNAs.

Table 4.3: Descriptive statistics of alignment with samtools of all samples. *TM*: Mean of Total Mapped (reads); *TU*: Mean of Total Unmapped (reads): *Mean M*: Mean Mapped (%); *Min M*: Min Mapped (%)

Group	TM	TU	Mean M	Max M	Min M	
mature	13166950	12194855	52.07	79.75	17.72	F
mature_hairpin	3644435	8836242	32.33	68.83	6.31	
mature_hairpin_genome	4334118	4565712	43.87	73.31	10.28	

### 4.5 IsomiR Annotation

Mirtop (v0.4.28) was used for the annotation of miRNAs and isomiRs. In figure Figure 4.8, the mean isomiR read counts are presented, which refer to an average calculation that helps describe how the reads of isomiRs are distributed within a dataset. Among the annotated isomiR sequences per sample, the reference miRNA accounts for 96.51%, with ranges varying from 88.70% to 98.13%. On the other hand, the distributions of isomiR variants are as follows: 3' Isoform (1.36%), 3' Addition (1.23%), 5' Isoform (0.64%), SNVs in the Central Offset Region (0.05%), SNVs in the Central Region (0.06%), Supported SNVs in the Central Region (0.05%), and SNV in Seed Region (0.05%), all of which are below 1.40%.

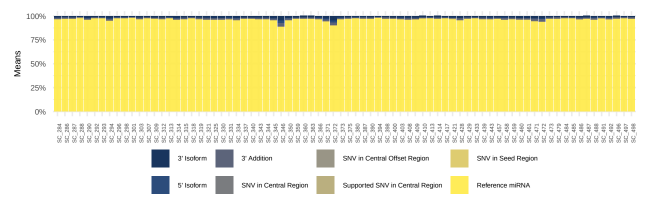


Figure 4.8: Annotation of miRNAs and isomiRs with mirtop (v0.4.28): Mean isomiR read counts

## 4.6 Differential expression analysis



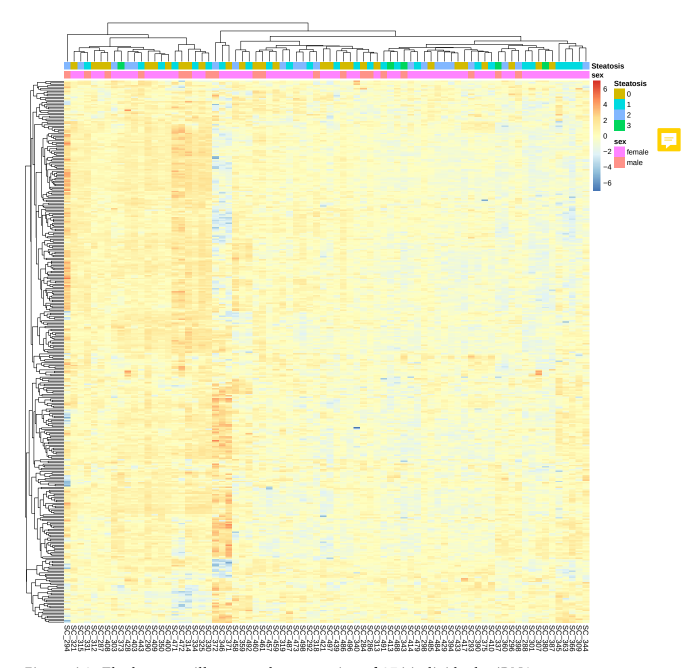


Figure 4.9: The heatmap illustrates the expression of 374 individual miRNA sequences across the analyzed samples. In this visualization, shades of red indicate increased miRNA expression, whereas shades of blue denote reduced or absent miRNA expression. Although a substantial number of miRNAs were identified, no distinct grouping patterns emerged among the analyzed samples, suggesting heterogeneity in miRNA expression profiles across the dataset.

Differential expression analysis is a bioinformatics and statistical technique used to identify genes, proteins, or other biomolecules that exhibit significant differences in expression levels between two or more biological conditions. For this analysis, the sequences of miRNAs of reference were

utilized annotated in the previous analysis.

Under the hood of *DESeq2*, we applied the *regularized logarithm* (*rlog*) transformation for normalization, which incorporates a prior on the sample differences<sup>[134]</sup>. This step is crucial because the standard logarithmic transformation can be sensitive to low expression values, whereas *rlog* addresses this issue by introducing regularization that helps stabilize variability between samples. This method facilitates seamless integration into subsequent analyses, as demonstrated in Figure 4.9.The figure shows a total of 374 miRNAs were found to be expressed differentially in the overall cohort when patients (samples) were categorized by sex and steatosis.

Since we were primarily interested in the transcriptional changes that may precede the initiation of the steatosis process, we focused on contrasting individuals across different levels of steatosis. In this experiment, the goal is to identify miRNAs that are expressed differentially at various levels of steatosis. In our differential expression analysis using *DESeq2* with the LRT model, we initially identified 169 miRNAs as upregulated and 205 downregulated in the subcutaneous white adipose tissue (scWAT) of individuals according their steatosis degree. Comparing these four groups and applying a false discovery rate (FIT) cutoff of less than 0.05, the number of statistically significant differentially expressed miRNAs was greatly reduced to 2 (Figure 4.10). Of these, we found *hsa-miR-372-3p* is upregulated and *hsa-miR-144-3p* is downregulated (Table 4.4).

Table 4.4: Differentially expressed miRNAs in subcutaneous white adipose tissue (scWAT). *FDR* : False Discovery Rate

	log2FoldChange	pvalue	FDR
hsa-miR-144-3p		p< 0.0002	0.04
hsa-miR-372-3p	1.085	p< 0.0002	0.04

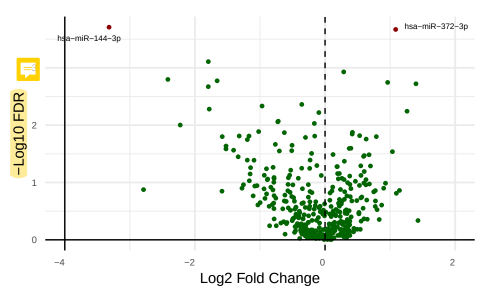


Figure 4.10: Plot showing differentially expressed miRNAs (red) in subcutaneous white adipose tissue (scWAT) according to the four groups of steatosis

Following their identification, we further analyzed the expression patterns of these two differentially expressed miRNAs by representing their normalized counts in boxplots across the four

F

steatosis groups Figure 4.11. This approach allowed us to observe how the expression of these significant miRNAs varies across different degrees of steatosis. The boxplots highlight the consistency of expression changes across groups, reinforcing the revance of these miRNAs in the context of steatosis severity. The miRNA hsa-miR-144-3p demonstrates a significant decrease in expression as steatosis levels increase, whereas hsa-miR-372-3p exhibits a trend of progressively higher expression across the varying levels of steatosis.

Despite observing a trend of increasing expression of hsa-miR-372-3p with rising levels of steatosis, the results from the Kruskal-Wallis test did not achieve statistical significance (p=0.28) when comparing the different states of steatosis. This suggests that, while there may be a tendency toward increased expression of the miRNA with the progression of steatosis, a larger sample size or an alternative approach is needed to draw firmer conclusions about its role in the progression of the disease.

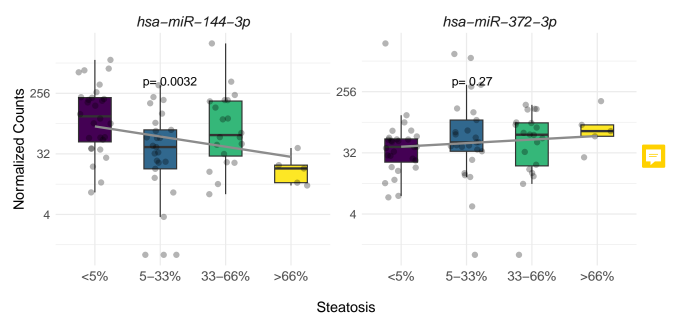


Figure 4.11: Boxplots of the differentially expressed miRNAs in subcutaneous white adipose tissue (scWAT) according to the four groups of steatosis. Each box represents the interquartile range (IQR) of the normalized counts, with the line inside the box indicating the median. The whiskers extend to show the range of the data, excluding outliers, which are displayed as individual points. p: p-value fot the Kruskal–Wallis test for the comparison between groups.

## 4.7 Target mRNA Selection and Validation

### 4.7.1 hsa-miR-372-3p

Potential target genes of the selected miRNAs were identified using the *multiMiR* package. This analysis included only validated interaction data, ensuring the biological relevance of the identified mRNAs. A subsequent filtering process was applied to select targets based on the type of experimental validation (e.g., luciferase assays, Western blot, or qRT-PCR) and functional support. As shown in Table 4.5, 24 target genes were identified after filtering.

Table 4.5: Selected interactions after filtering by database, experiment type (including luciferase assays, Western blot, or qRT-PCR), functional support (Functional MTI) and validated type for *hsa-miR-372-3p*.

Gene	Ensembl Identifier	Туре
LATS2	ENSG00000150457	validated
TGFBR2	ENSG00000163513	validated
NFIB	ENSG00000147862	validated
CDKN1A	ENSG00000124762	validated
VEGFA	ENSG00000112715	validated
TNFAIP1	ENSG00000109079	validated
TRPS1	ENSG00000104447	validated
MBNL2	ENSG00000139793	validated
RHOC	ENSG00000155366	validated
NR4A2	ENSG00000153234	validated
ERBB4	ENSG00000178568	validated
CDK2	ENSG00000123374	validated
LEFTY1	ENSG00000243709	validated
BTG1	ENSG00000133639	validated
WEE1	ENSG00000166483	validated
CCNA1	ENSG00000133101	validated
DKK1	ENSG00000107984	validated
PHLPP2	ENSG00000040199	validated
ATAD2	ENSG00000156802	validated
ADAMTS9	ENSG00000163638	validated
CADM2	ENSG00000175161	validated
ZBTB7A	ENSG00000178951	validated
TXNIP	ENSG00000265972	validated
KLF13	ENSG00000275746	validated

The target genes identified through multiMiR were subjected to functional enrichment analysis using KEGG (Kyoto Encyclopedia of Genes and Genomes) pathways<sup>[130]</sup>. The KEGG pathway enrichment analysis identified five significantly enriched pathways (p < 0.05) (Figure 4.12). The most prominent pathways include:

- *Cellular senescence* (KEGG ID: hsa04218): associated with cellular aging and the permanent arrest of the cell cycle in response to genetic damage, oxidative stress, or oncogenic signals.
- *Cell cycle* (KEGG ID: hsa04110): involved in the regulation and progression of the cell cycle, including critical checkpoints that ensure the fidelity of cell division.
- *Hepatitis B* (KEGG ID: hsa05161): related to infection by the hepatitis B virus and the inflammatory and carcinogenic processes it can trigger in the liver.
- *PI3K-Akt signaling pathway* (CGG ID: hsa04151): a key pathway regulating cellular processes such as proliferation, survival, and metabolism, frequently deregulated in cancer.
- *Pancreatic cancer* (KEGG ID: hsa05212): associated with molecular and cellular alterations characteristic of this cancer type, including aberrant signaling and apoptosis resistance.

These pathways are primarily linked to essential processes such as cell cycle regulation, intracellular

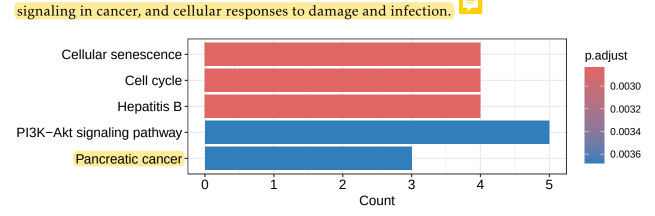


Figure 4.12: Top 5 Significant KEGG Pathways of hsa-miR-372-3p

### 4.7.2 hsa-miR-144-3p

Table 4.6 shows 22 target genes were identified for *hsa-miR-372-3p*, filtered by validated interactions and experimental evidence.

Table 4.6: Selected interactions after filtering by database, experiment type (including luciferase assays, Western blot, or qRT-PCR), functional support (Functional MTI) and validated type for *hsa-miR-144-3p*.

Gene	<b>Ensembl Identifier</b>	Type
NOTCH1	ENSG00000148400	validated
PLAG1	ENSG00000181690	validated
ZEB1	ENSG00000148516	validated
ZEB2	ENSG00000169554	validated
IRS1	ENSG00000169047	validated
MAP3K8	ENSG00000107968	validated
EZH2	ENSG00000106462	validated
APP	ENSG00000142192	validated
PTGS2	ENSG00000073756	validated
MET	ENSG00000105976	validated
ETS1	ENSG00000134954	validated
TGFB1	ENSG00000105329	validated
CFTR	ENSG00000001626	validated
FGG	ENSG00000171557	validated
MTOR	ENSG00000198793	validated
SMAD4	ENSG00000141646	validated
NFE2L2	ENSG00000116044	validated
PBX3	ENSG00000167081	validated
TTN	ENSG00000155657	validated
TUG1	ENSG00000253352	validated
PTEN	ENSG00000284792	validated
XIST	ENSG00000229807	validated

The target genes identified through multiMiR were subjected to functional enrichment analysis using KEGG where five significantly enriched pathways (p < 0.05) (Figure 4.13). The most prominent pathways include:

- *MicroRNAs in cancer* (KEGG ID: hsa05206): Highlighting the role of miRNAs in the regulation of gene expression, particularly in pathways associated with tumorigenesis and cancer progression.
- *Hepatocellular carcinoma* (KEGG ID: hsa05225): Specifically linked to liver cancer and the molecular mechanisms underlying its development and progression.
- FoxO signaling pathway (KEGG ID: hsa04068): A critical pathway regulating oxidative stress response, apoptosis, and metabolism, which plays a significant role in cancer and aging.
- *Gastric cancer* (KEGG ID: hsa05226): Associated with molecular and cellular alterations characteristic of gastric tumorigenesis.
- *Cellular senescence* (KEGG ID: hsa04218): Related to aging and the permanent arrest of the cell cycle due to stress signals or damage.

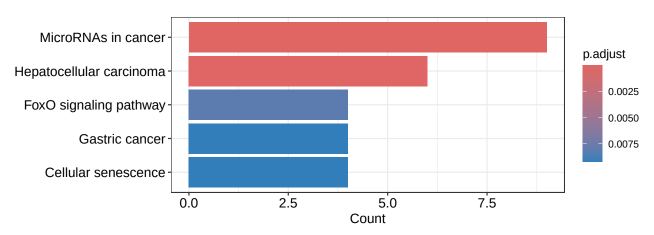


Figure 4.13: Top 5 Significant KEGG Pathways of hsa-miR-144-3p

These pathways reflect key biological processes, including tumorigenesis, stress response, and cell cycle regulation, underscoring the importance of miRNAs in these mechanisms. The findings provide valuable insights into potential molecular interactions and pathways relevant to the studied system.

# 4.8 The hsa-miR-372-3p and par-miR-144-3p Modulate Lipid Metabolism Genes in a HepG2 Steatosis Model.

The gene expression was evaluated in HepG2 cells transfected for 24 happens and then treated with Oleic Acid (OA) for an additional 24 hours to assess the impact of miRNA a cell model mimicking hepatic steatosis. Specifically, we investigated the relative expression of *PPARG* (Peroxisome Proliferator Activated Receptor Gamma), a key regulator of adipogenesis and insulin sensitivity; *PNPLA2* (Patatin Like Phospholipase Domain Containing 2), which encodes the Adipose Triglyceride Lipase

(ATGL) involved in triglyceride breakdown; DGAT2 (Diacylglycerol O-Acyltransferase 2), a critical enzyme for triglyceride synthesis; FAS (Fas Cell Surface Death Receptor), a mediator of apoptosis and cellular stress; and ACACA (Acetyl-CoA Carboxylase Alpha), the rate-limiting enzyme in de novo fatty acid synthesis. These genes play pivotal roles in lipid metabolism and, to a lesser extent, glucose metabolism, making them relevant targets for understanding the molecular mechanisms underlying hepatic steatosis.

The hsa-miR-372-3p mimic significantly downregulated the mRNA expression of the target gene ACACA (-91.40%  $\pm$  3.39; p < 0.05) in OA-treated HepG2 cells. A similar, although not statistically significant, decrease was observed in the mRNA levels of FAS (-90.50%  $\pm$  4.30; p < 0.093), suggesting a trend towards reduced expression. In contrast, the mRNA levels of DGAT2 (-73.7%  $\pm$  6.70; p = 0.462) and PNPLA2 (-55.10%  $\pm$  19.56; p = 0.243) did not show significant downregulation. Notably, the mRNA levels of PPARG remained unchanged compared to the negative control cells (Figure 4.14).

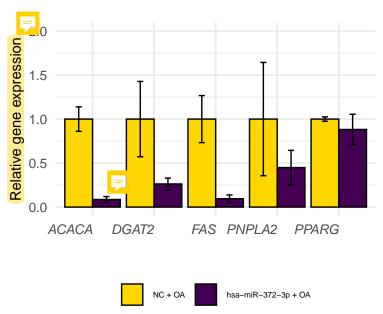


Figure 4.14: Gene expression analysis of HepG2 cells transfected with *hsa-miR-372-3p* mimic and treated with oleic acid to mimic in vitro hepatic steatosis was performed. The mRNA levels of protein-coding genes (*ACACA*, *DGAT2*, *FAS*, *PNPLA2*, *PPARG*), which are involved in glucose and lipid metabolism, were assessed. HepG2 cells were transfected with 50 nM of mirVana mimic *hsa-miR-372-3p* (5'-AAAGUGCUGCGACAUUUGAGCGU-3') along with a randomized sequence as a control (Negative Control 1). Twenty-four hours post-transfection, the cells were treated with oleic acid (0.5 mM) for 24 hours. The results are presented as mean relative gene expression ± standard error of the mean (SEM) (n = 3). Abbreviations: NC (negative control), OA (oleic acid), *ACACA* (Acetyl-CoA Carboxylase Alpha), *DGAT2* (Diacylglycerol O-Acyltransferase 2), *FAS* (Fatty Acid Synthase), *PNPLA2* (Patatin-like phospholipase domain-containing protein 2), *PPARG* (Peroxisome Proliferator Activated Receptor Gamma).

On the other hand, inhibition of hsa-miR-144-3p upregulated the expression of the putative target genes. The relative gene expression levels of the evaluated targets in HepG2 cells transfected and treated with Oleic Acid (OA) were as follows: ACACA (96.80%  $\pm$  53.79; p = 0.186), FAS (135%  $\pm$  160.34; p = 0.429), DGAT2 (32.74%  $\pm$  24.13; p = 0.128), PNPLA2 (48.83%  $\pm$  14.32; p = 0.140), and

PPARG (64.09% ± 41.25; p = 0.239) (Figure 4.14). Although the expression levels varied across genes, none of these changes were statistically significant (p > 0.05). These findings suggest that the regulatory effect of hsa-miR-144-3p on these genes may be limited under the experimental conditions employed.

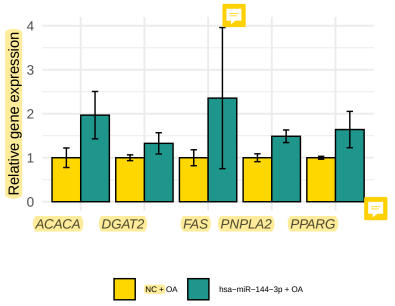
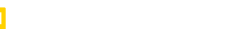


Figure 4.15: Gene expression analysis of HepG2 cells transfected with hsa-miR-144-3p inhibitor and treated with oleic acid to mimic in vitro hepatic steatosis was performed. The mRNA levels of protein-coding genes (ACACA, DGAT2, FAS, PNPLA2, PPARG), which are involved in glucose and lipid metabolism, were assessed. HepG2 cells were transfected with 50 nM of mirVana mimic hsa-miR-144-3p (5'-UACAGUAUAGAUGAUGUACU-3') along with a randomized sequence as a control (Negative Control 1). Twenty-four hours post-transfection, the cells were treated with oleic acid (0.5 mM) for 24 hours. The results are presented as mean relative gene expression  $\pm$  standard error of the mean (SEM) (n = 3). Abbreviations: NC (negative control), OA (oleic acid), ACACA (Acetyl-CoA Carboxylase Alpha), DGAT2 (Diacylglycerol O-Acyltransferase 2), FAS (Fatty Acid Synthase), PNPLA2 (Patatin-like phospholipase domain-containing protein 2), PPARG (Peroxisome Proliferator Activated Receptor Gamma).

## Chapter 5

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This study offers new insights about the role of miRNAs derived from subcutaneous adipose tissue in regulating metabolic processes related to hepatic steatosis. Through the analysis of smRNA sequencing data, differential miRNAs linked to key metabolic pathways were identified that may play a role in liver fat accumulation and the progression of hepatic steatosis in obese individuals. These findings address a critical gap in understanding the role of miRNAs in adipose liver metabolic crosstalk, which has implications for targeted interventions in hepatic steatosis.

Building on these insights, we specifically focused on identifying differentially expressed miRNAs associated with hepatic steatosis in subcutaneous adipose tissue. Although the preliminary analysis showed that hyperlipidemia had a significant *p* value, we decided not to include it in the final model for several reasons.

To maintain c ty in our analysis, our primary objective is to understand the direct relationship between hepatic steatosis and the expression of specific miRNAs in subcutaneous adipose tissue. The inclusion of hyperlipidemia, while relevant in other contexts, could complicate the interpretation of the results by introducing a variable that acts as a confounder. For hyperlipidemia, circulating miRNAs have been identified that are found in biological fluids (extracellular), while our goal is to identify miRNAs specific to subcutaneous adipose tissue (intracellular)<sup>[135,136]</sup>. This distinction is crucial, as it could divert attention from our central analysis and hinder the identification of miRNAs.

In this context, previous studies have identified miRNAs with TaqMan® MicroRNA Assays in adipose tissue from a cohort of fifteen overweight and obese individuals, investigating whether miRNA expression in human adipose tissue is specific to subcutaneous adipose tissue (scWAT) and intra-abdominal fat depots, as well as its association with parameters of obesity and glucose metabolism. However, this study has important limitations, such as the small sample size, which restricts the generalization of findings, and the inability to exclude that specific characteristics of the study population may have influenced the observed correlations. In contrast to these previous studies, to our knowledge, our study is the first to identify differentially expressed miRNAs in subcutaneous with sm-RNA seq in scWAT from 78 obese individuals within the FATe cohort, which encompasses different scales of hepatic steatosis, and being validated in vitro through a model of human hepatocyte cells. Our use of smRNA sequencing not only addresses the issue of sample size but also provides a more comprehensive profile of miRNA expression compared to TaqMan® assays.

Understanding the expression patterns of miRNAs across different tissues is crucial for gaining insight into both normal development and disease progression in those tissues<sup>[138]</sup>. miRNA expression profiles are altered in many diseases including steatosis. *hsa-miR-372-3p* has been identified as a potent miRNA associated with tumorigenesis<sup>[139]</sup>. It has been shown to promote tumor formation in testicular germ cell tumors<sup>[140,141]</sup>, lung cancer<sup>[142]</sup>, parathyroid carcinoma<sup>[139]</sup>, and osteosarcoma<sup>[143]</sup>.

Aditionally, recent studies have highlighted the role of *hsa-miR-372-3p* in the context of MASH. Specifically, hyperglycemia, lipid accumulation, and fructose treatment lead to increased expression of adipocyte enhancer binding protein 1 (*AEBP1*) in liver cells. *AEBP1* is a multifunctional protein involved in a variety of biological processes, including adipogenesis, cell differentiation, and macrophage cholesterol homeostasis<sup>[144]</sup>. Its emerging role in the formation of collagen-rich tissues and the development of tissue fibrosis is also significant in the context of MASH<sup>[145]</sup>. Also, *AEBP1* is a key component of a group of genes specifically dysregulated in the fibrotic processes associated with MASH<sup>[86]</sup>. Notably, the levels of *hsa-miR-372-3p* is found to be lower in patients with MASH and advanced fibrosis, where this miRNA interact functionally with *AEBP1* to reduce its expression. This suggests that the dysregulation of this miRNA may contribute to the progression of hepatic steatosis and its associated complications, further linking *hsa-miR-372-3p* to the metabolic dysregulation observed in MASH<sup>[145]</sup>.

Through transcriptomic and pathway analysis, we identified additional pathways already experimentally validated in which *hsa-miR-372-3p* is implicated. These include cell cycle regulation, intracellular signaling in cancer, and cellular responses to damage and infection. While these pathways support its role in tumorigenesis, they also suggest a broader implication of *hsa-miR-372-3p* in metabolic disorders, particularly in processes associated with liver fat accumulation and fibrosis progression.

One of the pathways identified is the PI3K-Akt signaling pathway, which plays a central role in regulating cell survival, metabolism, and inflammation. Dysregulation of this pathway has been linked to MASLD, particularly in the context of insulin resistance and abnormalities in lipid metabolism<sup>[146]</sup>. These findings align with prior studies showing that hsa-miR-372-3p enhances cell growth and proliferation by counteracting p53-mediated CDK inhibition and activating the Wnt/ $\beta$ -catenin signaling pathway<sup>[147,148]</sup>. Together, these pathways highlight a dual role for hsa-miR-372-3p in tumorigenesis and metabolic regulation.

Moreover, its downregulation in patients with advanced fibrosis suggests a potential regulatory role in the progression of hepatic steatosis to more severe conditions such as MASLD and hepatocellular carcinoma (HCC)<sup>[139,145,149]</sup>.

In this study, *hsa-mir-372-3p* was identified to be over-expressed in scWAT of obese patients with hepatic steatosis. However, it was significantly downregulated in MASH patients with advanced fibrosis compared to those with normal liver histology<sup>[145]</sup>. This change in expression underscores the potential role of *hsa-miR-372-3p* in the transition from simple steatosis to more severe liver conditions.

In our *in vitro* studies using HepG2 cells, the mimic of hsa-miR-372-3p effectively downregulated the expression of ACACA, a key gene involved in lipogenesis. This finding aligns with the observed trends towards reduced mRNA levels of FAS and indicates a potential regulatory role of hsa-miR-372-3p in lipid metabolism. Although genes such as DGAT2 and PNPLA2 showed downregulation trends, they did not reach statistical significance, nor did  $PPAR\gamma$ , which remained unchanged.





These results suggest that *hsa-miR-372-3p* may influence hepatic lipid metabolism and contribute to the progression of hepatic steatosis, indicating its potential relevance in MASH pathogenesis. Aditionally, the observed downregulation of key lipogenic genes, such as *ACCA* and *FAS*, reinforces the hypothesis that alterations in miRNA expression can significantly impact metabolic processes in the liver. *ACCA* is essential for converting acetyl-CoA into malonyl-CoA, a critical step in fatty acid synthesis, while FAS is responsible for synthesizing long-chain fatty acids.

Interestingly, the downregulation of *ACACA* and *FAS* observed in our study could be partially mediated by the PI3K-Akt pathway, as this signaling cascade is closely tied to lipogenesis and lipid metabolism. This suggests a possible interaction between *hsa-miR-372-3p* and key metabolic signaling pathways, further underscoring its central role in hepatic lipid regulation.

In conclusion, *hsa-miR-372-3p* has been proposed as a regulatory miRNA in MASH<sup>[150]</sup>, and the observed downregulation of critical lipogenic genes highlights the need for further investigation into the functional implications of this miRNA and its targets in the context of liver disease. These findings also open avenues for therapeutic strategies targeting *hsa-miR-372-3p*, potentially providing a novel approach to managing lipid accumulation and fibrosis in MASLD.

hsa-mir-144-3p belongs to the MIR-144 family, alongside its counterpart, the guide strand hsa-mir-144-5p. Although hsa-mir-144-3p is technically classified as the passenger strand, studies have demonstrated that it is often more abundant than the guide strand and plays a significant functional role in the organism<sup>[151]</sup>. This highlights the need to consider both strands when analyzing the biological impact of the MIR-144 family. This miRNAs is typically secreted into the extracellular space<sup>[152]</sup> and is frequently expressed in liver pathology<sup>[153]</sup>. Furthermore, MIR-144 has been implicated in the progression of various cancers<sup>[154]</sup>. In the context of liver-related metabolic diseases, these miRNAs have been associated with pathologies such as chronic hepatitis B and HCC<sup>[153,155]</sup>.

In our analyses, *hsa-miR-144-3p* emerged as a differentially expressed miRNA, suggesting its relevance in the context of hepatic steatosis. Previous studies have reported that this miRNA is involved in the progression of insulin resistance, which is closely linked to the development of hepatic steatosis and other metabolic disorders<sup>[86]</sup>. Additionally, *hsa-miR-144-3p* has been shown to inhibit nuclear factor erythroid 2-related factor 2 (*Nrf2*) expression and downstream antioxidant genes, which in turn reduces glucose consumption and exacerbates zinc-induced insulin resistance in HepG2 cells<sup>[156]</sup>.

Furthermore, previous studies found that the upregulation of *hsa-miR-144-3p* promotes hepatocyte proliferation by suppressing the expression of the *Txnip* gene (thioredoxin-interacting protein), which is involved in glucose metabolism and inhibition of cell proliferation. This suggests that this miRNA could play a crucial role in liver regeneration, making it a promising therapeutic target for hepatic steatosis and related conditions<sup>[157]</sup>. These findings indicate that *hsa-miR-144-3p* is involved in liver regeneration and gluconeogenesis, making it an interesting candidate for investigating its role in hepatic steatosis.

Through transcriptomic and pathway analysis, we identified additional pathways that have been experimentally validated in which hsa-miR-144-3p is implicated. These include pathways related to tumorigenesis, stress response, cell cycle regulation, and hepatic metabolism. Among these pathways, the FoxO signaling pathway has been established as a fundamental regulator in hepatic metabolism, not only for its role in gluconeogenesis and lipid balance but also for its interaction with CCAAT/enhancer-binding protein alpha ( $C/EBP\alpha$ ), a crucial regulator of adipogenesis. This highlights its relevance in metabolic homeostasis and preadipocyte differentiation [158].

Moreover, hsa-miR-144-3p is downregulated in patients with HCC<sup>[155]</sup>. Studies have reported that the suppression of this miRNA promotes proliferation and metastasis by targeting E2F transcription factor 3 (E2F3), an oncogene involved in the regulation of cell proliferation<sup>[154,159]</sup>. In our study, we found that the expression of hsa-miR-144-3p significantly decreases as the level of steatosis increases in scWAT, suggesting a potential link between this miRNA's regulatory function and the progression of hepatic metabolic disorders. These findings underscore the importance of hsa-miR-144-3p as a candidate for therapeutic targeting in HCC and related metabolic cortagions.

Furthermore, inhibition of *hsa-miR-144-3p* led to an upregulation of its putative target genes in HepG2 cells treated with oleic acid. While the expression levels of the evaluated targets displayed variations, the changes did not reach statistical significance, indicating that the influence of *hsa-miR-144-3p* on these genes might be constrained under the experimental conditions utilized. This suggests that, despite the potential regulatory role of hsa-miR-144-3p, the specific context and cellular environment can significantly impact its effectiveness in modulating gene expression.

The potential of hsa-miR-144-3p extends beyond its role in liver metabolism and cancer progression. It has been proposed as a promising therapeutic target and a diagnostic/prognostic tool in various cancers [[154];[160]]. Specifically, Rowe et al. [153] suggest that hsa-miR-144 could serve as a valuable biomarker in blood, plasma, or serum for chronic hepatitis B. This highlights the versatility of hsa-miR-144-3p not only in understanding disease mechanisms but also in facilitating early diagnosis and targeted therapies, reinforcing its importance in both clinical and research settings.

One of the main strengths of this study lies in the use of RNA-seq analysis to identify specific miRNAs scWAT, an innovative methodology that provides detailed and accurate information about their regulation compared to previous studies. Furthermore, this is the first study to differentially characterize miRNAs in a large cohort of 78 obese patients, significantly enhancing the generalizability of the results. The identification of hsa-miR-372-3p and hsa-miR-144-3p as key regulators in metabolic pathways related to lipogenesis and the progression of hepatic steatosis, along with their functional validation in in vitro models, reinforces the biological and clinical relevance of the findings. This approach also provides new insights into the metabolic interaction between scWAT and the liver, highlighting the potential of these miRNAs as biomarkers and therapeutic targets in the management of obesity and its hepatic complications.

Despite these strengths, the study has several limitations. First, the relatively small sample size may have reduced the statistical power to detect additional miRNAs, highlighting the need for larger cohorts to validate our findings and potentially uncover more miRNAs associated with liver steatosis. Second, while HepG2 cells served as a prevenient model for our in vitro studies, their limitations in fully mimicking adipocyte biology restrict the scope of our findings. An adipocyte model could have provided a more physiologically relevant system for studying the interplay between hsa-miR-144-3p and hsa-miR-372-3p and metabolic pathways. However, due to time constraints, this alternative was not feasible within the framework of this study. Future research should address these limitations by incorporating diverse cellular models and larger datasets to strengthen the generalizability and applicability of our conclusions.

This study highlights the role of miRNAs expressed in subcutaneous adipose tissue in the regulation of metabolic processes related to hepatic steatosis. We identified *hsa-miR-372-3p* and *hsa-miR-144-3p* as key regulators in metabolic pathways such as lipogenesis and confirmed their functional impact in in vitro models of hepatocytes. These findings reinforce their potential as biomarkers and therapeutic targets for the management of obesity and its hepatic complications.

# Chapter 6

# **Conclusions**



- 1. MiRNAs derived from subcutaneous adipose tissue play a fundamental role in regulating metabolic processes related to hepatic steatosis, suggesting that their expression may influence the progression of liver diseases.
- 2. Specific miRNAs, such as *hsa-miR-372-3p* and *hsa-miR-144-3p*, have been identified as potential biomarkers for the diagnosis and prognosis of hepatic steatosis and its complications, opening new opportunities for personalized medicine.
- 3. This study enhances the understanding of the relationship between adipose tissue and the liver, highlighting how miRNAs may be involved in regulating lipid accumulation and disease progression.
- 4. Further investigations are needed to validate these findings and explore the therapeutic potential of miRNAs in treating hepatic steatosis and other metabolic disorders.
- 5. The results of this study contribute to the current understanding of metabolic health, suggesting that modulation of miRNAs may offer novel strategies to address health issues related to obesity and liver diseases.

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