

# Bridging the gaps of microRNAs in obesity

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## **Chapter 5**

### Summary and General Discussion

In this thesis we have discussed multiple ways by which microRNAs emerged as powerful regulators in the path towards obesity and metabolic disease. Although we have not completely managed to bridge the gaps between the benches in the labs and clinical practice at the bedsides of patients, the efforts made in this thesis form the base of the first footsteps in this direction. Current treatment modalities, ranging from life style and dietary interventions to bariatric surgery with high risk of anastomosis leakage and dumping syndrome, did still not manage to dampen the rise in overweight and obesity. Curtailing the consequences of obesity for instance by antidiabetic, cholesterol lowering or antihypertensive therapy is merely an indication that the damage has already been done. As obesity is increasingly prevalent worldwide and known for its association with many comorbidities, such as insulin resistance, glucose intolerance, dyslipidemia and hypertension, the magnitude of the problem is expected to increase even more in the future underlining the need for the development of novel molecular paradigms revolutionizing its approach. To this end, the field of non-coding RNAs, in particular the small microRNAs, that regulate physiological functions in cells and tissues by binding to mRNAs inducing gene repression, accessible for external modulation, are tremendously attractive. The studies presented in this thesis were aimed at exploring the usage of circulating microRNAs to detect obesity and monitor progression and severity of metabolic alterations (**Chapter 2**) and deepening the knowledge of molecular mechanisms that are induced by microRNAs in the development of obesity (**Chapter 3 and 4**).

### **microRNA biomarkers, powerful tools to differentiate between health and disease that hit a brick wall**

In recent years, the discovery of microRNAs in extracellular fluids such as plasma and serum, opened up a field of minimally invasive biomarkers.<sup>1-5</sup> Beside the noninvasive accessibility from body fluids the presence of disease specific circulating miRNA signatures renders them attractive for diagnostic functions<sup>1</sup>. MiRNA expression profiling using gene arrays allows rapid identification of large populations of miRNAs in a single sample. In this work, micro-RNA expression profiling from pooled plasma samples led to the discovery of multiple microRNAs differentially expressed in obese versus normal weight women. Subsequent validation in a cohort verified that miR-216a allows differentiation between obese and non-obese women and therefore has biomarker capabilities. Beside this miR-216a also correlated negatively with dynamics of body composition such as body mass index, and waist circumference, as well as mean arterial pressure, triglycerides and high sensitivity-C reactive protein commonly known to be elevated in obese subjects. Although this microRNA showed very promising characteristics in its potential to differentiate between obesity and the ability to monitor metabolic alterations versus normal weight female subjects there are still some hurdles in the translation into routine clinical practice<sup>4</sup>. First the validation of this microRNA should

be performed in an even larger cohort, which can then be subdivided into different subgroups, for example normal weight with altered and unaltered lipid profile, obese with altered and unaltered lipid profile. This will allow more accurate differentiation between metabolic healthy and metabolic unhealthy obesity, and could clarify whether reduced levels of the miR-216a in obesity is specifically linked to BMI, or is rather linked to metabolic alterations. Multiple independent studies will give more insight into the reproducibility of the data. This approach will allow a more detailed analysis of diagnostic accuracy, determining detection thresholds and ranges, with a detailed analysis of sensitivity and specificity. The application of online biomarker repositories for the analysis and comparison of miRNA expression data between multiple cohorts and independent studies will aid in the translation of promising biomarkers into clinical practice. The standardization of isolation and detection methods is key for proper comparison. To date, there are many studies, and clinical trials in literature that identify circulating microRNAs as potential diagnostic tools in several diseases, but it seems as if all remained stagnated in this phase, none have reached diagnostic clinical guidelines and routine clinical practice yet.<sup>5-7</sup> The common problem addressed in these studies remains the need for larger-scaled cohorts and common guidelines prior to clinical implementation.<sup>6-9</sup> Conclusively, circulating microRNA- based biomarkers are powerful tools, with great potential that unfortunately hit a brick wall. Future studies should be directed at overcoming these translational hurdles, prior to generating more microRNA-based biomarkers.

### **microRNAs, fascinating epigenetic instruments capable of modulating energy homeostasis both centrally and peripherally**

The etiology of obesity and subsequent development of metabolic alterations and associated co-morbidities is multifold and includes contributions of genetic, epigenetic and environmental factors that involves concerted interactions between central and peripheral organ systems that regulate energy homeostasis. We show that single microRNAs are capable of modulating energy homeostasis both peripherally (**Chapter 3**) and centrally (**Chapter 4**).

On the one hand, adipose tissue has emerged as a major endocrine organ that plays a pivotal role in the regulation of energy homeostasis by coordinating energy storage in white adipose tissue (WAT), and energy combustion in brown adipose tissue (BAT). BAT, dense in mitochondria, could activate non-shivering thermogenesis in response to sympathetic signaling resulting in uncoupling of mitochondrial respiration and dissipation of energy in the form of heat. A mouse model with heterozygous ablation of *Twist1*, a transcriptional regulator in adipose tissue that negatively regulates brown fat metabolism, developed resistance to high fat diet induced obesity.<sup>10</sup> Consequently, repression of *Twist1* by a single miRNA, miR-337-3p enhanced the expression of several mitochondrial genes and proteins that are specific to BAT. This mechanism was substantiated by performing functional *in vitro* assays overexpressing miR-337 which resulted in a reduction of *Twist1* and increase in the main thermoregulatory uncoupling protein-1

UCP1 protein abundance. Beside the functional assays, the direct base-pairing between miR-337 and Twist1-3'UTR based on seed complementarity was confirmed by luciferase reporter assays. Therefore, the experiments performed in this study support the notion that miR-337 could induce browning by targeting Twist1, and therefore holds great potential in counteracting obesity and metabolic disease. Whether miR-337 truly is enough to counteract obesity with its capabilities to induce browning, could be further substantiated in follow-up studies, performed in an *in vivo* setting for instance by overexpressing miR-337 using adeno associated viral vectors followed by exposure to a high fat diet.

On the other hand, the central nervous system drives the coordination of energy balance by directing autonomic and behavioral responses to signals derived from peripheral organs. Metabolic perturbations arise from disruptions in dopaminergic signaling and variations in circadian rhythmicity that could alter timing of feeding, feeding behavior and result in hyperphagic obesity by overconsumption of food<sup>11</sup>. In analogy with every day practice, energy-rich midnight snacking and late-night eaters, are often refractory to weight loss therapy whereas early time restricted feeding which involves eating early in the day to be in alignment with circadian rhythms in metabolism improved insulin sensitivity in prediabetes<sup>12-14</sup>. Our results show that mice protected from high fat diet induced obesity with key differences in diurnal activity and diurnal feeding behavior show a baseline upregulation of dopaminergic rhythmicity. Beside the coordinated control of energy homeostasis by the central nervous system and the importance of circadian rhythmicity, there has been an increasing body of evidence that miRNAs play key roles in obesity-related diseases and the central control of energy homeostasis<sup>15</sup>. In this thesis we show that deletion of a single miRNA specifically in the brain could orchestrate beneficial effects for energy homeostasis while deletion of this miRNA gene in adipose tissue impairs insulin sensitivity in an *in vivo* setting. This data supports a model where miRNAs, particularly miR-216a has distinct functions in different tissues. Most importantly, we showed that brain-restricted miR-216a gene deletion led to protection from diet induced obesity and insulin resistance and found key differences in diurnal activity among these mice. The underlying mechanism that mediated the observed phenotypical differences were multifold. The prominent differences in diurnal activity orchestrate efficient energy handling shown by calorimetric measurements which is supported by a baseline upregulation in dopaminergic rhythmicity as previously described. In concert with the differences in monoaminergic rhythmicity, high fat diet exposure triggered a shift towards an anti-inflammatory environment mediated by IFN- $\beta$  signaling, providing protection against diet induced obesity. The explanation for the underlying mechanisms that orchestrate the observed phenotype stem from differential gene expression analysis. Although the phenotype arises from the combinatorial contribution of multiple mechanism, future studies could be directed towards systematic validation of the different components involved.

## Concluding Remarks and future perspectives

In this work we are fascinated by the comprehensive utility of small non-coding RNAs, miRNAs, that emerged as powerful regulators of physiological functions, and are accessible both as druggable disease targets as well as diagnostic and predictive diseases markers which could allow stratified health care. Although we have identified circulating miR-216a as a potential obesity marker, we recognize the fact that clinical implementation of microRNA-based biomarkers still has a long road ahead. Unless future studies address the common hurdles in the roads from identified biomarker towards implementation in clinical guidelines, the generation of novel miRNA-based biomarkers, will remain stuck in the “potential” biomarker phase and will not get off the ground.

Obesity and the associated metabolic alterations depend on epigenetic gene-environment interactions in both brain and adipose tissue in which microRNAs play significant roles. We provide evidence for single miRNA-based modulation of energy metabolism capable of protecting against the harmful effects of obesity. This could be accomplished either in the periphery by inducing browning of adipose tissue by miR-337, or centrally by modulating behavioral cues through monoaminergic rhythmicity and attenuation of the inflammatory response present in obesity, by brain specific deletion of miR-216a. Conclusively microRNAs are fascinating epigenetic tools that could modulate energy metabolism both centrally and peripherally to counter obesity and metabolic alterations. Ultimately, we have provided pre-clinical evidence for different levels of regulation accessible for miRNA modulation in obesity, recommendations for follow up studies, and recognize the need for subsequent clinical trials to develop a tailored approach using miRNA-based therapeutics.

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