Cross-anatomical single-cell definition and characterization of human adipose progenitor niche

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par

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Acceptée sur proposition du jury

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“To write a thesis is like cooking a rare steak; it takes time to get it done just right and even then, it's better served with a glass of red wine.”

Unknown

”I have not failed. I've just found 10,000 ways that won't work.”

Thomas Edison

“The most important thing is if you can find satisfaction in the little advances that you make, either on the problem itself or on your own skills and expertise.”

Bart Deplancke
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Not only this MD-PhD journey made me grow into the better person I am today but it also allowed me to meet my better half. Bianca, I love your way of thinking, your sensitivity, your humour and your enthusiasm. Your unwavering belief in me and my abilities has been a constant source of inspiration, and I am grateful for your love and support. I look forward to share my life with you.

Lausanne, the 8th March 2023
Abstract

Adipose tissue is a key metabolic and highly-dynamic organ whose dysregulation may cause clinical conditions of concern such as obesity and lipodystrophy. Its function varies based on the anatomy, in fact, visceral adipose tissue expansion carries a greater risk of life-threatening associated comorbidities than subcutaneous expansion. Among the responsible for adipose tissue plasticity are the adipose-derived stem and progenitor cells (ASPCs) which can commit to form new mature adipocytes even in a post-developmental adult organism. The advent of scRNA-seq techniques allowed to delineate a clear and unbiased picture of the murine ASPC landscape across depots, unraveling subpopulations with distinct functional properties and even non- and anti-adipogenic features, however a similar depth of understanding is still lacking in humans.

This work focuses on defining the human ASPC niche composition and equilibrium across four different depots (Subcutaneous (SC), Omental (OM), Perirenal (PR) and Mesocolic (MC)) and over more than 75 individuals of various BMIs. We took advantage of bulk and scRNA-seq techniques to explore hASPC heterogeneity then functionalize our findings in vitro over 30 donors.

We found that two main populations, the highly-proliferative adipose stem cells and highly-adipogenic pre-adipocytes, are ubiquitously present in all analyzed depots but their relative proportions display a depot-specific and BMI-dependent distribution. Despite their omnipresence, these subpopulations still exhibited depot-specific gene expression patterns, likely reflecting distinct AT properties. Five minor subpopulations are also shared across depots and have specific gene expression patterns resembling populations previously described in mice. We further identified two OM-specific mesothelial cell populations, cobblestone in morphology, out of which one highly expresses and secretes IGFBP2 (Insulin-like growth factor binding protein 2). This OM-specific IGFBP2+ population constitutes 2-5% of the non-immune, non-endothelial OM stromal vascular fraction depending on the donor’s BMI, appears to transition between mesothelial and mesenchymal cell states and inhibits the adipogenic capacity of hASPCs in a depot-specific manner through IGFBP2 secretion and integrin receptor signaling.

Altogether, our in-depth characterization of hASPC heterogeneity and function not only highlights the cellular uniqueness of different adipose niches but also identifies a new mechanism underlying the limited adipogenic capacity of OM hASPCs by uncovering an OM-specific IGFBP2+ mesothelial-like cell population that negatively regulates hASPC adipogenesis through IGFBP2 signaling. Further dissecting the precise mechanism of negative regulation in adipogenesis may lead to the discovery of new druggable targets to combat excessive adipose tissue expansion.

Keywords

Obesity, adipogenesis, human, adipose stem and progenitor cells, mesothelial cells, mesothelial to mesenchymal transition, anti-adipogenic, omentum, IGFBP2, TM4SF1, MSLN, scRNA-seq
Résumé

Le tissu adipeux est un organe clé et hautement dynamique du métabolisme dont la dérégulation entraîne des manifestations cliniques préoccupantes telles que l'obésité et la lipodystrophie. Sa fonction varie en fonction de l'anatomie, en effet, l'expansion du tissu adipeux viscéral comporte un plus grand risque de comorbidités associées potentiellement mortelles que l'expansion du tissu adipeux sous-cutanée. Parmi les responsables de la plasticité du tissu adipeux figurent les cellules souches et progénitrices dérivées du tissu adipeux (adipose-derived stem and progenitor cells = ASPC) qui peuvent se différencier en nouveaux adipocytes matures même dans un organisme adulte qui n’est plus en phase de développement. L’avènement des techniques de séquençage du transcriptome de cellule unique (single-cell RNA sequencing = scRAN-seq) ont permis de définir une image claire et non-biaisée de la configuration des ASPC chez la souris dans différents dépôts. Ceci a permis d’identifier des sous-populations ayant des propriétés fonctionnelles distinctes allant jusqu’à des comportements non et anti-adipogéniques. Une telle compréhension de la constellation des ASPCs fait toujours défaut chez l’Homme.

Ce travail se concentre sur la définition de la composition et de l’équilibre de la niche ASPC humaine dans quatre dépôts différents (sous-cutané (SC), omental (OM), périrénal (PR) et mésocolique (MC)) à partir de plus de 75 individus de divers IMC. Nous avons tiré parti d’analyses transcriptomiques pour explorer l’hétérogénéité des ASPC humaines, puis avons fonctionnalisé nos découvertes in vitro sur 30 donneurs.

Nous avons établi que deux sous-populations principales, les cellules souches adipeuses hautement prolifératives et les pré-adipocytes hautement adipogéniques, sont omniprésentes à travers tous les dépôts analysés, mais leur abondance relative est spécifique à chaque dépôt et dépend de l’IMC du donneur. Malgré leur omniprésence, ces sous-populations présentaient toujours des profils d’expression génique spécifiques au dépôt, reflétant probablement des propriétés propres à chaque tissu adipeux. Cinq sous-populations mineures sont également réparties entre les dépôts et ont des modèles d’expression génique spécifiques ressemblant à des populations précédemment décrites chez la souris. Nous avons en outre identifié deux populations de cellules mésothéliales qui sont spécifiques à l’OM, rôles de morphologie, dont l’une exprime et sécrète fortement l’IGFBP2, capable d’inhiber l’adipogénèse des hASPC d’une manière dépôt-spécifique grâce à la sécrétion d’IGFBP2 lui-même et sa signalisation via le récepteur d’intégrine.

Dans l’ensemble, notre caractérisation approfondie de l’hétérogénéité et de la fonction des ASPC humaines met non seulement en évidence l’unicité cellulaire des différentes niches adipeuses, mais identifie également un nouveau mécanisme sous-jacent à la reluctance adipogénique de ASPC omentales en découvrant une population cellulaire de type mésothélial IGFBP2+ qui inhibe l’adipogénèse des ASPC humains via la signalisation liée à IGFBP2 lui-même. Disséquer davantage le mécanisme de régulation négative du tissu adipeux peut conduire à la découverte de nouvelles cibles médicamenteuses pour lutter contre l'obésité.

Mots-clés

Obésité, adipogénèse, humain, cellules souches et progénitrices adipeuses, cellules mésothéliales, transition mésothé- liale à mésochymateuse, anti-adipogénique, omentum, IGFBP2, TM4SF1, MSLN, scRNA-seq
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<td>Adipose progenitor cells</td>
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<tr>
<td>ASC</td>
<td>Adipose stem cells</td>
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<tr>
<td>ASPC</td>
<td>Adipose stem and progenitor cells</td>
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<tr>
<td>AT</td>
<td>Adipose tissue</td>
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<tr>
<td>ATAC</td>
<td>Assay for Transposase-Accessible Chromatin</td>
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<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
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<tr>
<td>BAM</td>
<td>Binary Alignment Map</td>
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<td>BAT</td>
<td>Brown adipose tissue</td>
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<tr>
<td>BIOP</td>
<td>Bioimaging and optics facility</td>
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<td>BMI</td>
<td>Body mass index</td>
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<tr>
<td>BRB</td>
<td>Bulk RNA barcoding</td>
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<tr>
<td>BSA</td>
<td>Bovine serum albumine</td>
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<tr>
<td>CAC</td>
<td>Complete adipogenic cocktail</td>
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<tr>
<td>CER</td>
<td>Comité éthique pour la recherche</td>
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<tr>
<td>CHUV</td>
<td>Centre hospitalier universitaire vaudois</td>
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<tr>
<td>CP</td>
<td>Committed preadipocytes</td>
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<td>Ct</td>
<td>Cycle threshold</td>
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<tr>
<td>CT</td>
<td>Computerized tomography</td>
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<tr>
<td>D</td>
<td>Donor</td>
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<td>DE</td>
<td>Differentially expressed</td>
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<td>DEG</td>
<td>Differentially expressed gene</td>
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<tr>
<td>DEXA</td>
<td>dual energy X-ray absorptiometry</td>
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<tr>
<td>DEXA</td>
<td>dual energy X-ray absorptiometry</td>
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<tr>
<td>DMEM</td>
<td>Dulbecco’s modified eagle medium</td>
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<tr>
<td>DN</td>
<td>Double negative</td>
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<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
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<td>DP</td>
<td>Double positive</td>
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<tr>
<td>DPBS</td>
<td>Dulbecco phosphate buffered saline</td>
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<tr>
<td>ECM</td>
<td>Extracellular matrix</td>
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<tr>
<td>EDBB</td>
<td>École doctorale biotechnologie et bioingénierie</td>
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<tr>
<td>ELISA</td>
<td>Enzyme-linked immunosorbent assay</td>
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<td>EMT</td>
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<td>EPFL</td>
<td>École polytechnique fédérale de Lausanne</td>
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<tr>
<td>FACS</td>
<td>Fluorescence activated cell sorting</td>
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<td>FBS</td>
<td>Fetal bovine serum</td>
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<td>FC</td>
<td>Fold change</td>
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<td>FCCF</td>
<td>Flow cytometry core facility</td>
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<td>FCF</td>
<td>Flow cytometry facility</td>
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<td>FCS</td>
<td>Flow cytometry standard</td>
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<td>FDR</td>
<td>False discovery rate</td>
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<td>Abbreviation</td>
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<tr>
<td>FIP</td>
<td>Fibro-inflammatory progenitors</td>
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<td>FMO</td>
<td>Fluorescence minus one</td>
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<tr>
<td>GAM</td>
<td>Generalized additive model</td>
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<td>GB</td>
<td>Gallbladder</td>
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<td>GECF</td>
<td>Gene expression core facility</td>
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<td>GFP</td>
<td>Green fluorescent protein</td>
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<td>GLP-1</td>
<td>Glucagon-like peptide-1</td>
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<td>GO</td>
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<td>Genome Reference Consortium</td>
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<td>Histology core facility</td>
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<tr>
<td>HDL</td>
<td>High-density lipoprotein</td>
</tr>
<tr>
<td>HFD</td>
<td>High fat diet</td>
</tr>
<tr>
<td>HIV</td>
<td>Human immunodeficiency virus</td>
</tr>
<tr>
<td>HSC</td>
<td>Hematopoietic stem cell</td>
</tr>
<tr>
<td>HSD</td>
<td>Honestly significant difference</td>
</tr>
<tr>
<td>HSP</td>
<td>Heat shock proteins</td>
</tr>
<tr>
<td>IBI</td>
<td>Institute of bioinformatics</td>
</tr>
<tr>
<td>IBMX</td>
<td>3-Isobutyl-1-methylxanthine</td>
</tr>
<tr>
<td>IGF</td>
<td>Insulin-like growth factor</td>
</tr>
<tr>
<td>IGFBP2</td>
<td>Insulin-like growth factor binding protein-2</td>
</tr>
<tr>
<td>IHC</td>
<td>Immunohistochemistry</td>
</tr>
<tr>
<td>ILC2</td>
<td>Type 2 innate lymphoid cells</td>
</tr>
<tr>
<td>IMC</td>
<td>Indice de masse corporelle</td>
</tr>
<tr>
<td>IP</td>
<td>Interstitial progenitors</td>
</tr>
<tr>
<td>KD</td>
<td>Knock-down</td>
</tr>
<tr>
<td>KO</td>
<td>Knock-out</td>
</tr>
<tr>
<td>LAGB</td>
<td>Adjustable gastric banding</td>
</tr>
<tr>
<td>LOESS</td>
<td>Locally estimated scatterplot smoothing</td>
</tr>
<tr>
<td>MACS</td>
<td>Magnetic activated cell sorting</td>
</tr>
<tr>
<td>MAD</td>
<td>Median absolute deviation</td>
</tr>
<tr>
<td>MC</td>
<td>Mesocolon</td>
</tr>
<tr>
<td>MD</td>
<td>Medical doctor</td>
</tr>
<tr>
<td>MEM</td>
<td>Modified eagle medium</td>
</tr>
<tr>
<td>MMP</td>
<td>Matrix metalloproteinases</td>
</tr>
<tr>
<td>MMT</td>
<td>Mesothelial to mesenchymal transition</td>
</tr>
<tr>
<td>MRI</td>
<td>Magnetic resonance imaging</td>
</tr>
<tr>
<td>MSLN</td>
<td>Mesothelin</td>
</tr>
<tr>
<td>NA</td>
<td>Numerical aperture</td>
</tr>
<tr>
<td>NAFLD</td>
<td>Non-alcoholic fatty liver disease</td>
</tr>
<tr>
<td>NC1</td>
<td>Negative control 1</td>
</tr>
<tr>
<td>NGS</td>
<td>Next generation sequencing</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Definition</td>
</tr>
<tr>
<td>--------------</td>
<td>------------</td>
</tr>
<tr>
<td>NTP</td>
<td>Nucleotides triphosphate</td>
</tr>
<tr>
<td>OM</td>
<td>Omenum</td>
</tr>
<tr>
<td>OSAS</td>
<td>Obstructive sleep apnea syndrome</td>
</tr>
<tr>
<td>P</td>
<td>Passage</td>
</tr>
<tr>
<td>PB</td>
<td>Pacific blue</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate buffered saline</td>
</tr>
<tr>
<td>PC</td>
<td>Principal component</td>
</tr>
<tr>
<td>PCA</td>
<td>Principal component analysis</td>
</tr>
<tr>
<td>PCOS</td>
<td>Polycistic ovary syndrome</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>PDGF</td>
<td>Platlet-derived growth factor</td>
</tr>
<tr>
<td>PDGFR</td>
<td>Platlet-derived growth factor receptor</td>
</tr>
<tr>
<td>PFA</td>
<td>Paraformaldehyde</td>
</tr>
<tr>
<td>PR</td>
<td>Perirenal</td>
</tr>
<tr>
<td>RELM</td>
<td>Regional Earthquake Likelihood Models</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
</tr>
<tr>
<td>RT</td>
<td>Real-time</td>
</tr>
<tr>
<td>RYGBP</td>
<td>Roux-en-y gastric bypass</td>
</tr>
<tr>
<td>SARS</td>
<td>Severe acute respiratory syndrome</td>
</tr>
<tr>
<td>SC</td>
<td>Subcutaneous</td>
</tr>
<tr>
<td>SCA1</td>
<td>Stem cells antigen-1</td>
</tr>
<tr>
<td>SFM</td>
<td>Serum free medium</td>
</tr>
<tr>
<td>SFRP</td>
<td>Soluble frizzled-related proteins</td>
</tr>
<tr>
<td>SG</td>
<td>Sleeve gastrectomy</td>
</tr>
<tr>
<td>SNP</td>
<td>Single nucleotide polymorphism</td>
</tr>
<tr>
<td>SV</td>
<td>Science de la vie</td>
</tr>
<tr>
<td>SVF</td>
<td>Stromal vascular fraction</td>
</tr>
<tr>
<td>t-SNE</td>
<td>t-distributed stochastic neighbor embedding</td>
</tr>
<tr>
<td>T2DM</td>
<td>Type 2 diabetes mellitus</td>
</tr>
<tr>
<td>T2DM PA</td>
<td>Type 2 diabetes mellitus preadipocytes</td>
</tr>
<tr>
<td>TGF-β</td>
<td>Transforming growth factor beta</td>
</tr>
<tr>
<td>TM4SF1</td>
<td>Transmembrane 4 L Six Family 1</td>
</tr>
<tr>
<td>UCP1</td>
<td>Uncoupling protein 1</td>
</tr>
<tr>
<td>UMAP</td>
<td>Uniform Manifold Approximation and Projection</td>
</tr>
<tr>
<td>UMI</td>
<td>Unique molecular identifier</td>
</tr>
<tr>
<td>UNIL</td>
<td>Université de Lausanne</td>
</tr>
<tr>
<td>VD</td>
<td>Vaud</td>
</tr>
<tr>
<td>VSMP</td>
<td>Vascular smooth muscle progenitor cells</td>
</tr>
<tr>
<td>WAT</td>
<td>White adipose tissue</td>
</tr>
<tr>
<td>WC</td>
<td>Waist circumference</td>
</tr>
<tr>
<td>WHR</td>
<td>Waist to hip ratio</td>
</tr>
<tr>
<td>WHtR</td>
<td>Waist-to-height ratio</td>
</tr>
</tbody>
</table>
Chapter 1 | Introduction

**Note:** This chapter is based on a published review “Toward a Consensus View of Mammalian Adipocyte Stem and Progenitor Cell Heterogeneity” Trends Cell Biol 30, 937–950 (2020).

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* These authors contributed equally

**My contribution:** Conceptualized and wrote the manuscript.
1.1 The relevance of studying adipose tissue plasticity

Over thousands of years, humans have evolved leading a lifestyle where, getting food, mainly by hunting and gathering, was nearly as costly in energy as the energy that the food brought back. Throughout ages and civilizations, being overweight has been considered a privileged condition reserved only to the elite who could “afford it”. The recent revolution in human lifestyle as well as the increasing availability and degrading quality of food, have contributed to the dramatic change in the world’s weight distribution landscape. In 2016, more than 25% of the world’s population is overweight or obese\(^1\), regardless of gender, age, or social background\(^2\). What is alarming about excessive fat accumulation in the human body though is the consequences that it has for the metabolic health of patients.

Insulin resistance followed by type 2 diabetes, dyslipidemia, and high blood pressure together with obesity form the clinical features of the metabolic syndrome which facilitates the insurgence of cardiovascular diseases such as heart or brain stroke, as well as cancer\(^3\). Alone, cardiovascular diseases and cancer are the cause of a quarter of the world’s deaths\(^4\), and with it, obesity is thus one of the most widespread and preventable killers to be on earth. In addition, follow-up of these chronic and complex conditions such as metabolic syndrome, cardiovascular disease, and cancer drains numerous healthcare resources with a consequent negative impact on the financing of healthcare systems\(^5\). Despite long-term efforts from the scientific community to understand the mechanisms underlying fat accumulation and to develop effective therapies, no treatment is currently available\(^6\), and exercise and lifestyle change are not always applicable, especially in severely affected patients.

Efficient personalized medicine in the field of fat biology would permit a decrease in the excessive body fat burden both for patients and healthcare systems. If as shortly presented above, excess in adipose tissue mass can be highly detrimental to the human organism, it is not nevertheless a disposable organ as illustrated by the lipotoxic consequences of lipodystrophic conditions\(^7\). Overall, it is now well appreciated in the scientific community that adipose tissue constitutes a highly dynamic, endocrine-type organ playing an indispensable role in several biological processes including development, immunity, and energy homeostasis.

1.2 Clinical pathologies of the human adipose tissue

1.2.1 The overweight and obesity syndromes

Excess in body fat mass leads first to overweight followed by obesity. In an average men population, fat weight accounts for 15%-20% of the total weight, while for 25-30% of the woman's total weight\(^8,9\). To
diagnose overweight and obesity conditions, physicians require reliable and convenient measurements to use on a daily basis.

In clinical practice, the easiest metric to use is the Body Mass Index (BMI \([\text{kg/cm}^2]\)), defined as the weight of the individual divided by the square of their height. Empirically determined cutoffs define underweight, normoweight, overweight and obesity following the values shown in Table 1.1. BMI measurement however misses an important factor in the body weight distribution and can easily be biased by the ratio of “lean mass” = muscles versus “fat mass” = adipose tissue\(^1\). Moreover, it will not account for the anatomic distribution of the adipose mass, and for equal BMIs, a person can have its fat accumulated in the subcutaneous compartment versus the visceral one with the first case having a lower risk for developing obesity-associated comorbidities\(^10\). To partially overcome BMI limitations in discriminating between metabolically unhealthy visceral obesity versus subcutaneous one\(^11\) the waist circumference (WC) and the waist-to-hip ratio (WHR) are often used as complementary quick and affordable diagnostic tools, with the WC outperforming the BMI ratio when it comes to predicting intraperitoneal adipose tissue mass, and outperforming WHR when predicting posterior subcutaneous adipose tissue mass\(^12\). Combining BMI and WC measures allows for identifying normal risk, high risk and very high risk for developing associated lethal comorbidities as illustrated in Table 1.2. Finally, a recent study underly the importance of keeping waist circumference lower than half your height (Waist-to-height ratio (WHR) \(\leq 0.5\)) to prevent metabolic risk, out-performing the BMI and WC classification in terms of early detection of increased metabolic risk\(^13\).

With technological advances, and limited to research purposes for practical reasons, several more sophisticated methods have been used to precisely determine fat mass distribution in health and disease and to scientifically predict morbidity and mortality linked to overweight and obesity. Among those, bioelectric impedance, hydrodensitometry and air displacement plethysmography are non-invasive investigative techniques that rely on differences in conduciveness of hydrophobic adipose tissue and hydrophilic soft tissues\(^14\). The affordability and easy-to-use features of these devices justify their use in big population-based studies. However, the measure of adipose tissue being indirect and based on pre-set equations often calibrated on image-based quantification performed on white men reference population makes them inaccurate when transferred to different populations such as elders, children or women\(^15\). Dual-energy X-ray absorptiometry (DEXA)-scan relies on low-dose radiation exposure and attenuation of the electron beam based on the tissue it encounters in the body\(^16\). Although highly reproducible and relatively easy to perform, the output of the DEXA-scan measure is in two dimensions and thus cannot distinguish visceral fat from subcutaneous fat\(^17,18\). Once again, algorithms based on image-based adipose tissue measurements were developed to infer what percentage of the measured adipose mass is visceral\(^19\) and implemented in recent DEXA analysis software, however, the same limitation in transferability across populations described above apply in this case. Finally, image-based methods, such as computerized tomography (CT) and magnetic resonance imaging (MRI),
represent the highly technical albeit reference measuring tools to localize adipose tissue and accurately associate its anatomic distribution to morbidity\textsuperscript{20}.

*Table 1.1 – Clinical definition of weight status based on BMI\textsuperscript{21}*

<table>
<thead>
<tr>
<th>Weight</th>
<th>BMI (kg/m(^2))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Underweight</td>
<td>&lt; 18.5</td>
</tr>
<tr>
<td>Severe underweight</td>
<td>&lt; 16.0</td>
</tr>
<tr>
<td>Moderate underweight</td>
<td>16.0–16.98</td>
</tr>
<tr>
<td>Mild underweight</td>
<td>17.0–18.49</td>
</tr>
<tr>
<td>Normal range</td>
<td>18.5–24.99</td>
</tr>
<tr>
<td>Overweight</td>
<td>≥ 25</td>
</tr>
<tr>
<td>Pre-obese</td>
<td>25.0–29.99</td>
</tr>
<tr>
<td>Obese</td>
<td>≥ 30</td>
</tr>
<tr>
<td>Obese class I</td>
<td>30–34.99</td>
</tr>
<tr>
<td>Obese class II</td>
<td>35–39.99</td>
</tr>
<tr>
<td>Obese class III</td>
<td>≥ 40</td>
</tr>
</tbody>
</table>

*Table 1.2 – Risk identification of developing health comorbidities based on combined BMI and waist circumference\textsuperscript{13}*

<table>
<thead>
<tr>
<th>BMI classification</th>
<th>Waist circumference</th>
<th>No increased risk</th>
<th>No increased risk</th>
<th>Increased risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal weight</td>
<td>Low</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overweight</td>
<td>Low</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(25 to less than 30 kg/m(^2))</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Obesity I</td>
<td>Low</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(30 to less than 35 kg/m(^2))</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Obesity II</td>
<td>Low</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(35 to less than 40 kg/m(^2))</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Obesity III</td>
<td>Low</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(40 kg/m(^2) or more)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Waist Circumference: Low (men <94 cm, women <80 cm); High (men 94–102 cm, women 80–88 cm); Very high (men >102 cm, women >88 cm).

**Etiology**

The number one cause of the onset of overweight and obesity is behavioral. On the one hand, are poor dietary habits such as eating large amounts of processed food and drinking sugary drinks for a daily calorie intake higher than 2000 kcal, on the other hand, reduced physical activity results in reduced energy expenditure greatly contributing to weight gain over time.

Some rare forms of obesity accounting for less than 0.001% of the overall prevalence are driven by monogenic mutations (mainly in the leptin and melanocortin endocrine axis\textsuperscript{22}) and are at times associated with developmental delay *Table 1.3*, while several clinical syndromes with high familiarity prevalence including Polycystic Ovary Syndrome (PCOS), hypothyroidism and hypogonadism are characterized by early onset and resistant obesity\textsuperscript{23}.

Importantly, epigenetics seem to play a key role in obesity onset, and obese parents tend to give birth to obesity-predisposed offsprings\textsuperscript{24}. Finally, among iatrogenic causes of weight gain are medications including atypical antipsychotics, anticonvulsants, mood stabilizers, hypoglycemic drugs, glucocorticoids, and antidepressants\textsuperscript{25}.
In the context of excessive body weight, what is the most concerning are the linked comorbidities. Metabolically, insulin resistance followed by type 2 diabetes typically onset early in the context of obesity, together with dyslipidemia and high blood pressure they constitute the dreadful triad of the metabolic syndrome. The obstructive sleep apnea syndrome (OSAS) is typically linked to obesity and metabolic syndrome and contributes to an overall increase in the risk for cardiovascular diseases including stroke and heart failure. In the longer term, the metabolic imbalance also leads to non-alcoholic fatty liver disease and chronic kidney disease.

The osteoarticular system is also highly challenged by excessive weight. This results in a higher incidence of arthrosis, arthritis and sciatica events among obese patients. Osteo-articular pathogenic conditions also contribute to a vicious circle where pain impedes exercise which in turn worsens the cause of pain.

The onset of several cancers is facilitated by underlying obesity. Among the scientifically demonstrated ones are the esophagus adenocarcinoma, colorectal, pancreas, liver, gallbladder, kidney, thyroid, uterus, breast, ovary and blood (myeloma) cancers. In the attempt to explain such a striking correlation between obesity and cancers, several adipokines have been put forward as critical drivers.

Finally, not only the body is affected by obesity, but also the mind. Depression and anxiety are typically more prevalent in the obese population than the normoweight population. If the main explanation stems from the societal look that patients have to carry throughout their life, a physiologically detrimental impact of the high-sugar and high-lipids blood content on the brain is not to be excluded.

Treatments

The first line of management of overweight and obesity syndromes is nutritional and physical education accompanied by motivational support in the context of weight-management programs. With such a
conservative approach physicians can hope to achieve 5%-10% body weight loss which is often sufficient to improve the underlying metabolic dysbalance. The most challenging aspect of weight loss however is long-term maintenance. Five years after the weight-loss intervention, only 3% of the weight loss is usually retained.

When weight-management programs fail to significantly reduce the body weight of the patient, and the BMI remains above 30 kg/m², Swiss physicians can count on three anti-obesity medications to achieve a supplementary weight loss of 3 to 12% and improve the overall metabolic health (Table 1.4). Pancreatic lipase inhibitors (orlistat) induce malabsorption of free fatty acids and result in a median weight loss of 2.12 kg which is associated with a reduction of both type 2 diabetes mellitus and cardiovascular risk. The unpleasant side effect of malabsorption often limits the therapy to a couple of years. The agonist of the Glucagon-like peptide-1 (GLP-1) receptor (Liraglutide, Semaglutide) acts as an appetite suppressor and slows the gastrointestinal transit down. The resulting weight loss is an average of 3.2 kg and comes with positive effects on glycemia, blood pressure and cholesterol. Finally, a controverted combined medication (naltrexone/bupropion) analog of the amphetamine was approved by the European Medicine Agency in 2015 and at cost of several side effects such as headache, nausea and constipation has the potential of reducing the patients’ weight up to 5%. The American Food and Drug Administration approved in addition to the above-described drugs, several molecules acting on the brain (serotonin and opioid receptors), or physically taking space in the stomach to reduce caloric intake (Table 1.4).

The last bullet clinicians have to improve the metabolic health of morbidly obese patients is bariatric surgery, also known as metabolic surgery. The most common methods of bariatric surgery are adjustable gastric banding (AGB), sleeve gastrectomy (SG) and the Roux-en-y gastric bypass (RYGBP). The rationale stands on the one hand to restrict the stomach volume (AGB, SG and RYGBP), preventing excessive food intake, and/or on the other hand inducing malabsorption (RYGBP). If LAGB has been abandoned based on poor weight-loss outcomes in the long term, LSG and LRYGBP, being both laparoscopic surgeries, are safe and associated with low risks of perioperative complications and minimal mortality (0.1 to 0.2%). Long-term (> 10 years) follow-up reveals that on average 20-30% of the initial weight is lost without relapse. The weight loss is accompanied by significant improvement, at times complete remission, of several obesity-related comorbidities, coupled with an increase in patients’ survival up to 30-40%. 
1.2.2 Lipodystrophy

Opposite to overweight and obesity are lipodystrophic syndromes, characterized by the inability to generate and maintain functional adipose tissue. Lipodystrophy mainly concerns the subcutaneous adipose tissue and localizes differentially based on the type of syndrome\textsuperscript{7} (Fig. 1.1). The condition can be congenital, generalized (autosomal recessive mutation of AGPAT2, BSCL2 and CAV1 genes) or familial partial (autosomal dominant mutation of LMNA, CIDEC, LIPE or recessive missense mutation of adipogenesis master regulators PPARG and PLIN-1 as well as zinc finger protein ZMPSTE24), or acquired, once again generalized, partial or even localized in the case of subcutaneous drug injections (recombinant insulin treatment in the context of diabetes for example)\textsuperscript{52} (Fig 1.1). Lipodystrophy can also occur under long-term anti-retroviral therapy in the context of HIV infection\textsuperscript{53}. The prevalence of lipodystrophic syndromes is difficult to estimate but is in the range of extremely rare disorders (1-5 cases per million people)\textsuperscript{54}. The progressive loss of mature adipocytes leads to a number of metabolic and endocrinologic impairments including leptin and adiponectin defects as well as lipotoxicity and insulin resistance.

<table>
<thead>
<tr>
<th>Name (Trade Names)</th>
<th>Year Approved</th>
<th>Mechanism of Action / Clinical Effect</th>
<th>Average placebo-subtracted weight loss (%) at 14 months</th>
<th>Achieved ≥5% Weight Loss, Intervention vs. placebo (%)</th>
<th>FDA approval</th>
<th>EMA approval</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phentermine (Adipex, Lomaira)</td>
<td>1959</td>
<td>Sympathomimetic / Suppresses appetite</td>
<td>4.4 at 28 weeks</td>
<td>49 vs.16 at 28 wks</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Diethylpropion</td>
<td>1979</td>
<td>Sympathomimetic / Suppresses appetite</td>
<td>6.6 at 6 months</td>
<td>67.6 vs. 25.0</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Orlistat (Alli, Xenical)</td>
<td>1999</td>
<td>Intestinal lipase inhibitor / Reduces fat absorption by up to 30%</td>
<td></td>
<td></td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Phentermine-topiramate (Qsymia)</td>
<td>2012</td>
<td>Combination sympathomimetic and carbonic anhydrase inhibitor / Decreases appetite and binge eating behaviors</td>
<td>8.6</td>
<td>70 vs. 21</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Bupropion-naltrexone (Contrave)</td>
<td>2014</td>
<td>Combination of a dopamine and norepinephrine re-uptake inhibitor and mu-opioid receptor antagonist / Decreases appetite and cravings</td>
<td>4.8</td>
<td>48 vs. 16</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Liraglutide 3.0mg (Saxenda)</td>
<td>2014</td>
<td>GLP-1 receptor agonist / Decreases appetite, increases fullness, increases satiety</td>
<td>5.4</td>
<td>63.2 vs. 27.1</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Gelesis100 (Plenity)</td>
<td>2019</td>
<td>Superabsorbent hydrogel particles of a cellulose-citric acid matrix / Increases fullness. Considered a medical device but functions as a medication.</td>
<td>2.0 at 6 months</td>
<td>58.6 vs. 42.2</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Setmelanotide (Imciveree)</td>
<td>2020</td>
<td>Melanocortin-4-receptor agonist / Decreases appetite</td>
<td>N.A.</td>
<td>N.A.</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Semaglutide 2.4 mg (Wegovy)</td>
<td>2021</td>
<td>GLP-1 receptor agonist / Decreases appetite, increases fullness, increases satiety</td>
<td>12.4</td>
<td>86.4 vs. 31.5</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>
Clinics

The inability to store free fatty acids in mature adipocytes results in diffuse lipotoxicity. Non-specialized organs such as the liver, the pancreas and the heart will buffer the excessive circulating triglycerides at cost of having their function compromised\(^{455}\). Physically, patients have cadaveric facies, prominent muscles and veins as well as lytic skeletal changes. The associated hypo-leptinemia induces persistent hunger and voracious appetite. Metabolically, patients suffer from insulin-resistant diabetes mellitus coupled with hyperinsulinemia together with hypertriglyceridemia and low serum HDL levels. With disease progression, NAFLD followed by non-alcoholic steatohepatitis, pancreatitis, hypertrophic cardiomyopathy, nephropathy and organomegaly take over and undermine patients’ life expectancy. For the congenital forms of lipodystrophy, the average lifespan is 65 years old\(^{456}\), namely 30 years less than a healthy individual.

Treatments

The management of lipodystrophic syndrome is directed to limit the metabolic burden of lipotoxicity. First are dietary and exercise measures to keep a balanced diet and avoid excessive food intake. Leptin analog therapy allows for decreasing the appetite, the serum levels of glycosylated HbA1c and triglycerides within 4 months\(^{457,458}\) and is indicated for generalized lipodystrophic syndromes, while it is used off-label for the treatment of partial syndromes as it proved to be less effective in this context\(^{455}\).
1.3 The biology of the adipose tissue

If we want to combat obesity, there is an urgent need to better understand fundamental biologic processes involving adipose tissue (AT) expansion. In this work, I decided to focus on characterizing cells at the very root of the AT expansion, i.e. the stem cells that reside within the AT and are able to regenerate it. In order to be as close as possible to human biology, I decided to work with human samples and explored similarities and differences in the composition of the stem cells across four different anatomic depots. To date, the composition and behavior of the adipose-derived stem and progenitor cells (ASPCs) have mainly been studied in the mouse model while studies including human material have been limited to computational analysis and little to no functionalization of the findings was so far performed. In this section, I will present the state
of the art of ASPCs composition and function with a main focus on the murine model starting with a short overview of AT structure, function and anatomy.

1.3.1 A common structure different depots and functions

AT is the main center for long-term energy storage and lipids handling. Its conservation from simple invertebrates such as D. melanogaster all the way up to humans underlies its fundamental role for survival. It is defined as a connective tissue that arises from the mesenchymal layer of the embryo but, contrary to other connective tissues such as bone or cartilage it accounts for higher cellularity and a lower content of the extracellular matrix. Nested in a collagen net, adipocytes are the mature cells residing in the adipose tissue. Their size can vary from as little as 20 um to 300 um in diameter, and although they account for less than 30% of the absolute cell number of the adipose tissue they make up the biggest volume of the tissue and are therefore considered as the principal cell type in the tissue. Their main role is to store free fatty acids in the form of triglycerides, by doing so, they will increase their intracellular volume up to a point that becomes detrimental to the metabolic health of the entire individual. In fact, in the long term, this process, called hypertrophy, can lead to the exhaustion of the adipocyte itself which will start to secrete pro-inflammatory signals, eventually contributing to insulin resistance and metabolic syndrome.

Surrounding and supporting adipocytes stands the stromal vascular fraction (SVF), which not only hosts resident and transient immune cells and endothelial cells along blood vessels but is also the nest for a number of other poorly defined cell types, the so-called adipose-derived stem and progenitor cells (ASPCs). ASPCs are multipotent stem cells that are able to form a new functional adipose tissue when transplanted in lipodystrophic mice and are able to accumulate intracellular lipid-filled vesicles when cultured in vitro. Upon adipose tissue remodeling, ASPCs can commit and differentiate into mature adipocytes even in the adult organism. This process is called hyperplasia, and opposite to hypertrophy is associated with good metabolic health as it will allow to store free fatty acids in newly formed small and un-exhausted adipocytes. Hyperplasia likewise hypertrophy will eventually lead to the increase of the volume of the depot upon chronic excessive calory intake.

If the structure is common to all ATs, when it comes to function, it is important to distinguish between depots that store lipids and the ones that burn the lipids to generate heat. As such, three functional types of adipose tissue exist white, brown or beige ATs. Adipocytes of the white adipose tissue typically harbor a unique large lipid vesicle (unilocular) and are specialized in lipids storage. When it comes to brown AT, a vestigial adipose tissue in humans, but an important organ in mice, its main mission is to maintain body temperature homeostasis. It is in fact able to generate heat by highly expressing the uncoupling protein 1 (UCP-1). UCP-1 allows for the dissipation of the energy needed to produce ATP from the electron transport chain in the mitochondria and generate heat. Adipocytes of the brown adipose tissue are typically smaller in size than
the ones from the white adipose tissue and harbor more than one (multilocular) small lipid droplet in their cytoplasm. Finally, beige AT is an in-between and adaptative tissue type that upon external stimulation, develops within white adipose tissue and is able to generate heat. External stimulation is typically cold or adrenergic stimuli that result in the upregulation of the UCP-1 protein expression. Adipocytes of the beige adipose tissue are unilocular and express UCP-1. Whether beige adipocytes arise from an interconversion of white adipocytes or from a differential commitment of ASPCs is still a debated concept, but finding means to reprogram white adipose tissue to become thermogenic is a valuable research avenue in the context of improving the metabolic status of obese individuals.

1.3.2 Anatomy dictates the function of adipose tissue

A common way to define AT is based on its anatomy, which has been shown to correlate with metabolic health when overgrown. As such, “metabolically healthy” subcutaneous AT, using hyperplasia as a preferential mean of growth, is conventionally opposed to the “unhealthy” visceral one which typically expands through hypertrophy. However, the terms “visceral” and “subcutaneous” underlie several finer anatomic locations and, with it, potentially finer AT functions. For instance, visceral AT is the perirenal AT, which locates around the kidney and is attracting increasing attention being a site of dormant brown adipogenesis\textsuperscript{71,72}. It’s possible that crosstalk with the kidney itself to undermine kidney function is also an interesting detrimental mechanism linked to perirenal adipose tissue expansion\textsuperscript{71}. The mediastinal AT situated around the esophagus and the trachea is also considered as a visceral adipose tissue and a potential site for human brown-beige adipogenesis\textsuperscript{73,74}, yet it is but at the same time like the perirenal one is not confined within a mesothelial layer contrary to for example the pericardial, mesenteric or omental ATs, with the latter being considered as the “golden standard” visceral adipose depot\textsuperscript{71}. Pericardial adipose tissue and its anatomically close friend, epicardial adipose tissue are considered pro-inflammatory, proatherogenic, and cardiotoxic in the context of cardiometabolic disease, but also as an important source of energy for the myocardium when energy demand is increased\textsuperscript{75}. The mesenteric adipose tissue, a fold connecting the intestine to the abdominal wall, contributes to buffering the intestinal intake in many ways, both as an immune barrier and a lipidic reservoir to prevent lipid hepatotoxicity\textsuperscript{76}. Needless to say, that omental adipose tissue has consistently been linked to type 2 diabetes onset, cardiometabolic complications, and an overall proinflammatory status\textsuperscript{77}. The same heterogeneity holds between subcutaneous ATs. For example, the abdominal SC AT is slightly different from the gluteal one\textsuperscript{78}. So far, most of the functional analyses are restricted to subcutaneous and omentum, often taking advantage of the mouse model for practical reasons, however, evidence points to a regional specialization of every single adipose depot in the adult human body\textsuperscript{79}, there is, therefore, need to take these regional differences into account and better functionally characterize all existing adipose tissues.
1.4 The ASPC heterogeneity

In mouse, SCA1 surface protein, encoded by the Ly6a gene, is well established as the main marker to enrich ASPCs for a precursor cell population that has enhanced in vitro adipogenic capacity\(^8\). However, not all SCA1+ cells give rise to adipocytes in vitro, implying that an even finer granularity of cell states or even cell types exist. Therefore, much effort has been devoted to uncover additional surface markers. For example, mesenchymal markers, such as CD34, CD29, CD24, and platelet-derived growth factor receptor (PDGFR)α/β, were also shown to enrich for adipogenic precursors, or a fraction of them\(^8\)\(^-\)\(^4\). Our integration confirmed that Cd34, Cd29, and Pdgfra are expressed by virtually all ASPCs, while Cd24 and Pdgfrb exhibit a nonuniform expression, as discussed later (Fig. 1.2D-E). However, while SCA1 is widely used for enriching ASPCs in mouse, there is no SCA1 ortholog in human. Consequently, the in toto Lin– fraction of adipose SVF tends to be considered as the best representation of human ASPCs (hASPCs). Markers, such as CD29, CD34, CD13, CD44, CD73, CD90, CD142, CD9, CD10, and CD200\(^8\)\(^5\),\(^8\)\(^6\), have been used to further enrich for adipogenic cells in human, but there is no consensus yet on the exact molecular signature of hASPCs. Until recently, the identification of these markers was mostly based on flow cytometry and immunohistochemistry using hematopoietic, endothelial, and neural tissues as reference, which tends to introduce marker selection bias. However, over the past few years, the single-cell technology revolution has allowed researchers to delineate ASPCs at an unprecedented resolution (Fig. 1.2 and 1.3).

1.4.1 Three Main Subpopulations in White Adipose Tissue

Numerous studies\(^5\)\(^9\),\(^6\)\(^0\),\(^8\)\(^7\)-\(^9\)\(^1\) recently resolved ASPC heterogeneity in mouse subcutaneous (inguinal) white adipose tissue (WAT) and visceral (specifically epigonadal WAT) WAT (iWAT and eWAT, respectively). Despite differences in the utilized methodologies and/or mouse models (Table 1.5), all studies in mouse stratified ASPCs into two to three main subpopulations. Qualitative cross-comparison of the population-specific markers supports the notion that these populations are shared between subcutaneous and visceral depots, although intrinsic differences remain that are far from being fully understood. To empirically validate this ASPC classification, we integrated public data sets (highlighted in gray in Table 1.5) using the standard integration workflow of Seurat\(^9\)\(^2\),\(^9\)\(^3\). This large, integrated meta-data set provides a powerful means to explore the observed ASPC landscape across studies in an unbiased fashion. This is because it enables verification of the expression distribution and, thus, the specificity of previously established ASPC markers. In addition, it allows us to computationally explore parallels between the recently published scRNA-seq studies and to support the proposed population nomenclature. Specifically, our analyses revealed that the three populations (Adipose Stem Cells (ASC), Pre-Adipocytes (PreAs), and Adipogenesis regulators (Aregs)) are robust since they: 1) are characterized by specific gene expression signatures; 2) were first independently detected in each of the
utilized data sets; 3) can be projected onto one another upon data integration (Fig. 1.2 A, and Fig. 1.3A,G); and 4) are consistently clustered in various subclusters upon analysis of the integrated data set (Fig. 1.2B, and Fig. 1.3B,H). Finally, we were able to confirm that the three populations align with populations published across different studies (Fig. 1.2C, and Fig. 1.3C,D,I,J). Altogether, the integration shows that ASCs are a distinct population compared with PreAs and Aregs, which are closer in terms of gene expression but still feature distinct transcriptomic profiles. Specific to visceral fat are the mesothelial cells which also populate the Lin–fraction of visceral SVF together with ASPCs\textsuperscript{80,88,89,94,95}.

For human, only a couple of scRNA-seq studies have been published \textsuperscript{61,96,97}, and these do not necessarily reach the same conclusions. This renders it difficult to establish a clear picture of hASPC heterogeneity. However, a closer look at the reported findings does suggest that, based on shared gene signatures, at least two ASPC subpopulations are conserved between mouse and human, namely the ASCs and PreAs.
Table 1.5 – Methodological differences across scRNA-seq studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Model</th>
<th>Technique</th>
<th>Anatomical depot</th>
<th>SVF enriching strategy</th>
<th>Pop. names</th>
<th>Populations markers</th>
<th>Proposed nomenclature</th>
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<tbody>
<tr>
<td>Schwalie et al.</td>
<td>Adult Mouse</td>
<td>10x Genomics</td>
<td>Inguinal°</td>
<td>CD45−/CD31−/Tar19−/CD34+/Sca1+</td>
<td>P1, P2, P3</td>
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<td>ASCs, PreA, Aregs</td>
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<td>G1, G2, G3</td>
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</table>

Abbreviations: APC, adipose progenitor cells; Aregs, adipogenesis regulators; ASC, adipose stem cells; CP, committed preadipocytes; Fibro, fibroblasts; FIP, fibro inflammatory progenitors; HSC, hematopoietic stem cells; IP, interstitial progenitors; Mesos, mesothelial cells; PreA, preadipocytes; T2DM PA, type 2 diabetes-dependent preadipocytes; Gray boxes indicate data sets used in the integration analysis (see Fig. 1.2 and Fig. 1.3); Schwalie et al.87, Burl et al.88, Hepler et al.89, Merrick et al.90, Cho et al.91, Spallanzani et al.92, Vijay et al.93.
Figure 1.2 – Integrated scRNA-seq data of SVF isolated from eWAT and iWAT.

Publicly available scRNA-seq data of SVF that was isolated from eWAT (6645 cells) and iWAT (17608 cells) were integrated using the standard workflow of Seurat.

A. t-distributed stochastic neighbor embedding (tSNE) 2D cell map displaying the clustering of each dataset, analyzed individually.

B. tSNE 2D cell map colored by cell cluster identification.

C. Boxplot showing the distribution of scores of populations defined in different studies across cells. The score is calculated as the sum of scaled integrated expression of the top 100 DE genes of ASPC populations (Schwalie et al., Burl et al., Hepler et al.) or of a shorter list of specific markers (Merrick et al.) since the DE genes were not publicly available (Group1: Pi16, Dpp4, Dact2, Gpr1, Ptgs2, Bmp7, Wnt2, Stmn4, Wnt10b; Group2: Fabp4, Pparg, Icam1, Cd36, Dlk1, Gsc, Cyp1b1; Group3: F3, Fmo2, CleC11a)

D. Boxplot showing the distribution of log normalized expression of selected markers (Ly6a: Stem cell antigen 1 SCA1)

E. tSNE colored by the log normalized expression of selected markers. (Aoc3: VAP1; F3: CD142). The colors reflect population specificity (Green: ASCs, Red: PreAs, Blue: Aregs, Purple: Mesothelial cells).
1.5 The Adipose Stem Cell Population

1.5.1 ASCs in inguinal WAT

A first population of ASPCs that was uncovered using scRNA-seq (P1 in \(^{87}\), Group1 in \(^{60}\), and ASC2 in \(^{88}\)) was most stem-like in nature. This is why we propose to refer to this population as ASCs\(^{60,87,88}\) (Fig. 1.2A-C). Not only do ASCs express stem-related genes, such as \(\text{Ly6a}\)^{105}, \(\text{Dpp4}\)\(^{106-108}\), \(\text{Cd55}\)^{109,110}, and Wnt signaling pathway inhibitor \(\text{Sfrp4}\)^{111}, but they also lack others (e.g., \(F3\), encoding for CD142\(^{60,87,112,113}\)) (Fig. 1.2D-E), and are also highly proliferative \textit{in vitro}\(^{60}\). From a functional perspective, when ASCs \([\text{CD142}^–/\text{DPP4}^+\text{ (dipeptidyl peptidase 4)}]\) are \textit{in vitro} stimulated with insulin alone, they tend to be refractory to adipogenesis and barely express adipocyte-specific genes\(^{60}\). However, when stimulated with a complete adipogenic cocktail (CAC), ASCs \([\text{CD142}^–/\text{DPP4}^+\text{ or SCA1}^+/\text{CD55}^+\text{]}^{87}\) exhibit relatively high adipogenic potential, although its extent varied among studies\(^{60,87}\). This suggests that ASCs require key factors to unlock their commitment and differentiation potential and/or that they give rise to distinct cell states with different adipogenic potentials. Thus, ASCs likely represent a pool of mesenchymal stem cells that commit to adipogenesis only when exposed to the right mix of factors (i.e., CAC). In addition, their conserved capacity to form osteoblasts and induce osteoblast-specific markers (\(\text{Alpl, Bsp, Osx, and Ocn}\)) underscores their multipotency\(^{60}\).

Altogether, Lin–/SCA1+/CD55+ and Lin–/CD142–/DPP4+ cells, described in two different studies\(^{60,87}\), represent two cell populations with similar molecular and functional properties that are indicative of a stem-like nature. So far, the molecular mechanisms underlying the stemness of ASCs remain poorly understood. However, the known antiadipogenic factor, transforming growth factor (TGF)\(\beta\)^{70,114}, is able to regulate the identity and function of DPP4+ cells\(^{60}\). In fact, treatment with TGF\(\beta\) increased the proliferative capacity of ASCs, inhibited adipocyte formation (even when using a CAC), and upregulated the expression of ASC-specific markers, whereas the inhibition of TGF\(\beta\) signaling had opposite effects. Noteworthy, neither TGF\(\beta\) agonists nor antagonists had any effect on other ASPC subpopulations, underscoring the specificity of TGF\(\beta\) signaling to DPP4+ cells. The antiadipogenic Wnt signaling pathway\(^{70}\) also appears to have a role in regulating neoadipogenesis. This is because cells tend to maintain their stem state when \(\beta\)-catenin levels exceed those of TCF7L2, whereas the inverse balance induces adipogenesis\(^{115}\). Collectively, these findings point to the involvement of multiple signaling pathways in ASC maintenance, but the exact underlying molecular mechanisms remain poorly understood, including to which extent these pathways crosstalk, what triggers them, and how they influence overall cell state.

In human, even if a first low resolution study reported “homogeneity” amongst hASPCs\(^{97}\), it nevertheless showed, similar to mouse, that genes such as \(\text{CD55}\) and \(\text{THY1}\) were not uniformly expressed. Indeed, thereafter, it has been shown that one hASPC subpopulation expresses \(\text{DPP4, CD55 and MFAP5}\) and
transcriptionally resembles mouse ASCs. In addition and again similar to mouse, human ASCs showed full adipogenic potential only when stimulated by a combination of several adipogenic triggers, as opposed to insulin alone. Interestingly, a mesenchymal stem population characterized by THY1 expression has been described as responsible for adipogenic degeneration of the muscle in individuals suffering from type 2 diabetes. If this population transcriptomically resembles hASCs remains to be robustly proven.

### 1.5.2 ASCs in epigonadal WAT

A population with molecular and functional properties similar to ASCs has been uncovered in the scRNA-seq data sets from mouse eWAT (called ASC2 and Clusters 1 and 5) (Fig. 1.2B-E, and Fig. 1.3G-J). Visceral ASCs (Lin-/SCA1+/CD55high cells), for instance, showed higher proliferation and lower adipogenic differentiation capacity than the rest of visceral ASPCs. Furthermore, visceral ASCs (Lin-/CD142-/DPP4+ cells) exhibited lower adipogenic potential and responded to TGFβ treatment in a similar fashion as subcutaneous ASCs. Interestingly, the relative abundance of ASCs was lower in eWAT than in iWAT, suggesting that visceral WAT (visWAT) features a smaller early-stem cell pool.

From a mechanistic perspective, the PDGF pathway has a key role in adipose commitment and ASC pool maintenance and has been mostly studied in eWAT. PDGF receptors have been linked to stemness maintenance and need to be downregulated to initiate adipogenesis. It has been suggested that all mature adipocytes in major fat depots derive from Pdgfra+ cells and that the expression of PDGFRα precedes that of PDGFRβ. It has also been reported that adipocytes emerge from PDGFRβ+ preadipocytes in response to a high-fat diet (HFD). Based on our integration of scRNA-seq data sets, Pdgfra is virtually positive in all ASPCs, while Pdgfrb is only very lowly expressed by ASCs (compared with the rest of the ASPC pool) (Fig. 1.2D, Fig. 1.3E,K). These collective observations support the notion that ASCs are the ‘real’ ASCs that give rise to more committed progenitors, which in turn differentiate into adipocytes.

The implication of the PDGF pathway in stemness and adipogenesis prompted the community to specifically characterize the PDGFRβ+ progenitor pool using scRNA-seq (Hepler et al. 2018). A PDGFRβ+ population that shares many markers with ASCs was identified in such a way (Hepler et al. 2018) (Fig. 1.3I). Such cells were termed fibro inflammatory progenitors (FIPs) by the authors and were isolated as PDGFRβ+/LY6C+ cells. In line with previous observations in ASCs, analysis of the gene expression profile of FIPs revealed an active TGFβ signaling signature, and TGFβ treatment further upregulated FIP-specific markers, especially collagen (hence ‘Fibro’). At the same time, FIPs also exhibited a functional proinflammatory phenotype, expressing several cytokine genes, such as Il6, Cxcl2, and Cxcl10, and showing an ability to activate macrophages in vitro (hence ‘Inflammatory’). Finally, it was shown that FIPs are reluctant to undergo adipogenesis depending on
the culturing conditions, which supports the hypothesis that FIPs are similar to ASCs (Hepler et al. 2018). Nevertheless, the authors also revealed that FIPs exert antiadipogenic properties through a yet-to-be determined secreted factor. Thus, even if visceral FIPs share important markers with ASCs (Fig. 1.2A,C, and Fig. 1.3I), distinct phenotypic characteristics also appear to differentiate them from subcutaneous ASCs, making it difficult to establish a clear and definite analogy. Indeed, subcutaneous ASPCs cannot be stratified based on the markers that were used to sort FIPs since all subcutaneous PDGFRβ+ cells are also LY6C+. Therefore, FIPs currently constitute a rather mysterious cell population that may reflect an inherent ability of (visceral) ASCs to alter their own cell state and possibly function in response to specific immunological stimuli or physiological conditions.

1.6 The Pre-adipocyte Population

1.6.1 PreAs in inguinal WAT

The second and most abundant ASPC population uncovered through scRNA-seq, which we define here as PreAs, features cells that are marked by the expression of *Icam1* and *Aoc3* and several collagen and extracellular matrix remodeling factors (Fig. 1.2E, and Fig. 1.3F). So far, no study has primarily focused on PreAs. However, the molecular signatures as well as initial, functional characterization all point to a more committed adipogenic state compared with ASCs: first, several genes involved in adipogenesis-related functions, such as *Pparg*, *Fabp4*, *Lpl*, *Plin2*, or *Cd36*, are significantly upregulated in this population (P2 in 87, Group 2 in 60, and ASC1 in 88; Fig. 1.2E, and Fig. 1.3F); second, PreAs show lower proliferation compared with ASCs60, while exhibiting a high adipogenic capacity60,87, even when stimulated by insulin only, as opposed to ASCs. Taken together, Lin−/CD142−/ICAM1+ (intercellular adhesion molecule 1) cells60 and Lin−/SCA1+/VAP1+ (vascular adhesion protein 1) cells 87 appear to constitute similar cell populations that are in a committed adipogenic state. From a mechanistic perspective, PreAs are also the most refractory to the antiadipogenic effect of TGFβ60, which further supports the hypothesis that they are lineage-committed ASPCs.

In humans, a population resembling mouse PreAs, defined, among others, by the expression of *ICAM1*, *PPARg* and *GGT5*, was equally identified 96. Human PreAs seem to behave like mouse PreAs in vitro 96.

1.6.2 PreAs in epigonadal WAT

Burl et al.88 identified a population (ASC1) in both visceral and subcutaneous depots that resembles the PreA population (Fig. 1.2B-E, and Fig. 1.3G-J). In our analysis of this published data set, the fraction identified as ASC1 shares top markers with both PreAs and Aregs (CD142+) and could consequently further be stratified (Fig. 1.2C, and Fig. 1.3C,I). Although the latter study 88 was largely computational, a parallel study88 uncovered
a population defined by Lin−/PDGFRβ+/LY6C−/CD9− that, according to our data integration analyses, projects partially to PreAs (and to Aregs) (Fig. 1.2A, C, and Fig. 1.3G, H). Sorted Lin−/PDGFRβ+/LY6C−/CD9− cells exhibited a higher adipogenic potential compared with the Lin−/PDGFRβ+ total population, in line with the behavior of subcutaneous PreAs. Thus, it is likely that the authors analyzed a subset of PDGFRβ+ PreAs that were isolated from visWAT. Another independent study also revealed a higher adipogenic propensity in three clusters (Clusters 2, 4, and 6) that also project partially to PreAs and Aregs (integration by 90).
Chapter 1 | Introduction

![Image](image_url)

Figure 1.3 – Integrated scRNA-seq data of SVF isolated from iWAT or from eWAT.

Publicly available scRNA-seq data of SVF isolated from iWAT (17608 cells)\textsuperscript{96,100,102} (panel A-F) or eWAT (6645 cells)\textsuperscript{100,101} (panel G-L) were integrated using the standard workflow of Seurat\textsuperscript{103,104}.
A. tSNE 2D cell map displaying the clustering of each dataset from iWAT, analyzed individually.
B. tSNE 2D cell map from iWAT colored by cell cluster identification.
C. Percentage of the top 100 population markers that overlap with the top 100 population markers of published studies: Schwalie et al. (iWAT) 102, Burl et al. 100 (aggregate Mouse eWAT and iWAT), Hepler et al. 101 (eWAT, PDGFRβ+).
D. Boxplot showing the distribution of scores across cells for the Group1, Group2 and Group3 defined in Merrick et al. 96. The score is defined as the sum of the scaled integrated expression of a list of specific markers discussed in Merrick et al. 96 (Group1: Pi16, Dpp4, Dact2, Gpr1, Ptgs2, Bmp7, Wnt2, Stmn4, Wnt10b; Group2: Fabp4, Pparg, Icam1, Cd36, Dlk1, Gsc, Cyp1b1; Group3: F3, Fmo2, Clec11a).
E. Boxplot showing the distribution of log normalized expression of selected markers (Ly6a: Stem cell antigen 1 SCA1)
F. tSNE colored by the log normalized expression of selected markers. (Aoc3: VAP1; F3: CD142). The colors reflect population specificity (Green: ASCs, Red: PreAs, Blue: Aregs, Purple: Mesothelial cells).
G. tSNE 2D cell map displaying the clustering of each dataset from eWAT, analyzed individually.
H. tSNE 2D cell map from eWAT colored by cell cluster identification.
I. Percentage of the top 100 population markers that overlap with the top 100 population markers of published studies: Burl et al. 100 (eWAT), Hepler et al. 101.
J. Boxplot showing the distribution of scores across cells for the Group1, Group2 and Group3 defined in Merrick et al. 96. The score is defined as the sum of the scaled integrated expression of a list of specific markers discussed in Merrick et al. 96 (Group1: Pi16, Dpp4, Dact2, Gpr1, Ptgs2, Bmp7, Wnt2, Stmn4, Wnt10b; Group2: Fabp4, Pparg, Icam1, Cd36, Dlk1, Gsc, Cyp1b1; Group3: F3, Fmo2, Clec11a).
K. Boxplot showing the distribution of log normalized expression of selected markers (Ly6a: Stem cell antigen 1 SCA1)
L. tSNE colored by the log normalized expression of selected markers. (Aoc3: VAP1; F3: CD142). The colors reflect population specificity (Green: ASCs, Red: PreAs, Blue: Aregs, Purple: Mesothelial cells).
1.7 The Adipogenesis Regulators Population

1.7.1 Aregs in iWAT

A third population, closer in terms of gene expression to PreAs than to ASCs and defined by the expression of F3 (encoding CD142) (Fig. 1.2C,E), was first identified within iWAT of adult mice. Functionally, these cells were not only refractory to adipogenesis *in vitro*, but also had the capacity to inhibit the adipogenic differentiation of the remaining ASPCs, both *in vitro* and *in vivo*. Therefore, these cells were termed Aregs by the authors (Fig. 1.3F).

Given their distinctive transcriptomic clustering pattern, a second study also focused on CD142+ cells, but, contrary to the first, found CD142+ cells to be fully adipogenic. While there were clear methodological and mouse model differences, the exact reason why CD142+ ASPCs behaved differently between the two studies has yet to be resolved. Nevertheless, the overall consensus is that CD142+ ASPCs constitute a distinct population of cells featuring a transcriptomic signature that is clearly different from that of ASCs and PreAs (Fig. 1.2B,C, and Fig. 1.3C,F).

Going beyond adipose tissue, Areg-like cells have been recently identified in muscle, exhibiting molecular and functional (e.g., antiadipogenic) properties that are in line with those reported for subcutaneous Aregs. The authors proposed a mechanistic model that involves the secreted factor GDF10 as one of the main factors responsible for the inhibitory capacity of CD142+ cells, both in mouse and human. Interestingly, dystrophic mice were found to harbor fewer ‘muscle Aregs’ than their wild-type (WT) counterparts, raising the hypothesis that this change in cellular composition explains the increased accumulation of fat cells in dystrophic muscle.

1.7.2 Aregs in Visceral WAT

No scRNA-seq study of visWAT has so far specifically explored the existence of Aregs and, even though they both appear to display an antiadipogenic phenotype, FIPs and Aregs are clearly molecularly distinct (Fig. 1.2C). Even though a higher fraction of CD142+ cells in eWAT compared with iWAT has been reported using flow cytometry, to date, no one has ever sorted and characterized visceral Aregs fully. However, a recent study described an ASPC population (Lin-/Sca1+/PDGFRα+/PDPN+/THY1+/CD55– cells) that was transcriptomically similar to Aregs and that exhibited a low adipogenic propensity, even though it was not sorted using the CD142 surface marker itself. In line with this, our integration results revealed a population that express Areg-like markers in eWAT (Fig. 1.3H,I). Surprisingly, even though F3 remains a differentially expressed (DE) gene, it appears not to be the best marker for this population because it lacks specificity.
versus Fig. 1.3L). Such a broad, diffuse expression pattern could explain the higher CD142+ fraction observed in flow cytometry in visWAT without necessarily implying a higher proportion of actual Aregs. Nevertheless, using the relative fraction inferred from scRNA-seq data, and keeping in mind possible biases due to cell selection, the proportion of Areg-like marker-expressing cells in the visceral data set remains substantially higher than in the subcutaneous one (23.5% versus 7% when considering only ASPCs). A higher fraction of ASPCs featuring an antiadipogenic character, potentially comprising both ‘visceral Aregs’ and FIPs, would be consistent with the low adipogenic potential of total visceral ASPCs compared with their subcutaneous counterparts, a well-established notion in the field. These observations highlight once again not only the intrinsic similarities, but also differences between the different fat depots, and clearly emphasize the need to better functionalize all ASPC fractions.

And what about Aregs in humans? Markers of mouse Aregs (CD142, CLEC11A and FMO2) appeared broadly expressed across the two main human subpopulations: ASCs and PreAs. We conclude therefore that either orthologous human Aregs do not exist or may not be defined by this set of markers at the transcriptomic level. In accordance, no functional differences were found between sorted human CD142+ and ICAM1+ ASPCs. Nevertheless, when human CD142+ cells were sorted and compared to the corresponding CD142- and Lin- fractions, the same non-adipogenic character observed for mouse Aregs was seen for human CD142+ ASPCs. This underscores the general interest in providing a more in-depth molecular and functional characterization of human CD142+ cells.

1.8 Hierarchy of ASPCs

As implied by the names given to the different ASPC subpopulations in this introduction (ASCs, PreAs and Aregs), we propose a hierarchy between ASPCs (see arrows in Fig. 1.4). First of all, the expression pattern of Pdgfra and Pdgfrb, as already detailed above, indicates that the ASCs are the most stem-cell like ASPCs. The higher expression of Cd24 in ASCs further supports this hypothesis (Fig. 1.2E). Indeed, it has been proposed that CD24+ adipocyte precursors are stem cell-like ASPCs that become further committed by losing CD24 expression. Recently, this hierarchy has also been suggested in silico by applying a pseudotemporal trajectory analysis on scRNA-seq datasets, after which it was validated in vivo using fluorescence-based cell tracing. Specifically, when fluorescently-labelled subcutaneous ASCs (DPP4+ cells) were transplanted in a wild type subcutaneous fat pad, these cells acquired PreAs and Aregs markers (ICAM1 and CD142 respectively) within 7 days post-transplantation and a subset of them lost DPP4 expression after 14 days. These findings indicate that ASCs (DPP4+) give rise to both preAs (ICAM1+) and Aregs (CD142+). In contrast, fluorescently-labeled preAs (ICAM1+) and Aregs (CD142+) did not acquire DPP4 expression over time, underscoring the overall unidirectionality of ASC commitment and differentiation. However, PreAs and Aregs may
undergo immunophenotypic interconversion since a subset of mTomato-labeled ICAM1+ and CD142+ cells acquired to a certain extent CD142 and ICAM1 markers respectively \(^{96}\). How this interconversion is molecularly encoded, what determines the hierarchical equilibrium, and how different metabolic conditions affect this equilibrium, are important, yet outstanding questions (see section “The ASPCs and Obesity”).

1.9 Mesothelial cells

Mesothelial cells constitute the peritoneum both in mouse and human and belong exclusively to the visceral SVF. Not only do they line the walls of the abdominal cavity, but they are also found in small clusters within the visceral WAT itself, in the form of so-called ‘milky spot’ or ‘fat-associated lymphoid clusters’ \(^{125,126}\). Msln, Wt1, Lrrn4, and Upk3b are typical mesothelial markers and, in vitro, such cells display a characteristic epithelial/cobblestone morphology \(^{127–129}\). Not surprisingly, mesothelial cells were found among visceral ASPCs using scRNA-seq (in mouse \(^{80,88,95,98}\), in human \(^{59,60,94}\)) (Fig. 1.2A, B, E, and Fig. 1.3G, H, I). Thus, one should be aware that, for both human and mouse, cells of the mesothelial lineage exist within the visceral SVF and may cloud functional read-outs. Therefore, we argue that it may be best to sort these cells out before any downstream in vitro characterization of ASPCs.

However, what is interesting about mesothelial cells in the context of adipose biology, besides being a heterogeneous population themselves \(^{80,94,130}\), is their role at the crossroads between adipose tissue function and immunity as well as them being hotpots for cancer metastasis \(^{126,131,132}\). When peritonitis occurs, they recruit neutrophils for peritoneal fluid clearance \(^{80}\), whereas in physiological conditions, they support the interleukin (IL)-33 driven recruitment and differentiation of type 2 innate lymphoid cells (ILC2s) \(^{99,133,134}\). ILC2 cells secrete type 2 immunomodulatory cytokines \(^{130}\), which, together with regulatory T cells and eosinophils, enable the visceral cavity and adipose tissue to remain in a normal, uninflamed state, countering the development of type 2 diabetes \(^{135}\). In turn, loss of IL-33 results in weight gain, although the underlying mechanisms, as well as implications of ASPCs in this process remain poorly understood \(^{133}\).

In human, less is known about the interplay between mesothelial, immune, and stromal cells. However, it has been proposed that subclusters of mesothelial cells are implicated in beige adipogenesis based in the fact that these subclusters express a higher proportion of mitochondrial genes than any other ASPC subpopulation and that their relative cell number negatively correlated with glucose fasting levels \(^{94}\). Moreover, the beige adipocyte markers PLA2G2A \(^{136}\) and SOD2 \(^{137}\) were enriched among mesothelial subclusters, but not MYF5 and ZIC1, two brown adipocyte precursors cell markers \(^{137}\), providing additional support for the involvement of these human mesothelial cells in beige adipogenesis.
1.10 ASPCs and obesity

1.10.1 ASCs and obesity

We currently have a poor understanding of ASPC dynamics in obesity, and we know even less about the implication of each subpopulation. Nevertheless, scRNA-seq was performed, for example, on eWAT from both lean and diet-induced obese mice, revealing that obesity redistributes the prevalence of individual ASPC subpopulations. In addition, obesity specifically enriches for subgroups of ASPCs that exhibit enhanced extracellular matrix and immunomodulatory capacities as well as altered differentiation abilities. In line with this, the frequency of FIPs increased when mice were fed a HFD, while the DPP4+ stem fraction decreased. Together, these findings suggest that ASCs may play an important immunomodulatory role in WAT, consistent with previous observations, and that a diet-induced ASC imbalance toward a FIP-like phenotype may be one of the factors underlying obesity-induced adipose inflammation.

1.10.2 PreAs and obesity

One hypothesis is that intrinsic properties of ASPCs could explain the distinct impact of differential anatomical fat deposition during obesity on the development of metabolic disease (reviewed in). In line, subcutaneous ICAM1+ cells have a comparably high adipogenic capacity regardless of whether they are isolated from obese or lean mice, while visceral ICAM1+ cells exhibited a lower adipogenic potential when retrieved from obese mice compared to lean. This observation may explain why hypertrophic WAT expansion is favored over hyperplasia in the abdomen. Interestingly, a relative increase in the proportion of human PreAs seems to correlate with insulin resistance and type two diabetes based on a large scRNA-seq dataset from 25 obese patients. To our knowledge, this is the first time that a correlation has been found between a metabolic disease phenotype and differential ASPC composition in human. Whether this correlation, based on scRNA-seq cell proportions only, reflects causality remains to be established.

1.10.3 Aregs and obesity

The fact that WAT may harbor ASPCs such as Aregs or FIPs that could negatively modulate adipocyte formation has triggered a paradigm shift in our understanding of how adipose tissue plasticity could be mediated, pointing to potential novel therapeutic applications. For example, a reduction in adipogenesis-suppressing cells may be one of the mechanisms that lead to fat cell hyperplasia. Inversely, their increase may lead to fat cell hypertrophy due to a suppression of adipocyte formation. Interestingly, it has been proposed that a depletion of ASPCs may be responsible for the switch from hyperplasia to hypertrophy in the context of overfeeding. The fact that the abundance of CD142+ cells varies across depots and physiological state.
would support this possible shift \(96,102\). Indeed, while the increase of CD142+ cells in obese mice may at first glance seem counterintuitive given their possible anti-adipogenic phenotype, it may instead reflect a defensive strategy aimed at limiting adipose tissue (hyper)expansion. The main drawback of such a mechanism would be a balance favoring hypertrophy over hyperplasia upon excessive weight gain.

**A. Mouse**

**Figure 1.4 – Adipose derived stem and progenitor cells: toward a consensus view.**

**A.** Mouse white adipose tissue (WAT) \(96,100-102,139,143\): ASCs give rise to PreAs and Aregs. Aregs exert an inhibitory effect on adipogenesis. Such effect may in part be mediated by GDF10 based on Areg-like cells found in muscle \(120\). FIPs, CD9+ and mesothelial cells are exclusively present in visceral WAT. FIPs also exert an inhibitory action on adipogenesis. Mesothelial cells bridge ASPCs and immunoregulation. CD9+ cells are profibrotic \(144\).

**B. Human**

**Aregs**

**Hematopoietic stem cells**

Shadowed sections are based on (sc)RNA-seq analyses only, plain colors mean FACS isolation and characterization.
1.11 The scope of this thesis

This work attempts to redefine in an unbiased and cross-anatomical fashion the cellular composition of human adipose-derived stem and progenitor cells (hASPCs), a cellular compartment within the stroma of adipose tissue responsible for its renewal. We took advantage of cutting-edge single-cell RNA sequencing (scRNA-seq) techniques to identify cells that have common characteristics based on their RNA expression. I then characterized them molecularly and phenotypically with the ultimate goal of understanding why *in vitro* hASPCs isolated from intra-peritoneal depots are strikingly less adipogenic than the ones isolated from extra-peritoneal adipose depots.

At first, I performed scRNA-seq experiments on the non-endothelial, non-immune stromal vascular fraction (SVF) of the adipose tissue from four distinct anatomic origins: subcutaneous, omental, perirenal and mesocolic adipose tissues. This allowed to perform thorough bioinformatics analysis to identify new subpopulation and put them in the context of what was previously described in the literature about human and mouse ASPCs. I then identified surface proteins against which I designed and validated a sorting strategy that allowed to study each subpopulation separately with regard to their relative abundance in the SVF, adipogenic potential and proliferation ability in a systematic way across all adipose depots. Whenever possible I tried to link donor’s metadata including age, sex and BMIs to my findings.

At last, I explored the role of omentum-specific mesothelial cells in the context of adipogenesis. I demonstrated that mesothelial cells within the SVF of omental adipose tissue secrete inhibitory cues which are at least partially responsible for the observed low-adipogenic phenotype of intraperitoneal-derived SVF-adherent cells. I further proposed that the observed inhibition signaling originates specifically from mesothelial cells and relies on the IGFBP2 secreted protein which acts through an IGF-independent mechanism. Interfering with this signaling resulted in enhanced adipogenesis by the omentum.
Chapter 2 | A human omentum-specific mesothelial-like stromal population inhibits adipogenesis through IGFBP2 secretion

**Note:** This chapter is based on a manuscript under submission “A human omentum-specific mesothelial-like stromal population inhibits adipogenesis through IGFBP2 secretion”

* These authors contributed equally

**My contribution:** Conceptualized the study and wrote the manuscript. Conducted all experimental procedures and analyzed acquired images, flow cytometric measures, qPCRs, ELISAs and immunohistochemistry.
2.1 Abstract

Adipose tissue (AT) is a key metabolically dynamic organ that tends to have distinct functions dependent on its anatomical location. For example, subcutaneous (SC) AT tends to be more adipogenic compared to visceral AT whose expansion carries a greater risk for obesity-related co-morbidities. While significant progress has been made in mice to understand which cells mediate AT plasticity and how these are distributed across distinct adipose depots, a similar level of understanding is still lacking in humans.

Here, we probed the differentiation capacity of human Adipose-derived Stem and Progenitor cells (hASPCs, defined as Stromal Vascular Fraction (SVF) Lineage (Lin)-negative cells) from four adipose depots (SC, perirenal (PR), omental (OM), and mesocolic (MC)), revealing striking differences in their adipogenic potential. To explore the molecular and cellular origins of these differences, we performed bulk RNA-seq of 20 SC, 8 PR, 19 OM, and 4 MC primary ASPC samples and scRNA-seq of ~34'000 cells from these four depots, followed by functional characterization of the identified cell subpopulations.

On the one hand, these analyses revealed at least two major, ubiquitous hASPC subpopulations, adipose stem cells and pre-adipocytes, with distinct proliferative and adipogenic properties and whose proportions differed in function of adipose depot type and BMI. Moreover, despite their omnipresence, these subpopulations still exhibited depot-specific gene expression patterns, likely reflecting distinct AT properties. On the other, we identified an OM-specific, mesothelial-like stromal population that is defined by high expression of IGFBP2, constitutes 2-5% of the OM SVF Lin– fraction depending on the donor’s BMI, appears to transition between mesothelial and mesenchymal cell states, and inhibits the adipogenic capacity of hASPCs in a depot-specific manner through IGFBP2 secretion.

Altogether, our in-depth characterization of hASPC heterogeneity and function not only highlights the cellular uniqueness of different adipose niches, it also identifies a new mechanism underlying the limited adipogenic capacity of OM hASPCs by uncovering an OM-specific IGFBP2+ mesothelial-like cell population that negatively regulates hASPC adipogenesis through IGFBP2 signaling.

**Keywords**: obesity, adipogenesis, human, adipose stem and progenitor cells, mesothelial cells, mesothelial to mesenchymal transition, anti-adipogenic, omentum, IGFBP2, TM4SF1, MSLN, scRNA-seq
2.2 Introduction

Our understanding of key adipose tissue (AT) phenotypes, such as turnover and expansion dynamics in health and in response to altering metabolic conditions, is still limited, especially when it comes to human AT. This is further exacerbated by the fact that these AT phenotypes vary according to the anatomical location of the respective AT with for example the frequent opposition of the “metabolically healthy” subcutaneous (SC) AT to the “unhealthy” visceral one when overgrown. However, the terms “visceral” and “subcutaneous” underlie several finer anatomic locations and, with it, potentially more fine-grained AT characteristics. For instance, among visceral AT is the perirenal (PR) AT, which locates around the kidney and is attracting increasing attention as being a potential site of dormant brown adipogenesis. Its possible crosstalk with the kidney itself influences renal function and constitutes an independent risk factor for cardiovascular and chronic kidney disease. Even if PR AT can be considered as a visceral tissue, it is not confined within a mesothelial layer, contrary to for example the mesenteric (MC) and omental (OM) ATs. The latter tends to be considered the “golden standard” visceral adipose depot and is typically linked to type 2 diabetes onset, cardiometabolic complications, and an overall proinflammatory status. The MC AT, a fold connecting the intestine to the abdominal wall, contributes to buffering the intestinal intake, both as an immune barrier and a lipidic reservoir to prevent lipid hepatotoxicity. It serves as a scaffold to the mesenteric vasculature that nourishes the small and large intestines and, based on its anatomy, can further be stratified in the mesentery and the mesocolon, depending on whether it binds the small or large intestine, respectively.

While it is thus well-accepted that human ATs from distinct anatomical locations are linked to different metabolic risks when overgrown, little is known about what causes these phenotypic differences. One attractive hypothesis is that these differences could at least be partially driven by variation in the cellular composition of the stromal vascular fraction (SVF) across depots and specifically in that of the adipose-derived stem and progenitor cell (ASPC) pool. This hypothesis is supported by i) a recently published comprehensive single cell transcriptomic (scRNA-seq) atlas of whole human AT, as well as previously published studies, revealing the existence of several subpopulations amongst human ASPCs (hASPCs). However, these scRNA-seq studies focused on the two most commonly studied ATs: subcutaneous and omentum. Hence, a more overarching view on similarities and/or differences in hASPC composition beyond the SC and OM depots remains elusive. ii) While still scarce in humans, substantial evidence is mounting that in mice, ASPCs are also highly heterogeneous with the detection of at least three major ASPC subpopulations—Dpp4+ (or Ly6c+) cells being labeled as stem cells, the Icam1+ (or Aoc3+) ones as pre-adipocytes and the F3+ cells as adipogenesis-regulatory cells. Moreover, extensive downstream validation suggests that these subpopulations exhibit different functional properties with the Dpp4+ (or Ly6c+) cells being labeled as stem cells, the Icam1+ (or Aoc3+) ones as pre-adipocytes and the F3+ cells as adipogenesis-regulatory cells. A similar level of phenotypic characterization of hASPC populations is still lacking, likely reflecting the challenge of having access to and/or gathering enough human material to do so. Nevertheless, in one study, efforts were undertaken to functionally characterize
hASPC subpopulations that were similar to the ones found in mice, with the DPP4+ ASPCs being highly proliferative and less adipogenic than the ICAM1+ ASPCs\textsuperscript{96}. Together, these findings suggest that, similar to mouse, hASPCs may not only be highly heterogeneous but also functionally distinct. Yet, to date, no systematic, functional characterization of hASPC heterogeneity and behavior has been performed across several human adipose depots.

Here, we provide a comprehensive overview of gene expression profiles of SVF-adherent cells over 30 human donors in four major human depots: SC, PR, OM, and MC AT, supplemented with scRNA-seq data on ~34,000 non-immune (CD45−) and non-endothelial (CD31−) SVF cells (SVF/Lin−). We consistently detected two main hASPC subpopulations that are common to all depots and addressed similarities but also differences across these depots, as well as in comparison to the most commonly studied mouse ATs. Specifically, we found that pro-adipogenic/developmental genes are enriched in SC, non-adipogenic/inflammatory ones in OM, mitochondrial/thermogenic ones in PR, and protein folding/trafficking in MC. We isolated, quantified, and characterized different cellular subpopulations in SC, OM, and PR depots with regard to their adipogenic potential and proliferation abilities, validating two surface markers, CD26 and VAP-1, that enable the enrichment of highly proliferative and highly adipogenic cells, respectively, across all depots. Finally, we focused on resolving the mechanism underlying the lower adipogenic potential of OM-isolated SVF-adherent cells, compared to SC and PR ones. We identified a new and omentum-specific cell population that is susceptible to undergoing mesothelial-to-mesenchymal transition and negatively impacts the adipogenic potential of OM and SC hASPCs. We further linked the observed adipogenic inhibition to the secretion of IGFBP2, which potentially acts through the α5β1 integrin receptor.

2.3 Results

2.3.1 Human SVF precursor cells exhibit depot-dependent differences in their \textit{in vitro} adipogenic potential

To characterize the function of SVF-adherent cells, including hASPCs, across distinct human adipose depots, we differentiated cell lines from SC (20 donors), PR (8 donors), OM (19 donors), and MC (4 donors) AT (Table 2.1). As no consensus exists on the surface markers defining hASPCs and to not bias our strategy towards a potential ASPC (sub)population, we did not implement any enrichment strategy beyond plating SVF cells and culturing SVF-adherent cells, as is commonly done in the field. Once confluent, these distinct AT-derived primary cell lines were exposed to an adipogenic cocktail for 14 days (Fig. 2.1A, see Materials and Methods). Subsequent staining for lipid droplets revealed that only SVF-adherent cells that were isolated from white AT (WAT) situated outside the peritoneal cavity (i.e., SC and PR) are able to form mature adipocytes that are characterized by the effective accumulation of lipid droplets \textit{in vitro} (Fig. 2.1B-C, Supp. Fig. 2.1.1A). Conversely, cells that were isolated from intraperitoneal depots (i.e., OM and MC) barely formed any lipid
droplets under adipogenic differentiation conditions, or at most, tiny droplets that were difficult to distinguish from a background stain (Fig. 2.1B-C, Supp. Fig. 2.1.1B-C, see Materials and Methods). Interestingly, while both SC and PR hASPCs differentiated to a higher extent than intraperitoneal cells, PR lines showed the highest adipogenic potential in vitro, particularly when cells were differentiated straight after isolation (Fig. 2.1B-C, Supp. Fig. 2.1.1A). However, at longer times/passages, PR lines tended to become as adipogenic as SC ones (Supp. Fig. 2.1.2 and 2.1.3A-B). Furthermore, SC and PR lines showed high inter-individual variation in their ability to differentiate, which is observable as an adipogenic potential gradient for SC lines and a dichotomy for PR lines, as illustrated by very high versus very low adipogenic potential (Supp. Fig. 2.1.3A). In contrast, OM and MC lines were systematically resistant to adipogenic differentiation (Supp. Fig. 2.1.2), while also being the slowest growing lines (Supp. Fig. 2.1.1D).

We explored possible correlations between our experimental adiposcore (Supp. Fig. 2.1.2B) and physiological parameters such as BMI, age, and gender of the donors but found no correlations except for a tendency for PR cells to become less adipogenic in women and elderly people (Supp. Fig. 2.1.3C-F). However, we acknowledge that our cohort’s demographic characteristics can bias these observations (Table 2.1 and Table 2.2), i.e., included patients are mainly young and obese, while we also analyzed only a relatively small proportion of PR samples (n=8).

2.3.2 Human SVF-adherent cells exhibit transcriptomic differences that reflect their anatomical origin and adipogenic potential

To explore if the striking phenotypic difference between intra-peritoneal and extra-peritoneal cell lines is reflected in their respective transcriptomes, we set out to perform bulk RNA barcoding and sequencing (BRB-seq)150 of SVF-adherent primary cell lines from different individuals and depots both at the undifferentiated/expanding state (t0) and after 14 days of adipogenic differentiation (t14) (SC n=22, OM n=16, PR n=8, MC n=4, Supp. Fig. 2.1.4A). We found that the major source of variation is explained by the exposure to the adipogenic cocktail, followed by the anatomic origin of the cell lines (Fig. 2.1D, Supp. Fig. 2.1.4B-F). We observed that all samples at t0 highly express THY1, a well-known mesenchymal marker151, at similar levels, except OM samples in which it is slightly but significantly lower expressed (Supp. Fig. 2.1.4G). Upon exposure to a differentiation cocktail, cells from all depots induced genes related to extracellular remodeling, insulin response, and positive regulation of fat cell differentiation compared to their undifferentiated state. However, most of these adipogenic-related terms were more enriched in SC and PR compared to OM and MC (Fig. 2.1E-F, Supp. Fig. 2.1.4H-I). In addition, golden standard markers of adipogenesis and mature adipocytes such as FABP4, PPARG, CEBPA, ADIPOQ, PLIN1-2-4, LPL, and others (see Methods) were solely upregulated in PR and SC samples post-differentiation (Supp. Fig. 2.1.4J). The expression of the latter correlated with the lipid droplet accumulation of the corresponding lines as quantified by the image-based adiposcore (p=0.81,
Fig. 2.1G, Supp. Fig. 2.1.2, see Materials and Methods), showing that inter-individual variability in terms of adipogenicity is also reflected at the transcriptomic level. Overall, even if the differentiation medium induced an adipogenic response in OM and MC hASPCs, it did not activate key regulators leading to overt lipid accumulation under the form of lipid droplets as, for example, supported by the enrichment of the “lipid storage” term solely in PR and SC-derived cells (Fig. 2.1E).

Next to their clear molecular and phenotypic dissimilarities in adipogenic response, we further investigated potential transcriptomic differences between SVF-adherent cells from different depots, which could underlie their distinct adipogenic potential. As previously reported, developmental genes such as HOXC8-10, HOXA9, and HOXD8 were highly expressed in SC samples (Fig. 2.1H)\textsuperscript{78,152}, as further illustrated by the enrichment of numerous terms linked to morphogenesis and development compared to the other depots both at t0 and t14 (Fig. 2.1H, Supp. 1.5). Interestingly, at t14, SC samples also showed enrichment of cell differentiation-related terms compared to the other depots, even considering the highly adipogenic PR samples (Fig. 2.1I). In contrast, PR-enriched genes in differentiated hASPCs were related to thermogenesis, suggesting that these cells have brown-like or beige-like adipocyte characteristics (Fig. 2.1I)\textsuperscript{153,154}. In OM samples, we observed a non-adipogenic gene expression signature with positive and negative enrichment of the terms “negative regulation of differentiation” and “white fat cell differentiation” respectively, compared to differentiation medium-exposed SVF-adherent cells from the other adipose depots (Fig. 2.1I). Undifferentiated OM SVF-adherent cells also exhibited significantly higher expression of genes linked to an inflammatory response, which remained after exposure to an adipogenic cocktail (Fig. 2.1I, Supp. Fig. 2.1.5A, Supp. Fig. 2.1.6A). This is not entirely unexpected given that the OM samples that were analyzed using BRB-seq mainly originated from obese patients undergoing bariatric surgery (Supp. Fig. 2.1.4A, Table 2.1), whose OM fat has previously been reported to show signs of inflammation\textsuperscript{79,155–157}. Interestingly, in both t0 and t14 time points, OM cells showed an enrichment of expressed genes linked to the vasculature and epithelium/endothelium development (Fig. 2.1I, Supp. Fig. 2.1.6B). This is consistent with our observation that many Keratin-related genes such as KRT8, KRT9, KRT18, and also LRRN4 or UPK1B (Fig. 2.1H, Supp. Fig. 2.1.5B) were among the top differentially expressed genes in OM cells versus those from other depots. The latter may be reflective of a mesothelial cell signature\textsuperscript{158}, suggesting that the SVF-adherent cell pool of OM is composed of various cell types, other than hASPCs. Finally, genes that were specifically expressed in MC compared to other depots were linked to protein metabolism and trafficking (Fig. 2.1I).

Taken together, we found that each depot features specific gene signatures that can be linked to functional implications, highlighting the regional specialization of AT based on its anatomical location. In addition, the observed experimental adipogenic potential is mirrored by the up- or down-regulation of pro-adipogenic markers in extraperitoneal and intraperitoneal adipose depot-derived cells, respectively. Finally, mesenchymal markers are highly expressed in SVF-adherent cells from all depots, validating the high enrichment of
hASPCs in the SV-adherent fraction. However, OM-derived samples also express an epithelial/mesothelial gene signature, suggesting the presence of mesothelial cells within the Lin⁻ SVF of OM.

**Figure 2.1** – *Ex vivo* cultures of SVF-adherent cells feature an anatomic footprint in their phenotype and transcriptome.

(A) Schematic of the experimental setup; primary SVF-adherent cell lines from human subcutaneous (SC), perirenal (PR), omental (OM), and mesocolic (MC) adipose tissues were cultured in parallel and harvested at the undifferentiated (t0) and
differentiated (t14) states for transcriptomic (BRB-seq) analysis; the same lines were seeded in a separate assay plate to quantify their adipogenic potential using the adiposcore (see Materials and Methods).

(B) Representative fluorescence microscopy images of SVF-adherent cells directly after isolation expansion to confluence and adipogenic induction (t14); Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA, scale bar = 1 mm.

(C) Barplot showing the log (adiposcore + 1) quantification of SVF-adherent cells in B; n = 14-22, 4-5 donors, 3-5 independent wells.

(D) t-SNE map based on the transcriptomic (BRB-seq) data of SVF-adherent cells from the indicated adipose depots (SC - yellow, PR - brown, OM - purple, MC - blue) and time points (t0 - light, t14 - dark); n = 12-61, 4-20 biological replicates, 1-4 independent replicates for each.

(E) Dot plot showing enriched, representative terms found by GSEA performed on the differential gene expression analysis results of t0 versus t14 samples for each depot of the data shown in D.

(F) Boxplot displaying the “Positive regulation of fat cell differentiation score”, based on the scaled expression of the corresponding GO term (GO:0045600) of the data shown in D.

(G) Scatter plot showing the relationship between the image quantification-based experimental adiposcore (shown in Supp. Fig. 2.1.1) versus the “mature adipocyte score” based on the scaled expression of well-known adipogenic markers (see Methods) of the transcriptomic samples from the same donor. Samples are grouped by depots and donors. Spearman correlation and adjusted R² of y~log(x+1) (plotted orange line with 95% confidence interval) values are indicated.

(H) Heatmap of top differentially expressed genes when comparing the indicated depot versus the three others at t0 of the data shown in D.

(I) Dot plot showing representative, enriched terms found by GSEA performed on the differential gene analysis results of each indicated depot versus the others at t14 of the data shown in D.

*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, One-Way ANOVA and Tukey HSD post hoc test (C), unpaired two-sided t-test (F).
2.3.3 scRNA-seq of SVF Lin− fraction reveals both common and unique subpopulations across the subcutaneous, visceral, mesocolic, and perirenal adipose depots.

Next, we explored whether the afore-observed transcriptomic and phenotypic differences across SVF-adherent cells derived from distinct adipose depots could in fact be driven by cellular heterogeneity. To do so, and to be as representative as possible of the SVF-adherent cells of the bulk RNA-seq analysis, we performed scRNA-seq of SVF Lin− (i.e., CD45−/CD31−) cells that were isolated from SC (n=3), OM (n=3), MC (n=2, from the same donor), and PR (n=3) adipose samples (Table 2.2), analyzing a total of 34’126 cells (on average, ~8’500 cells per depot). We first analyzed each resulting dataset independently, i.e., per depot and per donor (Supp. Fig. 2.2.1A-B). In all datasets, we identified hASPCs, defined by the expression of THY1 and PDGFRA, well-known mesenchymal markers151, as well as a population expressing muscle-related markers such as MYH11 but also ACTA2 and TAGLN (Supp. Fig. 2.2.1C), resembling a transcriptomic signature of vascular smooth muscle progenitor cells (VSMPs)159. Consistent with our bulk transcriptomic analysis, we also identified mesothelial cells, defined by the expression of MSLN, UP3KB, LRRN4, and Keratin-related genes, exclusively among SVF Lin− cells that were derived from OM AT (Supp. Fig. 2.2.1C, Supp. Fig. 2.2.1A). Finally, we identified a small number of immune (high CD45) and endothelial (high PECAM1) cells in the datasets from donor D07 that likely escaped the magnetic-based (MACS) Lin− enrichment that was performed for the corresponding SVF Lin− cells, as opposed to FACS used for the remaining datasets, where no CD45+ nor CD31+ cells are detected (Supp. Fig. 2.2.1A-C, for the type of enrichment method used prior to scRNA-seq, see Table 2.2).

Consistent with recent reports61,96,145, hASPCs (THY1 and PDGFRA high) displayed high heterogeneity and could be further stratified into at least two main clusters (Supp. Fig. 2.2.1A). To explore if and to what extent the identified hASPC subpopulations share molecular features across adipose depots, we performed three independent analyses. First, we calculated the overlap of the top cluster markers between datasets (Supp. Fig. 2.2.1D). We found that, while the percentage of shared markers tends to be the highest within samples isolated from the same depot and donor (Supp. Fig. 2.2.1E-F), the overlap across depots and donors is, on average, over 50% for most of the identified subpopulations (Supp. Fig. 2.2.1D). This result was confirmed when projecting each dataset onto each other using scmap160, revealing that on average more than 75% of cells from one specific population projected onto the corresponding population in other datasets, regardless of the depot of origin (Supp. Fig. 2.2.1G). Finally, we integrated the data by considering each dataset as a different batch and correcting accordingly. Once again, we observed an excellent overlap of the depot-counterpart populations in the tSNE space (Fig. 2.2A, Supp. Fig. 2.2.1H-I), which was further confirmed by clustering analysis (Fig. 2.2B). Taken together, we can confidently stipulate that human adipose SVF contains at least two main hASPC (THY1 and PDGFRA high) subpopulations/states, a feature common to all studied...
depots: SC, PR, OM, and MC (Fig. 2.2C-D, Supp. Fig. 2.2.2A-B). To explore the universality of this finding, we assessed yet another unexplored adipose tissue, the AT which accumulates surrounding the gallbladder in a subset of morbidly obese patients, and even if relatively few hASPCs were ultimately captured, we still retrieved the two main hASPCs subpopulations (n=1, Supp. Fig. 2.2.1B).

Based on their respective gene expression signatures, we labeled those two hASPC subpopulations as adipose stem cells (ASCs) and pre-adipocytes (PreAs) (Fig. 2.2B). Indeed, ASCs from all depots shared a gene signature enriched for DPP4, CD55, and PI16, and showed enrichment in genes involved in proliferation, collagen synthesis and stemness (Fig. 2.2C, Supp. Fig. 2.2C-D), consistent with the corresponding ASC subpopulation from mouse epigonadal (visceral AT) and SC adipose depots\cite{96,100,102,146,161} (Supp. Fig. 2.2.3). On the other hand, PreAs differentially expressed known markers of committed adipogenic cells such as PPARG, FABP4, PDGFRA, APOC, and APOE and showed enrichment of terms linked to differentiation, commitment, and lipid transport (Fig. 2.2C, Supp. Fig. 2.2C-D), again aligning with the corresponding mouse PreA population\cite{96,100,102,146,161} (Supp. Fig. 2.2.3). Furthermore, our annotations are consistent with the two ASPC states observed in human SC AT\cite{96,145} and predicted for OM AT\cite{61,145} (Fig. 2.2E, Supp. Fig. 2.2.4). To our knowledge, these hASPC states have never been described for other human anatomical locations.

Upon integration, we identified five smaller populations of hASPCs defined by specific transcriptomic signatures that are most likely subpopulations of PreAs (Fig. 2.2B-C). One of them was defined by a high expression of HHIP (Supp. Fig. 2.2.5A-B). Among the top differentially expressed genes of this population, we recognized several key ortholog markers, such as F3, CLEC11A, GDF10, MGP, and INMT (Supp. Fig. 2.2.5A), of a mouse ASPC subpopulation that we have previously characterized as having non- and anti-adipogenic properties, and accordingly named Adipogenesis Regulators (Aregs)\cite{102,148}. To compare the transcriptomic signature of this population across species in a more unbiased way, we computed a score based on the orthologs of the top HHIP+ hASPC markers, or of Aregs (reported in Ferrero et al.\cite{146} and Zachara et al.\cite{148}, see Methods) and showed that in both species, the score was significantly enriched in the corresponding population (Supp. Fig. 2.2.3). Recently, Emont and colleagues identified a cluster of hASPCs that is characterized by enriched expression of EPHA3 in their human AT scRNA-seq atlas, which has substantial similarities to murine Aregs\cite{145}. Notably, EPHA3 is specifically expressed by the HHIP+ hASPCs that we identified in our analyses, further supporting its alignment with mouse Aregs (Supp. Fig. 2.2.5A). To solidify the point that the previously described EPHA3+ hASPCs are indeed similar to our HHIP+ hASPCs, we transferred our cell annotation onto the Emont et al. dataset\cite{145} and found that the EPHA3+ population has a significantly higher prediction score for our HHIP+ population than the rest of the hASPCs (Fig. 2.2F). Finally, given that HHIP is coding for a surface marker, we could confirm the existence of a human SVF Lin−/HHIP+ cell population in the SC adipose depot using flow cytometry (Supp. Fig. 2.2.5C-D).
Another small population of hASPCs, present in every depot and donor, which we refer to as IFIT+ hASPCs, is defined by an extremely specific expression of interferon-related genes such as IFIT3, IFI6, and IFI27, a gene signature that is reflective of a viral immune response (Supp. Fig. 2.2.6A-B). A mesothelial Ifit+ population has already been reported in mouse OM\textsuperscript{143}; yet, our IFIT+ population does not express mesothelial markers (Fig. 2.2C) but mesenchymal ones (Supp. Fig. 2.2.6C). However, we found that, based on the expression of ortholog genes between mice and humans, this population shared a very similar signature with Ifit+ cells that we have previously reported and that emerged when we integrated multiple mouse ASPC scRNA-seq datasets\textsuperscript{146} (Fig. 2.2G, Supp. Fig. 2.2.3).

Another small hASPC subpopulation that we detected was characterized by high expression of Secreted frizzled-related proteins 2 and 4 (SFRP2 and SFRP4) (Supp. Fig. 2.2.7A), and its gene expression patterns were similar to those of a mouse ASPC subpopulation identified using scRNA-seq\textsuperscript{146} (Supp. Fig. 2.2.3), and a hASPC subpopulation identified by Emont and colleagues\textsuperscript{145} (Supp. Fig. 2.2.7B). SFRPs are known to inhibit the Wnt signaling pathway, a key regulator of adipocyte differentiation\textsuperscript{162}, and SFRP2-4, in particular, were shown to be upregulated in obesity, especially in visceral WAT\textsuperscript{163}. While we identified this population as present in all depots, we observed a general higher expression of SFRP2, but not SFRP4, in hASPCs from OM adipose depots (Supp. Fig. 2.2.7C-D).

While most hASPC subpopulations seem to exist in all analyzed adipose depots, albeit at different proportions, we found two depot-specific cell clusters: FMO2+ cells were specific to PR and MC, while IGFBP2+ cells to OM adipose tissue (Fig. 2.2D, Supp. Fig. 2.2.2A-B). An enrichment of IGFBP2+ cell markers was also observed in our bulk transcriptomic datasets of OM samples compared to other depots, both at the undifferentiated and post-adipogenic induction states (Supp. Fig. 2.2.8A-B), thus confirming their specificity to OM. Moreover, when projecting our annotation onto the dataset by Emont and colleagues\textsuperscript{145}, our IGFBP2+ cluster seems to align with one of their clusters (hASPC6)(Fig. 2.2E, Supp. Fig. 2.2.4).

In conclusion, we found hASPCs to be constituted of two main subpopulations/states: the ASCs and PreAs, which are conserved across all depots. However, the heterogeneity of hASPCs is obviously more complex, with more (sub)cellular states. Among them, we identified an OM-specific subpopulation defined by high expression of IGFBP2. Finally, in line with the established definition of peritoneum covering visceral AT, we found mesothelial cells almost exclusively in the OM AT (Fig. 2.2D, Supp. Fig. 2.2.2A, Supp. Fig. 2.2.8C-E). Only (very few) cells originating from MC samples were also expressing mesothelial markers (Supp. Fig. 2.2.8E), in line with the MC being itself covered by the peritoneum.
Figure 2.2 – scRNA-seq reveals common and specific cell populations across adipose depots.

(A) t-SNE cell map of integrated scRNA-seq datasets across four depots and 6 donors (D) (Table 2.2): OM, n=3, SC, n=3, and MC, n=2 (same donor) from matched donors, and PR, n=3 colored by the clustering of each dataset analyzed individually (as shown in Supp. Fig. 2.2.1A).

(B) t-SNE cell map of the data introduced in A colored by the identified clustering: Adipose Stem Cells (ASCs) - green, Pre-adipocytes (PreAs) - red, HHIP+ ASPCs - light blue, IFIT+ ASPCs - gray, SFRP4+ ASPCs - light green, RBPS+ ASPCs - light-red, FMO2+ ASPCs - brown, mesothelial cells (Meso) - purple, vascular smooth muscle progenitor cells (VSMPs) - orange, endothelial cells (Endo) - yellow, and immune cells (Immune) - pink. The percentage of cells belonging to each cluster is shown by a dot plot, with the exact number of cells on the right.

(C) Dot plot of 10 of the main specific markers of each identified cluster shown in B.

(D) Barplot displaying the percentage of cells of each depot shown in B coming from each cluster, excluding immune and endothelial cells, see color legend below.

(E) UMAP computed on the integrated data of SC- and OM-derived hASPCs and mesothelial cells published in Emont et al. (2022)\cite{Emont2022}, colored by the predicted cell type/state when transferring our cell cluster annotation, see color legend for the predicted cell types below.
**F** Violin plot showing the distribution of the prediction score of the HHIP+ hASPC population when transferred onto the scRNA-seq atlas of hASPCs of Emont et al.\textsuperscript{145}, where they identified hASPC4 as being transcriptomically similar to murine Aregs\textsuperscript{102,148}.

**G** Boxplot showing the distribution of the murine "Ifit+ ASPC" scores across the detected, distinct human SVF cell populations. The scores were based on the human orthologs of the murine top markers of the Ifit+ ASPCs based on the integration of scRNA-seq datasets of subcutaneous and visceral murine adipose tissues described in Ferrero et al.\textsuperscript{146}
2.3.4 The Lin− fraction of SVF harbors three main subpopulations with specific functions that are ubiquitous across depots

After having characterized the heterogeneity of the cellular SVF Lin− landscape across depots, we aimed at refining our functional characterization between depots, now at the subpopulation level. We thereby first focused on the main cell populations that are ubiquitous across depots: the ASCs, the PreAs and the VSMPs (Fig. 2.2A). Based on our scRNA-seq analyses expression profiles, we developed a specific sorting strategy that would allow downstream stratification and characterization of each of the aforementioned main SVF Lin− populations that are shared across depots. This sorting strategy involves three layers: 1) the first layer involves CD26, encoded by the gene DPP4 and specifically expressed by ASCs (Supp. Fig. 2.3.1A). Consistent with previous studies61,96,102, Dpp4 expression is specific to the murine ASC cluster146. 2) The second layer involves Vascular-adhesion protein 1 (VAP1), encoded by the gene AOC3 and highly expressed in VSMPs (Supp. Fig. 2.3.1A). In mouse, Aoc3 expression has mainly been described as being enriched in the PreA population96,102,146. However, based on our scRNA-seq integration of murine data, Aoc3 is in fact also highly expressed by murine VSMPs (Supp. Fig. 2.3.1B). 3) The third layer aims to enrich for PreAs. Several candidate surface markers appear specific to the PreA population (i.e., GPC3 or ICAM1). However, we reasoned that a simpler PreA enrichment approach would be to select for low expression of CD26 and VAP1. This approach would hold true in every depot except for the OM adipose depot, where two additional OM-specific cell populations would first need to be excluded: the mesothelial and the IGFBP2+ cells. Based on our transcriptional analyses, we selected the transmembrane 4 L6 family member 1 (TM4SF1) as a marker to first exclude OM-specific populations from our analysis (Supp. Fig. 2.3.1A, C). In sum, our sorting strategy involves antibodies directed against CD26, VAP1, and TM4SF1 (see Materials and Methods) to enrich for human ASCs (SVF Lin−/TM4SF1+/CD26+) and VSMPs (Lin−/TM4SF1+/VAP1+) (later referred to as DN for “double negative”) enriched for PreAs (Fig. 2.3A-B).

As expected, and in line with the transcriptomic findings, only OM-derived SVF showed a clearly positive population when stained with anti-TM4SF1 antibody, confirming its high enrichment in OM depot (Fig. 2.3B, Supp. Fig. 2.3.1D). However, as in the scRNA-seq datasets, we did find a few TM4SF1+ cells among MC SVF Lin− cells as well (Supp. Fig. 2.3.1D). Analysis of the flow cytometry profiles gathered from up to 37 human donors (Table 2.1) allowed us to quantify the relative abundance of the targeted populations in each of the three adipose depots (Fig. 2.3C). We found that the ASC pool is less abundant in OM AT compared to that of PR and SC, while SC AT is dominated by PreAs and the OM and PR ones by VSMPs (Fig. 3D). In line with our scRNA-seq findings, we found the same three populations in MC AT with relative ratios that resemble those of OM AT (Supp. Fig. 2.3.1E-F).
Having confirmed the existence of these shared SVF Lin– subpopulations in each depot, we aimed to interrogate their phenotypic behavior in vitro. When sorted separately, the CD26+ population outpaced all other populations in terms of cell growth regardless of the depot of origin (Supp. Fig. 2.3.1G), a feature that confirms their stem-like nature and is consistent with previous observations in mouse and human96,164. While highly proliferative, CD26+ scored the lowest in terms of adipogenic potential (Fig. 2.3E-F), further supporting the hypothesis that they are located at the very root of the adipogenic lineage. The VAP1+ cells had the highest adipogenic potential, followed by DN cells (Fig. 2.3E-F). The latter populations also exhibited a lower proliferative capacity compared to the ASC population (Supp. Fig. 2.3.1G).

Taking advantage of the relatively large cohort of human donors (n=37, Table 2.1) from which the adipose tissue was sampled, we investigated potential correlations between the relative abundance of each of the hASPC subpopulations and corresponding metadata such as BMI, age, and gender of the donors. Interestingly, we found that while the proportion of CD26+ cells (enriching for ASCs) is not affected by BMI changes, the latter appears to be correlated with DN depletion (enriching for PreAs). This correlation is particularly high in the SC, but also in the OM AT and is accompanied by a slight increase in the proportion of VAP1+ cells (enriching for the VSMPs) (Fig. 2.3G). In contrast, the age or sex of the donor did not seem to affect the equilibrium of cell populations within the SVF Lin– pool of any of the three analyzed adipose depots (Supp. Fig. 2.3.2).

Despite high similarities in the transcriptomes of ASCs and PreAs across depots in the scRNA-seq data, we observed that OM CD26+ and DN cells are consistently and significantly less adipogenic than equivalent SC and PR cells. To determine if cell-intrinsic features could explain the low adipogenic capacities of the OM cells, we explored the depot-specific transcriptomic signatures of these subpopulations in our scRNA-seq dataset. We noticed that across depots, the transcriptomes of ASC cells are more related than PreA ones (Supp. Fig. 2.2.2D, Supp. Fig. 2.3.3A), supporting the hypothesis that depot-specific features accumulate along commitment. We then identified genes of ASCs or PreAs that were enriched in a depot-specific manner (Fig. 2.3H, Supp. Fig. 2.3.3B). In line with their high adipogenic potential, hASPCs from SC, and especially PreAs, showed significantly higher expression of well-known adipogenic genes and transcription factors such \textit{KLF4}, \textit{KLF6}, \textit{WISP2}, \textit{APOE}, \textit{APOC1}, and \textit{CD36}. The pro-adipogenic character of PR adipose depot-isolated cells was also reflected in their transcriptome (Supp. Fig. 2.3.3B-C). For example, \textit{PIK3R1} is the most up-regulated gene in PR compared to other adipose depots, with PI3K/Akt signaling playing a crucial role in adipogenesis of human mesenchymal stem cells165. In mice, PI3K/Akt signaling has also been linked to browning by regulating GDF5-induced \textit{Smad5} phosphorylation166. While \textit{GDF5} expression was virtually absent in our scRNA-seq data, \textit{SMAD5} expression was specific to PR PreAs and ASCs. Similarly, \textit{ZBTB16} is a PR-specific marker known to induce browning167. With respect to populations that showed limited adipogenic potential, MC cells overexpressed genes linked to unfolded protein or protein folding (Supp. Fig. 2.3.3C) such as Heat-shock-
proteins (HSPs) (Fig. 2.3H, Supp. Fig. 2.3B), a large family of molecular chaperones. HSPs have been reported to interact with PPARγ to either stabilize it and enhance adipogenesis (Hsp90)\(^{168}\) or to destabilize it and inhibit adipogenesis (Hsp20)\(^{169}\). OM cells once again showed an expression enrichment of genes linked to the inflammatory response (Supp. Fig. 2.3.3C). Among the candidates that were specific to OM were also a number of markers that were previously described in the literature as having a negative impact on adipogenesis (Fig. 2.3H, Supp. Fig. 2.3B, RARRES2, RSPO3, RPL7, PTN, GAL, ALDH1A1, IGFBP3\(^{152,170-172}\)). Taken together, we were not only able to validate the findings that we inferred from bulk transcriptomic data at the scRNA-seq level, we also provided evidence that the PreA subpopulations are likely the ones that contribute the most to the depot-specific transcriptomic signatures that we captured at the bulk level.
Figure 2.3 – The SVF Lin− composition differs between subcutaneous, omental, and perirenal adipose tissues, and shared sub-populations from different anatomical locations exhibit consistent but also intrinsic molecular and cellular phenotypes.

(A) Sorting strategy scheme to enrich for Adipose stem cells (ASCs), pre-adipocytes (PreAs), Vascular smooth muscle progenitors (VSMPs), and OM-specific cells.

(B) Flow cytometry-based representative profiles and gating strategy for SC, OM, and PR SVF from the same donor (D23) to isolate SVF Lin−/TM4SF1− cells.

(C) Flow cytometry-based analysis of the abundance of each cell subpopulation gated from the Lin−/TM4SF1− fraction of SVF cells; SC n = 37, OM n = 35, PR n = 17 donors.

(D) Bar plot to compare the flow cytometry-based abundance of the indicated SVF populations across depots. The three populations accumulate to 100% of Lin−/TM4SF1− gated cells by depot; SC n = 37, OM n = 35, PR n = 17 donors.

(E) Representative fluorescence microscopy images of SVF Lin−/TM4SF1−, CD26+, DN, and VAP1+ SVF populations from each depot after in vitro adipogenic differentiation (see Methods); Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA, scale bar = 100 um.

(F) Quantification of the adipogenic potential of the SVF Lin−/TM4SF1− populations shown in D; Values are normalized to average adiposcore of the reference Lin−/TM4SF1− population; n = 12-21, 3-7 donors, 1-4 independent wells each.

(G) Scatter plot showing the correlation between the % Lin−/TM4SF1− cells from each indicated SVF population and BMI across donors.

(H) Heatmap of the top 30 genes detected as significantly higher expressed in the indicated depot versus all other depots (only genes detected as differentially expressed in each pairwise comparison were retained) when focusing on ASCs (left) or PreAs (right); Average log normalized expression scaled by row.

*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, One-Way ANOVA and Tukey HSD post hoc test (C, D, F), and linear regression analysis with its relative goodness of fit, and the FDR-adjusted p-values of the Pearson correlations (G).
2.3.5 Mesothelial cells inhibit adipogenesis of omental hASPCs

We next questioned whether the presence of OM-specific cell populations (Fig. 2.4A) might influence the adipogenic capacity of the precursor cells themselves, as triggered by two key observations: 1) OM VAP1+ and DN cells, which are depleted of TM4SF1+ cells via the utilized sorting strategy, did show a discrete ability to differentiate (Fig. 2.3E-F); 2) several genes that were previously linked to the non-adipogenic phenotype of OM SVF-adherent cells were specific to mesothelial and/or IGFBP2+ cells (e.g., CD200, WT1, and ALDH1A2252, Supp. Fig. 2.4.1A).

Using TM4SF1 as a surface marker for OM-specific populations (mesothelial and IGFBP2+ cells, Supp. Fig. 2.3.1C), we separated TM4SF1+ cells from the total OM SVF Lin− fraction to study the adipogenic behavior of “pure” OM hASPCs (Fig. 2.4B). In line with our previous observation on the adipogenic potential of OM DN and VAP1+ subpopulations (Fig. 2.3E-F), we found that OM SVF Lin−/TM4SF1− cells, enriching for OM hASPCs and later referred to as TM4SF1− cells, are significantly more adipogenic than the total OM SVF Lin− fraction, which does contain the OM-specific mesothelial and IGFBP2+ cells. Not surprisingly, since mesothelial cells have previously been shown to be non-adipogenic, the OM SVF/Lin−/TM4SF1+ cells, here referred to as TM4SF1+ cells, did not accumulate any lipid droplets (Fig. 2.4C-D). Importantly, however, the increase in differentiation observed for TM4SF1− cells compared to the Lin− fraction was greater than expected by the simple, proportional removal of the non-adipogenic TM4SF1+ cells (accounting for roughly 20% of the total SVF Lin− fraction, Fig. 2.3C). Taken together, in vitro cultured OM hASPCs seem to be subjected to inhibitory cues that may stem from the OM-specific TM4SF1+ populations. Interestingly, the relative fraction of OM TM4SF1+ cells within the total SVF Lin− cell pool positively correlated with the BMI of donors, suggesting a possible functional role of TM4SF1+ cells in AT expansion (Supp. Fig. 2.4.1B). Morphologically, TM4SF1+ cells stood out from regular round and cobblestone-like OM hASPCs (OM SVF/Lin−/TM4SF1−)275,276 (Supp. Fig. 2.4.1C), as they had spindle-like shape characteristic of mesothelial cells.

To test whether the observed inhibitory cues within the OM SVF Lin− cell pool has a negative influence not only on the adipogenic potential of OM hASPCs but also on those of SC or PR, we set up a mixing experiment where SC Lin− or PR Lin− cells were co-cultured with increasing ratios of OM Lin− cells (Fig. 2.4E-F, and Supp. Fig. 2.4.2). We observed that despite a linear increase in the relative proportion of OM SVF Lin− cells among SC SVF Lin− ones, the resulting decrease in adipogenic potential was non-linear (Fig. 2.4F). In other words, the decrease in differentiation was greater than expected by the simple relative “dilution” of SC SVF Lin− cells. To control for the fact that SC cells were not overgrown by OM cells, we quantified the relative number of nuclei across conditions which was stable (Supp. Fig. 2.4.2A) and controlled that the expression of an SC-specific marker DKK2 was linearly increasing with the proportion of SC cells (Supp. Fig. 2.4.2B-C). Using a similar approach but this time mixing OM SVF Lin− cells with PR SVF Lin− cells did not reveal any regulatory effect, as we observed a relatively linear relationship between the increase in differentiation and the
proportion of PR cells per well (Supp. Fig. 2.4.2D-F). Thus, our findings suggest that the presence of OM TM4SF1+ cells results in lowered adipogenic potential, although this effect is not universal among hASPCs, hinting at depot-specific sensitivities to potential inhibitory cues stemming from OM SVF Lin– cells.

2.3.6 IGFBP2+ OM SVF Lin– cells can be found in situ and appear to transition between mesothelial and mesenchymal cell types

To better understand the cellular nature of the observed inhibitory effect, we set out to determine which TM4SF1+ population may be responsible for this adipogenic inhibition: the previously described mesothelial cells, and/or a small cluster of cells that highly express IGFBP2 (Fig. 2.4A and Supp. Fig. 2.3.1C). The latter appears a plausible candidate since IGFBP2 has previously been shown to exert an anti-adipogenic effect both in mice and humans\textsuperscript{177,178}, which is why we first aimed to better define this population. At the single-cell level, the IGFBP2+ population appeared to have an intriguing dual gene expression signature, sharing markers with both hASPCs and mesothelial cells (Supp. Fig. 2.4.3). Such expression signature may at first glance suggest a technical artifact known as doublets, when two cells are mistakenly co-captured and considered as a single one. However, IGFBP2+ cells did not display a larger library size or number of captured features (Supp. Fig. 2.2.1I), which would be expected for doublets due to a larger initial RNA content compared to singlets. More importantly, we found that these cells express, on the one hand, specific markers such as IGFBP2, RBP1, WNT4, or WNT6 and, on the other, markers to a higher level than in ASPCs or mesothelial cells alone (Supp. Fig. 2.4.3), which is technically impossible for randomly co-encapsulated cells. To validate the existence of this population in another independent dataset, we transferred our cell annotation onto the recently published scRNA-seq atlas of human SC and OM ATs\textsuperscript{145}. We found that, first, only cells from OM harbor a positive prediction score for IGFBP2+ cells (Supp. Fig. 2.4.4A), validating once more their specificity to the OM. Second, the cells predicted as IGFBP2+ aligned with a cluster that was independently identified by Emont et al.\textsuperscript{145} (Supp. Fig. 2.4.4B-D, Fig. 2.2F) and showed enrichment for IGFBP2+ cell markers, as illustrated by the marker-based expression score (Supp. Fig. 2.4.4E). Interestingly, the frequency of this population (relative to ASPCs and mesothelial cells) highly correlated with the BMI of the donors ($\rho=0.95$, Supp. Fig. 2.4.4F). Furthermore, and once again, aside from expressing their own specific markers (Supp. Fig. 2.4.4G-H), the predicted cells co-expressed mesothelial and ASPC markers (Supp. Fig. 2.4.4I) and aligned along a “bridge” between the two cell types. This duality in gene expression could reflect cells that are transitioning from one cell type to another. To computationally test this hypothesis, we performed trajectory inference on OM hASPCs (ASCs, PreAs), IGFBP2+ cells, mesothelial cells as well as VSMPs as a negative control. The trajectory was computed using PAGA as it can identify continuous and disconnected structures in the data\textsuperscript{179}. The inferred graph predicted branches connecting ASPCs to mesothelial cells through IGFBP2+ cells (Fig. 2.4G). As positive and negative controls of the validity of the graph structure, ASCs and PreAs were
also connected by a robust branch, as previously reported in mouse\textsuperscript{96,146}, and VSMPs were, as expected, not connected to the main trajectory. When ordering the cells by their pseudotime along the trajectory starting from ASCs (Supp. Fig. 2.4.5A), we observed a gradual decrease and increase of hASPC and mesothelial cell markers, respectively, along the connecting branch (Fig. 2.4H-I), as well as an up-regulation of IGFBP2+ cell markers during the transition (Fig. 2.4H, J, Supp. Fig. 2.4.5B). Altogether, these results indicate that IGFBP2+ cells might represent cells that transition between mesothelial and mesenchymal cell types. Accordingly, we found the GO term “epithelial-to-mesenchymal transition” (EMT) to be enriched among the IGFBP2+ cells’ differentially expressed genes (Supp. Fig. 2.4.5C-D). In addition to the genes enriched in the GO term, such as Slug (SNAI2), we also found several genes that are expressed by the transitioning cells that were previously linked with EMT, such as genes from the Wnt family, Matrix Metallopeptidase (MMPs), ZEB transcription factors, and others\textsuperscript{180–182} (Supp. Fig. 2.4.5D). TGF-β signaling, and especially TGF-β1, has also been described as a master regulator of EMT linked to wound healing and fibrosis\textsuperscript{183,184}. In line, we found that IGFBP2+ cells have an enriched expression linked to “response to TGF-β”, but not significantly to TGF-β1 in particular. These cells also express genes in relation to epithelial migration and proliferation. Finally, EMT in the peritoneum of mice has been shown to induce the following gene programs: angiogenesis, hypoxia, inflammatory responses, cell cycle markers, and downregulation of adhesion molecules\textsuperscript{185}. The corresponding GO terms were all significantly enriched among the IGFBP2+ cell markers (Supp. Fig. 2.4.5C). Thus, our findings point to the existence of cells that likely transition between mesothelial and mesenchymal cells, even under “steady-state-like” conditions.

Having identified IGFBP2-expressing and transitioning cells computationally, we next defined a sorting strategy to separate these cells and the mesothelial cells from the total human OM SVF. To do so, we retained TM4SF1 as a valuable marker to segregate OM-specific cells from OM hASPCs and added MSLN as a marker that is solely expressed by mesothelial cells (Supp. Fig. 2.3.1C). Hence, we defined IGFBP2+ cells as OM SVF Lin–/TM4SF1+/MSLN– and mesothelial cells as OM SVF Lin–/TM4SF1+/MSLN+ (Supp. Fig. 2.4.6A), each fraction accounting for 5 to 10% of the total OM SVF/Lin– fraction (Supp. Fig. 2.4.6B). While we were able to highlight a significant, positive correlation between the abundance of TM4SF1+ cells and the BMI of donors based on flow cytometry analysis (Supp. Fig. 2.4.1B), that between the abundance of TM4SF1+/MSLN– cells, enriching for IGFBP2+ cells specifically, and BMI was not significant (Supp. Fig. 2.4.6C). Using the same panel of markers, we then set out to localize IGFBP2+ cells \textit{in situ}. Interestingly, both MSLN and TM4SF1 highly stained the boundaries of the AT lobules (Supp. Fig. 2.4.7A), likely revealing a mesothelial mono-layer peritoneum-like structure that pads the OM itself. Accordingly, the majority of cells looked equally intense for both markers; we therefore defined the majority of cells as mesothelial cells (Fig. 2.4K, red arrows, Supp. Fig. 2.4.7B). However, intermingled among these mesothelial cells, we identified cells that were much more intense in the TM4SF1 channel than the MSLN one (Fig. 2.4K, and Supp. Fig. 2.4.7B, white arrows), reminiscent
of our IGFBP2+ cell gene expression signature. The absence of background staining was assessed by both unstained control and secondary-only staining (Supp. Fig. 2.4.7). Finally, and emphasizing their transitioning cell nature, we found that confluent Lin−/TM4SF1+/MSLN− cells harbor the specific mesothelial-cobblestone-like morphology, but when expanding, they tend to be spindle-like in shape, resembling mesenchymal cells (Fig. 2.4L, Supp. Fig. 2.4.8).
Figure 2.4 – OM SVF harbors cells and signals that mediate adipogenic inhibition, while IGFBP2+ cells appear in transition between mesothelial cells and ASPCs.

(A) t-SNE cell map of integrated scRNA-seq datasets highlighting the two OM-specific populations: Mesothelial cells in purple and IGFBP2+ cells in blue.

(B) Representative flow cytometry scatter plot of OM SVF Lin– cells (D05) stained with TM4SF1 antibody showing the gating strategy for sorting OM SVF Lin– specific subpopulations as Lin–/TM4SF1+ and Lin–/TM4SF1– cells.

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(C) Representative fluorescence microscopy images of OM SVF Lin−/−, Lin−/TM4SF1− and Lin−/TM4SF1+ cell populations after adipogenic differentiation (see **Methods**); Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA; Scale bars = 100 μm.

(D) Barplot showing the adiposcore of the cell populations that are indicated in C; n = 6-23, 4 donors, 1-6 independent wells for each.

(E) Representative fluorescence microscopy images of SVF Lin− cells in mixing experiments after 14 days of adipogenic differentiation, where SVF Lin− cells from OM and SC adipose tissues of donor 68 (D68) were mixed directly after cell isolation at the indicated proportions. Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA, scale bar = 100 mm.

(F) Quantification of the extent of adipogenic differentiation of the distinct, mixed OM and SC SVF Lin− cell populations, as presented in E. Values across biological replicates are normalized to the average adiposcore of the reference 100% SC Lin− condition. The relative proportion (0-100%) of SC SVF Lin− cells in each well is plotted on the x-axis. Error bars represent standard deviation from the average, the linear and exponential regression with corresponding R² coefficients are shown in red and blue, respectively. The black line represents the expected increase of adipogenesis for a linear dilution between 0 and 100% of SC SVF Lin− cells; n = 16, 4 biological replicates, 4 independent wells for each.

(G) PAGA-inferred trajectory superimposed on the PAGA-initialized ForceAtlas2 layout. The size of the dots is proportional to the number of cells in the cluster, and the thickness of the lines indicates the confidence of the obtained trajectory relationship (the thicker, the more confident).

(H) Heatmap showing the gene expression changes along pseudotime calculated on the trajectory shown in G. Genes decreasing from hASPCs (ASCs and PreAs) to Mesothelial cells are highlighted in red, genes increasing from hASPCs to Mesothelial cells are highlighted in purple, and genes specific to IGFBP2+ cells are highlighted in blue; log normalized gene expression scaled by row (quantile normalization).

(I) Scatter plot showing the average of quantile-normalized gene expression highlighted in red or purple on the heatmap shown in H for each cell along the pseudotime calculated on the trajectory shown in G. The plot focuses on the transition between PreAs (red) and Mesothelial cells (purple), passing by IGFBP2+ cells (blue). A locally estimated scatterplot smoothing (LOESS) smoothing with 95% confidence interval is shown.

(J) Scatter plot showing the average of quantile-normalized gene expression highlighted in blue on the heatmap shown in H for each cell along the pseudotime calculated on the trajectory shown in G. The plot focuses on the transition between PreAs (red) and Mesothelial cells, passing by IGFBP2+ cells (blue). A generalized additive model (GAM) fit with 95% confidence interval is shown.

(K) Confocal microscopy fluorescent images of the *in situ* immunohistochemistry-based localization of TM4SF1+ (green) and MSLN+ (pink) cells in OM adipose tissue in donor 67. Nuclei are stained with DAPI (blue). The arrows indicate TM4SF1+/MSLN− cells (white) and TM4SF1+/MSLN+ cells (red) in the periphery of the adipose tissue lobules. Scale bars, 50 μm. Experiments were repeated at least three times, yielding similar results.

(L) Brightfield microscopy images of OM SVF Lin−/TM4SF1+/MSLN− (i.e., IGFBP2+) cells from donor 67 reveal a mesothelial cobblestone-like morphology when confluent and fibroblast spindle-like morphology upon expansion; Scale bars, 10 mm.

*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, REML analysis with matched values for the same donor and Tukey HSD post hoc test (D).
2.3.7 IGFBP2+ SVF Lin− OM cells inhibit adipogenesis through IGFBP2

Having identified cells with low MSLN but high TM4SF1 expression both in situ by immunohistochemistry and ex vivo by FACS, we wanted first to validate that cells that we isolated as Lin−/TM4SF1+/MSLN− (Fig. 2.5A) correspond to the IGFBP2+ cells that we identified in the scRNA-seq data. We first confirmed enriched IGFBP2 expression in the sorted cells compared to total OM SVF Lin− or OM TM4SF1−/MSLN− ASPCs and SC SVF Lin− cells by qPCR (Fig. 2.5B). To assess whether IGFBP2 is secreted or intracellular186, we looked for the presence of the IGFBP2 protein in the supernatant of the OM-specific subpopulations as well as in SC and PR SVF Lin− cells as negative controls. Using ELISA, we measured the highest concentration of IGFBP2 in the supernatant of OM Lin−/TM4SF1+/MSLN− cells, quantified at approximately 35 ng/ml after 48h, followed by the mesothelial cells which secrete less than 20 ng/ml of IGFBP2. In contrast, we only detected low IGFBP2 levels in the supernatant of OM SVF Lin− cells, OM hASPCs (OM Lin−/TM4SF1−/MSLN−), or PR Lin− and no secretion by the SC SVF Lin− cells (Fig. 2.5C). To translate these IGFBP2 ELISA values to a more physiological model of IGFBP2 secretion by the OM AT, we incubated total OM AT in PBS and measured the secreted IGFBP2 amount after 24, 48, and 72 hours. The concentration of IGFBP2 increased linearly over time, leading to a secretion of ~5ng/mL for 100mg of tissue every 24h (Fig. 2.5D).

Having identified a sorting strategy to stratify the IGFBP2-secreting and mesothelial cells, we now aimed at identifying which cell fraction is responsible for the anti-adipogenic effect exerted by the OM-specific TM4SF1+ cells described above (Fig. 2.4C). More specifically, given that IGFBP2 is a well-known OM-specific adipokine that has been shown to have anti-adipogenic properties177,187,188, we wondered if the SVF Lin−/TM4SF1+/MSLN− cells could exert this effect in a paracrine fashion. To test this hypothesis, we used a transwell setup to expose receiving cells at the bottom to the secretome of either IGFBP2-secreting, mesothelial, or control cells, preventing cell-to-cell contact. At the bottom, we seeded the highly adipogenic SC SVF Lin− cells, since we already showed their sensitivity to the OM SVF Lin− mediated anti-adipogenic effect (Fig. 2.4E-F). By doing so, we observed the highest and most significant adipogenic inhibition on SC cells when they were exposed to OM SVF Lin−/TM4SF1+/MSLN− cells, while the adipogenic inhibition was milder and more variable when SC cells were exposed to the OM Lin−/TM4SF1+/MSLN+ fraction (Fig. 2.5E-F, Supp. Fig. 2.5.1A). To validate that the PR cells are less responsive to this inhibitory signal, as shown in direct co-culture experiments (Supp. Fig. 2.4.2D-F), we performed the same transwell experiment, but this time with PR SVF Lin− cells at the bottom. Consistent with our first results, PR hASPCs were rather insensitive to the inhibitory action of OM SVF Lin− cell subpopulations on adipogenesis (Supp. Fig. 2.5.1B-D).

To test whether IGFBP2 is at least partly regulating the high anti-adipogenic effect of IGFBP2-secreting cells, we knocked down (KD) IGFBP2 in the OM SVF Lin−/TM4SF1+/MSLN− cell population using siRNA probes. After validating the KD both at the mRNA and secreted protein levels (Fig. 2.5G-H), we used again a transwell set-up to expose SC SVF Lin− cells to their secretome as well as to that of OM SVF Lin−/TM4SF1+/MSLN− cells
treated with non-targeting siRNA control (NC1). We found that the SC cells exposed to the \( \text{IGFBP2} \) KD cells are significantly more adipogenic than those exposed to the control (Fig. 2.5I-J, Supp. Fig. 2.5.1E), further supporting the notion that Lin–/TM4SF1+/MSLN– cells exert an anti-adipogenic action via IGFBP2.

### 2.3.8 IGFBP2-mediated adipogenic inhibition occurs in an IGF-independent manner

Prompted by the evidence that IGFBP2 at least partially orchestrates the anti-adipogenic environment observed within OM SVF, we set out to better understand the mechanism underlying IGFBP2’s anti-adipogenic effect. First, we tested if exogenous recombinant IGFBP2 is itself inhibitory by treating SVF-adherent cells from SC or PR depots with increasing IGFBP2 concentrations ranging from 0.5 to 4nM (Supp. Fig. 2.5.2A-F). We observed a linear decrease of differentiation of SC cells with increasing IGFBP2 concentrations, while only a milder effect was observed on PR cells. Notably, we observed significant inhibition of SC cells starting from as low as 1nM IGFBP2 (Fig. 2.5K-L). This is a value close to that measured in the supernatant of IGFBP2 (Lin–/TM4SF1+/MSLN–)-secreting cells (Fig. 2.5C), which is why we used it for all further experiments.

IGFBP2 is known to act through two main mechanisms involving either IGF-dependent or IGF-independent signaling\(^{189}\). Under the first scenario, the presence of IGFBP2 in the extracellular environment of hASPCs would sequester IGF-I and/or IGF-II and interfere with their pro-adipogenic signaling\(^{190-193}\). Under the second scenario, IGFBP2 would activate a signaling cascade by binding to the \( \alpha 5\beta 1 \) integrin receptor, inducing cells to stay in their pre-adipocyte state\(^{194}\). Hence, we aimed to narrow down through which of these mechanisms IGFBP2 might influence adipogenesis of hASPCs from adipose depots to which we had recurrent access (i.e., SC, PR, and OM).

To test the first hypothesis of IGFBP2 sequestering IGFs, we co-treated SVF-adherent cells with both IGFBP2 and IGF-I or IGF-II, as well as with the three recombinant proteins alone. Based on the literature\(^{191,193}\), we decided to use a concentration of 10nM for both IGF-I and IGF-II, given that, rather surprisingly, we were unable to observe a significant effect on the adipogenic potential of hASPCs treated with IGFs at any concentration ranging from 2.5 to 40nM, neither on SC or PR SVF Lin– cells (Supp. Fig. 2.5.2A, G-J, D, K-N). For SC cells, the drop in adipogenic potential was comparable when cells were treated with IGFBP2 both in the presence or in the absence of IGFs (Fig. 2.5K-L, Supp. Fig. 2.5.3A), suggesting an IGF-independent action of IGFBP2. Once again, PR lines appeared to be less sensitive to the action of IGFBP2 and IGF treatments. In fact, even though we observed a similar trend to that observed for SC cell behavior when treating PR cells with IGFBP2 both in the presence or in the absence of IGFs, none of the observed decreases in adipogenic potential was significant when compared to the non-treated cells (Supp. Fig. 2.5.3B-D). Overall, this is consistent with our previous observations suggesting that PR SVF-adherent cells are less sensitive to the inhibitory effect of OM SVF Lin– cells in the cell mixing setup (Supp. Fig. 2.4.2D-F) and of OM SVF Lin–/TM4SF1+/MSLN– cells in the transwell setup (Supp. Fig. 2.5.1B-D).
Next, we explored to what extent OM TM4SF1– cells, enriching for OM hASPCs, can respond to IGFBP2 and IGF treatments, since these cells anatomically co-exist with the IGFBP2-secreting cells. Even if OM TM4SF1– cells are intrinsically lowly adipogenic, we observed a significant differentiation decrease when these cells were treated with IGFBP2 (Fig. 2.5M-N, Supp. Fig. 2.5.3F), which further supports the anti-adipogenic potential of IGFBP2-secreting cells in their depot of origin. Contrary to PR and SC cells, OM cells were more sensitive to the IGF-I and IGF-II treatments but with a high degree of variability between batches (Fig. 2.5M-N). However, when co-treated with IGFs and IGFBP2, the differentiation of OM TM4SF1– cells was again significantly lower than in non-treated cells (Fig. 2.5M-N, Supp. Fig. 2.5.3E). The fact that IGF treatment could not maintain the increased adipogenic potential when cells were treated with IGFs only, even if given in excess (10nM of IGFs versus 1nM of IGFBP2 for a stoichiometry of 1:1), further suggests an IGF-independent mode of action by IGFBP2.

We then tested the second hypothesis, namely that IGFBP2 may act in an IGF-independent fashion by activating the α5β1 integrin receptor. To do so, we used echistatin, a known antagonist of the integrin receptor, at a concentration of 100 nM for the first 48h of adipogenic induction, as longer treatment resulted in cell detachment. We therefore coupled echistatin to IGFBP2 treatment only during the first 48h of differentiation. Interestingly, we found that echistatin alone is able to significantly enhance the differentiation of SC SVF-adherent cells, while, when cells were co-treated with IGFBP2 and echistatin, the adipogenic potential of the treated cells was similar to that of non-treated control cells (Fig. 2.5K, O, Supp. Fig. 2.5.3A). Interfering with integrin receptor function in PR SVF-adherent cells yielded a similar trend in overall adipogenic potential as observed for SC cells (Supp. Fig. 2.5.3B-C, F). This result highlights the important role played by integrin receptor signaling in mediating the adipogenic potential of cells, as echistatin had a significant effect even on the highly adipogenic PR cells.

Finally, when treating OM TM4SF1– cells with echistatin, we observed a significant increase in the ability of these intrinsically non-adipogenic cells to accumulate lipid droplets (Fig. 2.5M, P, and Supp. Fig. 2.5.3E), in line with findings by Yau and colleagues. Furthermore, co-treatment with echistatin and IGFBP2, both competing for binding to the α5β1 integrin receptor, led to a significant increase in differentiation compared to non-treated cells, but less than echistatin-only treatments (Fig. 2.5M, P, and Supp. Fig. 2.5.3E).

Taken together, our observations point to the existence of an OM-specific and transitioning cell population that highly expresses and secretes IGFBP2, which negatively impacts the adipogenic potential of OM and SC hASPCs, by signaling through an IGF-independent mechanism involving the integrin receptor alpha. However, we cannot completely exclude that the restored adipogenic potential of the analyzed cells (as compared to non-treated control cells) may be driven by two independent and opposite effects, i.e., inhibition by IGFBP2 and enhancement by echistatin. Indeed, the observed significant increase in adipogenesis for example of PR
cells upon echistatin treatment (Supp. Fig. 2.5.B-C, F) suggests that the integrin receptor can also negatively regulate adipogenic potential in a manner that may be independent of IGFBP2 activation.
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**A**  
IGFBP2 high Lin- Mesothelial  
OM ASPCs - MSLN - CF568

**B**  
IGFBP2
Depot: OM SC

**C**  
IGFBP2
Depot: OM SC PR

**D**  
IGFBP2
Depot: SC SVF

**E**  
SVF
Depot: SC SVF

**F**  
Normalized adiposcore [a.u.]
Depot: OM SC

**G**  
IGFBP2
Depot: SC SVF

**H**  
IGFBP2
Depot: SC SVF

**I**  
IGFBP2
Depot: OM SC

**J**  
Normalized adiposcore [a.u.]
Depot: OM SC

**K**  
OM ASPCs D73  
Ctrl IGFBP2 Echistatin  
IGF-I IGF-II IGFBP2 IGF-I IGF-II

**L**  
IGFBP2
Depot: OM SC

**M**  
OM ASPCs D73  
Ctrl IGFBP2 Echistatin  
IGF-I IGF-II IGFBP2 IGF-I IGF-II
Figure 2.5 – IGFBP2+ cells secrete IGFBP2 to inhibit adipogenesis of neighboring cells through an IGF-independent, paracrine mechanism.

(A) Representative flow cytometry scatter plot of OM SVF Lin− cells (D53) stained with TM4SF1 and MSLN showing the gating strategy to enrich for specific SVF Lin− subpopulations: Lin−/TM4SF1−/MSLN− (OM ASPCs - Black border), Lin−/TM4SF1+/MSLN− (IGFBP2+ cells, Blue border), or Lin−/TM4SF1+/MSLN+ (mesothelial cells, Purple border); DP: Double Positive; DN: Double Negative.

(B) qPCR-based quantification of IGFBP2 expression. Ct values are normalized first to HPRT1 expression, then to the ΔCt of OM SVF cells; n=4, 2 donors, 2 technical replicates.

(C) ELISA-based quantification of secreted IGFBP2 (ng/mL) in the supernatant of the indicated cellular populations after 48h of secretion in a serum-free medium; n=8, 4 donors, 2 technical replicates.

(D) ELISA-based quantification of IGFBP2 levels (ng/mL), as secreted by 100mg of OM adipose tissue over the indicated time window in PBS; n=4, 2 donors, 2 technical replicates.

(E) Representative fluorescence microscopy images of “receiver” SC SVF adherent cells, at the bottom of a transwell set-up, after adipogenic differentiation when co-cultured with the indicated SVF populations on top of the transwell: paired SC SVF adherent cells, OM SVF adherent cells, OM SVF/Lin−/TM4SF1− (OM ASPCs), OM SVF/Lin−/TM4SF1+/MSLN− (IGFBP2+ cells), or OM SVF/Lin−/TM4SF1+/MSLN+ (mesothelial cells). Top row: SC cells from D25, OM cells from D54; bottom row: SC and OM cells from D65.

(F) Barplot showing the adiposcore quantification of “receiver” cells in E. Values are normalized to the average adiposcore of the reference top SC SVF adherent condition; n=12, 4 donors, 3 independent wells.

(G) qPCR-based quantification of IGFBP2 expression in SVF/Lin−/TM4SF1+/MSLN− cells subjected to either IGFBP2 siRNA or non-targeting siRNA control (NC1), as retrieved from the transwell set-up. SC SVF adherent cells are also used as negative control. Ct values are normalized first to HPRT1 expression, then to the ΔCt of NC1 control; n = 2, 1 donor, two technical replicates.

(H) ELISA-based quantification of IGFBP2 levels in the supernatant of OM SVF Lin−/TM4SF1+/MSLN− cells subjected to either IGFBP2 siRNA or non-targeting siRNA control (NC1). SC SVF/Lin− cells are used as negative control; n = 2, 1 donor, two technical replicates.

(I) Representative fluorescence microscopy images of “receiver” SC SVF adherent cells, at the bottom of the transwell set-up, after adipogenic differentiation when co-cultured, with the indicated cells on top of the transwell: paired SC SVF adherent control cells, OM SVF/Lin−/TM4SF1−/MSLN− cells treated with non-targeting siRNA control (NC1), OM SVF/Lin−/TM4SF1+/MSLN− cells treated with IGFBP2 siRNA. Top row: SC and OM cells from D74, bottom row: SC cells from D63, and OM cells from D75.

(J) Barplot showing the adiposcore quantification of “receiver” cells in I; n=16-20, 4 donors, 2-4 independent wells.

(K) Representative fluorescence microscopy images of SC SVF-adherent cells after adipogenic differentiation when treated with the indicated interfering compounds: IGFBP2 1nM, IGF-I 10nM, IGF-II 10nM, Echistatin 100nM.

(L) Barplot showing the adiposcore quantification of cells in K with a focus on the IGF-dependent signaling pathway of IGFBP2. Values are normalized to the average adiposcore of the untreated control cells (Ctrl); n=12, 4 donors, three independent wells.

(M) Representative fluorescence microscopy images of OM SVF/Lin−/TM4SF1−/MSLN− cells after adipogenic differentiation when treated with the indicated, interfering compounds: IGFBP2 1nM, IGF-I 10nM, IGF-II 10nM, Echistatin 100nM.

(N) Barplot showing the adiposcore quantification of cells in M with a focus on the IGF-dependent signaling pathway of IGFBP2. Values are normalized to the average adiposcore of the untreated control cells (Ctrl); n=9, 3 donors, three independent wells.

(O) Barplot showing the adiposcore quantification of cells in K with a focus on the IGF-independent signaling pathway of IGFBP2. Values are normalized to the average adiposcore of the untreated control cells (Ctrl); n=12, 4 donors, three independent wells.

(P) Barplot showing the adiposcore quantification of cells in M with a focus on the IGF-independent signaling pathway of IGFBP2. Values are normalized to the average adiposcore of the untreated control cells (Ctrl); n=9, 3 donors, three independent wells.

For images in E, I, K, and M: Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA, scale bar=100 um. *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, One-Way ANOVA and Tukey HSD post hoc test (B, C, F, L, O), REML analysis with matched values for the same donor and Tukey HSD post hoc test (J, N, P).
2.4 Conclusion

Despite significant efforts, our understanding of hASPC heterogeneity and function across human adipose depots is still limited, which has been further hampered by the lack of hASPC consensus markers. Consequently, hASPCs tend to be studied in form of total SVF Lin–, which blurs their cellular and functional characterization. To address this, we performed a comprehensive exploration of human SC, PR, OM, and MC AT SVF Lin– population structure and function. Our bulk analyses revealed extensive molecular and phenotypic variation among these depots (Fig. 2.1). On a global level, we confirmed earlier observations that only SVF-adherent cells from extraperitoneal ATs (SC and PR) displayed high adipogenic potential ex vivo, while their intraperitoneal counterparts (OM and MC) were refractory to adipogenesis. This is also reflected by the fact that SC and PR SVF-adherent cells featured a highly adipogenic transcriptomic signature compared to OM and MC ones (Supp. Fig. 2.1.4 H, J), which in contrast featured a more inflammatory and epithelial/mesothelial gene expression profile (OM) (Alvehus et al. 2010), or a protein trafficking (heat shock protein) signature (MC) (Fig. 2.1I). However, despite being highly adipogenic, we also found important molecular differences among extraperitoneal ATs, revealing that, contrary to SC, the gene expression profile of PR SVF-adherent cells was enriched for terms associated with the oxidative respiratory chain, thermogenic response, and mitochondrial activity (Fig. 2.1I). This suggests that PR hASPCs may be prone to being, potentially reflecting an influence of the nearby adrenal gland.

To better explore potential cellular mechanisms underlying the distinct adipogenic properties of the four analyzed depots, we resolved SVF Lin– heterogeneity by performing scRNA-seq on about 34'000 cells (an average of 8'500 cells per depot) and comparing the resulting data with publicly available datasets from both human and mouse ATs. These analyses allowed us to identify at least six hASPC populations that are shared across the Ats (Fig. 2.2B), including four relatively small ones, such as HHIP+ or IFIT+ hASPCs, as well as two main ones: i) the hASCs, which mapped to the mouse Dpp4+ population and the human DPP4+ cells, and ii) the hPreAs, which mapped to the mouse Icam1+/Aoc3+ population and human ICAM1+ clusters. Subsequent population sorting thereby revealed that the ASC pool is proportionally the smallest in OM AT (Fig. 2.3D), supporting the hypothesis that SC and PR ATs have a greater capacity to expand through hyperplasia compared to OM AT. A third cluster that was ubiquitous in all analyzed human depots is the VSMP cluster which highly expresses AOC3 (VAP1) (Fig. 2.2B, Supp. Fig. 2.3.1A). Although Aoc3 has mainly been described as being expressed by murine PreAs, murine VSMPs do exist and also highly express Aoc3 (Supp. Fig. 2.2.3, Supp. Fig. 2.3.1B). As hPreAs also exhibit basal AOC3 expression, we cannot completely rule out that VAP1 also enriches for a fraction of human AOC3-expressing PreAs. In our study, VAP1+ cells were the most adipogenic, but at the transcriptomic level, AOC3-high cells also expressed muscle-related markers (Fig. 2.2C), which seems contradictory. However, beige/brown AT progenitors have been described to upregulate muscle-related markers to become thermogenic. Thus, we cannot exclude that VSMP and/or
VAP1-enriched PreAs are in fact beige progenitors, although the fact that VAP1+ cell abundance was lower in SC compared to OM and PR ATs would in this regard be counterintuitive since SC is the preferential site for beige adipogenesis. Nevertheless, it seems worthy to elucidate the function of these VSMP cells given their greater abundance in high versus normal weight individuals across all analyzed adipose depots (Fig. 2.3G). This may reflect an attempt to either induce a thermogenic response to balance excessive energy take or to create new vasculature to support (excessive) adipose tissue expansion (or both). Overall, when comparing the human and mouse ASPC landscapes, there are clear similarities but also differences. For example, while F3+ ASPCs form a clearly distinct cluster in mouse, they appear to be rarer in humans, as they only emerged after aggregating all of our datasets (Fig. 2.2A-B). Moreover, while F3 is a specific marker for anti-adipogenic ASPC populations in mice, it is much less specific in humans, where HHIP appears to be a more specific marker for this cell population (Supp. Fig. 2.2.5A).

Next to AT-universal cell populations, we also found some that are exclusive to one adipose depot. A striking example is the mesothelial cells that we almost exclusively detected in OM AT (Fig. 2.2D). While the presence of such cells within the OM SVF fraction is well-established, our functional characterization revealed that the mesothelial cells, even if variable across replicates, can in fact mildly inhibit the differentiation of OM hASPCs (Fig. 2.5E-F), suggesting that the mesothelium surrounding the OM AT could have a regulatory impact on its plasticity. But more strikingly, we identified a mesothelial-like subpopulation that is also specific to the OM and that exhibits a robust anti-adipogenic capacity. We found that the latter OM SVF/Lin−/TM4SF1+/MSLN− cells highly secrete IGFBP2 (Fig. 2.5C) and can as such negatively regulate the adipogenic capacity of both SC and OM hASPCs (Fig. 2.5I-J), consistent with IGFBP2’s previously reported anti-adipogenic properties. Our findings thereby indicate that these cells’ anti-adipogenic properties are mediated through an IGF-independent mechanism, most likely through the activation of integrin receptor signaling in a paracrine fashion. Stromal populations that negatively regulate the adipogenic capacity of ASPCs have recently been discovered in mouse ATs by our and other labs. The discovery of the OM SVF/Lin−/TM4SF1+/MSLN− cells now suggests that also in humans, AT (here, OM) plasticity may be orchestrated by distinct regulatory features including not only (more systemic) endocrine signals but also specialized niche cells that are not necessarily mesenchymal in nature.

The identification of a TM4SF1+/MSLN−/IGFBP2-secreting cell population also provides new insights into the mechanisms underlying the well-established limited adipogenic capacity of OM hASPCs, even though IGFBP2 signaling could only partially explain the reduced capacity of OM hASPCs compared to other depots (Fig. 2.3E-F). This indicates that OM hASPCs still feature cell-intrinsic and transcriptomically independent mechanisms that render them refractory to differentiation and more specialized to mediating inflammation and vascular remodeling (Supp. Fig. 2.3.3C). Nevertheless, the identification of these IGFBP2-secreting cells raises questions as to their cellular origin and physiological relevance. We found that IGFBP2+ cells co-expressed
mesenchymal and mesothelial markers and showed enrichment of mesothelial to mesenchymal transition (MMT) markers (Fig. 2.4H-I, Supp. Fig. 2.4.5C). Moreover, when sorted as TM4SF1+/MSLN– cells, they exhibited a cobblestone-mesothelial morphology while, upon expansion, a spindle-mesenchymal one (Fig. 2.4L), suggesting that these cells are likely able to undergo MMT, a still poorly characterized process that has been described to also be driven by IGFBP2 itself\(^{210-213}\). While this cellular process is known, it has mainly been described in development, wound healing and cancer. Our results suggest however that MMT can also occur in adulthood at steady-like state. Interestingly, by projecting our annotation onto the recently published single-cell atlas of human AT\(^{145}\), we found not only that IGFBP2+ cells can be detected in the OM adipose depots of both lean and obese donors, which we confirmed through flow cytometry (Supp. Fig. 2.4.6), we also observed a highly positive correlation between inferred IGFBP2+ cell abundance and BMI (Supp. Fig. 2.4.4F). The latter observation appears to contrast with results from previous studies though, which found in fact an anti-correlation between BMI\(^{214-216}\), onset of metabolic syndrome\(^{217}\) including type 2 diabetes and NAFLD\(^{218}\) on the one hand and circulating IGFBP2 serum levels on the other. Since IGFBP2 is also secreted by other organs such as the liver\(^{187,218}\), additional research will therefore be required to reconcile IGFBP2’s role in seemingly controlling local OM adipose tissue plasticity versus acting as a systemic metabolic regulator.

While an important proportion of human visceral fat is contained in the OM, this depot is rather minimal in mouse\(^{219}\). It may therefore prove difficult to find an equivalent population in mouse. However, a very recent study by Zhang et al.\(^{91}\) of mouse epididymal AT did identify “mesothelial-like cells” that shared markers with both mesothelial and mesenchymal cells and that were also defined by high Igfbp2 expression. This suggests that OM IGFBP2+ cells may be cellularly and functionally conserved between mouse and human, which in turn may open new experimental avenues to study their relevance in mediating OM AT plasticity in distinct metabolic contexts. Longer-term, such studies may then lead to new therapeutic strategies to render OM hASPCs more adipogenic and less inflammatory, which could be a valuable novel approach to treat metabolic disorders linked to obesity, especially given the protective and anti-diabetic effect that has been attributed to IGFBP2\(^{216}\).
2.5 Materials and methods

Bioethics

All materials used in this study have been obtained from AT donors from two independent cohorts: the Co-hort of Obese Patients of Lausanne with ethically approved license by the commission of the Vaud Canton (CER-VD Project PB_2018-00119) and a control healthy cohort from renal transplantation donors with ethi-cally approved license by the commission of the Vaud Canton (CER-VD 2020-02021). The coded samples were collected undersigned informed consent conforming to the guidelines of the 2000 Helsinki declaration. Table 2.3 illustrates cohorts demographics.

Human ASPCs isolation and culture

2-3 cm³ biopsies from SC, OM, PR and MC ATs were washed in PBS to remove excess blood, weighted and finely minced using scissors. Minced adipose tissue was incubated with 0.28 U/ml of liberase TM (Roche #05401119001) in DPBS with calcium and magnesium (Gibco #14040091) for 60 min at 37 °C under agitation. Vigorous shaking was performed after 45 min of incubation to increase the yield of recovered SVF cells. The digested tissue was mixed with an equal volume of 1% human albumin (CSL Behring) in DPBS −/− (Gibco #14190094) to stop the lysis. Following a 5-min centrifugation at 400 g at room temperature, floating lipids and mature adipocytes were discarded by aspiration and the resuspended SVF pellet was sequentially filtered through 100-μm and 40-μm cell strainers to ensure a single cell preparation. To lyse red blood cells, pelleted SVF was resuspended in VersaLyse solution (Beckman Coulter #A09777) according to the manufacturer’s recommendations and washed once with 1% albumin solution. Obtained red blood cell-free SVF suspension was then either plated for experiments, expanded and cryoprotected or stained for sorting (see below). The SVF used for expansion or experiments was plated at a density of at least 100’000 cells per square centimeter in high glucose MEMalpha GlutaMax medium (Gibco #32561037) supplemented with 5% human platelet lysate (Sigma #SCM152) and 50 μg/ml Primocin (InvivoGen #ant-pm-2). For culturing human ASPCs, TrypLE Select reagent (Gibco #12563011) was used to collect the cells from the cell culture plates.

Bulk RNA barcoding and sequencing (BRB-seq)

All cells for BRB-seq were seeded in parallel in six 24-well plates. Cells from three wells were harvested undifferentiated (t0 time point) upon cell expansion in the 24-well plate. Cells from the three remaining wells were expanded until confluence and harvested in TRIzol (Sigma, #T3934) after 14 days of adipogenic differentation (t14 time point). RNA was extracted from all samples in parallel using the Direct-ZOL 96 well plate format (Zymo, #R2054), and BRB-seq libraries were prepared as previously described and further detailed by the Mercurius Protocol (Alithea Genomics). In brief, 7-200 ng of total RNA from each sample was reverse transcribed in a 96-well plate using SuperScriptTM II Reverse Transcriptase (Lifetech 18064014) with
individual barcoded oligo-dT primers, featuring a 12-nt-long sample barcode (IDT). Double-stranded cDNA was generated by second-strand synthesis via the nick translation method using a mix containing 2 μl of RNase H (NEB, #M0297S), 1 μl of E. coli DNA ligase (NEB, #M0205 L), 5 μl of E. coli DNA Polymerase (NEB, #M0209 L), 1 μl of dNTP (10 mM), 10 μl of 5x Second Strand Buffer (100 mM Tris, pH 6.9, (AppliChem, #A3452)); 25 mM MgCl₂ (Sigma, #M2670); 450 mM KCl (AppliChem, #A2939); 0.8 mM β-NAD (Sigma, N1511); 60 mM (NH₄)₂SO₄ (Fisher Scientific Acros, #AC20587); and 11 μl of water was added to 20 μl of ExoI-treated first-strand reaction on ice. The reaction was incubated at 16 °C for 2.5 h. Full-length double-stranded cDNA was purified with 30 μl (0.6x) of AMPure XP magnetic beads (Beckman Coulter, #A63881) and eluted in 20 μl of water.

The Illumina-compatible libraries were prepared by tagmentation of 10-40 ng of full-length double-stranded cDNA with 1 μl of in-house produced Tn5 enzyme (11 μM). After tagmentation, the libraries were purified with DNA Clean and Concentrator kit (Zymo Research #D4014) eluted in 20 μl of water and PCR amplified using 25 μl NEB Next High-Fidelity 2x PCR Master Mix (NEB, #M0541 L), 2.5 μl of each i5 and i7 Illumina index adapter (IDT) using the following program: incubation 72 °C—3 min, denaturation 98 °C—30 s; 15 cycles: 98 °C—10 s, 63 °C—30 s, 72 °C—30 s; final elongation at 72 °C—5 min. The libraries were purified twice with AMPure beads (Beckman Coulter, #A63881) at a 0.6x ratio to remove the fragments < 300 nt. The resulting libraries were profiled using a High Sensitivity NGS Fragment Analysis Kit (Advanced Analytical, #DNF-474) and measured using a Qubit dsDNA HS Assay Kit (Invitrogen, #Q32851) prior to pooling and sequencing using the Illumina NextSeq 500 platform using a custom primer and the High Output v2 kit (75 cycles) (Illumina, #FC-404-2005). The library loading concentration was 2.4 pM, and the sequencing configuration was as follows: R1 21c / index i7 8c / index i5 8 c/ R2 55c.

In parallel, the same cells were seeded in four independent 96well plates and imaged after 14 days of differentiation to quantify their adipogenic potential (see “In vitro adipogenic differentiation of hASPCs”).

Analysis of BRB-seq data

Preprocessing
After sequencing and standard Illumina library demultiplexing, the .fastq files were aligned to the human reference genome GRCh38 using STAR (Version 2.7.3a), excluding multiple mapped reads. Resulting BAM files were sample-demultiplexed using BRB-seqTools v.1.4 (https://github.com/DeplanckeLab/BRB-seqTools) and the “gene expression x samples” read, and UMI count matrices were generated using HTSeq v0.12.4.

General methods
Samples with a too low number of reads or UMIs were filtered out. Genes with a count per million greater than 1 in at least 3 samples were retained. Raw counts were then normalized as log counts per million with a pseudo count of 1, using the function cpm from EdgeR²¹ version 3.30.3. If the samples were from different
batches, the raw counts were first normalized using quantile normalization as implemented in voom from the package limma\textsuperscript{222} version 3.44.3 and then corrected for batch effects using combat from sva version 3.36.0. PCAs were computed using prcomp with the parameters center and scale set to TRUE. Differential expression analyses were performed using DESeq2\textsuperscript{223} version 1.28.1 and adding batch as a cofactor when necessary.

**Scores:**
Scores were calculated as the sum of the integrated gene expression scaled between 0 and 1 per gene of the mentioned gene lists. For Supp. Fig. 2.2.3A the top differentially expressed genes of murine fat depots populations identified based on the integration of 3 scRNA-seq datasets and published in Ferrero et al.\textsuperscript{146} were used to compute the cell type scores.

**Gene expression heatmaps:**
Heatmaps display row-normalized expression and were generated using pheatmap version 1.0.12. The columns and rows were clustered using the method “ward.D2” of hclust of the package stats.

**Gene set enrichment analysis:**
Gene set enrichment analysis was performed using the package clusterprofiler\textsuperscript{224} version 3.16.1.

**scRNA-seq of SVF Lin– cells**
SVF Lin– cells from different depots and donors were enriched with either FACS or MACS (Table 2.2) and resuspended in 1% human albumin in DPBS solution prior to be loaded into the Chromium Single Cell Gene Expression Solution (10x Genomics), following the manufacturer’s recommendations targeting a recovery of 4000 to 5000 cells per run. scRNA-seq libraries were obtained following the 10x Genomics recommended protocol, using the reagents included in the Chromium Single Cell 3’ v3 Reagent Kit. Libraries were sequenced on the NextSeq 500 v2 (Illumina) instrument using 150 cycles (18 bp barcode + UMI, and 132-bp transcript 3’ end), obtaining ~5 × 108 raw reads.

**Analysis of scRNA-seq data**

**Analysis of the datasets individually**
Raw fastqs were processed using the default CellRanger pipeline (v 2.1.0, 10X Genomics, Pleasanton, CA). The same transcriptome version was used to align all the datasets (GRCh38.92). All the data were then loaded on R (R version 3.6.1). Cells were filtered for the number of Unique Molecular Identifiers (UMIs) and genes using isOutlier from the package scater, which determines which values in a numeric vector are outliers based on the median absolute deviation (MAD) (nmads set between 3 and 4), and filters for too high a percentage
of UMIs mapping to mitochondrial RNA (~10%) or ribosomal RNA (~20%) or too low a percentage of UMIs mapping to protein-coding genes (~80%).

The datasets were first analyzed one by one using the Seurat pipeline\textsuperscript{103}. After cell filtering, only genes expressed in at least 3 cells were kept. The data were scaled for the number of UMIs and features using the function ScaleData and the remaining default parameters. The first 50 principal components of the PCA were computed using RunPCA, and then evaluated for significance using the JackStraw function of Seurat. Only the first PCs successively having a p-value < 0.05 among the top 50 PCs were selected for downstream analysis. Clustering was performed using FindNeighbors. The robustness of the clustering was assessed using clustree displaying the relationship between the clusters with increasing resolution. Differential expression analysis was computed using the FindAllMarkers function of Seurat for the selected clustering. Only genes detected as differentially expressed (log\textsubscript{2}FC > log\textsubscript{2}(1.2), p.adj < 0.05) for both the Likelihood-ratio test (test.use = “bimod”) and Wilcoxon Rank Sum test (test.use = “wilcox”) were selected.

Each sample was processed and sequenced individually, with the exception of the samples PR - D30 and PR - D61. The isolated cells of these two samples and donors were mixed. The cells were identified as belonging to each donor post-processing based on two criteria: the results of the clustering of the dataset, which clearly separated the cells from the two individuals, and the expression of \textit{XIST} as the two donors were of the opposite sex. Cells ambiguously assigned to a donor (i.e, having a positive expression of \textit{XIST} while clustering with the cells of the donor patient or the opposite) were filtered out.

Comparison of top markers of individual datasets
For each pair of subpopulations and dataset, the percentage of shared markers between their top 100 differentially expressed genes with the highest FC were calculated and displayed on Supp. Fig. 2.2.1D-F.

Scmap
The \textit{Scmap} package\textsuperscript{225} was used to project the cells of a dataset X onto the identified subpopulations of a dataset Y. Each pair of dataset X, Y and its inverse Y, X were computed. More precisely, the datasets were normalized using the “Single-cell Analysis Toolkit for Gene Expression Data in R” (\textit{scater} package). The data were log normalized using the logNormCounts functions using the size factor estimated with computeSumFactors. The 1000 most informative features of each dataset were selected using the selectFeatures function of \textit{scmap}, which is based on a modified version of the \textit{M3Drop} method. The centroids of each cluster for each dataset were calculated with the function indexCluster, and finally, the datasets were projected onto one another using the function scmapCluster.

Data integration
The datasets from each individual patient and depot, at the exception of GB-D07 (due to a very low number of captured ASPCs), were integrated following the standard workflow of Seurat pipeline. The datasets were
normalized in log scale with a scale factor of 10000. The top 2000 highly variable genes were selected using the FindVariableFeatures function with the parameter selection.methods set to “vst”. The anchors were identified using FindIntegrationAnchors. The top 2000 variable features identified by SelectIntegrationFeatures and the first 60 principal components of the PCA were used as input to perform canonical correlation analysis. The integrated data computed by IntegrateData were then used for dimensionality reduction and clustering based on the first 60 principal components of the PCA. Clustering was computed for different clustering resolutions. The final clustering result was based on the clustering results at different resolutions depending on the robustness of the clusters and the specificity of their differentially expressed markers. Top differentially expressed genes were identified using the FindConservedMarkers function of Seurat after setting the default assay to RNA, the adjusted p-values were combined using Tippett’s method as implemented by the function minimump from metap R package (meta.method = metap::minimump)²²⁶. Only groups of cells with at least 10 cells were tested (min.cells.group = 10). Specifically, for the IGFBP2+ cell cluster, as we found only a few cells per batch and we focused on that cell type in part of the manuscript, DEGs were further computed using EdgeR and correcting for batch. More precisely, genes not expressed in at least 2% of the cells were filtered out using the function filterByExpr. After converting the count matrix into a DGEList using DGEList, the data were normalized with calcNormFactors. The design matrix was defined following the formula ~0 + clust + batch, where clust corresponds to the cluster of every cell and batch to its dataset (as individually shown on Supp. Fig. 2.2.1A). The dispersion was estimated using estimateDisp. The quasi-likelihood negative binomial generalized log-linear model was fitted using glmQLFit, followed by the quasi-likelihood F-test glmQLFtest contrasting the IGFBP2+ cluster versus the other clusters (pondered by the number of clusters).

Identification of depot-specific markers for ASCs and PreAs

DEG analysis was performed on the integrated data, by selecting the cells of the population of interest (ASCs or PreAs) and contrasting between all possible pairs of depots using the function FindMarkers of Seurat. This is possible as we have 3 replicates for SC, OM, PR, and 2 for MC, however, for the latter, those were coming from two biological samples from the same donor. A set of markers was considered depot-specific when significantly differentially expressed in a depot versus any other depot. A gene was defined as differentially expressed when its average log Fold Change (defined as the average of the log Fold Change in each replicate) was positive and an adjusted p-value smaller than 0.05.

Comparison with murine ASPCs

a. Murine data integration

The integration of five datasets of adult mouse SC and OM ATs provided by Schwalie et al.¹⁰², Burl et al.¹⁰⁰, Hepler et al.¹⁴⁷ and Merrick et al.⁹⁶ was performed as described in Ferrero et al.¹⁴⁶. The clustering originally published in Ferrero et al.¹⁴⁶, focusing on ASPCs, merged the cells close to endothelial cells into one main cluster. The clustering was here revised to include vascular smooth muscle progenitor cells. For consistency
with the human data, the top markers of the subpopulation were computed as defined above. The top markers were ordered by the average of the log$_2$ Fold Change of each dataset.

b. Score

Scores of the mouse ASPC subpopulations, mesothelial cells, and vascular smooth muscle progenitor cells were based on their human orthologs and calculated as the sum of the gene expression scaled between 0 and 1 per gene of the top markers (average log$_2$ Fold Change across batches > 0 and adjusted p-value < 0.05) of each murine ASPC subpopulation (ASCs, PreAs, Aregs, Ifit+, and Cilp+ ASCs), mesothelial cells and vascular smooth muscle progenitor cells. The scores were then scaled by the number of genes on each list.

Comparison with the dataset from Emont et al.$^{145}$

The whole human single-nucleus/cell dataset (here reported as “scRNA-seq”) provided by Emont et al.$^{145}$ was downloaded on the single cell portal (study no. SCP1376, All cells). The dataset was then subsetted for the cells defined as ASPC or mesothelium by the authors (as defined in the metadata “cell_type2”), and the PCA was recomputed as well as clustering, tSNE and UMAP with the first 50 PCs as input. First, an IGFBP2 expression score was computed using the AddModuleScore function. The dataset containing only ASPCs, and mesothelial cells was then split by samples, and the symbol gene IDs were converted to Ensembl ID using the GRCh38 release 92 from the Ensembl gene annotation as reference. The few genes with no corresponding Ensembl IDs were filtered out, and, in the rare case of two corresponding Ensembl IDs, only one was kept. Each sample was log normalized with the default normalization of the Seurat package and then scaled for the features selected using SelectIntegrationFeatures with each of the samples of Emont et al.$^{145}$ and our generated single-cell SC and OM datasets as input. The first 50 PCs were computed based on the scaled data. Clustering was performed following the default Seurat clustering pipeline for resolutions spanning from 0.1 to 3. Each sample of the Emont et al.$^{145}$ dataset was then projected on our integration (see Analysis of single-cell RNA-seq, Data integration), using the FindTransferAnchors and TransferData functions of the Seurat package with the default parameters.

Trajectory analysis

Trajectory analysis was performed on the integrated normalized data subsetting for Epiploic samples. Potential doublets were excluded from the analysis using DoubletFinder$^{227}$ on each epiploic scRNA-seq dataset individually. Cells labeled as ASCs, PreAs, IGFBP2+ cells, Mesothelial cells, and VSMPs were selected. The first 50 PCs were computed using the pca function of scanpy$^{228}$ and the neighborhood graph was computed with the default parameters (pp.neighbors). The connectivity between our defined cell classifications was computed using the paga function$^{179}$, and low-connectivity edges were thresholded at 0.03. We computed the ForceAtlas2 (FA2) graph$^{229}$ using PAGA-initialization (draw_graph). The Dynverse package$^{230}$ was used to compute the most variable genes along the branch connecting PreAs and Mesothelial cells through IGFBP2+ cells (calculate_branch_feature_importance).
**FACS sorting of human SVF subpopulations**

SVF cells were resuspended in 1% albumin solution to the concentration of $10^5$ cells/ml, and the staining antibody panels (Table 2.4) were added in titration-determined quantities. The cells were incubated with the cocktail of antibodies on ice for 30 min protected from light, after which they were washed and stained with propidium iodide (Molecular Probes #P3566) for assessing viability, and subjected to FACS using a Becton Dickinson FACSAria II sorter or a MoFlo Astrios EQ, Cell Sorter - Beckman Coulter. Compensation measurements were performed for single stains using compensation beads (eBiosciences #01-2222-42).

The following gating strategy was applied while sorting the cells: first, the cells were selected based on their size and granularity or complexity (side and forward scatter), and then any event that could represent more than one cell was eliminated. Next, the live cells were selected based on propidium iodide negativity, and from those, the Lin−(CD31−/CD45−) population was selected. At first, all SC, OM, and PR cells were stained with the OM-specific panel, including mesothelial markers, but since SC and PR SVF cells were consistently negative for the TM4SF1 and MSLN markers over three consecutive experiments, SC and PR cells were only stained with the SC and PR panels, respectively (Table 2.4). For the SC samples, from the Lin− fraction of cells, Lin−/CD26+, Lin−/VAP1+, Lin−/DN, and Lin−/HHIP+ cells were defined against unstained controls and FMO controls. For the PR samples, from the Lin− fraction of cells, Lin−/CD26+, Lin−/VAP1+, and Lin−/DN cells were defined against unstained controls and FMO controls. For the OM samples, OM-specific subpopulations were first isolated from the Lin− gate as Lin−/TM4SF1+/MSLN− and Lin−/TM4SF1+/MSLN+ populations. From the remaining Lin−/TM4SF1− gate, we then isolated Lin−/TM4SF1−/CD26+, Lin−/TM4SF1−/VAP1+, and Lin−/TM4SF1−/DN cells. Acquired FCS files were analyzed using FlowJo software to infer population abundances that were plotted using GraphPad Prism.

**In vitro adipogenic differentiation and chemical treatments of hASPCs**

Cells were seeded for adipogenic differentiation at high density (65k cells/cm²) in 3-5 replicate wells of a 96-well black plate (Corning #353219). After 48h or when cells where confluent for at least 24h, cells were treated with induction cocktail (high glucose DMEM (#61965), 10% FBS, 50 μg/ml Primocin, 0.5 mM IBMX (Sigma #I5879), 1 μM dexamethasone (Sigma #D2915), 1.7 μM insulin (Sigma #I9278), 0.2 mM indomethacin (Sigma #I7378) for 7 days, followed by a maintenance cocktail (high glucose DMEM, 10% FBS, 50 μg/ml Primocin, 1.7 μM insulin) for another 7 days. No medium refreshment was performed between these two timepoints. For the chemical treatments, the above-mentioned differentiation and maintenance cocktails were supplemented with the recombinant IGFBP2 protein at 1nM (R&D, #674-B2-025), recombinant IGF-I protein at 10nM (Sigma, #I3769), recombinant IGF-II protein at 10nM (R&D, #292-G2-050), Blocking anti-human IGFBP2 antibody (scavanging) 1 mg/ml (R&D, #AF674) and Echistatin 100 nM (R&D, #3202). Chemicals were added to both induction and maintenance cocktails except for Echistatin which was added to the
induction cocktail only and withdrawn 48h after induction since inhibiting the integrin receptor resulted in cell detachment when Echistatin was kept in culture for longer periods than 48h. In the Echistatin mixed with IGFBP2 condition, only IGFBP2 was kept after 48h. IGFBP2, IGF-I and IGF-II were first titrated at the concentrations shown in Supp. Fig. 2.5.2.

**Cell proliferation assay**

Sorted cells were split into four and seeded in 4 different wells of a 12well plate and allowed to attach and start to proliferate for 7 to 10 days. One well of each cell population was trypsinized after this period. Cells were resuspended in 1 ml of medium, counted twice using a hematocytometer, and the mean count was used as the baseline number of cells from which cell increase was calculated. The same counting was performed on the remaining wells every two days. The expansion medium was refreshed every two days.

**Mixing and transwell experiments**

For the mixing experiments, unexpanded Lin– SVF cells were isolated with MACS using Miltenyi LD columns (Miltenyi, #130-042-901) on manual mono-MACS separators after staining with magnetic anti-human CD45 and CD31 microbeads (Miltenyi, #130-045-801 and #130-091-935) according to the manufacturer’s protocol. MACS-isolated Lin- cells from SC, OM, and PR samples were counted in duplicates and mixed at high density (65k cells/cm²) in 11 ratios from 0 to 100%. After 24h, the cells were induced to differentiate following the adipogenic differentiation protocol. For the transwell experiments, we used 96well plate format transwell inserts with 0.4 mm (Corning #CLS3391) pores to allow protein and small molecule diffusion through the membrane, but not cell migration. 96well transwell-receiving plates (Corning #3382) were first coated with type I collagen (Corning #354249) 1:500 in DPBS before use to facilitate cell adhesion. Sorted donor OM subpopulations and expanded receiver SC and PR SVF-adherent cells were plated and expanded separately onto the top transwell insert and the bottom receiving plate, respectively. When confluent, the transwell insert was put in contact with the receiver plate, and all cells were induced to differentiate following the listed differentiation protocol.

**Enzyme-linked immunosorbent assay (ELISA)**

For the supernatant measure, cells were expanded for two passages and seeded into a 6well plate. Once confluent, the expansion medium was aspirated, and wells were washed twice with PBS to ensure residual serum, dead cell and protein removal. 2ml of OPTI-Pro serum-free medium (Thermo, #12309050) was added to each well and incubated with the cells at 37°C for 48h. After incubation, SFM medium was harvested, spun for 10 min at 4°C max speed to clear potential cell debris. Cleared supernatant was aliquoted and stored at -80°C until further usage. For the whole AT IGFBP2 secretion assays, three times 200-400 mg of OM AT were put in 500ml of DPBS (Gibco #14190169) and incubated at 37°C for 24, 48 and 72 hours. After incubation,
DPBS was harvested, spun for 10 min at 4°C max speed to clear potential cell debris and stored at -80°C until further usage. The Anti-human IGFBP2 ELISA kit (Sigma, #RAB0233-1KT) was used to quantify IGFBP2 protein in the supernatants according to the manufacturer’s recommendations. Before loading samples on the ELISA membranes, the total protein concentration was quantified using the Qubit™ Protein Broad Range assay kit (Thermo, #A50669) and 300 ng of total protein was added per reaction. Incubation of samples with primary antibodies was performed O/N at 4°C. At the end of the assay, absorbance was read at 450 nm using a SPARK® Microplate reader.

**Immunohistochemistry**

Human AT biopsies were washed twice in PBS to remove excess blood and divided in 50 to 100 mg for fixation in 4% PFA (paraformaldehyde, electron microscopy grade (VWR #100504-858)) for 2 hours at 4°C with gentle shaking. Next, the tissue was washed with PBS and incubated with 30% sucrose O/N at 4°C with gentle shaking. Cryoblocks were prepared using Cryomatrix (Thermo Fisher Scientific #6769006), and 25-μm sections were generated using a Leica CM3050S cryostat at -30°C. The tissue was air-dried for 30 min at -20°C in the cryostat itself, then 1 h at RT. Slides were additionally fixed 10 min in 4% PFA at RT, washed two times 5 minutes with PBS, permeabilized at RT with 0.25% TritonX100 (Sigma #T9284) for 10 minutes, washed twice with PBS again and antigen blocking was performed at RT for 30 min with 1% BSA in PBS. Primary antibodies (anti-TM4SF1, anti-MSLN, anti-PLIN1) in 1% BSA were applied O/N at 4°C with gentle shaking following the titrations indicated in Table 2.4. The following day, after two PBS washes, and quick 1% BSA dip, the secondary antibody (anti-rabbit AF-647) in 1% BSA was applied for 40 min at RT following the titrations in Table 2.4. Nuclei were stained with 1mg/ml DAPI (Sigma #D9564) for 10 min and washed twice in PBS prior to mounting with Fluoromount G (Southern Biotech #0100-01). The slides were then imaged with a Leica SP8 Inverted confocal microscope (objectives: HC PL Fluotar 10x/0.30 air, HC PL APO 20x/0.75 air, HC PL APO 40x/1.25 glyc, HC PL APO 63x/1.40 oil). The results presented in Fig. 2.4K and Supp. Fig. 2.4.7 were replicated in at least three independent experiments. We note that we also verified that the signal we detected is not the result of autofluorescence of the AT or from unspecific binding of secondary antibodies (Supp. Fig. 2.4.7).

**Imaging and quantification of in vitro adipogenesis**

On the 14th day of differentiation, cells were either fixed with 4% PFA (EMS, #15710) and stained at a later timepoint or live-stained with fluorescence dyes: Bodipy 10 mg/ml (boron-dipyrromethene, Invitrogen #D3922) for lipids and Hoechst 1 mg/ml (Sigma, #B2883) for nuclei. Cells were incubated with the dyes in PBS, for 30 min in the dark, washed twice with PBS, and imaged. If the imaging was performed on live cells, we used FluoroBrite DMEM (Gibco # A1896701) supplemented with 10% FBS as acquisition medium. Given substantial variation in the extent of lipid accumulation by the tested cell fractions (within the same well but also across technical replicates), the imaging was optimized to cover the largest surface possible of the 96
well. Moreover, a z-stack acquisition in a spinning-disc mode and Z-projection were performed in order to capture the extent of in vitro adipogenesis with the highest possible accuracy. Specifically, the automated platform Operetta (Perkin Elmer) was used for imaging. First, 3–6 z-stacks were acquired for every field of view in a confocal mode of the microscope in order to produce high-quality images for downstream z-projection and accurate thresholding. Next, 25 images per well were acquired using a Plan Neofluar 10× Air, NA 0.35 objective for the transwell-receiving plates or 20x air objective NA 0.8 for normal 96w plates (Falcon, #353219), with no overlap for further tiling and with the aim of covering the majority of the well for an accurate representation of lipid accumulation (see Methods in \cite{148,231}). The lasers were set in time exposure and power to assure that in both the Hoechst and the Bodipy channels, the pixel intensity was between 500 and 4000, and in all cases at least two times higher than the surrounding background. The images, supported by Harmony software, were exported as TIFF files. They were subsequently tiled, and Z-projected with the maximum intensity method. To accurately estimate and represent differences in adipocyte differentiation, a quantification algorithm for image treatment was developed in collaboration with the EPFL BIOP imaging facility. In brief, image analysis was performed in ImageJ/Fiji, lipid droplets (yellow) and nuclei (blue) images were filtered using a Gaussian blur (sigma equal to 2 and 3, respectively) before automatic thresholding. The automatic thresholding algorithm selections were chosen based on visual inspection of output images. The area corresponding to the thresholded lipid signal was then divided by the area corresponding to the thresholded nuclei area and used to calculate the Adiposcore (totalLipidArea/totalNucleiArea). In the figures, representative blown-up cropped images of each sample are shown. To reduce technical variation across the biological replicates (different donors), adiposcores were normalized to the average adiposcore of the indicated control when we compared conditions within highly differentiating lines like SC and PR. Adiposcores were compared without normalization when we wanted to directly compare adiposcores across depots (i.e., Fig. 2.1C) or among poorly-differentiating samples like OM when the absolute values of adiposcores were < 0.01 (Fig. 2.4D, Fig. 2.5J, N and P, and Supp. Fig. 2.1.2B, and Supp. Fig. 2.1.3).

siRNA-mediated knockdown

To achieve knockdown of IGFBP2, direct transfection was performed on OM SVF Lin−/TM4SF1+/MSLN− cells using the IGFBP2 IDT, TriFECTA DsiRNAs kit using 3 pooled siRNAs: hs.Ri.IGFBP2.13.1, hs.Ri.IGFBP2.13.2, hs.Ri.IGFBP2.13.3. In brief, after sorting, cells were expanded for one or two rounds, then harvested and plated at mid-low density (45k cells/cm2) and allowed to adhere. The following day, transfection mix was prepared as Opti-MEM medium (Invitrogen #31985062), 1.5% Lipofectamine RNAiMAX (Invitrogen #13778150) and 20 nM of the pooled siRNAs. In the transfection mix, lipofectamine-siRNA transfection particles were allowed to form for 15 min at RT with gentle shaking. After incubation, the transfection mix was diluted 10 times (to a final concentration of siRNA of 2 nM) in MEMaAlpha GlutaMax medium (Gibco #32561037) supplemented with 2.5% human platelet lysate (Sigma #SCM152), w/o antibiotics and
exchanged to the plated cell medium. After 48h, medium was changed to differentiation medium (for the transwell assay), with serum free medium (for ELISA validation) or directly taken in TRIzol (for qPCR validation).

**RNA isolation and qPCR**

Expanded OM and SC SVF-adherent, OM SVF Lin—/TM4SF1—/MSLN—, OM SVF Lin—/TM4SF1+/MSLN— cells as well as cells subjected to siRNA-mediated knockdowns 48h post-transfection were collected into TRIzol (Sigma, #T3934). The direct-zol RNA kit (Zymo Research #R2062) was used to extract RNA, followed by reverse transcription using the SuperScript II VILO cDNA Synthesis Kit (Invitrogen # 11754050). Expression levels of mRNA were assessed by real-time PCR using the PowerUp SYBR Green Master Mix (Thermo Fisher Scientific #A25743). mRNA expression was normalized to the Hprt1 gene. Primer sequences used: *IGFBP2* – Fw CGAGGGCACTTGTGAGAAGCG, Rv TGTTCATGGTGCCTCCACGTC; *HPRT* – Fw CAGCCCTGGCGTGTGATTA, Rv GTGATGGCCCTCCTCCTT.

**Statistical methods**

The experiments were not randomized, and the investigators were not blinded in experiments. The paired Student’s t-test was used to determine statistical differences between two groups, with the null hypothesis being that the two groups are equal. Multiple comparisons were corrected using false discovery rate (FDR) correction. When specified, one-way ANOVA or RELM test followed by Tukey honest significant difference (HSD) post hoc correction was applied, the null hypothesis being defined so that the difference of means was zero. (Adjusted) *p*-value < 0.05, **p*-value < 0.01, ***p*-value < 0.001 were considered statistically significant. All boxplots display the mean as a dark band, the box shows the 25th and 75th percentiles, while the whiskers indicate the minimum and maximum data points in the considered dataset excluding outliers. All bar plots display the mean value and the standard deviation from the mean as error bar.
2.6 Miscellaneous

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Authors contributions

R.F., P.R. and B.D. designed the study and wrote the manuscript. R.F. conducted all experimental procedures and analyzed acquired images, flow cytometric measures, qPCRs, ELISAs and immunohistochemistry. P.R. conducted all analyses related to transcriptomics both at the single-cell and bulk levels. J.R. and M.Z. assisted with sample processing, cell culture and preparation of sequencing libraries. J.R. performed histological assays. J.P. provided assistance with flow cytometry-related procedures. D.A. and V.G. assisted with bulk transcriptomic-associated procedures and data processing. V.G. and W.S. assisted with the transcriptomic analyses. L.F., S.M., T. Z., N.P., M.S. and M.M. provided access to human samples. V.G., W.S. and C.C. provided extensive comments to the manuscript.

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Competing interest declaration

The authors declare to have no conflict of interest.
2.7 Supplementary Figures and Tables

Supplementary Figure 2.1.1 – Across depot comparisons of SVF-adherent cells at low passage.

(A) Total number of nuclei in each well (see Methods for more details) of images in Fig. 2.1B; n=14-22, 4-5 donors, 3-5 independent wells.

(B) Representative confocal images of intraperitoneal OM- and MC-derived SVF-adherent cells after 14 days of differentiation; Top: lines that form very few mature lipid droplets, Bottom: lines that form small lipid droplets that are barely distinguishable from background; Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA.

(C) Image-processing steps to define what signal from the lipid channel is considered as fully formed lipid droplets and which is not. Red squares indicate the thresholding parameters chosen for the final analysis to exclude background from the low adipogenic lines and to not underestimate the lipid area of the highly differentiating lines (PR). In the first two columns, green denotes lipids, red nuclei, and in the last two columns, red is the signal measured above the defined threshold (see Methods).
(D) Relative cell number increase over time of culture for SC, OM, and PR SVF-adherent cells; n=12, 4 donors per depot; Black stars compare SC versus OM, Red: PR versus OM, Blue: SC versus PR.

*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, One-Way ANOVA and Tukey HSD post hoc test (A), and RELM analysis and Tukey HSD post hoc test (D).
Supplementary Figure 2.1.2 – Adipogenic differentiation potential of cell lines is highly variable across individual donors.

(A) Full well images of four independent wells after image processing (25 merged Z-projected tiles taken with a 20x objective, see Materials and Methods) enabling us to quantify the adipogenic potential of each cell line that was analyzed by BRB-seq18 after 14 days of adipogenic differentiation. The cells were differentiated after 2 to 6 passages. The blue color represents the signal above threshold for the DNA stain, and the yellow color represents the signal above threshold for the lipid stain. Subcutaneous, yellow (SC): n=104, 4 independent wells, 26 cell lines, 20 donors (D); Perirenal, brown (PR): n=36, 4 independent wells, 9 cell lines, 8 donors; Omentum, purple (OM): n=88, 4 independent wells, 22 cell lines, 18 donors;
Mesocolon, blue (MC): n=16, 4 independent wells, 4 cell lines, 4 donors; Scale bar=1 mm. B2 indicates the second biological replicate of a certain line, and 1y indicates the few samples from patient 1 year post bariatric surgery.

(B) Distribution of the ratios between the total lipid area and the total DNA area measured in the images shown in A, i.e., adiposcore (see Methods for more details), ordered by increasing values across all cell lines shown in A.

(C) Total number of nuclei in each well quantified by segmenting the DNA area above threshold of images in A and plotted in the same order as in B.
Supplementary Figure 2.1.3 – The adipogenic potential of SVF-adherent lines from different depots and their correlation with metadata.

(A) Quantification of the adipogenic potential derived from images in Supp. Fig. 2.1.2A representing differentiated SVF-adherent cells after expansion (passaged 2 to 6 times). On the y axes, the log(adiposcore+1) is plotted. SC: n=104, 4 independent wells, 26 cell lines, 20 donors; PR: n=36, 4 independent wells, 9 cell lines, 8 donors; OM: n=88, 4 independent wells, 22 cell lines, 18 donors; MC: n=16, 4 independent wells, 4 cell lines, 4 donors.

(B) Total number of nuclei in each well (see Methods) of images shown in Supp. Fig. 2.1.2A.

(C) Scatter plot showing the correlation between the adiposcore of highly adipogenic lines (SC - left and PR - right) and the BMI of respective donors. The line represents the linear regression analysis.

(D) Scatter plot showing the correlation between the adiposcore of highly adipogenic lines (SC - left and PR - right) and the age of respective donors. The line represents a linear regression analysis.

(E) Barplot showing the distribution of the adiposcore of SC lines between men and women.

(F) Barplot showing the distribution of the adiposcore of PR lines between men and women.

*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, and linear regression analysis with its relative goodness of fit, and the FDR-adjusted p-values of the Pearson correlations (C, D), One-Way ANOVA and Tukey HSD post hoc test (E, F).
Supplementary Figure 2.1.4 – Human SVF-adherent cells from different depots differentially upregulate the adipogenesis response upon exposure to an adipogenic cocktail.

(A) Plot showing the number of donors and distribution of their BMI and sex included in the BRB-seq analysis across the different depots and time points.

(B) PCA based on the BRB-seq data of SVF-adherent cells from the indicated adipose depots (SC - subcutaneous, PR - perirenal, OM - omentum, MC - mesocolic) and indicated time points (t0 - undifferentiated, t14 - 14 days post-adipogenic induction).

(C) PCA, as described in B, colored by donors.
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(D) t-SNE map shown in Fig. 2.1D computed on the 10 first PCs of the PCA displayed in B, colored by donors.
(E) t-SNE map shown in Fig. 2.1D colored by the number of detected genes.
(F) t-SNE map shown in Fig. 2.1D colored by the BMI of the donors.
(G) Boxplot displaying the expression distribution of THY1, a known mesenchymal cell marker, across samples from the indicated depots at t0.
(H) Boxplots displaying the “white fat cell differentiation score” based on the scaled expression of the genes from the GO term “white fat cell differentiation” (GO:0050872) for the indicated depots and time points (see Methods).
(I) Boxplots displaying the “response to insulin score” based on the scaled expression of the genes from the GO term “response to insulin” (GO:0032868) for the indicated depots and time points (see Methods).
(J) Boxplots displaying the “mature adipocyte score” based on the scaled expression of the following markers: FABP4, PPARG, ADIPOQ, LIP, LPL, PLIN1, PLIN2, PLIN4, CEBPA, CEBPB, CIDEC, CIDEA, for the indicated depots and time points (see Methods).

*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, unpaired two-sided t-test (G-J).
Supplementary Figure 2.1.5 – SVF-adherent cells from different depots have distinct transcriptomic signatures both when undifferentiated and when differentiated.

(A) Dotplot showing enriched, representative terms that were found by GSEA performed on the differential gene expression analysis results of each indicated depot versus all others at t0 (i.e., undifferentiated state) of the transcriptomic data shown in Fig. 2.1D.

(B) Heatmap of top differentially expressed genes when comparing the indicated adipose depot versus the three others at t14.
Supplementary Figure 2.1.6 – The expression of genes related to “inflammatory response” and “vasculature and epithelium development” is enriched in OM adipose depot-derived cells compared to those from other depots.

(A) GSEA plot of selected inflammatory response GO terms (“humoral response” GO:0006959, “cellular response to lipopolysaccharides” GO:0071222, “neutrophil chemotaxis” GO:0030593), based on the differential expression analysis of SVF-adherent cells derived from OM adipose depots versus those from other depots (SC, PR, MC) at t0 (i.e., undifferentiated state) or t14 (i.e., post-adipogenic induction).

(B) GSEA plot of the GO terms “epithelial cell development” (GO:0060429) and “morphogenesis of a branching epithelium” (GO:0048754), based on the differential expression analysis of SVF-adherent cells derived from OM adipose depots versus those from other depots (SC, PR, MC) at t0 (i.e., undifferentiated state).
Supplementary Figure 2.2.1 – scRNA-seq reveals common cell populations across adipose depots and donors.

(A) t-SNE cell maps of individual scRNA-seq datasets of SVF Lin– cells isolated from four adipose depots (SC, OM, MC, and PR) and six different donors (D, as indicated in the corner of each t-SNE, see Table 2.2), visualizing the identified subpopulations of hASPCs: adipose stem cells (ASCs; green), pre-adipocytes (PreAs; red), OM-specific cells (blue), as well as mesothelial
cells (purple), vascular smooth muscle progenitor cells (VSMPs; dark orange), endothelial (light orange) and immune cells (pink). The number of cells per dataset from top to bottom and by row: 3929, 4169, 2162, 4262, 2042, 2670, 8583, 600, 509, 2650, 2550.

(B) left - t-SNE cell map of a scRNA-seq dataset of SVF Lin− cells isolated from gallbladder-associated adipose tissue from one donor, visualizing the identified populations: immune cells (pink) and hASPCs (ASCs in green and preAs in red). The high proportion of immune cells in the dataset likely reflects a technical artefact where SVF cells may have been under-stained prior to MACS separation; right - PCA of the highlighted hASPCs of the left panel visualizing the two hASPC subpopulations: adipose stem cells (ASCs; green) and pre-adipocytes (PreAs; red), a total of 54 cells.

(C) Boxplot showing the log normalized gene expression distribution of selected markers in the different hASPC subpopulations depicted in panels A and B.

(D) The percentage of shared top 100 differentially expressed genes between each subpopulation and sample shown in panels A and B; each point represents the number of shared markers between the indicated subpopulations (x-axis) of an individual dataset X and a subpopulation (coloring) in another individual dataset Y.

(E) The percentage of shared top 100 differentially expressed genes as shown in panel D but limited to the comparison of the same populations (ASCs with ASCs, PreAs with PreAs, and VSMPs with VSMPs) across depots and donors, split by the type of depot of the comparison pairs; comparisons between samples originating from the same depots are highlighted in shades of green, and comparison of samples originating from different depots are in shades from yellow to red.

(F) The percentage of shared top 100 differentially expressed genes as shown in panel D but limited to the comparison of the same populations (ASCs with ASCs, PreAs with PreAs, and VSMPs with VSMPs) across depots and donors, stratified according to whether the pairs of compared samples are originating from the same donor (green) or not (red).

(G) The percentage of cells of a subpopulation projected onto each subpopulation and sample based on scmap results (see Methods). Each point represents the percentage of a subpopulation (x-axis) of an individual dataset X projected onto a subpopulation (coloring) of another dataset Y. Projections of subpopulations (x-axis) non-existing in the reference data are highlighted as shaded circles.

(H) t-SNE cell map of the integration of all scRNA-seq datasets described in Fig. 2.2A and B colored by sample.

(I) Violin plot showing the distribution of library size (left) or the number of detected genes (right) across the different clusters shown in Fig. 2.2B.
Supplementary Figure 2.2.2 – Human ASCs and PreAs feature different signaling pathways.

(A) Barplot displaying the percentage of cells of each cluster shown in Fig. 2.2B coming from each batch.
(B) Barplot displaying the percentage of cells of each cluster shown in Fig. 2.2B coming from each depot (shades of colors indicate the different batches). The values were corrected for the number of cells per batch and the number of replicates per depot (n=3 for SC, PR, and OM, versus n=2 for MC).

(C) Dot plot showing enriched biological process GO terms based on differentially expressed genes of ASCs or PreAs of the integrated scRNA-seq data shown in Fig. 2.2A and B.

(D) Heatmap of the differentially expressed genes between the Adipose Stem Cell (ASC) population and the Pre-adipocyte (PreA) one across depots.
Supplementary Figure 2.3 – Several cell populations that were identified in the human SVF Lin– fraction across adipose depots correspond to cell populations identified in mice.
(A) Boxplot showing the distribution of the indicated murine WAT cell population scores across the detected, distinct human SVF cell populations. The scores were based on the human orthologs of the murine top markers, which were computed in the analysis presented in Ferrero et al. integrating scRNA-seq datasets of subcutaneous and visceral adipose tissue.

(B) Boxplot showing the distribution of the indicated human cell populations across the different mouse cell populations defined in Ferrero et al. The scores were based on the mouse orthologs of the human top markers of the clusters shown in Fig. 2.2B.

(C) t-SNE cell map of integrated scRNA-seq datasets from OM and SC fat depots visualizing the identified subpopulations of mouse subcutaneous cells: adipose stem cells (ASCs), pre-adipocytes (PreAs), Aregs, Ifit+ ASPCs, Cilp+ ASPCs, mesothelial, endothelial, and immune cells.

*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, unpaired two-sided t-test (A, B), only the statistics for the highest mean(s) are shown. The indicated adjusted p-value categories hold for all comparisons with the corresponding category, except if stated otherwise.
Supplementary Figure 2.2.4 – Metadata of the scRNA-seq atlas of hASPCs and mesothelial cells published by Emont et al.59.

(A) UMAP of hASPCs and human mesothelial cells from scRNA-seq data provided by Emont et al.59 colored by the depot of origin.

(B) UMAP of hASPCs and human mesothelial cells from scRNA-seq data provided by Emont et al.59 colored by the clustering published in the latter study.
Supplementary Figure 2.2.5 – A HHIP+ hASPC population shares key marker genes with the murine Adipogenesis regulators (Aregs).

(A) Dot plot displaying the average expression and percentage of expressing cells of the top 20 markers of HHIP+ hASPCs across the clusters shown in Fig. 2.2B. The orthologs of the murine top Areg markers (as defined in Zachara et al., 2022) are highlighted in bold.

(B) t-SNE cell map of the integrated data of human SVF cells shown in Fig. 2.2A and B colored by the log normalized expression of HHIP. The HHIP+ hASPC cluster is highlighted with a red circle.

(C) Representative flow cytometry-based gating of HHIP+ events on SC adipose depot-derived SVF Lin– cells. The HHIP+ staining is highly variable across donors: donor D34 harbors a more distinct population, D25 more a continuum of positive events.

(D) Flow cytometry-based quantification of Lin–/HHIP+ events in SC SVF Lin–, n=11.
Supplementary Figure 2.6 – A population of IFIT+ hASPCs features enriched viral immune response gene expression.

(A) Dot plot showing the average expression and percentage of expressing cells of the top 20 markers of IFIT+ hASPCs across the clusters shown in Fig. 2B.

(B) Dot plot showing representative GO terms that are enriched based on the differentially expressed genes of IFIT+ hASPCs.

(C) Boxplot showing the distribution of the log normalized expression of PDGFRA and PTPRC (CD45) across the cluster shown in Fig. 2B.

In order from left to right:
- ASCs
- PreAs
- IGFBP2+
- SFRP4+
- IFIT+
- HHIP+
- RBPS5+
- Meso
- Endo
- Immune
Supplementary Figure 2.7 – A hASPC population shared across depots is defined by high SFRP2 and SFRP4 expression.

(A) Dot plot displaying the average expression and percentage of expressing cells of the top markers of SFRP4+ hASPCs across the clusters shown in Fig. 2B.
(B) UMAP of hASPCs and human mesothelial cells from scRNA-seq data provided by Emont et al. colored by the prediction score of SFRP4+ cells when transferring our cell cluster annotation.
(C) Box plot showing the distribution of SFRP2 (top) and SFRP4 (bottom) expression in the different cell types from the integrated scRNA-seq data, colored by depot of origin.
(D) Box plot showing the distribution of SFRP2 (top) and SFRP4 (bottom) expression in the different hASPC subpopulations from the integrated scRNA-seq data, colored by depots of origin.
Supplementary Figure 2.8 – Mesothelial and IGFBP2+ cell markers are enriched in SVF cells from the OM adipose depot compared to other depots, both pre- and post-adipogenic induction.

(A) Volcano plot displaying differential gene expression results based on the BRB-seq\textsuperscript{150} data of expanded SVF-adherent cells from the OM adipose depot versus SVF-adherent cells from other depots (SC, PR, and MC fat depots). Top IGFBP2 markers identified using scRNA-seq datasets are highlighted in blue, while significantly differentially expressed genes (log2FC > 1, adjusted p-value < 0.01) are highlighted in darker colors.

(B) Volcano plot displaying differential gene expression results based on the BRB-seq\textsuperscript{150} data of SVF-adherent cells from the OM adipose depot versus SVF-adherent cells from other depots (SC, PR, and MC fat depots) post-adipogenic induction. Top IGFBP2+ cells markers identified using scRNA-seq datasets are highlighted in blue, significantly differentially expressed genes (log2FC > 1, adjusted p-value < 0.01) are highlighted in darker colors.

(C) Volcano plot displaying differential gene expression results based on the BRB-seq\textsuperscript{150} data of SVF-adherent cells from the OM adipose depot versus SVF-adherent cells from other depots (SC, PR, and MC). Top mesothelial markers identified using scRNA-seq datasets are highlighted in purple, while significantly differentially expressed genes (log2FC > 1, adjusted p-value < 0.01) are highlighted in darker colors.
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(D) Volcano plot displaying differential gene expression results based on the BRB-seq\textsuperscript{150} data of SVF-adherent cells from the OM adipose depot versus SVF-adherent cells from other depots (SC, PR, and MC) post-adipogenic induction. Top mesothelial markers identified using scRNA-seq datasets are highlighted in purple, while significantly differentially expressed genes (log2FC $> 1$, adjusted p-value $< 0.01$) are highlighted in darker colors.

(E) Volcano plot displaying differential gene expression results based on the BRB-seq\textsuperscript{150} data of SVF-adherent cells from the OM adipose depot versus SVF-adherent cells from other depots (SC, PR, and MC) post-adipogenic induction. Top mesothelial markers identified using scRNA-seq datasets are highlighted in purple, while significantly differentially expressed genes (log2FC $> 1$, adjusted p-value $< 0.01$) are highlighted in darker colors.

(F) Box plot displaying the log normalized expression of MSLN (top) and UPK3B (bottom) across hASPCs (ASCs, PreAs, HHIP+, IFIT+, SFRP4+, RBP5+ ASPCs), IGFBP2+ cells, Mesothelial cells (Meso) and Vascular Smooth Muscle progenitors (VSMPs), grouped by the depot of origin indicated on the x-axis.
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Supplementary Figure 2.3.1 – The selection of scRNA-seq-inferred surface markers enables the enrichment of main SVF cell populations across all analyzed adipose depots.
(A) t-SNE cell map of integrated scRNA-seq datasets described in Fig. 2.2A and B colored by the expression of genes corresponding to the surface markers that were used to isolate each subpopulation experimentally.

(B) Violin plot showing the distribution of the log normalized expression of Aoc3 based on the scRNA-seq integration of murine datasets (see Methods).

(C) Boxplot showing the distribution of the log normalized scRNA-seq-based expression for IGFBP2, MSLN, and TM4SF1 (color) across the different human cell populations (x-axis).

(D) Flow cytometry profiles of Lin-/TM4SF1+ populations after TM4SF1 staining from five different adipose depot-derived SVF cells from the same donor.

(E) Flow cytometry-based analysis of the abundance of each indicated cell population gated from the Lin− fraction of MC SVF cells. Bar plots indicate mean, error bars standard deviation; n=2 donors.

(F) Bar plot to compare flow cytometry-based abundances of the indicated cell populations across SC, PR, OM, and MC adipose depots within the SVF Lin− fraction. The three populations accumulate to 100% by depot. For OM, each cell population is also TM4SF1− to deplete for OM-specific populations; SC n=37, OM n=35, PR n=17; MC n=2 donors.

(G) Relative cell number increase over time of culture for CD26+, DN, and VAP1+ cells in each depot. For OM, the three cell populations were gated from the Lin−/TM4SF1− population and the growth speed of Lin−/TM4SF1+ OM-specific cells were also recorded; n=12, 3 donors, 3-4 populations per depot.

SC - Subcutaneous, PR - Perirenal, OM - Omentum, MC - mesocolic/mesenteric; *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, One-Way ANOVA and Tukey HSD post hoc test (E) and RELM analysis and Tukey HSD post hoc test (F). Black compares CD26+ versus DN, Red CD26+ versus VAP1+, Blue DN versus TM4SF1+, Green VAP1+ versus TM4SF1+, Pink DN versus VAP1+.
Supplementary Figure 2.3.2 – Correlation between the FACS-based abundance of the indicated SVF Lin– subpopulations shared across depots and physiological data of the donors.

(A) Grouped bar plots showing subpopulation abundance in male versus female donors (SC n=37, OM n=35, PR n=17 donors).

(B) Scatter plot showing the correlation between FACS-based subpopulation abundance and donor age. The line represents a linear regression analysis.

SC - Subcutaneous, PR - Perirenal, OM - Omentum; *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, Unpaired Two-Way ANOVA (mixed model) multiple comparisons and Tukey HSD post hoc test (A) and linear regression analysis with its relative goodness of fit, and the FDR-adjusted p-values of the Pearson correlations (B).
Supplementary Figure 2.3.3 – ASCs and PreAs, even if detected across depots, exhibit distinct gene expression profiles.

(A) Heatmap of the correlation between 2000 randomly selected ASCs and PreAs based on the first 30 principal components of the PCA space of integrated scRNA-seq data shown in Fig. 2.2A; a similar number of cells was selected for each depot and population.

(B) Heatmap of the top 30 genes detected as significantly higher expressed in the indicated depot versus all other depots (only genes detected as differentially expressed in each pairwise comparison were retained) when focusing on ASCs (left) or PreAs (right); log normalized expression.

(C) Dot plot of enriched, representative GO terms based on the differentially expressed genes specific to the indicated depot, as explained in A.

SC - Subcutaneous, PR - Perirenal, OM - Omentum, MC - Mesocolic.
Supplementary Figure 2.4.1 – Known anti-adipogenic markers are overrepresented in OM-specific cell populations that express TM4SF1, and the proportion of TM4SF1+ cells positively correlates with BMI. Mesothelial and IGFBP2+ cells also display different morphologies.

(A) Boxplot showing the distribution of log normalized expression of the scRNA-seq data of WT1, ALDH1A2, and CD200 (x-axis) across the indicated cell populations (defined by the colors).

(B) Scatter plot showing the correlation between the OM SVF Lin−/TM4SF1+ fraction based on flow cytometry analysis and the BMI of donors; the line represents a linear regression analysis with its relative goodness of fit; the p-value was computed performing a Pearson correlation.

(C) Bright-field transmission light microscopy images of spindle-like OM (Omentum) ASPCs (OM SVF/Lin−/TM4SF1−) and cobblestone-like OM-specific TM4SF1+ populations.
Supplementary Figure 2.4.2 – Mixing OM SVF with PR SVF does not affect the adipogenic potential of PR cells.

(A) Total number of nuclei in each well (see Methods for more details) of images in Fig. 2.4E, n=16, 4 biological replicates, 4 independent wells for each.

(B) Boxplot showing the distribution of batch normalized expression of DKK2 of BRB-seq data of SVF-isolated cells from the indicated depots and treatment conditions, n=12-61, 4-20 biological replicates, 1-4 independent wells for each.

(C) qPCR-based gene expression levels of DKK2 (a subcutaneous depot-specific gene), normalized by HPRT1 expression and 0% subcutaneous (SC) to control for correct mixing ratios in the experiment shown in Fig. 2.4E. The linear regression and corresponding R2 coefficient values are shown in red; a black line links the lowest value to the highest value; n=4, 2 biological replicates, 2 independent wells for each.

(D) Representative fluorescence microscopy images of SVF Lin– cells in mixing experiments after 14 days of adipogenic differentiation, where SVF Lin– cells from OM and PR of Donor 68 were mixed directly after cell isolation at the indicated proportions. Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA, scale bar=100 mm.

(E) Total number of nuclei in each well (see Methods for more details) of images in C; n=12, 4 biological replicates, 3 independent wells for each.

(F) Quantification of the extent of adipogenic differentiation of the distinct, mixed OM and PR SVF Lin– cell populations, as presented in E. Values across biological replicates are normalized to the average adiposcore of the reference 100% PR Lin– condition. The relative proportion (0-100%) of PR SVF Lin– cells in each well is plotted on the x-axis. Error bars represent standard deviation from the average, linear and exponential regression with corresponding R2 coefficients shown in red and blue, respectively. The black line represents the expected increase of adipogenesis for a linear dilution between 0 and 100% of PR SVF Lin– cells; n=16, 4 biological replicates, 4 independent wells for each.

SC - Subcutaneous, PR - Perirenal, OM - Omentum, MC - Mesocolic.
Supplementary Figure 2.4.3 – IGFBP2+ cells exhibit a specific gene expression profile that also shares signatures with both ASPCs and mesothelial cells.

(A) Dot plot displaying the average expression and percentage of expressing cells of the top IGFBP2+ cell markers across the clusters shown in Fig. 2B.

(B) Boxplot showing the distribution of the score based on the top mesothelial cell markers (purple) or the top ASC and preA markers (green) in OM hASPCs (ASCs and PreAs), IGFBP2+ cells, and mesothelial cells.
Supplementary Figure 2.4.4 – Cells from the sc and snRNAseq atlas of WAT published by Emont and colleagues are predicted as IGFBP2+ cells.

(A) Boxplot showing the distribution of the prediction score of IGFBP2+ cells when transferring our cell cluster annotation on the data published by Emont et al. for the indicated adipose depots (x-axis); SC - Subcutaneous, OM - Omentum.
(B) UMAP computed on the integrated data of hASPCs and human mesothelial cells reported by Emont et al.\textsuperscript{59} colored by the clustering provided in the same study.

(C) Bar plot displaying the number of cells predicted as IGFBP2+ cells into each of the clusters of mesothelial cells and ASPCs originally reported by Emont et al.\textsuperscript{59}, shown in B.

(D) UMAP described in B colored by the prediction score of IGFBP2+ cells when transferring our cell cluster annotation on the data reported by Emont et al.\textsuperscript{59}.

(E) UMAP described in B colored by the score based on the top IGFBP2+ cell markers.

(F) Correlation between every donor’s BMI and the percentage of hASPC6 cells (IGFBP2+-like) for each donor based on the scRNA-seq dataset provided by Emont et al.\textsuperscript{59}; the percentage was calculated for each donor as the fraction of mesothelial cells and ASPCs combined.

(G) tSNE cell map of our integrated scRNA-seq data colored by the log-normalized expression of the indicated IGFBP2+ cell markers.

(H) UMAP described in B colored by the log-normalized expression of some IGFBP2+ cell markers as in G.

(I) UMAP described in B colored by the log-normalized expression of the indicated markers shared by predicted IGFBP2+ cells and ASPCs or Mesothelial cells.
Supplementary Figure 2.4.5 – IGFBP2+ cells are cells that transition between mesothelial and mesenchymal cell states.

(A) PAGA-inferred trajectory\(^{179}\) superimposed on the PAGA-initialized ForceAtlas2 layout between ASCs, PreAs, IGFBP2+ cells, mesothelial cells, and VSMPs as shown in Fig. 2.4G, colored by the inferred pseudotime (starting from ASCs).
(B) PAGA-inferred trajectory as shown in A colored by the log normalized expression of IGFBP2.

(C) Dot plot of key GO terms enriched based on IGFBP2+ cell markers.

(D) Heatmap showing the change of gene expression along the trajectory pseudotime shown in A for EMT-related genes (top: genes found as enriched when performing GO enrichment analysis, bottom: other EMT-related genes found in the literature). For visualization purposes, the number of cells was downsampled proportionally along pseudotime (see Methods).
Supplementary Figure 2.4.6 – TM4SF1 and MSLN markers allow distinguishing between the two OM-specific subpopulations.

(A) Representative flow cytometry plots of SC (Subcutaneous), OM (Omentum), and PR (Perirenal) SVF/Lin– from donor 53 stained with TM4SF1 and MSLN and gating strategy to enrich for ASPCs (Lin–/TM4SF1–/MSLN–), IGFBP2+ cells (Lin–/TM4SF1+/MSLN–) or mesothelial cells (Lin–/TM4SF1+/MSLN+) exclusively in OM SVF. Similar profiles were obtained from at least three donors.

(B) Quantification of the relative abundance of indicated cell populations based on Flow cytometry profiles; n=23-47 donors.

(C) Scatter plot showing the correlation between the OM SVF Lin–/TM4SF1+/MSLN– fraction based on flow cytometry analysis and the BMI of donors; the line represents a linear regression analysis with its relative goodness of fit; the p-value was computed performing a Pearson correlation.
Supplementary Figure 2.4.7 – Detection of TM4SF1+/MSLN+ and TM4SF1+/MSLN− cells by in situ immunohistochemistry.
(A) Confocal microscopy fluorescent images after TM4SF1 (Green), Perilipin (PLIN1) (Yellow), and MSLN (Pink) immunohistochemistry staining of whole OM AT cryo-cuts. The top row is the unstained control. DAPI staining for nuclei is colored in Cyan. The experiment was repeated three times, yielding similar results.

(B) Confocal microscopy fluorescent images after TM4SF1 (Green), Perilipin (PLIN1) (Yellow) and MSLN (Pink) immunohistochemistry staining of whole OM AT cryocuts. The top row is the unstained control. DAPI staining for nuclei is colored in Cyan. The experiment was repeated three times, yielding similar results. The white arrows point to TM4SF1+ cells, the red arrows point to TM4SF1+/MSLN+ cells.
Supplementary Figure 2.4.8 – TM4SF1+/MSLN- cells change morphology upon expansion *in vitro*.

(A) Bright-field transmission light microscopy images of confluent or expanding spindle-like OM (Omentum) ASPCs (OM SVF/Lin-/TM4SF1-) and cobblestone-like Mesothelial cell (OM SVF/Lin-/TM4SF1+/MSLN+) and IGFBP2+ (defined as OM SVF/Lin-/TM4SF1+/MSLN-) cell populations.
Supplementary Figure 2.5.1 – PR SVF-adherent cells are insensitive to the inhibition exerted by IGFBP2-secreting cells in a transwell setting.

(A) Total number of nuclei in each well (see Methods for more details) of images in Fig. 2.5E; n=12, 4 donors, 3 independent wells.

(B) Representative fluorescence microscopy images of “receiver” PR SVF-adherent cells, at the bottom of the transwell set-up, after adipogenic differentiation when co-cultured with the indicated SVF fractions at the top: paired PR SVF-adherent cells, OM SVF-adherent cells, OM SVF/Lin-/-TM4SF1- (OM ASPCs), OM SVF/Lin-/-TM4SF1+/MSLN- (IGFBP2-secreting cells), or OM SVF/Lin-/-TM4SF1+/MSLN+ cells (mesothelial cells). First row: PR and OM cells from D54, Second row: SC and OM cells from D65.

(C) Barplot showing the adiposcore quantification of bottom cells in B. Values are normalized to the average adiposcore of the reference top PR SVF-adherent condition; n=12, 4 donors, 3 independent wells.

(D) Total number of nuclei in each well (see Methods for more details) of images in B; n=12, 4 donors, 3 independent wells.

(E) Total number of nuclei in each well (see Methods for more details) of images in Fig. 2.5I; n=16-20, 4 donors, 2-4 independent wells.

SC Subcutaneous - Yellow, OM Omentum - Purple, PR Perirenal - Brown.
Supplementary Figure 2.5.2 – Experimental titration of IGFBP2, IGF-I, and IGF-II recombinant proteins to test their effect on adipogenic differentiation.

(A) Representative fluorescent microscopy images of SC SVF-adherent cells treated with the indicated concentrations of interfering compounds. Scale bar=100 mm.

(B) Barplot showing the adiposcore quantification of IGFBP2-treated cells in A. The adiposcores are normalized to the non-treated cells (Ctrl); n=4, 2 donors, 2 independent wells.

(C) Total number of nuclei in each well (see Methods for more details) for IGFBP2-treated cells in A; n=6, 3 donors, 2 independent wells.

(D) Representative fluorescent microscopy images of PR SVF cells treated with the indicated concentrations of interfering compounds. Scale bar=100 mm.
Barplot showing the adiposcore quantification of IGFBP2-treated cells in D. The adiposcores are normalized to the non-treated cells (Ctrl); n=4, 2 donors, 2 independent wells.

Total number of nuclei in each well (see Methods for more details) for IGFBP2-treated cells in D; n=6, 3 donors, 2 independent wells.

Barplot showing the adiposcore quantification of IGF-I-treated cells in A. The adiposcores are normalized to the non-treated cells (Ctrl); n=4, 2 donors, 2 independent wells.

Total number of nuclei in each well (see Methods for more details) for IGF-I-treated cells of panel A; n=6, 3 donors, 2 independent wells.

Barplot showing the adiposcore quantification of IGF-II-treated cells in A. The adiposcores are normalized to the non-treated cells (Ctrl); n=4, 2 donors, 2 independent wells.

Total number of nuclei in each well (see Methods for more details) for IGF-II treated cells of panel A; n=6, 3 donors, 2 independent wells.

Barplot showing the adiposcore quantification of IGF-I-treated cells in D. The adiposcores are normalized to the non-treated cells (Ctrl); n=4, 2 donors, 2 independent wells.

Total number of nuclei in each well (see Methods for more details) for IGF-I treated cells of panel D; n=6, 3 donors, 2 independent wells.

Barplot showing the adiposcore quantification of IGF-II-treated cells in D. The adiposcores are normalized to the non-treated cells (Ctrl); n=4, 2 donors, 2 independent wells.

Total number of nuclei in each well (see Methods for more details) for IGF-II treated cells of panel D; n=6, 3 donors, 2 independent wells.

For each image: Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA. *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001 One-Way ANOVA and Tukey HSD post hoc test.
Supplementary Figure 2.5.3 – Treatment with IGFBP2 did not significantly affect the adipogenic differentiation of PR SVF-adherent cells, while treatment with Echistatin increased the accumulation of lipid droplets.

(A) Total number of nuclei in each well (see Methods for more details) for cells in Fig. 2.5K. n=12, 4 donors, 3 independent wells per replicate.

(B) Representative fluorescence microscopy images of PR SVF cells after adipogenic differentiation when treated with the indicated interfering compounds. IGFBP2 1nM, IGF-I 10nM, IGF-II 10nM, Echistatin 100nM.

(C) Total number of nuclei in each well (see Methods for more details) for cells in A. n=12, 4 donors, 3 independent wells per replicate.

(D) Barplot showing the adiposcore quantification of cells in A focusing on the IGF-independent signaling pathway of IGFBP2. The adiposcores are normalized to the non-treated cells (Ctrl); n=12, 4 donors, three independent wells.

(E) Total number of nuclei in each well (see Methods for more details) for cells in Fig. 2.5N. n=12, 4 donors, 3 independent wells per replicate.

(F) Barplot showing the adiposcore quantification of cells in A focusing on the IGF-dependent signaling pathway of IGFBP2. The adiposcores are normalized to the non-treated cells (Ctrl). n=12, 4 donors, three independent wells.

SC Subcutaneous - Yellow, OM Omentum - Purple, PR Perirenal - Brown; For each images: Yellow - Bodipy stains for lipids, blue - Hoechst stains for DNA *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001, One-Way ANOVA and Tukey HSD post hoc test.
## Table 2.1 – Donors’ information

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**Legend**

AS: Adiposcore
BF: Brightfield images
BIB: Barcoded bulkRNA sequencing
CE: Chemical experiments
CI: Confocal imaging
ELISA: Enzyme-linked immunosorbent assay
FC: Flow cytometry
IR: Immunohistochemistry
KD: Small hairpin RNA knockdown experiments
Mixing experiments
Profiles: Flow cytometry scatter plots
scRNAseq: Single-cell RNA sequencing
TW: Transwell

**Notes**

- Table data includes donors’ information such as height, weight, BMI, age, gender, and type of surgery.
- The table is structured to compare various measurements and surgical outcomes.
- The data is presented in a clear, organized manner, facilitating easy comparison and analysis.

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Table 2.2 – Donors’ specifications for scRNA-seq

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Table 2.3 – Cohorts specifications

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Table 2.4 – Antibody specifications for FACS

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<tr>
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<td>Thermo</td>
<td>A31573</td>
<td>Donkey</td>
<td>AF487</td>
<td>1:200</td>
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Table 2.5 – Antibody specifications for IHC

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<th>Host species</th>
<th>Fluorophore conjugate</th>
<th>Biotium Mix n Stain fluorofore conjugate</th>
<th>Working titration</th>
<th>Primary (I) or Secondary (II)</th>
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<td>Thermo</td>
<td>A31573</td>
<td>Donkey</td>
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Chapter 3 | Conclusion and future perspectives
3.1 The adipose-tissue SVF and its heterogeneity across species

3.1.1 The ASPC heterogeneity in mouse

The advent of scRNA-seq techniques allowed for unprecedented and unbiased revelation of new cell sub-populations within heterogeneous cell populations. If the scientific community has been able to produce whole body atlases of simple research models such as C. elegans and D. melanogaster, when it comes to higher mammals accounting for billions of cells, we have no other ways than exploring organ by organ, tissue by tissue the underlying cell heterogeneity. The Deplancke lab undertook the scRNA-seq-based exploration of adipose stem and progenitor cells (ASPCs) in 2015 and since then contributed to a number of groundbreaking discoveries, among which was resolving the heterogeneity of mouse subcutaneous (SC) ASPCs. Shortly after, several other labs corroborated our findings and added depth to general knowledge by resolving epigonadal or omental mouse ASPCs heterogeneity, all concluding on the fact that the ASPC pool is heterogenous regardless of the anatomic depot of origin. Specifically, the mouse ASPC landscape across depots is populated by two canonical mesenchymal cell types/states, the adipose stem cells (ASC) and the pre-adipocytes (PreA). Downstream functionalization of these cell sub-populations corroborated the choice of nomenclature, given that ASCs as opposed to PreAs showed greater proliferation abilities, while the latter the highest adipogenic potential. Transcriptomically close to ASCs and PreAs stood an enigmatic cell type characterized by a high and specific expression of F3 and which robustly clustered apart from the two main populations. Technically, this mouse F3+ population should not be considered as an ASPC as it is unable itself to form mature adipocytes, instead, exerts a negative regulation on neighboring adipogenic cells like ASC or PreAs, hence the proposed nomenclature of Aregs (for Adipogenesis regulators). Lineage tracing analysis further validated the newly established niche landscape, in fact it was shown that ASCs stand at the very root of adipogenesis, and can give rise to both PreAs and Aregs in vivo. By integrating all publicly available datasets in 2020, a fourth, final mesenchymal population showed consistent clustering. This is the Clip+ population. Although no functional characterization was performed on the Clip+ population itself, their transcriptome is somewhat close to both ASCs and Aregs and an interesting hypothesis would be that they could be specialized Aregs precursor cells. Trajectory inference analysis and targeted lineage tracing experiments would help lift the veil on this aspect. Interestingly, a few cells from the integrated dataset (lfit+ cluster) are characterized by high expression of interferon-related markers. The gene expression signature of these cells is reminiscent of fibroblast-inflammatory cells that are highly specialized in innate immune response directed against viruses that locate preferentially in the red pulp of the spleen. Therefore, these cells may act as specialized sensors for viral infection of the adipose tissue, and it would be interesting to interrogate their role in the tropism of specific viruses, such as SARS-CoV2, towards adipose tissue. Besides the two canonical ASPC clusters, the Areg and its potential progenitor
Clip+ and the specialized anti-viral cluster Ifit+, which are identifiable in both subcutaneous and epigonadal mouse AT, is a cell cluster that is not mesenchymal in nature, but rather epithelial, and that exclusively stems from epigonadal- and omental-derived datasets: the mesothelial cells. Considering the newly established subpopulation landscape of the ASPCs in the adult mouse, further work should focus on interrogating how the equilibrium of the niche changes upon physiological conditions in health and disease. As such, it would be useful to develop reporter mouse models for each of the uncovered cell subpopulation. This could be achieved, for example, by fluorescently labeling one of the top differentially expressed markers of each subpopulation. Once these tools ready, we would be able to interrogate changes in the relative abundance of each cell subpopulation in different biological scenarios (i.e. early development and pre-pubertal time, upon aging and post-menopausal time, across genders, as well as under high-fat diet or high-glucose pulse conditions) and test the biological limits of adipose tissue plasticity.

The anti-adipogenic effect of Aregs is one of the most intriguing aspects that arose from studying the mouse ASPC niche. Despite that the non- and anti-adipogenic character of these cells have been challenged by other labs, we and others have proved the phenotype consistency of Aregs (CD142+ (=F3+)) cells across a number of different conditions (i.e. mouse strains, FACS gating strategy, differentiation cues) in recent studies. Out of all tested conditions, two appeared to be crucial to ensure the correct sorting and characterization of non- and anti-adipogenic Aregs. On the one hand, in adult mice, it is important to exclude contamination of Aregs by Vap1+ ASPCs. Hence Dong and colleagues proposed a refined sorting strategy, where Aregs are defined as SC SVF Lin−/Sca1+/VAP1−/CD55−/CD142+ ASPCs. On the other hand, the animal age from which cells were isolated turned out to be critical to ensure Aregs phenotype. In fact, Aregs isolated from SC adipose tissue of pup mice, specifically from the pre-weaning age, appeared to be highly adipogenic – to at least the same extent as the other ASPCs (CD142- (=F3-) ASPCs) – and were incapable of negatively influencing the adipogenic potential of neighboring cells. Weaning is a critical event in the metabolism of mammals including humans, where the gut of the offspring has to learn to digest complex sugars and nutrients that were absent in the mother's milk. As such, weaning is accompanied by a drastic change in the gut microbiota. It is by now well established that some nutrients in the gut can only be absorbed following pre-metabolization by the microbiome, and a number of vitamins and essential nutrients for the organism, including retinoic acid, are subjected to this bacterial pre-processing step before being at the host organism disposal.

We showed that retinoic acid, together with secreted proteins CD142 and MGP play a key role through an auto-paracrine axis to ensure the non-adipogenic character of the Aregs on one side, and on the other side in the inhibitory mechanism of Aregs toward neighboring ASPCs. Hence, it would be tempting to hypothesize that changes in the gut microbiome upon weaning might be influencing the phenotype of Aregs. This could be tested by exploring Aregs behavior in germ-free mice, mice subjected to selected depletion of microbiota through antibiotic treatment, or by altering the timings of the diet-transition enforced by weaning.
Chapter 3 | Conclusion and future perspectives

3.1.2 The ASPC heterogeneity in human

Mice are certainly a great resource when it comes to expanding our general knowledge of how fundamental biology mechanisms operate at the whole-body scale in a mammalian organism. However, findings in mice might not be always translated in humans. A relevant example is the discovery of the leptin hypothalamic-pituitary endocrine axis in the mid ‘90s. The leptin axis is one of the major food intake regulation axes in mice. Knocking out the \textit{Lep} gene in the wild-type mouse is responsible by itself for the onset of severe obesity as a consequence of excessive food intake by the animal (\textit{Ob/Ob} mouse model). However, when it comes to humans, perturbations in the leptin signaling axis were described in only sporadic cases of severe obesity, as a consequence of homozygotic mutation in the leptin-coding region of \textit{Lep}. In addition, except for the patients with a non-sense mutation in the leptin gene itself, where the leptin analog therapy rescued a normal body weight, no drug-discovery-related attempt to enhance the leptin axis was effective for weight management therapy, and to date, the only drug-development attempts concerning the leptin axes are carried out in the field of cancer biology. Several other important differences exist between the mouse and human adipose tissue. For example, the brown adipose tissue is a fully functional and important organ in mice, both in early life and adulthood, responsible for maintaining body temperature homeostasis throughout the entire lifespan of the animal. In humans, brown adipose tissue is mainly present in newborns, while during adulthood, only vestigial depots are found in the supraclavicular, mediastinal, cervical, axillary, paraspinal and perirenal regions and might only marginally contribute to the regulation of body temperature. Another striking difference in the AT distribution in mice versus humans is the anatomy of the main visceral AT. The omentum, the main human visceral adipose tissue is nothing more than a tiny line of tissue above the stomach of the mouse, barely distinguishable from the pancreas, inversely, the mouse epigonadal AT occupies the vast majority of the animal abdominal space, while is barely distinguishable from the human gonad itself. Given these differences in adipose tissue biology and the possibility that findings in mouse AT might not extrapolate to humans, there was a need to understand the heterogeneity and functionality of human ASPCs.

Throughout my Ph.D., I got access to adipose tissue biopsies from three anatomic locations (Subcutaneous = SC, Perirenal = PR, Omentum = OM) and sporadic from two of them (Mesocolon = MC and the Gallbladder-associated AT = GB) and was able to collect samples from up to 85 donors (Table 2.1). This allowed to depict a comprehensive snapshot of similarities and differences in the hASPC composition in function of the anatomic depot of origin both at the single-cell (~34’000 SVF/Lin– cells analyzed) resolution (Fig. 2.2) and at the bulk transcriptomic one (20 SC, 8 PR, 19 OM, and 4 MC primary cell lines analyzed) (Fig. 2.1). Like in mice and in other publicly available work, two main hASPC populations, the hASCs (DPP4+, CD55+) and the hPreAs (ICAM1+, APOD+) appeared to be present in all four canonical adipose tissues (SC, PR, OM. And MC) as well as in the GB AT, which is at times present around the gallbladder of morbidly obese donors exclusively. We
experimentally proved that hASC-enriched SVF cells (Lin−/TM4SF1−/CD26+) are more proliferative and less adipogenic than hPreA-enriched SVF cells (Lin−/ TM4SF1−/DN) regardless of the anatomic depot of origin (Fig. 2.3E-F and Supp. Fig. 2.3.1G). In silico trajectory analysis that we performed, also hints at the lineage dependency of these two cell populations (Fig. 2.4G) as was previously described96. Aside hASCs and PreAs stand another transcriptomically well-distinct cell subpopulation common to all anatomic locations. These cells that resemble Vascular Smooth Muscle Progenitors (VSMP)159 highly express muscle-related markers such as actin and myosin (Fig. 2.2.1C) and represent between 2 to 10% of the SVF Lin− cells depending on the depot(Fig. 2.2D). Using AOC3 marker encoding for VAP-1 surface protein, we were able to sort VSMP-enriched SVF cells (Lin−/TM4SF1−/VAP1+) (Fig. 2.3E-F) which surprisingly showed the highest adipogenic potential in vitro (even higher than the PreAs defined as Lin−/TM4SF1−/DN) (Fig. 2.3E-F) coupled to the slowest growing pace (Supp. Fig. 2.3.1G). If at the first glance is contradictory that muscle-primed cells can be so adipogenic, this could be explained by the fact that VSMPs are in fact beige adipocytes progenitors 69,204-207. Another more technical explanation for the observed high-adipogenic phenotype of VSMPs would be that despite the fact that they significantly express to the highest extent AOC3, a number of PreAs also express AOC3. It would thus be advisable to investigate if the high adipogenicity of VAP1+ cells come from in fact a “contamination” by highly adipogenic PreAs.

Like in mouse, we identified a number of other minor clusters including one expressing HHIP that resemble mouse Aregs (Supp. Fig. 2.2.5), one expressing SFRP4 that resembles Clip+ cells in mouse (Supp. Fig. 2.2.7) and even one that expresses IFIT like mouse Ifit+ cells that may be involved in innate immunity response (Supp. Fig. 2.2.6)236. While the overall transcriptomic signatures are close to the ones of the mouse populations, the top expressing markers of human cells are slightly different and this difference should be taken into account for further investigations. Interestingly, one of the minor clusters from the human merged dataset was specific to the PR and MC AT (Fig. 2.2D, Supp. Fig. 2.2.1H, Supp. Fig. 2.2A-B), however, >80% of the cells from this cluster come also from one single donor (Supp. Fig. 2.2A), it is thus difficult to draw conclusions on if this cluster is truly PR-MC-specific or patient-specific. Aside from the above-mentioned pan-anatomic clusters, stand out two depot-specific ones: the mesothelial and the IGFBP2+ cells, which are exclusively retrieved in the OM AT (Fig. 2D, Supp. Fig. 2.2A-B) and whose new functional properties are discussed in the paragraph below. Overall, mounting evidence point to the fact that we should no longer ignore the underlying human nor mouse ASPCs heterogeneity for future work aiming at understanding the neo-adipogenesis process. For future perspectives, it would be interesting to include even more adipose tissue types (for example the pericardial, epicardial, mediastinal or orbital) to study not only their ASPC composition but also the heterogeneity of two important cellular players of the adipose tissue that were missed by the current study: the mature adipocytes themselves and the resident immune cells.
3.2 Mesothelial cells are key effectors of the stromal vascular fraction

Mesothelial cells build up the peritoneum, a monolayer of mesothelial cells upholstering the abdominal cavity. However, the functional role of mesothelial cells in the abdominal cavity has not been extensively studied so far. In this work, we identified mesothelial cells (OM SVF Lin−/TM4SF1+/MSLN+) and mesothelial-like IGFBP2+ cells (OM SVF Lin−/TM4SF1+/MSLN−) (Fig. 2.5A, Supp. Fig. 2.3.1C) exclusively in the human SVF of the OM AT (Supp. Fig. 2.4.6A). In mice, mesothelial cells can be found in the murine epigonadal and omental ATs. Therefore, future works studying visceral AT ASPCs should make sure to exclude mesothelial cells from the “true” ASPCs for any transcriptomic and functional analyses. To do so, we used TM4SF1 as a pan-mesothelial marker, and we observed that the OM SVF/Lin− fraction depleted of TM4SF1+ cells (OM SVF/Lin−/TM4SF1− cells), was significantly more adipogenic than the total fraction of OM SVF Lin− cells (Fig 2.4C-D), hinting to the fact that mesothelial cells and mesothelial-like cells might play a negative role in regulating the OM hASPC adipogenesis. Moreover, we demonstrated, that TM4SF1+ cells can further be segregated into two functional subpopulations, one that is MSLN+ (OM SVF Lin−/TM4SF1+/MSLN+) and another that is MSLN− (OM SVF Lin−/TM4SF1+/MSLN−) (Fig. 2.5A, Supp. Fig. 2.3.1C). While both populations were able to inhibit the adipogenesis of hASPCs to some extent, only the MSLN− fraction did so in a consistent and robust way (Fig. 2.5E-F). We were further able to prove that the inhibition stems from the fact that TM4SF1+/MSLN− highly secrete and express IGFBP2 (Fig. 2.5B-C), which inhibited adipogenesis through the activation of the integrin receptor α5β1 (Fig. 2.5K-P)186. During the length of this thesis, we demonstrated this point by chemical and recombinant protein interference assays in primary cultures (Fig. 2.5K-P). Future works could further complement this line of research by better understanding the intracellular downstream effects of the integrin receptor activation. This could be done also in primary human cultures, by knocking down the integrin receptor and comparing the phosphorylation status of intracellular proteins in the presence or absence of the IGFBP2 protein treatment. Similarly, to further understand the transcriptional reprogramming occurring in OM SVF cells upon IGFBP2 treatment, ChIP-seq or ATAC-seq assays could be performed.

In our samples, the percentage of OM-specific TM4SF1+ cells (mesothelial and IGFBP2+ together) measured by flow cytometry positively correlated with the donor’s BMI (Supp. Fig. 2.4.1B). Taking advantage of the Emont et al. atlas145, we were able to corroborate in silico a positive correlation between IGFBP2+ cells specifically and the donors’ BMIs (Supp. Fig. 2.4.4F). While such findings hint at a functional role of the IGFBP2+ cells in vivo, IGFBP2 serum levels in humans negatively correlated with BMI214–216, and characteristic features of the onset of the metabolic syndrome217, including type 2 diabetes and NAFLD218. Circulating levels of IGFBP2, however, are highly determined by hepatic IGFBP2 secretion218,253, and, therefore, might not reflect its regulation in OM AT. To further characterize the role of IGFBP2 in human physiology it would also be
interesting to interrogate whole genome sequencing biobanks to look for SNP in the IGFBP2 locus and study what is the phenotypic trait associated to it, with a specific focus on metabolic traits.

IGFBP2-secreting cells are rare among the OM SVF (2-5% of Lin− cells) (Fig. 2.4.6B) and once in culture are slow-growing cells, difficult to expand to reach useful cell numbers. This could constitute a critical limitation for studies aiming to unravel their biological function. The inner limitations of human biopsy collections might also limit our ability to better understand the effects of IGFBP2 signaling in vivo. In this sense, it would be useful to investigate whether IGFBP2-secreting cells exist in the epigonadal AT of the mouse. A possible avenue to do so is to validate antibodies directed against the mouse TM4SF1 and MSLN antigens, and to use the same sorting strategy described in humans, however, given the described fundamental differences that we systematically observed across human and mouse AT biology, it would be more appropriate to favor an unbiased approach and first look for Igfbp2+ cells in publicly available scRNAseq datasets obtained from mouse epigonadal SVFs directly91,100,101,145. This could allow us to identify new mouse-specific surface markers to eventually validate through flow cytometry. In that direction, Zhang and colleagues91 recently identified through single-cell transcriptomics a mesothelial cell population that highly expresses Igfbp2 in the adult mouse epWAT. In contrast, at a younger age (P3) Igfbp2 was differentially expressed by the smooth muscle cell cluster (likely resembling the human VSMPs). Ultimately, genetically modified animal tools would greatly help to explore the role of Igfbp2 in vivo. Transgenic human Igfbp2-expressing mice, as well as Igfbp2-KO and eGFP-tagged mice, are commercially available and could provide new avenues to evaluate the role of IGFBP2 in adipose tissue biology upon metabolic challenges. Similarly, the eGFP-tagged-IGFBP2 transgenic mouse would allow to easily sort Igfbp2+ cells as well as to visualize them in situ.

In this work we highlighted a new functional role of mesothelial cells with regard to adipogenesis regulation, however they might also be functionally relevant with regard to the cross-talk with the immune system143,201,254. We showed that SVF-adherent cells from the OM AT highly express pro-inflammatory markers and upregulate pathways related to immune response (Fig. 2.1, Supp. Fig. 2.1.5A). Interestingly, the mesothelial cluster in the scRNA-seq specifically expresses chemokines and cytokines like IL-33, IL-1b and IL-18 as well as WT1 that all have been shown to contribute to the low-grade inflammation often linked to visceral obesity252,254–256. Further effort should be put into understanding if the inflammatory signals we captured among OM SVF-adherent cells are secreted by mesothelial cells, the ASPCs or if it is a crosstalk between the two cell types that induces an overall inflammatory response. The cross-talk between mesothelial cells and ASPCs could be studied using co-culture transwell systems, where non-inflammatory cells like SC or PR ASPCs would be harvested for transcriptomic analysis after being exposed to paracrine signaling cues of mesothelial or IGFBP2-secreting cells, or vice-versa. Similar experiments could be designed to evaluate the interaction of IGFBP2-expressing cells with immune cells. Another valuable and interesting approach would be to use
spatial transcriptomics and histochemistry techniques to study the physical interaction between immune and mesothelial cells.

3.3 The emerging concept of negative regulation of adipogenesis

3.3.1 Do Aregs exist in human?

Aregs is a new functional cell population that was first described by our lab in 2018\textsuperscript{102}. In mice, they are defined as subcutaneous SVF Lin–/Sca1+/CD142+ cells, which appear to inhibit adipogenesis both \textit{in vitro} and \textit{in vivo}. Their relative abundance among SVF cells increases upon HFD feeding\textsuperscript{102,148,149}, which may seem counterintuitive at a first glance, but in fact could reflect either (1) a compensatory mechanism to counteract the adipose tissue expansion, or (2) the development of intrinsic resistance to the regulatory mechanism of Aregs upon weight gain. Finding an equivalent population in humans was among the fundamental endeavors of the present work. Interestingly, while a clear and consistent cluster of \textit{F3}-expressing cells was identified in both subcutaneous and visceral SVFs of the mouse\textsuperscript{145,257}, it looks like they are an extremely rare cell type in humans. In fact, we were able to detect an Areg-like cluster only after aggregating all our datasets together. In addition to their rareness, \textit{F3}, the highly specific mouse marker is much less specific to human Aregs and is expressed instead at low levels by all the hASPCs. Based on our analysis \textit{in silico}, \textit{HHIP} should be considered a more valuable marker in the context of human Aregs biology given its specific enrichment in what we identified as the Areg-like cluster (\textbf{Supp. Fig. 2.2.5}). Validating a sorting strategy based on HHIP staining would therefore be an appropriate approach to better characterize human Aregs.

Despite human Areg-like cells are yet to be fully defined and characterized, understanding the mechanism of action of mouse Aregs is highly relevant since it is likely conserved across species. In fact, recent work from our lab showed that when SC hASPCs (SC SVF-adherent cells) are exposed to mouse Aregs (SC SVF Lin–/Sca1+/CD142+) in a transwell system, this resulted in significant impairment in their adipogenic potential, similar to the observed inhibitory action of Aregs on mouse adipogenic CD142– ASPCs (SC SVF Lin–/Sca1+/CD142–)\textsuperscript{148}. Using the same rationale, it would be interesting to interrogate the effect of the inhibitory power of mouse Aregs on hASPCs derived from OM and PR AT as well, aiming at understanding if Aregs’ inhibitory power is universal. The human-mouse hybrid setup could also be a useful tool to better understand the mechanism of action of Aregs taking advantage of the genetic differences between species.

In an interesting twist, our lab recently reported that when mouse adipogenic CD142– ASPCs are exposed to the Aregs secretome in the transwell setup, not only they ceased to be adipogenic, but their transcriptome became close to the one of Aregs. This is suggestive of a snowball effect where Aregs presence would induce an Areg-like state in their surrounding cells\textsuperscript{148}. This could stem from a canonical protein-receptor cell-signaling mechanism\textsuperscript{148,149} but also through signaling based on extracellular vesicles/exosomes “transmission”.

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Mesenchymal cells are known heavy producers of extracellular vesicles that carry anti-inflammatory and therapeutic properties. The hypothesis that Aregs would therefore signal through exocytosis of cytoplasmatic vectors that would fuse to the cytoplasm of receiving cells and carry Aregs’ mRNA with them could be interesting to pursue. Our findings showing that hASPCs can be influenced by Aregs could be helpful for this purpose, as one could look for contaminating mouse mRNA into the receiving hASPCs transcriptome as a way to elegantly test if Aregs signal through an extracellular vesicle flow. This approach, however, would not detect peptides, lipids or other metabolites that could mediate actions through extracellular vesicle communication, which cannot be ruled out.

In the present work we show for the first time that negative regulators of adipogenesis exist not only in mice but also in humans. Mesothelial cells and specifically IGFBP2+ cells, residing within the OM AT are able to negatively regulate the neoadipogenesis of OM ASPCs but also SC ASPCs. This finding opens a number of research avenues to understand how these cells modulate the plasticity of OM AT in vivo. This could have important therapeutic consequences, as OM AT expansion is linked to metabolically unhealthy obesity. In order to advance in a more unbiased way toward this understanding, it would be great to undertake a multiomics approach, where we would perform combined proteomics, lipidomics and metabolomics on IGFBP2-secreting cell supernatant to identify in a more direct way what are the cues that are effectively secreted by the cells. Another interesting approach in both Aregs and IGFBP2+ cell biology would be to test the universality of their regulatory mechanism on whole new systems such as the muscle and the bone marrow as well as on bone or cartilage morphogenesis. In all these systems to have a precise means to control tissue renewal would allow precious advances in tissue engineering and transplantation.

### 3.3.2 Adipogenesis is differentially regulated across anatomic locations

Despite high similarity across hASPCs derived from different anatomic regions, we and others were able to demonstrate, that cells from different depots carry over a transcriptomic footprint from their anatomic origin (Fig. 2.1H-I). As such, SC SVF-adherent cells highly express several developmental related HOX genes, PR cells have several pathways linked to oxidative respiratory chain and thermogenesis upregulated while OM cells are extremely inflammatory and carry an epithelial signature compared to MC which are characterized by heat-shock-protein (HSP) expression. It is therefore reasonable to think that different regulatory processes reign in different anatomic depots. Interestingly IGFBP2 signaling seems to act differentially across depots. While OM and SC hASPCs adipogenesis is effectively blunted by IGFBP2 at the concentration of around 30 ng/ml (1nM) (Fig. 2.5K-P), PR cells seem to be overall less sensitive to IGFBP2 (Supp. Fig. 2.5.1B-C, Supp. Fig. 2.5.3B, D-F). Interestingly, OM SVF-adherent cells compared to all other depots differentially express IGFBP2 both at the undifferentiated state and the differentiated one. In contrast, PR ASPCs upregulate IGFBP2 exclusively upon differentiation (low IGFBP2 expression at PR t0, high IGFBP2 expression at PR t14) (Fig. 3.1).
Taken together, not only does the concentration of the inhibitory cue seem to play a key role in determining selectively which adipose depots to regulate, but determinant could also be the timing of secretion and the signaling mode. In this sense, it is important to note that IGFBP2 might not only act as a secreted protein, but it can also have intracellular actions, for example, by interacting with tumor-suppressing genes such as PTEN. It would therefore be interesting in the case of PR to identify if the IGFBP2 upregulation results in its secretion or not.

IGFBP2 certainly contributes to the refractoriness of OM ASPCs to form lipid droplets in vitro but it still cannot be fully explained why intraperitoneal-derived ASPCs are so strongly refractory to adipogenesis upon cell culture. Even the most highly adipogenic OM ASPC population (OM SVF Lin−/TM4SF1−/VAP1+) is by far not as adipogenic as the SC or PR corresponding populations (Fig. 2.3E). This observation hints at the fact that there must be some cell-intrinsic mechanism that prevents OM ASPCs to become adipocytes. Several OM-specific markers aside from IGFBP2 could potentially be involved in the cell-intrinsic non-adipogenic character of OM ASPCs. For instance, retinoic acid, GAL, PTN, SLPI, BCHE and CD200 are all differentially expressed by OM ASPCs and have previously been described as being anti-adipogenic factors. Interfering with these markers, by knocking them down or overexpressing them as well as dissecting their signaling pathways may lead to new discoveries in the field of adipogenesis regulation. Rspo2 has been reported to play a key role in the inhibitory mechanism used by Aregs to inhibit. To test how hASPCs from different depots respond to RSPO2 treatment would be interesting to evaluate its relevance in human adipose tissue biology.

![IGFBP2 expression at t0 and t14 across depots.](image)

Boxplot showing the distribution of batch normalized expression of IGFBP2 of BRB-seq data of SVF-isolated cells from the indicated depots and treatment conditions, n=12-61, 4-20 biological replicates, 1-4 independent wells for each.
3.4 Understanding the adipogenic lineage

3.4.1 The mesothelium as a possible origin for OM adipocytes

Aside from being able to negatively regulate adipogenesis, we showed the IGFBP2+ cells are susceptible to transition between mesothelial and mesenchymal cell states. Based on scRNA-seq data and pseudotime analysis, we showed that these cells order along a trajectory connecting ASPCs to mesothelial cells (Fig. 2.4G). Along the connecting branch, mesothelial and ASPCs markers increase and decrease, respectively, while some genes including IGFBP2 and others previously linked to mesothelial to mesenchymal transition (MMT) peak at the center of the trajectory (Fig. 2.4J). While MMT acts similarly to the Epithelial to Mesenchymal Transition (EMT), its role in physiopathology is still unclear and has mainly been described in lung, liver and kidney fibrosis as well as in the context of peritoneal fibrosis during dialysis. MMT is however not necessarily a detrimental event. Due to these transitional properties, the omentum is routinely used in visceral surgery as a living patch to reconstruct perforated intestines. MMT is uttermost important during organogenesis, where mesothelial cells - expressing Msln - give rise to various cell types, such as fibroblast, or smooth muscle, of the visceral organs. Our work put forward the hypothesis that MMT would not only occur during early developmental stages but would happen in adulthood in a steady-like state. Whether the MMT in adulthood would be the root of adipocyte formation in the omentum is a controverted subject that, if proven valid, would revolutionize our vision of OM AT plasticity. A lineage tracing study performed on murine epididymal fat originally, showed that eWAT but not scWAT nor BAT adipocytes express Wt1, likely reflecting an origin from Wt1+ mesothelial cells, while mesothelial cells were also described to be adipogenic in vitro. A very recent study confirmed that mature eWAT adipocytes do express Wt1, but also demonstrated that only Wt1 Pdgfra+ cells were able to give rise to mature adipocytes, while canonical Krt19+ Wt1+ mesothelial cells were unable to do so both at the developmental level and under forced adipose tissue expansion via high-fat diet. Interestingly, our IGFBP2+ cells express PDGFRa but not KRT19. We never directly tested the adipogenic abilities of our IGFBP2-secreting cells but this would be certainly an interesting experiment to perform. Similarly, it would be interesting to see if overexpression of known MMT transcription factors in mesothelial cells isolated from human omentum, would trigger a gene expression profile similar to the IGFBP2+ cells.

3.4.2 Redirecting the adipogenic lineage towards beiging

In the present work, we were able for the first time to determine the relative abundance of different cell subpopulations in the SVF Lin− niche of human SC, OM and PR AT of over 30 donors (Fig. 2.3B-C). This analysis showed that the ASC population is the least abundant in the OM AT, compared to SC and PR (Fig. 2.3D, Supp. Fig. 2.3.1F). This likely reflects exhaustion of the OM ASC pool in adulthood, which could explain the
preferential hypertrophic mode of expansion of OM AT. Conversely, PreAs predominate the ASPC landscape of SC ASPCs, while VSMP (or VAP1+-expressing PreAs) predominate the visceral ones (i.e. OM, PR and MC) (Fig. 2.3D, Supp. Fig. 2.3.1F). With increased BMI, and regardless of the anatomic depot, we highlighted an overall shift of the niche equilibrium from the PreAs (whose abundance negatively correlates to BMIs) to the VSMP pool (whose abundance positively does) (Fig. 2.3G). If we couple this hypothesis with the cell hierarchy model proposed by Merrick and colleagues96, where mouse ASC would give rise to ICAM1+ cells and CD142+ cells, it would be interesting to explore if human ASC could give rise to both DN PreAs and VAP1+ VSMPs/PreAs as an early commitment to white or beige adipogenesis respectively. If this was the case, the observed increase in the relative proportion of VAP1+ cells in all ATs may reflect a desperate attempt to induce a thermogenic response to balance the excessive energy intake or to create new vasculature with the AT expansion.

Sprouting literature highlights how beige/brown adipogenesis is often coupled to the upregulation of myogenic-like markers necessary to dissipate energy under the form of heat69,204–207. In this regard, it would be interesting to test the ability of VAP1+ cells specifically to upregulate UCP1 under adrenergic stimulation or to perform respirometry or mitochondrial activity tests on these cells and compare their behavior with the other cell subpopulations as this would mount evidence supporting the hypothesis that VAP1+ cells could be thermogenic.

3.5 The Omentum AT as a hotspot for abdominal cancer metastasis

More than a bare energy reservoir, the omentum is a fascinating crossroad between connective and lymphatic tissue able to secrete endocrine signals, combat infections and store lipids. Clinically many visceral malignancies, including ovarian, gastric and colorectal cancers preferentially metastasize to the omentum263. The exact signals that mediate this process are still subject to research and future efforts will be needed to dissect if the pro-metastatic signals come from a particular stromal cell population. In mice, metastatic cells are found in so-called “milky spots”, of the epigonadal WAT264, which are specialized niches of immune and mesothelial cells reminiscent of small lymph nodes143,265,266. Milky spots do not exist in the human omentum, but, in line with them, mesothelial cells rather locate on the periphery of the tissue267. Nevertheless, with the whole new resolution we brought to the cell composition of the stromal vascular fraction of the omentum, it would be relevant to question the role that mesothelial as well as IGFBP2+ cells play in the nesting of tumoral cells in the omentum. Mesothelial cells notably secrete IL-6 and IL-8 involved in gastric cancer metastasis268. TM4SF1, our validated pan-mesothelial surface marker, has been described as a bad prognosis marker in pancreatic ductal cancer, as it might facilitate metastasization of the tumor269. Consequently, current efforts focus on TM4SF1 as a potential new therapeutic target to prevent the invasion and metastasization of ovarian cancer270. Finally, IGFBP2 itself has been described as “the dark horse” in...
metabolism\textsuperscript{209} and constitutes a marker for tumor progression in gliomas, prostatic cancer and breast cancer. Among the proposed mechanisms by which IGFBP2 could influence tumor prognosis are the interaction with extracellular matrix (ECM) components, such as surface proteoglycan receptors and integrin receptors. By affecting ECM components, IGFBP2 would allow tumor cells to detach from their native matrix and migrate. In addition, IGFBP2 could also influence tumor progression by downregulating the expression of $\text{PTEN}$ or $\text{CDKN2A}$, which are known tumor-suppressing genes\textsuperscript{271} as well as preventing the translocation of p21 or PAPA1 to the\textsuperscript{72}.

It would be extremely insightful to study if the metastatic process would be altered in genetically modified mice that have their mesothelial cell function impaired either by KO of IGFBP2 or TM4SF1. It would also be interesting to use in situ techniques such as immunohistochemistry or spatial transcriptomics in human metastasis resections. Macrophages also seem to play a key role in facilitating tumor metastasis to the omentum\textsuperscript{273}, which could be influenced by the crosstalk with mesothelial cells, creating the perfect microenvironment for cancer proliferation. Therefore, mesothelial cells in adipose tissue open a whole new field for the identification of therapeutic approaches to improve the prognosis of such aggressive diseases.

Altogether we provided the first systematic and cross-anatomical functionalization of hASPC subpopulations proposed a new mechanism explaining why visceral ASPCs are reluctant to adipogenesis in vitro and discussed universal and unique characteristics of the adipose tissue in the function of its anatomic location and species of origin. Interfering locally with the IGFBP2 signaling in the visceral adipose tissue may contribute to promoting a healthy neoadipogenesis axed on hyperplasia instead of hyper-trophy and ease the inflammatory status linked to visceral obesity.
References


References


References


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2 months Internal medicine – HJB, St-Imier, Switzerland
1 month Infectious Diseases – Hôpital Tenon, Paris, France
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2018-2020 BIO-378 Travaux pratiques de physiologie, Prof. Pexieder – The electrocardiogram
2018-2020 BIO-377 Physiologie par systèmes, Prof. Roy
2015-2017 Physiscs, physiology and histology support lessons to first year medical school students (EtuPrep)
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2020 Super sandwich seminar, SV faculty seminar, EPFL, Talk
2021 IBI-EDBB Grad Student Mini-Symposia, EPFL, Talk
2019-2020 Joint UNIL-EPFL PhD retreat, Talk and Poster
2019 Summer school “Shaping the future of medicine”, Poster
2019 Stem cell retreat, Münsingen, Switzerland
2018-2022 MD-PhD retreat Unil, Poster and Talk (2022)

Prizes and grants
2021 Prize for the best poster, “An omentum specific mesothelial population inhibits adipogenesis”,
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Transplantation immunology – HUG, Geneva, Switzerland

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1 month
General surgery – Clinica Luganese, Laugano, Switzerland

1 month
Otolaryngology – HFR, Fribourg, Switzerland

2 months
Internal medicine – HJB, St-Imier, Switzerland

1 month
Infectious diseases – Hôpital Tenon, Paris, France

2 months
Emergencies – Hôpital Tenon, Paris, France

Other work experiences
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