Proteomic signatures of human visceral and subcutaneous adipocytes

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Declaration of Interests

The authors declare that no conflict of interest exists.
Abstract

Objective: Adipose tissue distribution is a key factor influencing metabolic health and risk in obesity-associated comorbidities. Here we aim to compare the proteomic profiles of mature adipocytes from different depots.

Methods: Abdominal subcutaneous (SA) and omental visceral adipocytes (VA) were isolated from paired AT biopsies obtained during bariatric surgery of 19 severely obese women (BMI > 30 kg/m²) and analysed using state-of-the-art mass spectrometry. Differential expression analysis and weighted gene co-expression network analysis (WGCNA) were performed to investigate proteome signature properties and to examine a possible association of the protein expression with the clinical data.

Results: We identified 3,686 protein groups and found 1,140 differentially expressed proteins (adj. p-value < 0.05), of which 576 proteins were upregulated in SA and 564 in VA samples. We provide a global protein profile of abdominal SA and omental VA, present the most differentially expressed pathways and processes distinguishing SA from VA, and correlate them with clinical and body composition data. We show that SA are significantly more active in processes linked to vesicular transport and secretion, and to increased lipid metabolism activity. Conversely, the expression of proteins involved in the mitochondrial energy metabolism and translational or biosynthetic activity is higher in VA.

Conclusion: Our analysis represents a valuable resource of protein expression profiles in abdominal SA and omental VA, highlighting key differences in their role in obesity.

Keywords: Adipose tissue; Visceral adipose tissue; Subcutaneous adipose tissue; Obesity; Adipocytes; Metabolism
Abbreviations:

AT – adipose tissue; VAT – visceral adipose tissue; SAT – subcutaneous adipose tissue; SA – subcutaneous adipocytes; VA – visceral adipocytes
1. Introduction

As the major adipose tissue cell type, adipocytes are essential for their physiological and pathophysiological processes, which include energy storage and release, metabolic homeostasis, and immune response regulation (1). They store energy in the form of triglycerides, predominantly in two anatomical depots: visceral and subcutaneous (2). Both sites share similar properties; however, the functional differences between subcutaneous adipose tissue (SAT) and visceral adipose tissue (VAT) with respect to insulin sensitivity, fatty acid composition, and bioactive molecule secretion are critical to health outcomes (3). Especially in the context of obesity, evidence supports a distinct pathological contribution related to the anatomical location of adipose tissue (4). Hence, SAT is generally considered metabolically less detrimental compared to VAT, which is strongly associated with the risk of cardiovascular disease, diabetes, cancer, and other comorbidities (5,6). However, detailed molecular features related to the functional differences of adipose depots remain obscure. In particular, little is known about alterations at the protein level.

Developments in proteomics technology provide us with a powerful tool for identifying the differences between the adipose tissue depots and enable us to reveal cell-specific expression patterns (Brockman and Chen, 2012). While the human adipose proteome has already been investigated, studies are significantly limited by intrinsic adipose tissue heterogeneity (7–10), lack comparison of different depots (11,12) or rely on a targeted approach limited to monitoring only selected proteins (13). To overcome these issues, we used isolated mature adipocytes from paired SAT and VAT biopsies of 19 morbidly obese female subjects (BMI ~ 40) and performed label-free comparative liquid chromatography-mass spectrometry (LC-MS) suitable for in-depth proteome analysis.

Our goal was to characterize the proteomic features of SAT and VAT adipocytes and their potential contribution to the function of adipose tissue in obesity. By employing the combination of the extreme phenotype of our cohort, which further amplifies differences between adipose locations,
and state-of-the-art proteomics, our study unveiled complex depot-specific profiles of isolated adipocytes. This is, to date, the most comprehensive proteomic analysis of adipocytes, newly highlighting the distinct role of each depot in obesity and thus advancing our understanding of adipocyte biology.

2. Methods

2.1 Patients

This study included 19 women (mean age = 47 ± 8 years, obesity grade II and III, BMI = 41.32 ± 4.23 kg/m²) who underwent bariatric/metabolic surgery at the Department of Surgery, Vitkovice Hospital, Ostrava. Patient selection followed the guidelines of the International Federation for the Surgery of Obesity (IFSO) (www.ifso.com/body-mass-index). All patients were white of Central European origin, 10 suffered from arterial hypertension, 5 from hyperlipidemia, 3 from type II diabetes, 4 from asthma. The study originally included 20 women and 4 men; however, the male samples were under-represented and thus excluded from the study – data available at the Pride repository. One female patient was excluded due to the outlying data pattern of visceral adipocyte protein intensities.

Anthropometric assessment (body weight, height, waist, and hip circumference) was performed prior to surgery. DXA measurements were performed to determine body composition and body fat distribution using a Hologic densitometer (Discovery A; Hologic, Waltham, MA, USA) calibrated according to the manufacturer’s recommendations. The precision error of the densitometer and coefficient of variation were established and the value of the least significant change was calculated as described in previous studies (14–16). Fasting blood glucose, cholesterol, triglycerides, HDL, LDL, and HbA1c concentrations were also measured prior to surgery. Subjects characteristics are shown in Table 1.
All participants signed written informed consent before participation in the study and the design of the study was approved by the Multicentric Ethics Committee of Vitkovice Hospital, Ostrava (No. EK/3/17) according to the principles of the Declaration of Helsinki.

### 2.2 Adipocyte isolation and culture

SAT was obtained from the subcutaneous abdominal area between the xiphoid process and umbilicus on the left side of the patient. The incision was then used as a laparoscopic port and all procedures were performed laparoscopically in the standard manner as described in previous studies (17,18). VAT was taken from the front part of the omentum majus. Utilized bariatric procedures included e.g. laparoscopic greater curvature plication (LGCP), laparoscopic sleeve gastrectomy (LSG) and Roux-en-Y gastric bypass (RYGB). The type of surgery did not influence the way of AT biopsy.

Obtained biopsies were placed into a tube with 20 ml of Dulbecco’s Modified Eagle’s Medium/Nutrient Mixture F-12 Ham (DMEM/F12, Sigma-Aldrich, Cat# D6421) and transferred at room temperature for adipocyte isolation. The following procedures were carried out under a laminar flow hood using sterile equipment. Adipocytes were isolated using a previously described method (19), briefly described below: Adipose tissue was washed three times with sterile PBS and visible blood veins were removed from samples. After mincing into approximately 10 mg fragments using scissors, tissue was digested with Standard Grade NB 4 Collagenase (Serva, Cat# 1745401), 1 mg/ml in DMEM/F12 for 2 hours at 37 °C with shaking (100 rpm). The enzymatic reaction was stopped by adding DMEM/F12 supplemented with 10 % foetal bovine serum (FBS, Biosera, Cat# 1001). The mixture was filtered through a 500 μm plastic mesh and centrifuged at 300×g for 5 minutes. This step was repeated twice. Floating adipocytes were then collected, washed twice in DMEM/F12 and seeded in a T75 flask in DMEM/F12 supplemented with 10% FBS. The cells were allowed to recover overnight before they were washed three times in PBS and frozen at -80 °C.
2.3 Protein extraction and on-filter digestion

All adipocyte samples were completely thawed and vortexed with an SDT lysis buffer (4% Sodium dodecyl sulphate (SDS; Sigma-Aldrich, Cat# 436143), 0.1 M Dithiothreitol (DTT; Thermo Fisher Scientific, Cat# R0862), 0.1 M Tris/HCl (Sigma-Aldrich, Cat# 10812846001), pH = 7.6) in a Thermomixer® (Eppendorf) for one minute. The mixtures were subsequently heated at 95 °C for 30 min while mixing at high speed. The supernatant was cleared by centrifugation (20,000×g, 15 min) and the solution below the lipid layer was collected for further processing. Protein extract quality and its concentration were assessed by 1D SDS-PAGE (20) to ensure comparable conditions for all samples during filter-aided sample preparation (FASP) (21,22). Protein lysates (50 µg) were mixed with 8 M UA buffer (8 M urea (Sigma-Aldrich, Cat# U1250) in 0.1 M Tris/HCl (Sigma-Aldrich, Cat# 10812846001), pH 8.5) on Microcon-30kDa cut-off filters (Merck Millipore, Cat# MRCF0R030) and centrifuged at 14,000×g for 30 minutes at 20 °C. The retained proteins were washed with 200 µl UA buffer. The final protein precipitate in the Microcon devices was mixed with 100 µl of UA buffer containing 50 mM iodoacetamide (Sigma-Aldrich, Cat# I1149) and incubated in the dark for 20 minutes at room temperature. Subsequently, the samples were washed three times with 100 µl UA buffer and three times with 100 µl of 50 mM NH₄HCO₃ (Sigma-Aldrich, Cat# A6141). Finally, trypsin (enzyme:protein ratio of 1:50; sequencing grade, Promega, Cat# V5111) was added onto the filter and the mixture was incubated for 18 hours at 37 °C. The tryptic peptides were then eluted by centrifugation followed by two additional elution steps with 50 µl of 50 mM NH₄HCO₃. Peptides were then cleaned by liquid-liquid extraction (3 iterations) using water saturated ethyl acetate (23). Cleared FASP eluates were evaporated completely in a SpeedVac concentrator (Thermo Fisher Scientific). The resulting peptides were extracted into LC-MS vials by 2.5 % formic acid (FA) in 50% acetonitrile (ACN) and 100% ACN with an addition of polyethylene glycol (20,000; final concentration
0.001 %) (24) and concentrated in a SpeedVac concentrator (Thermo Fisher Scientific). The samples were processed in two batches (12 sample pairs per batch).

2.4 Liquid chromatography-mass spectrometry (LC-MS/MS)

Peptide mixtures were measured using an LC-MS/MS system consisting of an Ultimate 3000 RSLCnano system (SRD-3400, NCS-3500RS CAP, WPS-3000 TPL RS; Thermo Fisher Scientific) combined with an Orbitrap Fusion Lumos system (Thermo Fisher Scientific) with Digital PicoView 550 nanospray ion source. Tryptic digests (~2 µg/injection) were online concentrated and desalted on a trapping column (100 µm × 30 mm, filled with 3.5 µm X-Bridge BEH 130 C18 sorbent (Waters, Milford)) using 0.1% formic acid (FA) in water. The peptides were eluted from the trapping column onto an Acclaim Pepmap100 C18 analytical column (3 µm particles, 75 µm × 500 mm; Thermo Fisher Scientific, Cat# 164570). Peptides were separated using a 120-minute gradient (mobile phase A – 0.1% FA in water; mobile phase B – 0.1 % FA in 80% (v/v) acetonitrile. Peptides were eluted with a linear 75 min gradient of 1–30 % of mobile phase B, followed by a 30 min increase to 56 % of mobile phase B and a 5 min increase to 80 % of mobile phase B, and a 10 min wash of 80 % mobile phase B at a flow rate of 300 nL/min.

MS data were acquired using data-dependent acquisition. Target values for full scan MS spectra were 4 × 10⁵ charges in m/z range 350–2,000 with a maximum injection time of 54 ms and a resolution of 60,000 at m/z 200. MS/MS scans after HCD fragmentation with 30% collision energy were performed at a resolution of 30,000 at m/z 200 with an ion target value of 5 × 10⁴ and a maximum injection time of 50 ms. All peptide mixtures were analysed separately in a randomized manner; VA and SA samples from each patient were analysed together.
2.5 MS data analysis

MS raw data files were analysed using MaxQuant software (v. 1.6.1.0) (25) using default settings unless otherwise noted. MS/MS ion searches were performed against the UniProtKB Human FASTA database (Homo sapiens, taxon ID 9606, version 2018-05 from 28.05.2018) and cRAP contaminant database (version 170518; thegpm.org/crap) using the Andromeda search engine (Cox et al., 2011). Carbamidomethylation on cysteine was set as a fixed modification and N-terminal acetylation, methionine oxidation, and asparagine/glutamine deamination as variable modifications. Trypsin/P enzyme with 2 allowed missed cleavages and minimal peptide length of 6 amino acids were set. The false discovery rate (FDR) was 0.01 for both protein and peptide levels. The match-between-runs algorithm was used across the whole dataset to improve peptide matching. The mass spectrometry proteomics data and the search results have been deposited to the ProteomeXchange Consortium via the PRIDE (26) partner repository with the dataset identifier PXD024734.

2.6 Statistical and bioinformatics analyses

Protein intensities reported in the proteinGroups.txt file (output of MaxQuant) were further processed using the software container environment (https://github.com/OmicsWorkflows) version 3.7.2a and R software (27). Processing workflow is available upon request. Briefly, it included the following: a) removal of decoy hits and contaminant protein groups; b) exclusion of 4 male sample pairs and an outlying sample pair; b) protein group intensities log₂ transformation, c) LoessF normalization, and d) missing values imputation using the imp4p package. Protein groups were only considered for subsequent analysis if they contained measured intensity value in at least 8 samples of VA or SA and were identified to at least 2 peptides. The imputed normalized protein intensities (S File 1 (28)) were used for differential expression analyses using the LIMMA R package (29). The linear
model used to compare paired differences between SA and VA samples was adjusted for batch effect by adding batch number as a variable in the model. The correlation between sample pairs was included in the linear model using appropriate functions from the LIMMA package (30). Subsequently, the results were adjusted for multiple hypothesis testing using the Benjamini and Hochberg procedure (31) implemented in the LIMMA package. Limma test results are provided in Supplementary File 2 (28).

Global correlation analysis was performed on normalised imputed log2-transformed intensity data. Two separate correlation matrices were constructed, i.e. for SA and VA intensities. The Pearson correlation coefficient was used to calculate the correlations. The correlations were then subtracted from 1, resulting in values in the 0 to 2 range. These values were then used as distances in unsupervised hierarchical clustering using Ward’s algorithm (32). The resulting correlation matrices present the correlations ordered into clusters obtained using this method.

The relationship between SA and VA protein expression and clinical traits was assessed using a Weighted Gene Co-expression Network Analysis (WGCNA) R package (33,34). The log2-transformed normalised and imputed intensity data was used to construct the biweighted mid-correlation signed co-expression network. The adjacency matrix satisfying the approximate scale-free topology network was used to calculate Topological Overlap Measure, and the corresponding dissimilarity was used to produce a hierarchical clustering tree of proteins. Modules of highly interconnected proteins were defined using the Dynamic Tree Cut of the dendrogram branches, and subsequent merging of highly similar modules. Each module was then assigned a designated colour. For each module, an eigengene was defined as the first principal component of the module, and module membership (MM) was determined by calculating the Pearson correlation between each module eigengene and each protein, together with corresponding p-values (35). Module-trait relationships were calculated as Pearson correlations between each module eigengene and clinical traits. Modules with a significant trait correlation p-value < 0.05 and absolute correlation value > 0.5 were selected for STRING(36).
functional enrichment analysis. Hub proteins with the highest MM and proteins with the most significant trait correlation based on gene significance (GS) values within the selected modules were further investigated in the context of AT and obesity.

Gene ontology (GO) enrichment analyses were performed in Cytoscape 3.8.0 using ClueGO (v. 2.5.7) and the CluePedia (v. 1.5.7) plugin (37) showing only terms where the p-value < 0.05. The enrichment of biological processes, molecular function and the cellular compartment for the whole adipocyte proteome was set to show the top 2 levels of GO terms. Default search parameters were used for the enrichment of significant proteins. The pathway analyses were performed using the open-source peer-reviewed pathway database Reactome (38) with subsequent pathway visualization using ClueGo + CluePedia including the import of STRING database scores (36) into the network. This approach facilitated the visualization of complex pathway networks with protein-protein interactions among the most differentially expressed proteins between SA and VA.

The prediction of putative secreted proteins was conducted using the SignalP-5.0 Server, enabling the prediction of the secretory signal peptide sequence for transport by the Sec translocon and cleaved by Signal Peptidase I using a deep neural network (Almagro Armenteros et al., 2019). This analysis was performed using the FASTA sequence of the most differentially expressed proteins with log₂FC > 1.

3. Results

3.1 Overview of global adipocyte proteome

In this study, we used mature adipocytes derived from abdominal SAT and omental VAT biopsies collected from 19 obese female patients during bariatric surgery. Adipocytes from two distinct adipose tissue depots were used for label-free quantitative LC-MS profiling. We generated a dataset of 38 adipocyte proteomes with a total of 3,686 identified proteins; we provide this proteome dataset in an easily accessible and minable format for the benefit of the entire community (PRIDE accession PXD024734). In order to minimize the chance of false positivity, we excluded protein
groups identified only by a single peptide match, and protein groups quantified in less than 8 of the 19 SA or VA samples for subsequent analysis. The final dataset with 2,939 quantified protein groups was used to compare measured intensities in SA and VA samples (S File 1 (28)). The whole dataset was comparable in the means of identified proteins between the SA and VA. The variability between the replicates was minimal with Pearson correlation coefficients between measured intensities of [0.81 – 0.93] for not normalized log2 transformed data, and [0.83 – 0.94] for loess normalized log2 transformed data (data not shown).

First, we annotated the whole proteome with gene ontology (GO) terms and performed an enrichment analysis to describe the proteome structure, and subsequently compared the functional distribution of differentially expressed proteins. The overall adipocytes proteome enrichment analysis showed the distribution of all proteins within the adipocytes' cellular components (CC), biological processes (BP), and molecular function (MF) (Figure 1). The most enriched GO CC terms included cytoplasm, cytosol, extracellular space, and the endomembrane system, as well as inner cell complexes like the mitochondrial protein complex, actin filament bundle, and ribonucleoprotein complex. The most enriched biological processes in our adipocyte proteome are the establishment of localization, small molecule metabolic process, oxidation-reduction process, cellular metabolic process, and cellular component organization or biogenesis. Among others, the ATP metabolic process is also significantly enriched with over 55% of associated genes for a given term. The proteome enrichment analysis based on the molecular function shows mainly catalytic activity, protein binding, small molecule binding, followed by proteins involved in hydrolase activity, oxidoreductase activity, structural molecule activity, cofactor binding, or lipid binding.
3.2 Global adipocyte proteome correlation maps reveal cell-specific functional expression patterns

Next, we hypothesized that proteins are present in functionally related networks with a close relation to protein abundance levels. Therefore, we used a global correlation map with a pairwise relationship of all 2,939 proteins. The cross-correlation of all measured protein intensities generated a matrix of 8,637,721 Pearson’s correlation coefficients which were clustered using unsupervised hierarchical clustering into the global correlation map for SA and VA samples respectively. The global correlation maps show distinct expression patterns between SA and VA proteomes. Therefore, the apparent protein clusters were submitted to GO enrichment analysis, and we revealed several functionally related protein clusters in both adipocyte proteomes (Figure 2a, b). Although most of the cellular processes and compartments formed clusters in both proteomes, some clusters were larger with a stronger linear relationship in one of the proteomes than in the other. The SA proteome creates highly correlated clusters enriched especially in peptide metabolic and biosynthesis processes, transport, the establishment of protein localization, extracellular exosome, and vesicular transport. On the other hand, the VA proteome creates a large cluster of strong correlations enriched by proteins of cellular respiration, generation of precursor metabolites and energy, and mitochondrion and clusters involved in lipid metabolic process, as well as protein transport and localization, and peptide metabolic processes.
3.3 Bioinformatic analyses of differentially expressed proteins reveal adipocytes’ functional differences

To reveal differences between SA and VA protein expression, we compared both adipocyte cell lines at protein abundance level. A comparison of the measured protein MS intensity values between VA and SA proteomes revealed 1,140 differentially expressed proteins (adj. p-value < 0.05) (S file 2 (28)). Where 576 proteins were upregulated in SA, and 564 in VA samples, respectively. To investigate what biological processes, molecular functions, and cellular components are enriched by the SA and VA upregulated proteins, all significantly upregulated proteins were submitted to ClueGO (v.2.5.7) with a significance threshold of 0.05 and merging all redundant groups with > 50% overlap. All proteins were annotated based on current knowledge of protein domain molecular-level activities, locations relative to cellular structures, or participation in biological processes, to reveal the major biological differences between SA and VA.

Based on the cellular component enrichment analysis, most of the upregulated proteins in SA are associated with membrane-related terms such as organelle membrane, vesicle, whole membrane, bounding membrane of organelle, etc. In contrast to SA, VA upregulated proteins are enriched mainly by proteins localized to the cytosol, extracellular vesicle, mitochondrion, and mitochondrial respiratory chain complex I (Figure 3a).

The enrichment analysis of biological processes (Figure 3b) displayed most of the SA upregulated proteins in the cellular component organization, the establishment of localization in cell, vesicle-mediated transport, response to organic substance, or cellular catabolic process. Generally, the most significantly enriched groups were related to transport activities like vesicle-mediated transport, the establishment of localization in cell or Golgi vesicle transport, but also to metabolic processes associated with lipid metabolism and storage, cellular catabolic processes or carbohydrate derivative metabolic processes. On the other hand, the most VA upregulated proteins were associated with the
organonitrogen compound metabolic process, organelle organization, organic substance catabolic process, cellular catabolic process, or organic acid metabolic process. Taken together all enriched groups in VA, the most significant enrichment was observed in metabolic processes especially in relation to the cellular catabolism including macromolecules and small molecules. Moreover, a significant enrichment was observed in terms related to mitochondrial energy metabolism such as oxidoreductase activity, mitochondrial respiratory chain complex I assembly, electron transport or generation of precursor metabolites and energy.

In terms of molecular function enrichment analysis (Figure 3c) SA upregulated proteins showed only a few significantly enriched functional groups in comparison to VA. Here, most of the SA upregulated proteins act in cytoskeletal protein binding, cell adhesion molecule binding, purine ribonucleotide binding or coenzyme binding. In contrast, the upregulated proteins of VA are functionally enriched especially in oxidoreductase activity, coenzyme binding, nucleotide binding or RNA binding.

**3.4 Pathway enrichment analysis reveals the most profound SA and VA differences**

In order to explore processes and pathways enriched by the upregulated proteins in SA and VA respectively, we analysed the lists of upregulated proteins using the Reactome database. The Reactome database allows over-representation pathway analysis of curated pathways and reactions in human biology. Reactome pathways are arranged in a hierarchy and the over-represented pathways score is corrected for false discovery rate using the Benjamani-Hochberg method. This resulted in 115 significantly over-represented terms in SA and 96 in VA respectively (S File 3a, b (28)). Since the results showed significant over-representation at multiple levels within Reactome event hierarchy, we compared both adipocyte types at the root terms and subsequently focused on the most significantly enriched pathways relevant to the adipocytes or AT physiology/pathophysiology.
The over-representation analysis showed the most pronounced differences between both adipocyte types, where the upregulated proteins of SA are involved in wider spectra of cell processes in comparison to VA. SA are enriched especially with pathways and processes involved in extracellular matrix organization (FDR: 8.84E-04), vesicle-mediated transport (FDR: 7.44E-03), protein localization (FDR: 3.90E-02), or programmed cell death (FDR: 4.33E-02). Less significant over-representation was also observed for processes and pathways in the metabolism (FDR: 1.08E-01), small molecule transport (FDR: 1.54E-01), and the protein metabolism (FDR: 1.72E-01) (Figure 4a). In contrast to SA, upregulated proteins in VA are significantly over-represented mainly in the metabolism (FDR: 3.56E-14) and cellular response to external stimuli (FDR: 3.10E-06) (Figure 4b).

In order to select the most relevant Reactome pathways and reactions, both lists of significantly upregulated proteins were analysed together in one analysis using Cytoscape ClueGO, showing only significant results and merging redundant groups with > 50% overlap (S file 4a (28)), but also separately for each adipocyte type (S file 4b, c (28)). Proteins often operate as a protein complex or interacting partners in complex networks. The identification of these protein networks and interactions is essential for understanding the functional properties of SA and VA proteins; however, not all interactions may have been recognized by the ontology enrichment analysis itself. Therefore, to extend the analysis of Reactome pathways and reactions, we also performed an interaction analysis using the STRING database scores. This created a large protein-protein interaction network combined with the most significantly enriched pathways by the upregulated and downregulated proteins of SA, and vice versa for VA (Figure 5). Some enriched groups of associated pathways, processes and interactions share proteins of both groups; however, in order to simplify the complex network, we evaluated the pathways and processes separately for each adipocyte type.
3.4.1 SA pathways – vesicular transport and lipid metabolism

SA upregulated proteins enrichment analysis showed associations with chylomicron assembly and remodelling, together with other processes related to apolipoproteins, metabolism of fat-soluble vitamins, especially retinoid metabolism and transport, and ligand binding and uptake by scavenger receptors. Other metabolism-related processes include peroxisomal lipid metabolism with beta-oxidation of very long chain fatty acids. Another significantly enriched term was glycosphingolipid metabolism.

The enrichment of collagen proteins with several integrins and laminins implies ECM interactions. Further, the integrins were linked to proteins involved in caldesmon calcium binding process, which was also enriched by upregulated tropomyosin responsible for the regulation of glucose uptake.

Further, we observed proteins associated with proteasome mediated degradation, Hedgehog (Hh) ligand biogenesis and ERAD degradation. One of the largest SA upregulated protein clusters was formed by proteins of RAS-like family of small GTPases and other proteins involved in vesicle mediated transport. The most enriched processes by these proteins were RAB geranylgeranylation, and RAB8A, 10, 13, 14 exchange GDP for GTP, together with insulin regulated trafficking by GLUT4 and vesicle translocation and docking at the plasma membrane. Other processes related to this group of proteins include COPI vesicle uncoating at the ER, dissociation of cis-SNARE at the ER by the NSF ATPase activity, or proteins involved in antigen peptide transport and other transport processes.
3.4.2 VA pathways – energy metabolism and protein synthesis activity

Among the VA pathways we observed protein clusters involved especially in the energy metabolism. The most significant enrichment was observed in the case of the protein cluster forming mitochondrial complex I. The complex I was also linked to proteins of the pyruvate metabolism and citric acid cycle. The enrichment analysis also shows proteins of glucose metabolism, glycolysis or gluconeogenesis, and other energy metabolism related terms. In the context of the significant enrichment of processes associated with mitochondrial metabolism, we also observed proteins associated with glutathione conjugation. Another, and one of the largest, protein cluster displays VA upregulated proteins involved in ribosomal assembly, translation initiation and translation. This process is also connected to the group of protein phosphatases, which are associated with a regulation of translational activity by nonsense-mediated decay. Along with the elevated expression of translational apparatus proteins, we observed clusters of proteins involved in protein folding, activation and stabilization like chaperonins and heat shock proteins in ATP-dependent manner. Further analysis showed several upregulated signalling proteins in VA. The network is centred around MAPK3, which is linked to several other signalling proteins such as AKT2, SRC, STAT5A, STAT5B, GNAI1, GNB1, and GNB4. Correspondingly, this was accompanied by the upregulation of several phosphatases, plausibly regulating the kinases in a negative feedback loop.
3.5 The most differentially expressed proteins

The top 20 most upregulated proteins from SA and VA are shown in Table 2 and Figure 6a. We also ranked all proteins by mean intensity values to reveal the position of the most differentially expressed proteins within the entire proteome. The most profound differences are observable within the least abundant proteins (Figure 6b). However, the majority of the most differentially expressed proteins by Log₂FC – especially GLIPR2, and ASPN, but also EZR, ALOX15, OGN, GPD1L, SLC38A10, PEOLO, CORO2B, and ANXA8L2 – are among the more abundantly expressed proteins, which might suggest a major physiological role in distinguishing between SA and VA. Additionally, we plotted the intensities of the top 20 most differentially expressed proteins by Log₂FC, distinguishing experimental and missing imputed values in individual samples (Figure 6c). This revealed several proteins potentially expressed exclusively either in SA (ALPL, CLIC6, TLCD2) or in VA (AGPAT9, ANXA8L2, PPAD1).

3.6 Prediction of putative secreted proteins

AT is an important endocrine organ which secretes several signalling molecules such as adipokines, which may contribute to the pathogenesis of obesity comorbidities. We submitted FASTA protein sequences of significant proteins with Log₂FC > 1 to the SignalP-5.0 Server. This server enables the prediction of the presence of secretory signal peptides transported by the Sec translocon and cleaved by Signal Peptidase I using deep neural networks (39). This analysis predicted 46 secreted proteins out of 175 SA upregulated proteins but only 20 among the 167 upregulated proteins in VA (S File 5 (28)). The 20 most differentially expressed SA proteins included 7 potentially secreted proteins and therefore possibly detectable in peripheral blood, namely ALPL, NOTCH3, FKBP9, GLB1, SIL1, RCN3,
CD248. Similarly to SA, we identified 6 proteins out of the 20 most upregulated proteins in VA with a prediction to be secreted, i.e. OGN, ASPN, LUM, APCS, PPRELP, and BGN.

We also performed an over-representation analysis of a predicted secreted proteins combined list using a combination of STRING together with Reactome database enrichment, and ClueGO biological processes enrichment analysis. This analysis revealed several interacting protein groups mainly among SA proteins. The largest enriched group involved in chylomicron assembly, triglyceride-rich lipoprotein particle remodelling, cholesterol transfer activity, regulation of cytokine production in immune response and other related processes contained upregulated SA proteins APOA1, APOA2, APOB, APOE, ADAM10, CLU, DAG1, RCN, and TINAGL1 and the upregulated VA protein APOA4. A significant association was also observed for SA upregulated NENF and NUCB2 which were associated with negative appetite regulation. Additional significantly enriched terms worth mentioning included dermatan sulphate (VA: BGN, DCN) and keratan sulphate biosynthetic process (SA: GNS, GLB1; VA: LUM, OGN, PRELP).

3.7 Association of clinical traits with the protein expression

To investigate the association of clinical traits with the SA and VA proteome signatures, we adapted the WGCNA package in R which facilitates the identification of highly similar expression patterns and the subsequent correlation of these patterns with clinical data. Using the WGCNA co-expression network analysis, we obtained 11 and 16 modules of highly correlated proteins for SA and VA respectively (S File 6 (28)). The modules were assigned colours independently for the SA and VA proteomes. The analysis of VA also contained a grey module of proteins which were not assigned to any module. The module eigengenes (ME) were then correlated with the clinical data to identify modules with significant associations for SA (Figure 7) and VA (Figure 8) respectively. The functional enrichment of significantly correlated modules was performed using the STRING database and GS
was used to measure the degree of association between the traits and individual proteins (SA – S File 7, VA – S File 8 (28)).

SA module-trait correlation analysis showed that several modules were significantly associated with the clinical traits (Figure 7). A positive correlation of waist circumference measured at the inferior margin of the ribs (waste1) was observed for the SA Purple and Salmon module, and a negative correlation was associated with the Red module. The SA Purple module (n = 76) eigengenes also correlated with the hip circumference, BMI, and DXA lean mass index. Although the Purple module was enriched by proteins involved in immune response and proteins associated with a response to infectious disease, the subsets of proteins significantly associated with the traits were enriched mainly by proteins of extracellular space and extracellular matrix disassembly, e.g. CD44, CTSL, LCP1 or CAPN2. The SA Salmon module (n = 38) enriched by proteins in the RNA binding process was also positively correlated to the DXA visceral fat mass. The Red module (n = 200) associated with the negative correlation was enriched by proteins involved in vesicle-mediated transport and in the transport of small molecules, along with G protein signalling pathways. In addition to the Purple module, hip circumference and DXA lean mass index also correlated with the SA Green-yellow module (n = 75) enriched by proteins involved in the RNA metabolism, and the SA Black module (n = 177) enriched by proteins involved in immune system processes and stress response. The SA Black module was also associated with a positive correlation with HDL levels. A negative correlation with HDL levels and conversely a positive correlation with glycemia was linked to the SA Turquoise module (n = 719) enriched by proteins involved in vesicular transport and the protein metabolism. The subsets of significantly associated proteins with HDL levels and glycemia were associated especially with proteins involved in proteasomal protein catabolic process and regulation of G2/M transition of the mitotic cell cycle. Likewise, a positive correlation to glycemia and cholesterol levels was observed for the SA Brown module (n = 406) enriched by proteins involved in the translation and metabolism of proteins. Conversely, a negative correlation was observed with the SA Green module (n = 541) enriched significantly by mitochondrial proteins and proteins involved in the citric acid and
respiratory electron transport. The most significant positive correlation of SA proteins with glycemia was observed for FAM129B (GS = 0.82), PIN1 (GS = 0.81). Conversely, the most significant negative correlation was observed for PXN (GS = 0.80), or HIBADH (GS = -0.80) accompanied by other dehydrogenases, e.g. ACAD11 (GS = -0.79), ACADM (GS = -0.74), or HSDL2 (GS = -0.74) involved in fatty acid beta-oxidation. An inverse correlation to cholesterol level was observed for the SA Blue module (n = 433), enriched by proteins of the endomembrane system, especially proteins of the endoplasmic reticulum, and the Pink module (n = 163) enriched by ribosomal proteins and proteins involved in intracellular transport and localization. The overall most significant correlation to cholesterol was associated with the RAB22A (GS = -0.84), LGALS12 (GS = -0.78) and STX-8 (GS = -0.74).

The VA module-trait correlation analysis (Figure 8) showed different correlation patterns compared to the SA proteome. A positive correlation with waist measured at the umbilical level and total body fat was associated with the VA Greenyellow module (n = 124) enriched by organelle organization and translational initiation. A negative correlation with both waist circumferences, but also with DXA lean mass and DXA visceral fat mass, was associated with the VA Turquoise module (n = 600). The VA Turquoise module is enriched largely by processes in mitochondria, such as the citric acid cycle and respiratory electron transport or fatty acid metabolism. The most significant association with the DXA visceral fat mass was observed for NADH dehydrogenases, especially NDUFB8 (GS = -0.82) and NDUFB4 (GS = -0.79). A negative correlation with the DXA visceral fat parameters was also observed for the VA Grey60 (n = 42) module that is enriched by proteins in carboxylic acid and small molecule metabolic processes. Conversely, a positive correlation was associated with the VA Pink (n = 159) and Red module (n = 194). The VA Pink module is enriched by proteins in actin cytoskeleton organization and metabolism of RNA, whereas the VA Red module is enriched by proteins involved in vesicle-mediated transport and exocytosis. A positive correlation was also observed for cholesterol and the VA Blue, Green, and Cyan modules. The VA Blue module (n = 516) is enriched by proteins involved in the cellular metabolic processes, whereas, the VA Green module (n = 200) is enriched by proteins
involved in antigen processing and the presentation of peptide antigens, and proteasome degradation. The VA Cyan module (n = 60), which is also associated with the waist1 to hip ratio, is enriched by cytosolic proteins. Similar to the Cyan module but with a negative correlation, the VA Salmon module (n = 80) enriched by proteins in oxidative phosphorylation was associated with cholesterol but also with LDL levels. Here, the most significant association with cholesterol was observed for RAB43 (GS = -0.82) and ECHDC3 (GS = -0.81). Also, a negative correlation to cholesterol and LDL levels was observed for the VA Brown module (n = 258). This module is significantly enriched by proteins involved in the vesicle-mediated transport and lipid metabolism. The most significant negative correlation with cholesterol (GS = -0.81) and LDL (GS = -0.82) was observed for ANTXR2.

4. Discussion

In our study, we focused on the comparison of protein expression profiles from mature adipocytes isolated from abdominal SAT and omental VAT collected from severely obese women in order to explore major depot-specific differences. We identified 3,686 proteins groups and found 1,140 differentially expressed proteins (adj. p-value < 0.05), of which 576 proteins were upregulated in SA and 564 in VA samples. Our analysis revealed cell-specific protein expression patterns of coordinated processes in both adipocyte proteomes. Most of the protein clusters observed in the global correlation mapping contained functionally or mechanistically related protein clusters also observed in the enrichment analysis of differentially expressed proteins. Our study demonstrated that SA are more active in processes related to vesicular transport and secretion, together with increased lipid metabolism activity. Conversely, the expression of proteins involved in translational or biosynthetic activity and energetic metabolism was higher in VA. Moreover, the functional differences were further highlighted by distinct correlations with the clinical and body composition data.

Oxidative phosphorylation is essential in the metabolism by supplying ATP for cellular processes. Our results indicate that oxidative phosphorylation, and processes related to mitochondria and the
mitochondria energy metabolism, represent one of the most pronounced differences between SA and VA. The global correlation cluster mapping of the VA proteome showed more balanced and significantly higher correlations of the mitochondrial proteins and proteins involved in mitochondrial processes in comparison to SA. The pathway analysis of differentially expressed proteins then indicated that especially the proteins of mitochondrial electron transport chain Complex I, pyruvate metabolism, and citric acid cycle are more abundantly expressed in VA. Our results are in agreement with several lines of evidence indicating that the visceral compartment is generally characterized by a relatively higher mitochondrial oxidative metabolism (40,41). This could be further underlined by obesity-induced changes that differentially affect mitochondria in SA and VA. More specifically, previous studies showed a negative correlation between oxidative phosphorylation capacity and BMI, and between electron transfer system capacity and BMI in SA but not in VA (42,43). Since our analysis investigated the proteome of adipocytes derived from the extreme phenotype of body composition, we can hypothesize that observed differences might to some extent serve as an indicator of SA mitochondria impairment, which could in turn contribute to the enhanced energetic and metabolic activity of VA. Further, underlining the relevance of bioenergetics metabolism, number of proteins involved in mitochondrial regulation showed significant negative correlation with clinical and body composition parameters including glycemia, triglycerides level, waist to hip ratio, LDL, and cholesterol. Notably, the systematic downregulation of OXPHOS capacity and mitochondrial reorganization as a general alteration of adipocytes in obesity was also recently reported in the diet-induced obesity murine model, affecting both depots, with larger changes observed in SA (44).

In addition to OXPHOS, anaerobic glycolysis also contributes to ATP synthesis; previous findings suggest that in adipocytes it is in fact a more relevant energy source in adipocytes (45). In this study, we observed a group of proteins involved in glycolysis to be more abundantly expressed in VA (e.g. ALDOA, ENO1, GPI, HK2, PFKL, PFKM, PFKP, PGAM1, PGK1, PPP2CA), which might reflect the overall higher energy demand of VA, as well as a partial compensatory mechanism for overloaded mitochondria. This is also in agreement with previous findings that the OXPHOS capacity of VA
adipocytes remained unaffected with higher BMI, while glycolysis increased (43). The upregulated expression of glycolytic pathway proteins might also be the consequence of inferior capillary density and limited angiogenic growth capacity of visceral tissue, potentially leading to local hypoxia and a shift towards an anaerobic metabolism in less vascularized areas (46). Notably the glycolytic pathway yields intermediates which are further utilized in aerobic cellular bioenergetics, which also appeared to be upregulated in VA in our study. This includes enzymes directly involved in TCA and the pyruvate metabolism (e.g. ACO2, ADHFE1, GLO1, HAGH, L2HGDH, LDHA, LDHB, ME1, PDPR, SUCLA2, SUCLG1), as well as in the above-mentioned upregulation of several Complex I proteins. Furthermore, in VA we revealed a higher abundance of ATP-citrate lyase (ACLY), the enzyme responsible for the cytosolic synthesis of acetyl-CoA from citrate, and higher protein expression of other lipogenic enzymes including ACACA, ACACB, FASN, and HACD3. Altogether, our results support the notion that glycolytic and lipogenic processes are augmented in VA. This is further clarified by the fact that several enzymes involved in the regulation of these processes are amongst the most expressed proteins of VA (e.g. AGPAT9, GPD1L). This is in agreement with previous reports of increased glucose uptake rates in VA, particularly in obese individuals, where VAT acts as an essential glucose sink (47).

Metabolic activity requires newly synthesized proteins to be complemented by molecular chaperones which assist their folding into native three-dimensional states. This process should be further accompanied by intracellular trafficking in order to deliver the synthesized proteins into different organelles. The potential demand for novel proteins in the VA is supported by the overexpression of protein groups including multiple chaperones, ribosomal proteins and elongation factors, and dynactin complex subunits. This correlates with previous studies reporting the superior upregulation of genes involved in protein synthesis in the visceral compartment (41). Moreover, the increase of several members of the COP9 (Constitutive photomorphogenesis 9) signalosome involved in the regulation of ubiquitin-proteasome pathway, along with the ESCRT (endosomal sorting complexes required for transport) complex, also argues for tightly orchestrated proteostasis in VA. In comparison, in SA we observed an overrepresentation of components of 26S proteasome machinery.
(e.g. PSMB8, PSMC4, PSMC5, PSMD11, PSMD12, PSMD13, PSMD2, PSMD4, PSMD7), and proteins involved in the endoplasmic reticulum-associated degradation (ERAD) pathway (e.g. DERL2, ERLEC1, ERLIN1, RNF185, SEL1). Further, we revealed the upregulation of the Calnexin/Calreticulin chaperone system dedicated to N-glycosylated proteins folding, which, in line with previous studies, also suggests enhanced protein traffic along the secretory pathway in SA (48,49). This is also well illustrated by the upregulation of SIL1 and RCN3, involved in secretory trafficking, which are among the most overexpressed SA proteins. This system plays a vital role in Ca2+ homeostasis and ensures that assembled glycoproteins leave the endoplasmic reticulum, or orchestrate their degradation via ERAD, when correct protein folding is unachievable (50). Importantly, the expression of several members of this pathway is elevated as a result of ER stress and is a part of unfolded protein response. For instance, UGGT – significantly overrepresented in SA in our study – plays the role of a checkpoint allowing misfolded proteins to rebind to the chaperones, thus preventing their exit from the ER (50).

This study also clarifies depot-specific differences between numerous modes of vesicular trafficking systems. Trafficking between subcellular compartments involves transport containers whose formation is orchestrated by multiple conserved coat complexes. These encompass the coat protein complexes (COPI and II) involved in endoplasmic reticulum export, clathrin and caveolin coat families involved in the subsequent steps defined by the exocytic and endocytic pathways (51). Additionally, cargo capture is necessarily connected to cellular components that direct the transport of vesicles to their unique destinations. Two large protein families that contribute significantly to vesicle targeting are the Rab GTPases and the soluble N-ethyl-maleimide-sensitive factor attachment protein receptor (SNARE) family of docking/fusion proteins (51). In our study, upregulation in the above-mentioned trafficking networks was observed in SA, reflecting either increased intracellular transport of various cargo or the extensive regulation of lipid droplet formation and interactions with other organelles such as endosomes, peroxisomes, and mitochondria, where lipids and signalling molecules are exchanged (52). Remarkably, this is well in line with our data that showed the enrichment of
predicted secreted proteins in SA, compared to VA. It is also noteworthy to underline the enrichment of RABs which are also involved in the transport pathway, including the trafficking of high-affinity glucose transporter GLUT4 (S3). In agreement with our results, multiple proteins involved in vesicular trafficking were previously reported by Xie and colleagues who characterized the SA proteome in lean men (49). Highlighting the variability in transport mechanisms, it is also worth noting that the number of VA proteins associated with the vesicle-mediated transport is negatively correlated with cholesterol and LDL levels, and positively with visceral fat mass. In addition, the largest SA module (Turquoise) is enriched in proteins involved in vesicular transport and positively correlated to glycaemia and negatively to HDL level. Altogether, our data suggest distinct features of protein synthesis, transport, and turnover in adipocytes from different depots.

In adipocytes, triglycerides (TG) are synthesized from free fatty acids (FFAs) esterified to a glyceride-glycerol backbone. The majority of FFAs are delivered through lipoprotein lipase (LPL) breakdown of triglyceride-rich plasma lipoproteins. This process is partially complemented by the direct uptake of circulating FFA (5). In the current study, we observed the overexpression of several lipid homeostasis regulators in SA, encompassing the main gatekeeper enzyme for the entry and FFA re-esterification in adipose tissue – LPL, crucial long-chain fatty acid transporter in adipocytes CD36, and proteins of the apolipoprotein family (e.g. APOA1, APOA2, APOB, APOD, APOL2, APOM, and APOE). Notably, the expression of hormone-sensitive lipase (LIPE), the rate-limiting step for lipolysis of TG stored in adipocytes, was higher in VA. Apolipoproteins regulate the assembly of lipoprotein particles, maintain their structure and affect their metabolism by binding to membrane receptors (S4). In agreement with our results, higher mRNA expression of APOE was previously reported in SA (S5). Conversely, another study revealed a higher level of APOA1 and APOE in a visceral depot in a mixed-sex cohort (41). Nonetheless, further research is needed, as this is – to the best of our knowledge – the first proteomic study to unveil such pronounced depot-specific differences in multiple apolipoproteins in mature adipocytes. In addition to LPL-catalysed hydrolysis of triglycerides, it is worth mentioning that lipoproteins may also be whole-particle internalized directly by cells through
LDL receptor family member proteins such as VLDLR, LDLR, and LRP1 (56) which were, in comparison, upregulated in VA. Notably, several other lipid metabolism regulators were overexpressed in SA, including ACOX1, a rate-limiting enzyme in peroxisomal fatty acid β-oxidation, and a cluster of proteins involved in the glycosphingolipid metabolism (e.g. ARSA, ARSD, ASAH1, CTSA, ESYT1, GLB1, GLTP, HEXB, STS, SUMF2). This is of particular importance, as disturbances in glycosphingolipid content regulation emerged as an essential contributor to the development of insulin resistance (57).

Altogether, our data imply large lipid flux and the prominent role of the lipoprotein metabolism in a subcutaneous compartment, and propose distinct depot-dependent features of TG utilization.

Obesity is characterized by considerable adipose tissue expansion, and ECM reorganization creates space for adipocyte enlargement. In line with this, proteins associated with cytoskeletal and ECM remodelling were in our study significantly correlated with increased visceral fat mass (VA, Pink module). ECM is a complex structure composed of different proteins, proteoglycans, and polysaccharides (58). In the current study, we revealed prominent upregulation in ECM components including collagens (e.g. COL3A1, COL4A1, COL4A2, COL6A1, COL6A2, COL6A3), laminin (LAMB2, LAMC1), and integrin (e.g. ITGA1, ITGA6, ITGAV, ITGB5) in SA. The importance of ECM remodelling in a distinctive pathological contribution of adipose tissue depending on its anatomical site was previously demonstrated by CILAIR-based comparative secretome analysis of obese VAT and SAT (59). Remarkably, we found that the majority of the VAT most over-secreted proteins mentioned in this study is in fact enriched in SA in our analysis (e.g., COL6A3, LAMC1, HSPG2, CD14). By contrast, two of the most over-secreted SAT proteins (COL1A2, THBS1) identified by Roca-Rivada et al. were found enriched in VA in our study. This negative correlation might suggest protein depletion in cellular fraction due to high secretory activity. Nonetheless, these discrepancies might also be attributed to dissimilar experimental set-up, including differences in cohorts, methodology, and analysis of whole tissue vs. isolated mature cells. Further, the SA expression of proteins involved in extracellular matrix disassembly, i.e. CD44, CTSL, LCP1 or CAPN2, correlated with the clinical data including waist to hip ratios, total weight or BMI. Significantly, several studies implicate a causal role
of CD44 in adipose tissue inflammation and insulin resistance (60,61). In a similar manner, another emerging regulator of ECM interactions involved in insulin resistance, CD248, is among the most expressed proteins of SA (62).

In our study, five of the most upregulated proteins of VA were small leucine rich repeat proteoglycans, ubiquitous ECM components involved in matrix structural organization (OGN, ASPN, LUM, PRELP, and BGN). Indeed, SLRPs have previously been reported in insulin resistance-associated visceral secretome, showed a positive correlation with BMI and central obesity, and play overall substantial roles in mediating metabolic inflammation (63–65). Together, these proteins may be considered to constitute promising targets for following research of the depot-specific imprint in the peripheral system. Altogether, ECM components emerge as an additional essential player in the pathophysiology of obesity, and it is plausible that the depot-specific composition of ECM and its interactions are directly implicated in the processes associated with adipose tissue dysfunction.

To conclude, our study characterized the proteome of mature adipocytes from subcutaneous and visceral adipose depots. Of special interest are proteins that are predicted or annotated as secretory, and could be utilized as promising depot-specific biomarkers for advanced obesity monitoring. Previously published data concerning obesity emphasize the activity and deleterious role of the visceral compartment. In addition to this widely accepted concept, our results highlight the role of subcutaneous adipocytes in the regulation of fundamental processes associated with obesity, including the lipid metabolism, vesicular transport, and ECM composition.

This is to date the most comprehensive comparative proteomic analysis of isolated adipocytes; however, some limitations must be considered. Our study focused on female patients with clinically severe obesity, as they represent the primary group of bariatric surgery patients (66). Extrapolations of our data should thus be approached with caution; the differences reported here must be confirmed in future mixed cohorts studies employing a combination of functional and metabolomics approaches. Further, while the extreme phenotype of our cohort is useful in the enhancement of
differences between studied depots, any extrapolation of our results to non-obese subjects remains questionable, as well as any inference of causality based on observed data. On the other hand, to the best of our knowledge this is the first large-scale non-targeted proteomics analysis of abdominal SA and omental VA derived from paired biopsies of a relatively homogenous and large (n = 19) cohort, performed together with a bioinformatic analysis of the whole proteome, and differentially expressed proteins. Our effort provides insight into the functional role of adipocytes in obesity and may thus serve as a roadmap for future research. Hopefully, a better understanding of the distinct roles of adipocytes will help improve the characterization of the molecular pathophysiology of obesity and its associated diseases and facilitate relevant therapeutic interventions.
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Author Contributions

Conceptualization, PHr, JK, and JBV; Clinical assessment and sample biopsies, MP, PHo, MM, and MB; Cell handling, JK, and PL; LC-MS, DP, and ZZ; Data analysis, PHr, DK, and DP, Investigation, and data interpretation, PHr, JK, and JFK; Writing - original draft PHr, and JK; Writing – Review & Editing, PHr, JK, MP, PHo, MM, MB, DK, PL, JFK, DP, ZZ, and JBV; Visualization, PHr, DK, Project administration and funding acquisition, JBV

Data Availability

The mass spectrometry proteomics data and the search results have been deposited to the ProteomeXchange Consortium via the PRIDE (26) partner repository with the dataset identifier PXD024734. Supplementary data is available at figshare.com - Hruska P. et al. Proteomic signatures of human visceral and subcutaneous adipocytes - Supplementary files DOI: 10.6084/m9.figshare.14626341 (28). The R scripts are available upon request.
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**Figure title and legends**

**Figure 1 – Global proteome GO term enrichment**

GO enrichment analysis of all identified proteins (n = 2939) in the SA and VA adipocytes. The enrichment analysis was retrieved using ClueGO limited to top 2 GO term levels for biological processes (BP), cellular components (CC), and molecular function (MF). The figure shows the percentage of associated genes from the adipocyte proteome represented by significantly enriched GO terms (adj. P-value <0.05). The GO terms are sorted from the most significant on top to less significant on the bottom for each GO term group, respectively.

**Figure 2 – Global correlation map of SA and VA proteome**

The global correlation maps show cross-correlations of all quantified proteins using Pearson correlation coefficient and subsequent unsupervised hierarchical clustering for a) SA and b) VA proteome, respectively. This analysis revealed extensive maps of co-regulated proteins in respective clusters. In the map, proteins with a strong correlation or anti-correlation to each other cluster together in red or blue areas, respectively. The protein lists of the main correlation clusters were submitted to ClueGO enrichment analysis of biological processes (BP), and cellular components (CC). The most significantly enriched terms of two cluster levels can be found on the right from the maps.
Figure 3 – SA and VA GO term enrichment

GO enrichment analysis of all significantly upregulated SA (n = 576) and VA (n = 564) proteins. The enrichment analysis was retrieved using ClueGO with a significance threshold of 0.05 and merging all redundant groups with > 50% overlap for a) cellular components (CC), b) biological processes (BP), and c) molecular function (MF). The figure shows the most significantly enriched GO terms (adj. P-value < 0.05) for SA, and VA upregulated proteins, respectively.

Figure 4 – Pathway and reaction analysis of differentially expressed proteins

Reactome pathway and reactions enrichment analysis of all significantly a) SA and b) VA upregulated proteins. Figures show the most significant terms (p-values < 0.05) using the Reactome build-in Voronoi diagram visualization tool. The terms are visualized based on the Reactome event hierarchy showing the most significant terms in bright yellow.

Figure 5 – SA and VA upregulated proteins pathway and interaction analysis

The figure shows the most significant Reactome pathways and reactions (p-values < 0.05) together with associated proteins and protein-protein STRING interaction links visualized using Cytoscape GlueGO+CluePedia plugin. SA and VA Reactome pathway and reaction term nodes are shown as red and blue circles, respectively. The size of the nodes indicates the significance of the node - the bigger, the more significant. Related terms connect edges of a gray color. Nodes sharing upregulated proteins of both groups are transparent. Bright red and green edges indicate the ability of the protein to activate or inhibit the coupled protein. The most significant terms of each cluster group
were visualized and manually adjusted to obtain a reader-friendly pathway network. All pathways and networks with group cluster affiliation are accessible in S File 4 (28).

**Figure 6 – The most differentially expressed proteins**

a) Volcano plot for differential protein expression analysis showing the top 20 most upregulated proteins of SA (green) and VA (red).

b) Protein abundance rank of the most differentially expressed proteins. The mean intensity values of all measured proteins were ordered from the larger to the lowest values and ranked accordingly. The plot shows the relative representation of the most differentially expressed proteins within the whole adipocyte proteome.

c) The box plot shows the differences between SA and VA intensity values of the most differentially expressed proteins. Cross stands for imputed values and dot for experimental intensity values.

**Figure 7 – SA module-trait relationships**

SA module-trait relationship and corresponding p-values (Pearson) between the module eigengenes and selected clinical traits. Strong positive correlations are in red, while strong negative correlations are blue, numbers in individual cells are correlation coefficient (p-value). Significant module-trait correlations (p-value < 0.05) with the absolute correlation value > 0.5 are in bold. The size of the modules is in brackets. Waist1 stands for the waist circumference measured at the inferior margin of the ribs, and waist2 measured at the umbilical level.
Figure 8 – VA module-trait relationships

VA module-trait relationship and corresponding p-values (Pearson) between the module eigengenes and selected clinical traits. Strong positive correlations are in red, while strong negative correlations are blue, numbers in individual cells are correlation coefficient (p-value). Significant module-trait correlations (p-value < 0.05) with the absolute correlation value > 0.5 are in bold. The size of the modules is in brackets. Waist1 stands for the waist circumference measured at the inferior margin of the ribs, and waist2 measured at the umbilical level.
Table 1 – Subjects description

| Mean ± SD                                      |
|-----------------------------------------------|
| Age [years]                                   | 47 ± 8                        |
| Height [cm]                                   | 164 ± 4                       |
| BMI [kg/m²]                                   | 41.32 ± 4.23                  |
| Waist1 [cm] – the inferior margin of the ribs | 110 ± 9                       |
| Waist2 [cm] – umbilical level                 | 125 ± 12                      |
| Hip [cm]                                      | 130 ± 9                       |
| Waist1 to hip ratio                           | 0.84 ± 0.04                   |
| Waist2 to hip ratio                           | 0.96 ± 0.07                   |
| **DXA**                                       |                               |
| Body weight [kg]                              | 112 ± 12                      |
| Lean mass [kg]                                | 56 ± 5                        |
| Body fat mass [kg]                            | 53 ± 8                        |
| Visceral fat mass [g]                         | 1001 ± 236                    |
| Visceral fat volume [cm³]                     | 1082 ± 255                    |
| Visceral fat area [cm²]                       | 208 ± 49                      |
| Android to gynoid ratio                       | 1.07 ± 0.09                   |
| Fat mass index [kg/m²]                        | 30 ± 3                        |
| Lean mass index [kg/m²]                       | 21 ± 1                        |
| **Clinical laboratory data**                  |                               |
| Glycemia [mmol/l]                             | 5.9 ± 0.9                     |
| Cholesterol [mmol/l]                          | 5.3 ± 0.7                     |
| Triglycerides [mmol/l]                        | 2.1 ± 0.9                     |
| HDL [mmol/l]                                  | 1.2 ± 0.2                     |
| LDL [mmol/l]                                  | 3.4 ± 1.0                     |
| HbA1c [mmol/l]                                | 3.9 ± 0.4                     |

The table shows average body composition and clinical data of the 19 women included in this study before the bariatric surgery.
| UniProt ID | Protein name                                      | Gene name | Log2 FC | adj. P value |
|-----------|--------------------------------------------------|-----------|---------|-------------|
| P05186    | Alkaline phosphatase, tissue-nonspecific isozyme | ALPL      | 5.30    | 2.81E-13    |
| Q9HBR0    | Putative sodium-coupled neutral amino acid transporter 10 | SLC38A10 | 4.08    | 3.97E-16    |
| Q5T5P2    | Sickle tail protein homolog                      | KIAA1217  | 4.04    | 2.47E-09    |
| Q9H4G4    | Golgi-associated plant pathogenesis-related protein 1 | GLIPR2   | 3.97    | 4.76E-13    |
| Q96NY7    | Chloride intracellular channel protein 6         | CLIC6     | 3.72    | 3.78E-13    |
| Q14558    | Phosphoribosyl pyrophosphate synthase-associated protein 1 | PRPSAP1  | 3.48    | 1.21E-08    |
| A6NGC4    | TLC domain-containing protein 2                  | TLCD2     | 3.44    | 8.14E-15    |
| Q9UM47    | Neurogenic locus notch homolog protein 3         | NOTCH3    | 3.29    | 9.14E-14    |
| Q9U003    | Coronin-2B                                       | CORO2B    | 3.16    | 1.80E-12    |
| Q9BRX2    | Protein pelota homolog                           | PELO      | 3.03    | 3.39E-03    |
| O95302    | Peptidyl-prolyl cis-trans isomerase FKBP9        | FKBP9     | 3.00    | 7.93E-12    |
| Q9NR28    | Diabolo homolog, mitochondrial                   | DIABLO    | 2.95    | 2.77E-04    |
| P08473    | Nephrin                                          | MME       | 2.93    | 7.14E-08    |
| P16278    | Beta-galactosidase                               | GLB1      | 2.89    | 9.13E-11    |
| P22413    | Ectonucleotide pyrophosphatase/phosphodiesterase family member 1 | ENPP1    | 2.83    | 2.59E-07    |
| Q9H173    | Nucleotide exchange factor SIL1                  | SIL1      | 2.80    | 3.35E-05    |
| P21926    | CD9 antigen                                      | CD9       | 2.76    | 1.41E-11    |
| Q96D15    | Reticulocalbin-3                                 | RCN3      | 2.75    | 2.47E-09    |
| Q9Y4F1    | FERM, RhoGEF and pleckstrin domain-containing protein 1 | FARP1    | 2.71    | 6.91E-10    |
| Q9HC00    | Endosalin                                        | CD248     | 2.50    | 1.57E-12    |
| P02786    | Transferrin receptor protein 1                   | TFRC      | -2.39   | 6.09E-05    |
| Q8N6C5    | Immunoglobulin superfamily member 1             | IGSF1     | -2.45   | 4.60E-06    |
| O75891    | Cytosolic 10-formyltetrahydrofolate dehydrogenase | ALDH1L1   | -2.46   | 2.54E-12    |
| O95671    | N-acetylserotonin O-methyltransferase-like protein | ASMTL    | -2.48   | 1.60E-04    |
| P27487    | Dipetidyl peptidase 4                            | DPP4      | -2.58   | 3.41E-05    |
| P21810    | Biglycan                                         | BGN       | -2.61   | 3.15E-05    |
| P51888    | Prolargin                                        | PRELP     | -2.64   | 4.22E-04    |
| O95433    | Activator of 90 kDa heat shock protein ATPase homolog 1 | AHS1A    | -2.74   | 3.39E-03    |
| P02743    | Serum amyloid P-component                        | APCS      | -2.84   | 3.10E-09    |
The table provides a list of the 20 most differentially expressed proteins for SA and VA, respectively.

Fold-change and significance of changes of the Limma differential expression analysis are demonstrated as log₂ fold-change (Log₂FC) and adjusted P-value for multiple hypothesis testing.

Potentially secreted proteins are in bold.

| P51884 | Lumican | LUM | -2.93 | 1.83E-04 |
| Q9BXN1 | Asporin | ASPN | -3.03 | 9.39E-08 |
| Q8N335 | Glycerol-3-phosphate dehydrogenase 1-like protein | GPD1L | -3.05 | 1.53E-08 |
| P20962 | Parathymosin | PTMS | -3.12 | 3.39E-05 |
| P15311 | Ezrin | EZR | -3.13 | 6.70E-07 |
| P49915 | GMP synthase [glutamine-hydrolyzing] | GMPS | -3.21 | 9.14E-14 |
| P16050 | Arachidonate 15-lipoxygenase | ALOX15 | -3.30 | 2.20E-09 |
| Q53EU6 | Glycerol-3-phosphate acyltransferase 3 | AGPAT9 | -3.50 | 9.14E-14 |
| P20774 | Mimecan | OGN | -3.57 | 6.25E-06 |
| Q5VT79 | Annexin A8-like protein 2 | ANXA8L2 | -3.81 | 7.41E-12 |
| Q13442 | 28 kDa heat- and acid-stable phosphoprotein | PDAP1 | -4.00 | 3.80E-08 |
Figure 7

SA – Module–trait relationships

![Figure 7 Image](https://academic.oup.com/jcem/advance-article-doi/10.1210/clinem/dgab756/6406610)
