Mass spectrometry-based absolute quantification of 20S proteasome status for controlled *ex-vivo* expansion of Human Adipose-derived Mesenchymal Stromal/Stem Cells

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Running Title: LC-SRM quantification of 20S proteasome for ADSCs expansion

Glossary:

sP20S: standard 20S proteasome

iP20S: immunoproteasome

β5t P20S: thymoproteasome

 $\alpha 4s$ P20S: spermatoproteasome

ADSCs: Adipose-derived Mesenchymal Stromal/Stem Cells

AT: Adipose Tissue

FBS: Fetal Bovine Serum

IDMS: Isotope Dilution Mass Spectrometry

IFN γ : interferon γ

MSCs: Mesenchymal Stem Cells

SILAC: Stable Isotope Labeling by Amino acids in Cell culture

SRM / MRM: Single Reaction Monitoring / Multiple Reaction Monitoring

SVF: Stromal vascular fraction

Summary:

The proteasome controls a multitude of cellular processes through protein degradation and has been identified as a therapeutic target in oncology. However, our understanding of its function and the development of specific modulators are hampered by the lack of a straightforward method to determine the overall proteasome status in biological samples. Here, we present a method to determine the absolute quantity and stoichiometry of ubiquitous and tissue-specific human 20S proteasome subtypes based on a robust, absolute SILAC-based multiplexed LC-Selected Reaction Monitoring (SRM) quantitative mass spectrometry assay with high precision, accuracy, and sensitivity. The method was initially optimized and validated by comparison with a reference ELISA assay and by analyzing the dynamics of catalytic subunits in HeLa cells following IFNγ-treatment and in range of human tissues. It was then successfully applied to reveal IFNγ- and O₂-dependent variations of proteasome status during primary culture of Adipose-derived-mesenchymal Stromal/Stem Cells (ADSCs). The results show the critical importance of controlling the culture conditions during cell expansion for future therapeutic use in humans. We hypothesize that a shift from the standard

proteasome to the immunoproteasome could serve as a predictor of immunosuppressive and differentiation capacities of ADSCs and, consequently, that quality control should include proteasomal quantification in addition to examining other essential cell parameters. The method presented also provides a new powerful tool to conduct more individualized protocols in cancer or inflammatory diseases where selective inhibition of the immunoproteasome has been shown to reduce side effects.

Introduction:

Some hematopoietic malignancies can be successfully treated by inhibition of the catalytic core 20S proteasome complex, and more recent findings indicate that the proteasome is a promising target for the treatment of other cancer types or other pathologies including inflammatory diseases (1–3). Although the cylindrical $\alpha7\beta7\beta7\alpha7$ barrel-like structure of the 20S catalytic core proteasome has been preserved throughout evolution, the oligomeric protease has evolved, resulting in a higher heterogeneity of subunit compositions in mammals. As schematically represented in Figure 1A, there exist at least six distinct forms of 20S proteasomes in human cells and tissues. The standard 20S proteasome (sP20S) is composed of constitutive ($\alpha 1-\alpha 7$ and $\beta 3$, $\beta 4$, $\beta 6$, and $\beta 7$) and catalytic subunits $(\beta 1, \beta 2 \text{ and } \beta 5)$. It is the most abundant 20S subcomplex in most cell types, but significant amounts of other 20S forms have been observed in some human tissues and cells in their basal state, or are induced in specific environmental conditions (4). For instance, in response to pro-inflammatory cytokines or in immune cells, the three catalytic subunits of the sP20S can be replaced in a highlyregulated way by their immuno counterparts to form the immunoproteasome (iP20S), which has nonidentical cleavage specificities (5). Two intermediate proteasomes harboring a mixed assortment of standard and immunocatalytic subunits (β1i P2OS, β1iβ5i P2OS) have also been observed in various human tissues and cells, and their existence is consistent with the rules of cooperative assembly of inducible catalytic subunits (6). Some other 20S subtypes are much more tissue-specific, such as the thymoproteasome (containing β_{1i} , β_{2i} and β_{5t} catalytic subunits) and the spermatoproteasome (containing a specific isoform of $\alpha 4$ called $\alpha 4s$) which are found in the thymus and male germ cells, respectively. The environment- or tissue-specific subunit composition of human 20S proteasome has been shown to fulfill specialized functions that the standard proteasome can only exert suboptimally (4). For example, immune cells contain a significant proportion of immunosubunit-containing 20S proteasome subtypes which display specific proteolytic preferences thanks to which they produce antigens for presentation to CD8 T cells (6). In the thymus, the unique cleavage preference of B5t explains the essential role played by the thymoproteasome in the positive selection of developing T lymphocytes (7). Thus, knowledge of 20S proteasome composition will help us to better understand proteasome function in a given tissue or cell type. It is also crucial when developing novel therapeutic strategies targeting specific 20S proteasome variants. Indeed, the immunoproteasome is a valuable target in several ongoing oncology trials as well as in the treatment of inflammatory and autoimmune diseases (2, 3, 8). To assess proteasome status, by determining the absolute quantity and stoichiometry of all subtypes, thus requires precise and accurate absolute quantification of several subunits from the same biological sample. As mRNA and protein levels reported from proteomic and transcriptomic analyses show a low correlation (9), quantification must be performed at the protein level to allow an accurate and complete description of the 20S proteasome status. Several ELISA protocols have been published or are commercially available to determine absolute levels of both 20S and 26S proteasome complexes, but only at the global level, *i.e.*, they make no distinction between the different subcomplexes (6, 10–14). Only Guillaume et al. (6) considered the heterogeneity of 20S subtypes when developing their ELISA assay by using different in-house produced antibodies directed against four different standard and immunocatalytic β subunits. More recently, standard and immunoproteasome subtypes were determined by surface plasmon resonance imaging (SPRI) using specific inhibitors (15). However, the multiplexing capacity of these methods is insufficient to fully assess proteasome heterogeneity in a single assay.

To overcome this limitation, we propose to use Selected Reaction Monitoring (SRM), an isotope dilution mass spectrometry (IDMS)-based technique which has been approved for the quantification

of multiple biomolecules. In addition to its robustness, precision and accuracy, the main advantage of SRM is its capacity to quantify many analytes simultaneously. SRM has recently been successfully applied in protein assays (16, 17). In these studies, the proteins themselves were not directly detected and quantified, but were analyzed at peptide level, after enzymatic proteolysis. This is a critical step because, in most cases, the absolute quantification relies on the addition of an isotopically-labeled peptide standard (called AQUA peptide) (18) late in the experimental workflow, which might introduce a high variability due to differences in sample preparation. Thus, for accurate and robust absolute quantification of proteins, and in particular to determine protein stoichiometry within protein complexes, the use of isotope-labeled peptides as internal standards requires careful optimization of experimental conditions (19). An alternative approach based on artificial genes coding for concatenated proteotypic peptides (QconCAT strategy) (20) may be used to decrease quantification biases. For robust absolute quantification, an isotope-labeled equivalent of the full-length target protein is considered the ideal internal standard. Several approaches based on the synthesis of heavy versions of the protein of interest have emerged recently. Most of them rely on metabolic labeling and purification of a recombinant version of the protein of interest, or a shorter specific protein sequence, like the Protein Standards for Absolute Quantification (PSAQ) (21), absolute SILAC (22), PrEST (23), FlexiQuant (24), and TAQSI (25) methods. Another approach consisting in the chemical labeling of proteins using isobaric stable isotope tagging technology (26) has recently been described to reduce costs and the time required to grow cells in heavy amino acid-containing media. In all these approaches, because the internal standard is processed together with its endogeneous analog throughout the whole workflow, accuracy is markedly better than achievable with AQUA and QconCAT approaches when determining absolute quantities of proteins in various matrices (17). However, to our knowledge, the absolute quantification relying on the isotopic dilution of labeled proteins has been limited so far to monomers, and no reports of assays to determine the stoichiometry of macromolecular complexes have been published.

Mesenchymal stem/stromal cells (MSC) hold great potential in regenerative medicine because of their multi/pluripotency and immunosuppressive properties. Over the last decade, the clinical use of MSC has rapidly increased, and more than 800 clinical trials assessing MSC therapy in multiple clinical settings are currently registered (https://www.clinicaltrials.gov/). Adipose-derived Stem Cells (ADSCs) are a subclass of mesenchymal stem/ stromal cells initially derived from the bone marrow. However, to obtain the critical number of cells before transplantation, ADSCs must be expanded *in vitro*. This essential step has raised important concerns about the quality of adult stem cells, and inter-patient variability is a challenge when seeking to define ADSCs. The development of uniform protocols for both preparation and characterization of MSCs, including standardized functional assays to assess their biological potential, will be critical in contributing to their clinical utility.

Here, we applied the SILAC method to engineered cell lines expressing either the standard 20S proteasome or the immunoproteasome to produce and purify isotope-labeled, endogeneous versions of standard proteasome and immunoproteasome in vivo. These isotope-labeled proteasome complexes were used as internal standards to quantify the absolute concentration of all 20S catalytic and non-catalytic subunits in biological samples of various origins. The method developed was applied to determine the absolute concentration of total 20S proteasomes and the exact stoichiometry of six ubiquitous and tissue-specific 20S proteasome subtypes in a multiplexed LC-SRM assay with high precision (>92%), accuracy (>90%), and sensitivity (<1 fmol on column). Our results show that the absolute quantity and stoichiometry of the proteasome are challenged both by IFNy stimulation and O₂ levels during ex vivo expansion of primary ADSCs. Thus, determining proteasome status, which is a central contributor to maintaining stem cell homeostasis - characterized by stemness, capacity for self-renewal and cell differentiation (27–29) – might constitute an additional relevant quality control parameter for the production of ADSCs for clinical applications, which is of interest as the number of quality markers currently available is limited (30). Furthermore, accurate and precise assessment of proteasome abundance and heterogeneity could also help when seeking to achieve selective inhibition of a particular proteasome subtype, like the immunoproteasome, for personalized therapies in cancer

or autoimmune diseases. This is the first study to report the simultaneous determination of absolute quantity and stoichiometry for macromolecular complexes based on the isotopic dilution of labeled proteins in numerous human tissues and primary ADSCs culture.

Experimental Procedures

Cell lines, Culture conditions, SILAC, human samples

HEK 293T, HCT116, HeLa and RKO cell lines were grown in DMEM medium supplemented with 10% fetal bovine serum (FBS). U937, HeLa S3 and NB4 cell lines were grown in RPMI 1640 medium supplemented with 10% FBS. KG1a cell line was grown in RPMI 1640 medium supplemented with 20% FBS. MRC5 cell line was grown in MEM-α medium supplemented with 10% FBS. All cultures were supplemented with 2×10^{-3} M glutamine, 100 units/mL penicillin, 100 µg/mL streptomycin, and maintained at 37 °C under 5% CO₂. Unsynchronized cells were harvested at 80% confluence for adherent cells or at a concentration of 1.10⁶ cells per ml of culture for suspension cells. HeLa cells were treated with interferon-γ (R&D Systems, Minneapolis, MN) at 100 ng/ml in fresh medium.

Human 293-EBNA cells, "HEK-EBNA sP2OS" (mainly expressing sP2OS), and 293-EBNA cells engineered to express iP2OS, "HEK-EBNA iP2OS" (by transfecting 293-EBNA cells with cDNAs encoding the three immunocatalytic subunits β 5i, β 1i and β 2i) were obtained as previously described (6). HEK-EBNA sP2OS cells were cultured in SILAC medium which is composed of DMEM supplemented with 10% dialyzed FBS, 4 mM L-glutamine, 200 mg/L L-Proline, 100 mg/mL L-arginine ($^{13}C_6$), and L-lysine ($^{13}C_6$) (Cambridge Isotope Lab.), 100 IU/mL penicillin and 100 µg/mL streptomycin in 150 cm² culture plates and maintained at 37°C under 5% CO₂. HEK-EBNA iP2OS were cultured in the same SILAC medium as HEK-EBNA sP2OS, but further supplemented with 5 µg/mL Puromycin and 600 µg/mL Hygromycin to maintain selective pressure. Ten cellular doublings were performed in this medium to achieve an incorporation rate of 95% heavy amino acids in proteins (assessed by MS). Standard 20S proteasome and iP2OS were then purified as described earlier (31). Absolute quantities and purities of both purified proteasome subtypes were then assessed as described in Supplementary information I-1. Isotopelabeled sP20S and iP20S were stored as 10- μ L aliquots at 1.158 and 0.980 pmol/ μ L, respectively, in 20 mM Tris/HCl, pH 7.2, 1 mM EDTA, 1 mM DTT, 10% glycerol, and at -80°C to ensure stability over time.

Whole-cell lysates from various human tissues were supplied by AMSBio (HT-201 (Brain); HT-311 (Colon); HT-804 (Heart); HT-314 (Liver); HT-601 (Lung); HT-406 (Ovary); HT-102 (Skeletal muscles); HT-701 (Spleen); HT-401 (Testis); HT-704 (Bone Marrow); P1234264 (Thymus)).

Human ADSC were isolated from subcutaneous adipose tissue (AT) obtained from non-obese human donors (body mass index <26) undergoing elective abdominal dermolipectomy (Plastic Surgery Department, Rangueil Hospital, Toulouse, France). No-objection certificates were obtained to comply with bioethics law no. 2004-800 of 6 August 2004. The stromal vascular fraction (SVF) was obtained by enzymatic digestion of adipose tissue (AT) with collagenase NB4 (Roche Diagnostics, Indianapolis, IN). Cells were then seeded at 4,000 cell/cm² (P0) and cultivated in α -minimal essential medium supplemented with 10% FBS (Gibco), 100 µg/ml streptomycin, 100 U/ml penicillin, 25 µg/ml amphotericin (Thermo Fisher Scientific Life Sciences, Waltham, MA). Medium was changed twice a week. Cells from the same patients were cultured under normoxic (20% O₂, 5% CO₂) or hypoxic (1% or 5% O₂, 5% CO₂) conditions in an Xvivo System (BioSpherix, Paris, NY) to maintain cells in hypoxic conditions at all culture steps. Interferon- γ (100 ng/ml) was added to the medium in some experiments.

Our experimental protocols were approved by French research ministry's institutional ethics committee of (N°: DC-2015-23-49) and informed consent was obtained from all subjects in line with current regulations (no subjects age under 18 were included).

Adipocyte differentiation

ADSCs obtained from patients were seeded at 20,000 cell/cm² and were exposed to an adipogenic cocktail containing IBMX (3-isobutyl-1-methylxanthine) 0.45 mM, Dexamethazone 1 μ M and Indomethacine 60 μ M. An immunoproteasome inhibitor, 100 nM ONX-0914 (solubilized in 0.001% DMSO), was also added to the cell medium in some conditions; 0.001% DMSO was added to the control condition in these cases. Cells were grown at 37 °C under normoxic (20% O₂) or hypoxic conditions (5% and or 1% O₂). The medium was changed every 2–3 days throughout the culture process. At the end of experiment, the cell lineage was determined using Oil Red O which stains for adipocytes.

20S Proteasome purification.

When they reached 80% confluence, HEK-EBNA cell lines (HEK-EBNA sP20S or HEK-EBNA iP20S) were harvested in HKMG buffer (10 mM Hepes pH 7.9, 10 mM KCl, 5 mM MgCl₂, 10% glycerol, 10 mM ATP, 1% NP40, protease and phosphatase inhibitor (Roche)) and centrifuged for 10 min at 10,000 g. Supernatants were kept and used for proteasome immuno-purification as previously described (31).

ELISA-based absolute quantification of the 20S proteasome

ELISA-based absolute quantification of the 20S proteasome was performed as previously described (14). Briefly, ELISA assays were performed in 96-well plates (IMMULON HBX 4, Thermo Scientific, Milford, MA, USA). The plate was coated with 100 µL MCP21 monoclonal antibody (European Collection of Cell Cultures) at 5 µg/mL by incubation at 4 °C overnight. Wells were then washed 3 times with PBS/T-buffer. Non-specific sites were blocked by incubation with 2% BSA in PBS for 1 h at room temperature with slow shaking. The plate was washed 3 times with PBS/T-buffer. The samples (cell lysates) were then deposited in triplicate and incubated for 2 h at room temperature. The plate was washed with PBS/T-buffer and then incubated with the polyclonal rabbit anti-20S antibody (PW 8155, ENZO LIFE Sciences) for 1 h at room temperature under slow shaking. The plate was washed 3 times

with PBS/T-buffer and antibody binding was revealed using horseradish peroxidase-conjugated antirabbit antibody and 2 mg/mL ABTS substrate (2,2-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid)). The reaction was monitored by measuring the optical density at 416 nm (μQuant; Bio-Tek instruments, Inc., Winooski, VT, USA). The amount of proteasome in the sample was calculated by comparison with the calibration curve produced with purified 20S proteasome purified from human erythrocytes (ENZO LIFE SCIENCE). A linear dose–response was observed between 0 and 20 ng.

Proteasome activity assay

Cultured cells were harvested in HKMG buffer and sonicated in ice with a Bioruptor Plus (Diagenode) (15 min, cycle 45 s/15 s (ON/OFF), position High). Protein concentration was determined by detergentcompatible protein assay (DC Assay – BioRad) according to the manufacturer's recommendation. Proteasome activity was assayed in 96-well black plates (Greiner Bio-One, Frickenhausen, Germany). 10 µL of each lysate fraction were added to 40 µL of Tris-HCl 100 mM and 50 µL of Suc-LLVY-AMC (for chymotrypsin-like activity), Boc-LRR-AMC (for trypsin-like activity) and Z-LLE-AMC (for PGPH activity) substrate in 200 mM Tris-HCl, pH 8 (Enzo Life Science) at a final concentration of 400 µM/well. Kinetic assays were performed at 37 °C in a FLX-800 spectrofluorometer (BIOTEK, Winooski, VT, USA) over 90 min, reading fluorescence every 5 min, at 460 nm following excitation at 360 nm.

Samples preparation and in-gel digestion for mass spectrometry analysis

Samples were heated to 95 °C for 5 min in Laemmli buffer to denature proteins; 100 mM chloroacetamide was then added to the sample followed by incubation for 30 min at room temperature in the dark. Proteins were loaded onto a 12% acrylamide SDS-PAGE and concentrated in a single band, visualized by Coomassie staining (Instant Blue – Expedeon). The gel band containing the whole sample was cut and washed several times in 50 mM ammonium bicarbonate, acetonitrile (1:1) for 15 min at 37 °C. Trypsin (Promega) digestion was performed over night at 37 °C in 50 mM ammonium bicarbonate at a trypsin/total protein ratio of 1/50. Peptides were extracted from the gel by two incubations in 10% formic acid, acetonitrile (1:1) for 15 min at 37 °C. Extracts were dried in a Speed-Vac, and resuspended in 2% acetonitrile, 0.05% trifluoroacetic acid prior to LC-MS/MS analysis.

For LC-SRM analysis, 1 pmol of labeled iP20S and 1 pmol of labeled sP20S were added to 25 µg of total proteins before the protein denaturation step. When used, AQUA peptides (Thermo Scientific Pierce Protein Research) were spiked into samples just before mass spectrometry analysis at a concentration resulting in injection of a final quantity of 70 fmol.

Whole proteome analysis of extracts from human tissues and ADSCs grown under normoxic and in hypoxic conditions (1% O₂). Peptide mixtures were analyzed by nano-LC-MS/MS using an Ultimate3000 system (Dionex) coupled to an LTQ-Orbitrap Velos mass spectrometer (Thermo Fisher Scientific, Bremen, Germany) when analyzing human tissue extracts or to a Q-Exactive Plus mass spectrometer (Thermo Fisher Scientific, Bremen, Germany) when analyzing ADSCs. Five microliters (human tissue) or two microliters (ADSCs) of each peptide sample at $1 \mu g/\mu L$ were loaded onto a C18 precolumn (300 μm inner diameter x 5 mm; Thermo Scientific) at 20 μL/min in 5% acetonitrile, 0.05% trifluoroacetic acid. After 5 min of desalting, the precolumn was switched online with the analytical C18 column (75 µm inner diameter x 50 cm; home-made) equilibrated in 95% solvent A (5% acetonitrile, 0.2% formic acid) and 5% solvent B (80% acetonitrile, 0.2% formic acid). Peptides from human tissues were eluted using a 5-50% gradient of solvent B over 105 min at a flow-rate of 300 nL/min and peptides from ADSCs were eluted using a 5–25% gradient of solvent B over 165 min and a 25-50% gradient of solvent B over 135 min at a 300 nl/min flow rate. Both mass spectrometers were operated in data-dependent acquisition mode. During the analysis with the LTQ-Orbitrap Velos, survey scan MS spectra were acquired in the Orbitrap over the 350–1800 m/z range with resolution set to 60,000 (these parameters were 350-1500 m/z and 70,000 with Q-Exactive Plus, respectively). On the LTQ-Orbitrap Velos, the twenty most intense ions per survey scan were selected for CID fragmentation, and the resulting fragments were analyzed in the linear trap (LTQ). On the Q-Exactive Plus, the ten most intense ions per survey scan were selected for HCD fragmentation, and the resulting fragments were analyzed in the Orbitrap. Dynamic exclusion was used with a 60-s or a 30-s window (on the LTQ-Orbitrap Velos or the Q-Exactive Plus, respectively) to prevent repeated selection of peptides.

Raw mass spectrometry files were processed using MaxQuant (version 1.5.5.1) and Andromeda was used to match MS/MS spectra against the Human SwissProt database (March 2017 release – 20,181 entries for human tissues analysis / release of August 2018 – 20,386 entries for ADSCs analysis) and a list of potential contaminant sequences provided in MaxQuant1.5.5.1., with Carbamidomethylation of

cysteines set as fixed modification. Oxidation of methionine and protein N-terminal acetylation were set as variable modifications. The digestion specificity of trypsin was defined as cleavage after K or R, and up to two missed trypsin cleavage sites were allowed. The precursor mass tolerance was set to 20 ppm for the first search and 4.5 ppm for the main search. The mass tolerance in MS/MS mode was set to 0.8 Da for tissue analysis and to 20ppm for ADSC analysis. Minimum peptide length was set to seven amino acids, and minimum number of unique peptides was set to 1. A target-decoy approach was used to validate hits, using a reverse database and applying a peptide and protein false-discovery rate of 1%. The "match between runs" option in MaxQuant was enabled, with a time window of 0.7 min, to allow cross-assignment of MS features detected in different runs. Only unique peptides were used for quantification when analyzing human tissues, whereas both unique and razor peptides were used for the quantification when dealing with ADSCs data. Quantitative proteomic analysis was performed on the normalized LFQ intensities from the "proteinGroups" table in the MaxQuant output. Protein entries identified as potential contaminants by MaxQuant were eliminated from the analysis, as were proteins identified by fewer than two peptides. Tissue-specific 20S proteasome subunits (α 4s, β1i, β2i, β5i, and β5t) were compared across the different tissues based on their normalized LFQ intensities (Figure 3B). To compare the proteomes of ADSCs grown under normoxic or hypoxic conditions, the LFQ intensity values were used for the quantitative analysis. Fold-changes were log₂ transformed and thresholds calculated based on their distribution. The upper threshold was calculated as Q3 + 1.5*IQ, and the lower threshold as Q1 - 1.5*IQ, where IQ is the interquartile, and Q1 and Q3 are the first quartile and the third quartile, respectively. Proteins with fold-changes outside these thresholds were considered outliers from the global distribution. Outlier proteins with a p-value of less than 0.05 were considered as differentially expressed between normoxic and hypoxic conditions. GO terms enrichment analysis was performed on these proteins using GOrilla (32).

Multiple Reaction Monitoring to quantify 20S proteasome subunits

Dried peptide samples were resuspended in a solution containing 2% acetonitrile, 0.05% TFA, to obtain a final concentration of 1 μ g/ μ L. To obtain data for the whole set of peptides/transitions, samples were injected twice as the list of transitions was split into two methods. The sample (2.5 μ L, about 2.5 μ g protein equivalent) was loaded onto the system and analyzed on a hybrid triple quadrupole-ion trap mass spectrometer 6500 QTrap (AB Sciex) equipped with a nanoelectrospray ion source coupled to an Ultimate 3000 system (Dionex) for chromatographic peptide separation. Separation was achieved using a 60 min gradient from 0 to 50% of solvent B (80% acetonitrile, 0.2% formic acid) at a flow-rate of 300 nL/min. Spray voltage was set to 2500 V, curtain gas to 35 psi, nebulizer gas to 5 psi, interface heater temperature to 75 °C, and cycle time to approximately 3 s (3.1620 s and 3.0380 s for the two methods). Peptides were loaded onto a C18 precolumn (300 μ m inner diameter x 5 mm; Thermo Scientific) at 20 μ L/min in 2% acetonitrile, 0.05% trifluoroacetic acid. After 5 min of desalting, the precolumn was switched online with the analytical C-18 column (75 μ m inner diameter x 50 cm; inhouse-packed with Reprosil C18) and equilibrated in solvent A (5% acetonitrile, 0.2% formic acid).

Proteotypic peptide sequences were selected based on results of previous discovery experiments (33– 35) and by referring to the golden rules (16) (*i.e.*, considering isoforms, variants, PTMs reported on protein sequence or possibly artefactually-induced by sample handling, or missed cleavages, observed in Protein (UniProt) or MS databases (PeptideAtlas)). To achieve maximal sensitivity, collision energies (CEs) were optimized within a 6 V window around the CE value recommended by Skyline software and based on the m/z of the precursor. Final SRM transitions are given in Supplementary Data 3. Transitions could be unambiguously assigned thanks to the co-injected isotope-labeled peptides. Samples were run in a blinded fashion except for calibration curves, for which the lowest concentrations were injected first. To check system suitability and performance before injecting each batch of samples, Total Ion Current was tuned (chromatographic solvent ions of m/z 50 to 1000) at 2000 Da/s scan speed in Q1. Quality controls (QC) (20 fmol of tryptic digest of betagalactosidase; transitions are given in tryptic digests of 100 fmol, 500 fmol or 1000 fmol of isotopically-labeled sP20S and iP20S spiked in 2.5 µg of HeLa, then one QC sample, and finally a blank sample (the maximum carry-over observed was less than 0.01%). Quantitative reproducibility over time was checked based on signal intensities for heavy transitions (maximum deviation of 50% allowed); injection batches were generally carried out over periods of less than one week. Quantitative data analyses were performed using Skyline-Daily open-source software (36). Area values

for all transitions were first extracted automatically by the software, then checked manually and adjusted if necessary (*i.e.*, exclusion of data points if S/N < 10). Light transition (L) peak area signals were then normalized with respect to their labeled counterparts (H) (after correction by considering the 95% incorporation rate for $R(^{13}C_6)$ and $K(^{13}C_6)$ in labeled sP20S and iP20S). For each transition, technical replicates of injection (typically three) were averaged. Ratios (L/H) of all the transitions used to assay a given protein or for total 20S proteasome (details in Supplementary Data 3) were then averaged. The amount of labeled reference mix (containing equimolar concentrations of isotope-labeled sP20S and iP20S) spiked into the biological sample before sample preparation was then used to determine the absolute quantity of each proteasome subunit or total 20S proteasome. For each transition, LOD and LLOQ were experimentally determined by injecting heavy-isotope-labeled sP20S and iP20S spiked at increasing concentrations in a HeLa protein lysate and processed by the same method as applied to the other biological samples. LOD and LLOQ values were calculated using QuaSAR (37), which was implemented through the Skyline interface (see Supplementary Data 5).

Supplementary Data 4) were injected between each sample. Carry-over was checked by injecting

Targeted MS measurements were highly multiplexed, and used to quantify proteasome subunits across cell lysates, tissues and primary cells of human origin. The assay developed used internal standards for each analyte, to confidently detect and precisely quantify the proteins of interest. Thus, the analyses meet the expectations of Tier 2 level.

Determining the stoichiometry of the six major 20S proteasome subtypes

The concentrations and stoichiometries of all proteasome subtypes were determined as explained hereafter.

Total proteasome absolute quantity was calculated by averaging the quantities of all 20S non-catalytic subunits (α 1-7; β 3, β 4, β 6, β 7) (mean (α 1,2,3,5,6,7; β 3,4,6,7) \rightarrow Total P20S). A stoichiometry of two non-catalytic subunits per 20S proteasome and a molecular weight of 700,000 g/mol were used in these calculations.

As $\beta 5$ is integrated into the standard proteasome, its fraction exclusively represents the fraction of standard proteasome ($\beta 5 \rightarrow sP20S$). Similarly, the level of $\beta 5t$ corresponds to the fraction of thymoproteasome " $\beta 5t$ P20S", and can be obtained by subtracting the sum of $\beta 5$ and $\beta 5i$ levels from the total amount of 20S proteasome (Total P20S - $\beta 5 - \beta 5i = \beta 5t \rightarrow \beta 5t$ P20S). As the $\beta 2i$ subunit is integrated into both the immunoproteasome and the thymoproteasome, the difference in its quantity compared to $\beta 5t$ can be used to determine the fraction of immunoproteasome ($\beta 2i - \beta 5t \rightarrow iP20S$). The $\beta 1i$ -containing proteasome subtypes correspond to immunoproteasome, thymoproteasome, and intermediate proteasome $\beta 1i$ - $\beta 5i$. Thus, the quantity of $\beta 1i$ - $\beta 5i$ P20S is equal to the difference in abundance of $\beta 1i$ and $\beta 2i$ proteins since both immunoproteasome and thymoproteasome contain $\beta 2i$ ($\beta 1i - \beta 2i \rightarrow \beta 1i\beta 5i$ P20S). $\beta 5i$ is contained in the immunoproteasome and in both types of intermediate proteasomes ($\beta 5i \rightarrow iP20S + \beta 5i P20S + \beta 1i\beta 5i P20S$). Thus, the amount of $\beta 5i P20S$ can be calculated as follows: $\beta 5i + \beta 5t - \beta 1i = Total P20S - \beta 1i - \beta 5 \rightarrow \beta 5i P20S$.

Finally, the spermatoproteasome level can be determined from the quantity of the α 4s isoform (PSA7L protein _PSMA8 gene) by subtracting the abundance of the α 4 subunit from the total 20S proteasome content (using specific peptides of the PSA7 isoform_PSMA7 gene as detailed in Sup Figure 8A) (sP20S – α 4 $\rightarrow \alpha$ 4s P20S).

Thus, the absolute amounts of non-catalytic subunits, as well as those of the catalytic β_{2i} , β_{5} , β_{5i} and β_{1i} subunits determined by the SRM method were used to calculate the proportions of the six main 20S proteasome subtypes, as summarized below:

mean (α1,2,3,5,6,7; β3,4,6,7) **→** Total P20S

 $\beta 5 \Rightarrow sP20S$ Total P20S - $\beta 5 - \beta 5i = \beta 5t \Rightarrow \beta 5t P20S$ $\beta 2i - \beta 5t \Rightarrow iP20S$ $\beta 1i - \beta 2i \Rightarrow \beta 1i\beta 5i P20S$ $P20S - \beta 1i - \beta 5 \Rightarrow \beta 5i P20S$ $sP20S - \alpha 4 \Rightarrow \alpha 4s P20S$

Experimental design and statistical rationale

SRM analyses: All statistical analyses were performed on at least three independent biological replicates. For each biological replicate, results from at least two injection replicates were averaged. Probability values (*p*) were determined by one-way and two-tailed analysis of variances (ANOVA). Differences were considered to be statistically significant at confidence levels of 95% (*), 99% (**), or 99.9% (***).

Coefficients of variation were calculated as ratios of the standard deviation over the mean of the values and are expressed as a percentage. Accuracy was determined as the ratio of the difference of the experimental value and the reference value over the reference value, and is expressed as a percentage.

Label-free MS analyses (Figure 3B and Supplementary Figure 11): The "human tissue extract" dataset contains mass spectrometry results from the analysis of 11 different human tissues analyzed in

triplicate, corresponding to 33 raw files. The "ADSC" dataset contains mass spectrometry results from the analysis of three different patients under two different oxygen percentages used for cell culture, samples were analyzed in triplicates, and thus produced 18 raw files. Quantitative proteomic analysis was performed on the normalized LFQ intensities from the "proteinGroups" table in the MaxQuant output.

The data sets corresponding to the mass spectrometry analyses presented in this study have been deposited in the following repositories: PRIDE (Project accession: PXD011894) for label-free MS analyses, and PeptideAtlas (Dataset identifier: PASS01219) for targeted MS analyses. The detailed descriptions of all analyses (raw and processed file names, sample name, biological replicate number, MS technical replicate number, corresponding figure) are summarized in Supplementary Data 8.

Results

Design and validation of the SRM assay to determine 20S proteasome status

The method aimed to simultaneously determine the absolute 20S proteasome quantity and subtype stoichiometry in a single assay. To achieve this goal, we designed a workflow combining IDMS and SRM to exploit its multiplexing capacity, sensitivity and robustness. In terms of total 20S proteasome absolute quantification, preliminary optimizations were used to select the most appropriate mode of internal standardization to correct for a range of experimental biases (Figure 1B; Supplementary Information I-1). As detailed in Supplementary Information I-1 and Supplementary Figures 1-5, the isotope-labeled whole proteasome complex added to biological preparations clearly provide a much more robust absolute quantification method than spiking with individual isotope-labeled "AQUA" proteotypic peptides for each 20S subunit. Indeed, despite good analytical performance in terms of reproducibility and linearity, absolute quantification using AQUA peptides technology lacked accuracy. This critical issue was overcome by using the absolute SILAC quantification method. We produced and affinity-purified isotope-labeled standard proteasome and immunoproteasome to obtain highly pure

standards with a high level of subtype purity (min 99%) and with an excellent isotopic incorporation rate (min 95%) (Supplementary Information I-1, Sup Figures 3 & 4). The optimized quantification method then consisted in the distribution of a reference mixture of 20S proteasome subunits with carefully-controlled stoichiometry into protein lysates (Figure 1B). After digestion of proteins using trypsin, an SRM method including at least one peptide sequence (with three associated transitions for each light and labeled surrogate) for the 11 non-catalytic subunits (α 1-7; β 3, β 4, β 6, β 7) (Supplementary Data 3) was applied. With this method, absolute quantification of total 20S proteasome was achieved in protein lysates of eight different human cell lines of diverse origins (HEK 293T, HCT116, RKO, U937, HeLa S3, NB4, KG1a, and MRC5). 20S proteasome concentration determined by the SRM method was compared to those obtained with the reference ELISA method (Figure 2A). A high correlation was found between the two methods (coefficient of determination = 0.98; slope = 1.01). Altogether, these data demonstrate that the newly-developed method is suitable for absolute quantification of 20S proteasome in human cell lysates.

Next, we implemented the method to detect the six standard and immune-specific catalytic subunits of the 20S proteasome (β 1, β 2, β 5, β 1i, β 2i, β 5i), as the absolute quantification of these subunits is required to determine the stoichiometry of the different 20S proteasome subtypes present in biological samples (Figure 1). Two to three proteotypic peptide sequences (and three or four associated transitions for each light and labeled surrogate) were carefully chosen, paying attention to favoring wide distribution over the protein sequence. Transitions and detection were then optimized for the six catalytic subunits. In total, the final method to quantify all α and β subunits comprised 206 independent MS transitions (103 "light" transitions and 103 "heavy" transitions corresponding to heavy surrogate peptides) associated with optimized dwell times and voltages for detection by the mass spectrometer (Supplementary Data 3). The method to quantify of the absolute levels of all six 20S proteasome catalytic subunits into the 20S proteasome (Supplementary Information I-2). Based on the excellent accuracy (97 ± 2%) and variability (CVs below 15%) of the method to measure the

absolute levels of catalytic subunits (Supplementary Information I-2 and Supplementary Figure 7B&D), these data were further used to determine changes in the stoichiometries of the four 20S proteasome subtypes observed in a model of IFN γ -treated HeLa cells (sP20S, β 5i P20S, β 1i β 5i P20S, and iP20S – no β 5t P20S nor α 4s P20S detected) (as detailed in Experimental Procedures) (Figure 2B). As expected, iP20S was highly and significantly increased (by a factor of 12), becoming the major proteasome subtype after 96 h stimulation with IFN γ , whereas sP20S levels dropped by nearly 3-fold (Figure 2B). Strikingly, the two intermediate 20S immunoproteasome subtypes, β 5i P20S and β 1i β 5i P20S, which each represented around 10% of the total 20S present, did not vary significantly after the cytokine stimulation. This interesting result shows that the main impact of IFN γ stimulation on the composition of proteasome complexes is the progressive replacement of the sP20S by the iP20S subtype.

Determining the stoichiometry of 20S subtypes in human tissues of broad origins

Next, we applied the method to determine the absolute quantities and precise stoichiometries of 20S proteasome subtypes in 11 protein lysates extracted from human tissues of broad origins (Figure 3). In most tissues, the proteasome represented 0.2 to 1% of the total protein present (Figure 3A). A shotgun label-free LC-MS/MS proteomics analysis exclusively identified and quantified α 4s and β 5t in testes and thymus samples, respectively, confirming their tissue-specificity (4), whereas the immunocatalytic subunits (β 5i, β 1i, and β 2i) were much more widely distributed across tissues (Figure 3B). The stoichiometries of the six major 20S proteasome subtypes (Figure 1A) were determined based on the absolute quantities of α and β subunits measured using the optimized method (as explained in Experimental Procedures). The absolute level of the catalytic β 5t subunit provided the abundance of the thymoproteasome (Figure 1A). Because this protein is not incorporated into labeled sP20S and iP20S internal standards, the absolute quantity of β 5t was deduced from absolute amounts of total P20S, β 5 and β 5i (Figure 3C). Similarly, the α 4s subunit (PSMA7L protein) which is representative of the spermatoproteasome could not be directly quantified. Although α 4 and α 4s are very similar in

sequence, three peptide sequences that are specific to the PSMA7 protein (α 4) were detectable and were used to distinguish between the two isoforms (Supplementary Figure 8A). The absolute level of α 4s was deduced from standard P20S and α 4 contents (Figure 1B, Experimental Procedures). As in the shotgun proteotypic peptides corresponding to the thymoproteasome assay, and spermatoproteasome were only observed in thymus and testes tissues, respectively, (Figure 3B) and these two 20S proteasome subtypes represented only 20% and 12% of the total proteasome pool in these organs, respectively (Figure 3C). A principal component analysis (PCA) biplot based on catalytic subunit composition (Figure 3D) grouped brain, muscle, heart, and testes tissues as they almost exclusively contain standard proteasome β subunits (β 1, β 2, and β 5). Bone marrow, spleen, and thymus, were observed on the opposite side of the graph because they contain high levels of β_{1i} , β_{2i} , and β5i immunosubunits. Interestingly, intermediate 20S subtypes (β5i P20S and β1iβ5i P20S) were only detected in low abundance (less than 10% of the total 20S proteasome pool) in bone marrow and spleen, suggesting that the immunoproteasome is the most important proteasome subtype for antigen processing among immunosubunit-containing subtypes. Liver, colon, ovary and lung tissues were placed at an intermediary position on the PCA graph (Figure 3D), with 50 to 70% of immunosubunitcontaining 20S proteasome subtypes (Figure 3C).

Interestingly, the absolute amount of the α 4 subunit, measured using peptides common to PSMA7 and PSMA7L proteins (Sup Figure 8A-B), was stoichiometric and highly correlated with the total 20S proteasome amount (measured using the α and β non-catalytic subunits, apart from α 4) both in the panel of eight cell lines (R² = 0.97; slope = 0.99) and in tissues of diverse origins (R² = 1.00; slope = 1.01) (Sup Figure 8 C-D). This result once again confirms the accuracy and precision of the method developed while also emphasizing the need for absolute quantification to detect perturbations in proteasome subunit expression or incorporation of subunits into mature proteasomes. It also indicates that this broad panel of normal and tumor human cells do not contain the non-canonical alternative α 4- α 4 20S proteasome complex in which a second copy of α 4 occupies the position normally held by α 3 (38, 39).

The abundance of some proteasome subunits measured using our optimized MRM assay was compared with mRNA expression data retrieved from the human protein atlas database (40) in corresponding tissues (Supplementary Figure 9). Although positive correlations were observed between some proteins and mRNA molecules abundances, these results suggest that measuring mRNA levels are not suitable when seeking to predict protein expression levels for 20S proteasome subunits in human tissues.

Proteasome status is affected by priming ADSCs with IFNy during their ex-vivo expansion

Thanks to their immunosuppressive properties, MSCs are considered a promising tool for cell therapy. Much effort has been directed toward enhancing MSCs activity by treatment with IFNγ. However, any ex vivo modifications may also fundamentally alter the cells, and understanding determinants that affect their immunomodulatory activity is essential if we are to develop effective MSC strategies. MSCs obtained from patients must be expanded ex vivo if they are to be used in the clinic, and this process must be carefully controlled (41). Recently, we reported a decrease in the immunosuppressive properties of ADSCs cultured up to the senescent stage and demonstrated that this effect was the result of proteasome-mediated indoleamine 2,3 dioxygenase (IDO) degradation (42). Priming of MSCs with IFN γ has been shown to extensively potentiate their therapeutic activity (43, 44) but, to our knowledge, and although proteasome functions cover a broad spectrum of biological functions, the proteasome status has never been assessed in this context. When used to measure the consequences of IFN_Y-stimulation of *ex vivo*-expanded patient-derived ADSCs on proteasome status (Figure 4A), our method revealed no major changes in the total 20S proteasome quantity (Figure 4B). However, a massive replacement of standard catalytic subunits by their immunological counterparts was clearly observed (Figure 4C), resulting in a strong shift from the standard to the immunoproteasome subtype (Figure 4D). Indeed, while the standard subtype represented 75% of the 20S proteasome pool when ADSCs were expanded in an IFN_Y-free medium, this proportion was decreased to 25% after four days'

culture in the presence of the cytokine (Figure 4D). The massive induction of the immunoproteasome by cytokine stimulation is likely to have a considerable impact on the repertoire of degraded proteins and antigenic peptides loaded onto major histocompatibility complex class I (MHCI) molecules.

Expansion of ADSCs under different O_2 concentrations affects 20S proteasome status and their capacity to differentiate

Conventionally, MSC culture for clinical applications is performed under normoxic conditions (21% oxygen tension), even though oxygen levels within tissues are typically much lower (hypoxic) than these standard culture conditions. Therefore, oxygen tension represents an important environmental factor that may affect how MSCs perform *in vivo*. However, the impact of hypoxic conditions on distinct mesenchymal stem cell characteristics, such as the proteasomal status, still remains unclear.

We applied our method to analyze the 20S proteasome status after 10 days of ADSCs expansion under three different oxygenation concentrations: 1%, 5%, and 20%. Interestingly, proteasome abundance, proteolytic activity and composition were all affected by dioxygen levels (Figure 5 A-C). Indeed, when comparing the effect of ADSC expansion in hypoxic or normoxic conditions (1% vs. 20% O₂ levels), we detected a significant increase in total 20S proteasome abundance alongside a nearly 2-fold significant upregulation of the three immunoproteasome catalytic subunits (Figure 5A&C). Both changes probably account for the higher 20S proteasome chymotrypsin-*like* activity measured following culture in normoxic conditions (Figure 5B). Indeed, structural differences in active sites account for the higher chymotrypsin-*like* activity of the β 5i subunit of the immunoproteasome compared to its β 5 standard counterpart (5, 45). Moreover, the relative abundance of 20S proteasome-associated PA28 $\alpha\beta$ activator is decreased in hypoxia compared to normoxia (p = 0.02) while the O₂ level had no significant impact on the level of association of the other 20S CP regulators (Supplementary Figure 10B). This last observation further supports the increase measured in immunoproteasome subtype in normoxic compared to hypoxic conditions as PA28 $\alpha\beta$ is known to preferentially associate with the 20S immunoproteasome variant *in cellulo* (34).

Overall, in the absence of IFNγ in the culture medium, the major proteasome subtype observed in ADSCs was the standard proteasome, which represented 60 to 80% of the total proteasome pool, depending on patient and O₂ level (Sup Figure 9A). Given the massive induction of immunoproteasome upon exposure to the pro-inflammatory cytokine (Figure 4C) and the absence of immune cells in the ADSC culture medium, the inter-individual differences observed in immunoproteasome content might result from variability in the inflammatory context before surgery. Immunoproteasome subunits have been shown to be required for adipocyte differentiation (29). Accordingly, our data indicate that high levels of immunoproteasome correlate with high adipogenic potential at 20% O₂, and the reverse under hypoxic conditions (Figure 5D). Strikingly, pharmacological inhibition of the immunoproteasome using the ONX0914 inhibitor (46) leads to a marked decrease in the capacity of ADSCs to differentiate into adipocytes, which confirms previous observations in the PSMB8 KO mice (29) using an orthogonal method.

To get a broader picture of the effect of O_2 levels on the proteome of ADSCs, a whole proteome analysis was performed on ADSCs grown under normoxic or in hypoxic conditions (1% of O_2). Importantly, a new cohort of three patients was used to obtain this new biological material. Out of 3624 proteins identified and quantified, 67 proteins were found at increased levels in hypoxic conditions and 129 were over-represented in normoxic conditions (Supplementary Figure 11 and revised Supplementary Data 7). All three immunocatalytic subunits were increased under normoxic conditions (fold increase of 1.6 to 1.9) although statistical significance was only reached for β_{11} and β_{21} (p-value = 0.05). As discussed above (supplementary Figure 7), this apparent discrepancy can easily be explained by the lower precision of label-free quantification results compared with SRM quantification data.

Functional analysis based on GO Biological processes revealed the most significant pathway related to the proteins for which abundance was modified both under hypoxic and normoxic conditions to be the

"oxidation-reduction" pathway (FDR = 1.2E-06 and 5.9E-03 for proteins up-regulated in hypoxic and in normoxic conditions, respectively). This result was to be expected following a change in O_2 levels from 20% to 1%. Other enriched GO biological pathways included "response to oxidative stress" (FDR = 4.2E-02) in the proteins that were more abundant in normoxic conditions. This pathway is particularly interesting as the immunoproteasome has been associated with responses to oxidative stress (47–49). ADSCs were also shown to negatively regulate oxidative stress in vivo (50) and notably through Nrf2 which is one of the main transcription factors involved in controlling oxidative stress. In our wholeproteome analysis, we observed that NQO1 (two-fold increase, p-value = 0.02), one of the best-known targets of Nrf2 (51), was increased under normoxic compared to hypoxic conditions. Reactive oxygen species seem to play an important role in adipocyte differentiation (52-54). A function of the immunoproteasome related to ADSC differentiation could be the clearance of proteins having undergone oxidative damage, when this type of damage accumulates it could be toxic for the cell. Significantly, 25 of the proteins for which overall abundance differences were detected are functionally related to the "cell differentiation" GO Biological Process (FDR = 1.4E-02 for proteins over-represented in cells grown under hypoxic conditions). Among these, 16 were increased in the hypoxic cells whereas 9 were decreased. These proteins are listed in Supplementary Data 7 (4th sheet). Given the importance of the immunoproteasome in regulating ADSC differentiation when stimulated with dexamethasone, and because our results clearly demonstrate that the immunoproteasome is increased under normoxic compared to hypoxic conditions, it is tempting to speculate that some of these deregulated proteins might be direct or indirect targets of the immunoproteasome. In particular, Transforming growth factor-beta-induced protein ig-h3(TGFBI/BGH3) and Plasminogen activator inhibitor 1 (PAI1) were both increased in our hypoxic cells (4.1- and 2.9-fold increase, respectively, with p-values of 8.5E-4 and 0.03, respectively), and these proteins are known to be target genes of TGF β 1, a pathway required for hypoxia-mediated inhibition of adipocyte differentiation (55). Of course, further experiments will be necessary to confirm that these proteins are downstream players of the immunoproteasome affecting the adipogenic potential of ADSCs.

In conclusion, after as few as 10 days of ADSCs expansion under different dioxygen levels, the method developed could detect mild but still significant changes in 20S proteasome status, and in particular variations in the level of 20S immunoproteasome. This change in proteasome composition might be causal for the observed O₂-induced variations in differentiation potential of ADSCs even though further investigation will be required to determine the precise mechanisms.

Discussion

This study aimed to develop an assay to determine subunit absolute quantity and stoichiometry within a highly heterogeneous macromolecular protein complex, the 20S proteasome. To achieve this goal, a robust, accurate and sensitive absolute quantification method with multiplexing ability had to be developed to allow the absolute quantification of the 19 different subunits of the 20S proteasome in a single assay.

Absolute quantitation based on IDMS using the SRM scanning mode suits the requirements for multiplex detection and is also recognized for its specificity, sensitivity, and accuracy (16, 17). Because many proteasome subunits had to be analyzed, we first used the most straightforward IDMS proteomics strategy, the AQUA (18) method, which involves spiking various isotope-labeled synthetic peptides into samples as internal standards. Although mainly used for the absolute quantification of monomeric proteins, the AQUA method has also been successfully applied to assess the substoichiometric incorporation of the Rpn13 ubiquitin receptor within the 26S proteasome (56). Our AQUA peptide sequences were carefully chosen to meet the requirements for IDMS analysis of proteins (16) and, when tested in simple or more complex biological matrices, they provided good linearity in signal response, low variability, and adequate sensitivity. These analytical performances explain why the AQUA technology was so rapidly adopted for the LC-SRM validation of protein biomarkers in various biological matrices (16, 17). However, we noted a defect in trueness that was easily detected in our system because non-catalytic proteasome subunits are expected to be present

at equimolar stoichiometry. Responses were observed to be highly dependent on peptide sequences, probably because of incomplete enzymatic digestion of the protein of origin; other reasons for the discrepancy observed could be incomplete peptide solubilization, peptide instability, or artefactual chemical modifications. Thus, isotope-labeled peptides could not be used to accurately assess proteasome subtype stoichiometry unless laborious optimizations were undertaken (19). Several IDMS methods based on the use of isotope-labeled protein standards like PSAQ (21), absolute SILAC (22), PrEST (23), or TAQSI (25) were developed to overcome the issues of the AQUA methodology. To the best of our knowledge, none of these has yet been applied to determine protein complex stoichiometry. To achieve this goal on highly heterogeneous 20S proteasome complex, we optimized a workflow relying on SILAC-based absolute quantification. A reference mixture containing equimolar concentrations of both isotope-labeled and purified 20S standard- and immuno-proteasome subtypes was produced, gualitatively and guantitatively assessed, and spiked into protein lysates extracted from cell cultures or tissues. After protein digestion, the SRM method was optimized for carefully chosen proteotypic peptides. In this approach, use of a whole isotope-labeled protein as internal standard represents a clear advantage over the PrEST approach (23) because all proteotypic peptides can theoretically be used for quantification. This method was validated by comparison with the reference ELISA method for total 20S proteasome absolute quantification in a broad range of biological samples.

Although 20S and 26S proteasome complexes can be routinely quantified using different ELISA assays (6, 10–14, 33), this technique lacks the multiplexing capacity to assess the complete diversity of proteasome subtypes because up to 19 different protein chains can be incorporated into the 20S proteasome macromolecular assembly (Figure 1A). Moreover, to our knowledge, the precise stoichiometries of tissue-specific β 5t- and α 4s-containing 20S proteasome subtypes have never yet been determined. Using our optimized assay, for the first time, we accurately quantified the stoichiometry of the tissue-specific thymoproteasome and spermatoproteasome; their low levels in thymus and testes tissues (20% and 12% of the total 20S proteasome pool, respectively) support

findings in previous reports showing that β 5t and α 4s are detected in a subpopulation of cells in these two organs (4, 58).

Then, we demonstrated that our method could also precisely monitor changes in 20S proteasome β catalytic subunit composition following IFN γ -activation of HeLa cells. We took advantage of the known stoichiometry of β catalytic subunits in the 20S proteasome to assess the trueness of the method, which exceeded 96%. In comparison, the TOP3 label-free quantification approach barely achieves 78% accuracy after discarding outliers arising from unrepresentative peptides. Converting peptide abundance into protein concentration requires careful selection of peptide sequences and optimization of SRM transitions (19) or, alternatively, the removal of incoherent peptides from global MS1-based quantification datasets, for instance using covariation of peptide abundances (59). Thus, even if label-free MS1-based quantification methods can be used to obtain a rough estimate of protein complex stoichiometry, as previously reported (60–62), these approaches are most appropriate for the high-throughput determination of changes in the relative abundances of protein complex subunits (34, 35) or estimation of protein copy-number without requiring spike-in standards (63).

MSCs obtained from bone marrow (BM-MSCs) or AT (ADSCs) are promising tools for cell therapy in regenerative medicine, to treat severe inflammatory and autoimmune diseases, or to prevent transplant rejection (64). Cells must be expanded *ex vivo* to obtain sufficient numbers for use in therapy. Thus, culture conditions must be carefully controlled both for safety issues and to optimize therapeutic effectiveness. Using our method, we showed that pre-stimulation of ADSCs with IFNγ and increasing O₂ levels both affect the status of the 20S proteasome during *ex vivo* expansion of primary ADSCs. In both cases, changes in proteasome composition were observed, in particular an increase in immunoproteasome stoichiometry. These results emphasize the high plasticity of the 20S proteasome when exposed to external stimuli, but also the multiple biological roles played by the immunoproteasome subtype. IFNγ is known to induce the immunoproteasome through the formation of new proteasome particles incorporating immunocatalytic subunits in place of the standard ones.

Pre-stimulation of both BM-MSCs and ADSCs with pro-inflammatory cytokines, in particular IFN γ , has been shown to increase their immunosuppressive properties (43, 64, 65), by increasing release of several soluble immunosuppressive factors, in particular kynurenine, a product of indoleamine 2,3dioxygenase (IDO) activity (65). Moreover, both BM-MSCs and ADSCs are antigen presenting cells (64) and, when exposed to IFN γ , they up-regulate expression of HLA class I molecules on their surface. This response protects them from NK-mediated lysis (43, 66), and promotes their immunomodulatory effects. Strikingly, both mechanisms linked to the immunosuppressive properties of MSCs are affected by proteasome status. Indeed, a shift from standard to immunoproteasome considerably increases the production of MHC class I-binding peptides in vivo (67) and IDO is known to be a proteasome substrate (68). Proteasome-mediated degradation of IDO was recently reported to explain the reduction in immunosuppressive potential observed in clinical-grade expanded MSCs which had reached replicative senescence (42). Mechanistically, IFN_Y is known to increase IDO protein via transcriptional activation (69) but it is also responsible for a strong induction of the immunoproteasome-PA28 $\alpha\beta$ complex, a proteolytic system functioning in the absence of ubiquitination (67). As IDO is degraded through the ubiquitination-dependent SOC3-proteasome pathway (68), the increase in ubiquitin-independent degradation triggered by IFNy might constitute a mechanism further promoting IDO stabilization in this physiological context.

In addition to its major role in antigen processing, the immunoproteasome seems to degrade oxidized proteins more efficiently than the standard 20S CP (70, 71). *De novo* synthesis of 20S proteasome and in particular of immunoproteasome is crucially important for maintaining efficient proteostasis in oxidative stress conditions (49). The evidence provided here of low total proteasome content and an absence of immunoproteasome in brain probably accounts for the previously reported higher sensitivity of brain tissues to oxidative stress (72). Activation of the immunoproteasome and autophagy occur during the early stages of ESC differentiation, to allow degradation of damaged proteins and avoid their transmission to differentiated cells (27). Moreover, recent findings indicate that the immunoproteasome is required for differentiation of adipocytes (29) and skeletal muscle (28).

In line with this requirement, our results demonstrate that ADSCs grown under hypoxic conditions, had a lower immunoproteasome content and exhibited a lower differentiation potential than their counterparts grown under 20% O₂. In addition, our results suggest that immunoproteasome activity is causal for the change in adipogenic potential in normoxic conditions. Thus, the significant increase observed in all three immunocatalytic subunits and in overall proteasome activity in normoxia-cultured primary ADSCs compared to the cells cultured in hypoxic environments could reflect modulation of their therapeutic capacities. Importantly, ADSCs are adult pluripotent stromal stem cells isolated from white AT where O₂ levels are below 5% but they are routinely expanded *ex vivo* under 20% O₂. Manufacturing practices for ASC expansion must therefore be carefully optimized and controlled to maintain their therapeutic capacity. The results from the approach developed here applied to primary ADSCs grown in conditions close to real clinical production clearly demonstrate that IFNγ and dioxygen levels could be key parameters in this process.

To conclude, the method presented here allows robust and rapid determination of the complete status, *i.e.*, absolute quantity and stoichiometry, of a highly heterogeneous macromolecular protein complex, the 20S proteasome, in various human tissues and cells. When applied to primary ADSCs expanded in different culture conditions, our results highlighted a high plasticity in proteasome composition and abundance which might be related to modulation of the ADSCs' immunosuppressive and differentiation properties. The method developed thus constitutes a sound approach to complement immunophenotyping or other methods to monitor protein markers (73) for the optimization and control of manufacturing processes for ADSCs expansion. Knowledge of proteasome composition is also of major interest for therapeutic purposes. Indeed, upregulation or dysregulation of immunoproteasome catalytic subunits have been observed in several human diseases and disorders, such as inflammatory and autoimmune diseases, cancer, diseases of the central nervous system, and aging (74). For instance, mutations reported in immunocatalytic subunits and the resulting defects in 20S immunoproteasome assembly and activity observed in PRAAS (Proteasome-Associated Autoinflammatory Syndrome) patients could be monitored using our method (reviewed in (75)). Our

assay could also be of great interest when seeking to assess patients before instigating immunotherapy as the efficacy of tumor antigen processing and presentation is closely linked to levels of 20S immune-type proteasomes in antigen presenting cells (75). Detection of increased concentrations of 20S proteasomes or changes in subtype profiles could also be monitored in extracellular body fluids to diagnose and/or as a prognostic marker of various diseases (76). In this context, a more specific targeting of the different heterogeneous forms of the proteasome will lead, in the long run, to more specific treatments generating fewer side effects and less chemoresistance than caused by broadspectrum proteasome inhibitors (77). However, to design such targeted therapies, tools to precisely determine patients' proteasome status, like the one presented here, will need to be developed.

More generally, the developed strategy could be extended to assess the absolute level, dynamic, and heterogeneous nature of many other biologically-relevant macromolecular systems such as the human spliceosomal hprp19/CDC5L complex (19, 61), the nuclear pore complex (78), core ribosomal proteins (62), or even host-pathogens interactions (79). These adaptations would be more accurate than the peptide-based mass spectrometry methods currently used (61). The sole requirement for this adaptation is that it must be possible to ectopically produce and purify each individual subunit in carefully-controlled absolute quantities with heavy-isotope incorporation.

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Footnotes :

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Figures legends

Figure 1: workflow for determination of total 20S proteasome absolute quantity and stoichiometry by LC-SRM

A- Schematic representation of the six main 20S proteasome subtypes with different sets of subunits. The sP20S is the most abundant 20S subtypes in most cell types. The 20S proteasome subtypes containing at least one immunocatalytic subunit are predominantly observed in immune cells but can also be induced in response to inflammatory cytokines. Other 20S proteasome subtypes are tissuespecific, like the thymoproteasome (β 5t P20S) and the α 4s proteasome (α 4s P20S) which have been observed in the thymus and in reproductive cells like testes, respectively.

B- Determination of total 20S proteasome quantity and 20S subtypes stoichiometry in Human cell lysates or tissues using SRM. Endogeneous proteasomes contained in biological samples were digested with trypsin. The SRM approach was designed to quantify in a multiplexed manner a set of proteotypic peptides corresponding to α subunits, as well as β catalytic and non-catalytic subunits. Isotope-labelled standard and immunoproteasome complexes were used as internal standards in the final method. They were spiked in the biological sample at equimolar quantities before the tryptic digestion. Alternatively, heavy « AQUA » peptides were added to the endogeneous peptides lysate after tryptic digestion. At least three endogeneous peptides, and three transitions per peptide were analyzed. Thanks to internal normalization with the isotopically-labeled peptides transitions, the absolute quantity of selected α and β non-catalytic subunits as well as of β catalytic subunits were obtained and used to compute the absolute quantities of total 20S proteasome and to assess the stoichiometries of the six main 20S proteasome subtypes in biological samples. Details are provided in the Experimental Procedures section and in Experimental section.

Figure 2: Final validation of the LC-SRM method to determine absolute quantity of total 20S proteasome and to precisely and accurately monitor the dynamics of 20S subtypes stoichiometries

A- Correlation curve between the 20S proteasome concentration measured by the ELISA reference method and the one determined by LC-SRM in a panel of eight human cell lines (HEK 293T, HCT116, RKO, U937, HeLa S3, NB4, KG1a, and MRC5). The LC-SRM method was based on the isotopic dilution of equimolar amounts of in-house isotopically-labeled sP20S and iP20S (R($^{13}C_6$) and K($^{13}C_6$)) in each cell line protein lysate. One peptide sequence (and three transitions for the light and the heavy surrogate) and three peptide sequences were analyzed for non-catalytic subunit (α1-7; β3,4,6,7) and catalytic subunits (β1,2,5,1i, 2i, 5i), respectively (see Supplementary Data 3 for more details on peptides and transitions). An equivalent of 2.5 μg total proteins containing the sP20S and iP20S heavy internal standard mix were injected on column and analyzed by SRM, as detailed in the Experimental Procedures. Three biological replicates (and three technical replicates) were analyzed to obtain statistics. The total proteasome concentrations measured in the initial protein lysates are presented.

B- Dynamics of 20S subtypes stoichiometries upon IFN γ stimulation of HeLa cells. The absolute quantities of each of the six catalytic subunits measured by the LC-SRM method were computed to calculate the stoichiometries of 20S proteasome subtypes as detailed in Experimental Procedures. HeLa cells were stimulated for 0, 24, 72, or 96 h with IFN γ and the SRM method used to quantify proteasome subunits in each total cell lysate obtained at each time point of cytokine stimulation. Peptides sequences, transitions and applied voltages are detailed in Supplementary Data 3. Three biological replicates were analyzed to obtain statistics. The sum of the stoichiometries of the four 20S proteasome subtypes is very close to the expected value of 100 (obtained from independent measurement of total 20S proteasome absolute quantity).

C- Method variability (CV %) and accuracy (%) calculated with data obtained from IFN γ -stimulated HeLa cells. Accuracy was calculated from comparison of total 20S proteasome quantity obtained from non-catalytic subunits (reference values) and from $\beta 1/\beta 1i$, $\beta 2/\beta 2i$, and $\beta 5/\beta 5i$ couples of catalytic subunits (experimental values) (n = 36). Accuracy were computed as follow: accuracy = (experimental value – reference value) / reference value * 100. Method variability: Coefficients of variation on all

independent measurements (n = 24). Details on stoichiometry determination of 20S proteasome subtypes are provided in Experimental Procedures.

Figure 3: The method is able to precisely and accurately monitor 20S proteasome absolute quantities and subtypes stoichiometries in a panel of human tissues

A: Proteasome 20S absolute quantity (% w/w total proteins) measured by the LC-SRM method and 20S Proteasome chymotrypsin-like activity measured by degradation of LLVY-AMC peptide, in a panel of 11 human tissues.

B: Label-free relative quantification of tissue-specific 20S proteasome subunits by label-free LC MS/MS reveals the tissue-specificity of α 4s and β 5t subunits but not the one of immunocatalytic subunits. Relative abundances of each subunits across the tissues are represented. The highest abundance is arbitrarily set to 1.

C: Stoichiometries of the six proteasome 20S subtypes in different Human tissue determined by the LC-SRM method. Details on stoichiometry determination of 20S proteasome subtypes are provided in Experimental Procedures.

D: Biplot Principle Component Analysis of 11 human tissue samples (3 replicates per tissue) based on 20S proteasome catalytic subunits composition. The dashed circles represent the main clusters observed depending on content on standard (β 1, β 2, β 5) or immuno (β 1i, β 2i, β 5i) subunits.

Figure 4: Proteasome status is affected by priming ADSCs with IFNy during their ex-vivo expansion

A: Workflow for the determination of the effect of priming of ADSCs with interferon- γ on their 20S proteasome status, using the LC-SRM method. Primary cells obtained from two patients were used for the study.

B: Proteasome 20S absolute quantity (% w/w total proteins) measured by the developed LC-SRM method in ADSCs obtained from patients (n = 2 biological replicates – 3 technical replicates).

C: Fold change in the six catalytic subunits of the 20S proteasome during IFNγ stimulation. ADSCs were cultivated in the presence of the cytokine (100 ng/ml). Absolute quantities of the catalytic subunits were determined by the developed LC-SRM approach and fold changes are calculated relative to time 0h (n=2 biological replicates – 3 technical replicates).

D: Dynamics in the stoichiometries of 20S proteasome subtypes during the course of IFN γ stimulation of primary ADSCs derived from patients (n = 2 biological replicates – 3 technical replicates). Calculations are detailed in Experimental Procedures.

Figure 5: Expansion of ADSCs in different O₂ levels affect 20S proteasome status and ADSCs' capacity to differentiate

A: Proteasome 20S absolute quantity (% w/w total proteins) measured by the LC-SRM method in five patients derived primary ADSCs cultivated at three different levels of O_2 (1%, 5%, and 20%).

B: Proteasome 20S chymotrypsin-like activity measured by the *in vitro* degradation of the LLVY-AMC peptide by proteasome in ADSCs lysates after cultivation in three different levels of O_2 (1%, 5%, and 20%) (n = 5 patients – 3 technical replicates).

C: Relative content of standard and immuno catalytic 20S proteasome subunits measured by the method in five patients derived primary ADSCs cultivated at three different levels of O_2 (1%, 5%, and 20%).

D: Differentiation potential of human ADSCs from different patients under different levels of O₂ (1%, 5%, and 20%). The immunoproteasome inhibitor ONX-0914 alters the differentiation capacity of ADSCs at 5% and 20% O₂ concentrations. Adipogenic differentiation is indicated by Oil red O staining.

Figure 1









Figure 4





