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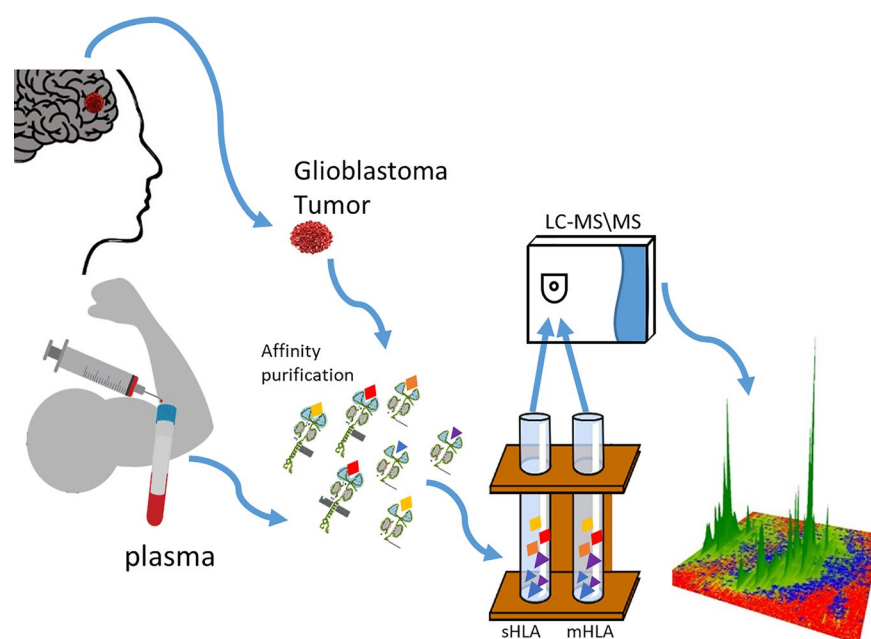
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In Brief

Glioblastoma is an aggressive brain tumor, thus early detection and immunotherapy may improve survival. This study includes large-scale analyses of the peptidome of the plasma-soluble HLA molecules of Glioblastoma patients and healthy controls, and the membranous HLA of the patients' tumors. These HLA peptidomes contain many HLA peptides derived from tumor antigens, providing potential opportunities for early diagnosis, and possibly also for personalized immunotherapy. The results emphasize the usefulness of the plasma-soluble HLA peptidomes as a source for biomarkers.

Graphical Abstract



Highlights

- The plasma-soluble HLA peptidome of Glioblastoma contains many tumor antigens.
- This sHLA peptidome may serve as a rich source for disease biomarkers.
- The sHLA peptidomes are highly correlated with the tumor's membranous HLA peptidomes.
- The HLA peptidomes differ significantly from the proteomes of the tumors.

Identification of Tumor Antigens Among the HLA Peptidomes of Glioblastoma Tumors and Plasma*

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Glioblastoma multiforme (GBM) is the most aggressive brain tumor with poor prognosis to most patients. Immunotherapy of GBM is a potentially beneficial treatment option, whose optimal implementation may depend on familiarity with tumor specific antigens, presented as HLA peptides by the GBM cells. Further, early detection of GBM, such as by a routine blood test, may improve survival, even with the current treatment modalities. This study includes large-scale analyses of the HLA peptidome (immunopeptidome) of the plasma-soluble HLA molecules (sHLA) of 142 plasma samples, and the membranous HLA of GBM tumors of 10 of these patients' tumor samples. Tumor samples were fresh-frozen immediately after surgery and the plasma samples were collected before, and at multiple visits after surgery. In total, this HLA peptidome analysis involved 52 different HLA allotypes and resulted in the identification of more than 35,000 different HLA peptides. Strong correlations were observed in the signal intensities and in the repertoires of identified peptides between the tumors and plasma-soluble HLA peptidomes of the individual patients, whereas low correlations were observed between these

HLA peptidomes and the tumors' proteomes. HLA peptides derived from Cancer/Testis Antigens (CTAs) were selected based on their presence among the HLA peptidomes of the patients and absence of expression of their source genes from any healthy and essential human tissues, except from immune-privileged sites. Additionally, peptides were selected as potential biomarkers if their levels in the plasma-sHLA peptidome were significantly reduced after the removal of tumor mass. The CTAs identified among the analyzed HLA peptidomes provide new opportunities for personalized immunotherapy and for early diagnosis of GBM. *Molecular & Cellular Proteomics* 18: 1255–1268, 2019. DOI: 10.1074/mcp.RA119.001524.

Glioblastoma multiforme (GBM)¹ is the most common and aggressive primary brain tumor, with a median survival of about 15 months (1, 2). The grim prognosis facing most GBM patients calls for improvements in targeted therapeutic treatments, as well as in discovery of proper biomarkers for early

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detection of the disease. Recently, immunotherapy of different cancers, including GBM, has drawn significant attention (3, 4), reviewed in (5–13). Much of the recent excitement about cancer immunotherapy stems from the success in treating patients with immune checkpoint modulators, which induce anti-cancer T cell immune reactions that can break tolerance and bring about complete responses in increasingly larger percentages of patients (14, 15). Identification of tumor antigens is needed for development of effective cancer immunotherapy, including GBM, and a good source for such antigens are the pools of HLA-bound peptides presented preferentially, or even exclusively, by the tumor cells (16), reviewed in (17–20). Tumor-specific antigens (TSA) and tumor-associated antigens (TAA) that can serve as candidates for cancer immunotherapeutics have been searched for extensively. Indeed, many such antigens were already identified, yet none has induced sufficiently strong anti-cancer immune reaction to eradicate the tumors in large cohorts of patients (18, 20, 21). One subset of preferred TAAs are Cancer/Testis Antigens (CTA), which are abnormally expressed in the malignant cells, and are normally expressed only in fetal tissues and in immune privileged sites, such as the male germ cells, placenta and ovary, but are absent from the normal somatic cells of any healthy tissue (22), reviewed in (23). Another special groups of tumor antigens are neoantigens, which are attracting significant attention as potential targets for cancer immunotherapy (24). These are more likely common in tumors with higher mutational load and therefore less frequently found in GBM.

HLA class I molecules are predominantly expressed as membrane anchored, cell-surface proteins (mHLA). Yet, the HLA molecules are also released from cells (25, 26), reviewed in (27), and can be recovered from the soluble HLA molecules in human plasma (sHLA) for analysis of their bound peptidome (28–30). Confounding factors that limit the use of the sHLA peptidome as a source for biomarkers are the presence of small amounts of sHLA molecules in the plasma of all healthy individuals (27, 31), and in addition, some of the sHLA alleles are released to the circulation in larger amounts than others (32, 33).

It has been suggested that shedding of the sHLA molecules represents a mechanism for evasion of immune recognition of tumor cells, which release larger amounts of sHLA molecules relative to the healthy cells (33). Because the sHLA molecules carry with them their original load of bound peptides, the immunopeptidomes of the sHLA molecules of cancer patients include disproportionally large fractions of peptides originating from the tumor cells, even though the tumors constitute

just small fraction of the body mass. It is therefore intuitive to assume that biomarkers of disease can be found among those sHLA peptides. Such plasma-sHLA molecules could then be a useful source of cancer biomarkers for early detection of recurrence during follow-up (28).

Analyses of the HLA peptidomes, based on immunoaffinity purification of the mHLA molecules from cell lines, tumor tissues or patients' plasma, followed by analysis of the bound peptides by chromatography and tandem mass spectrometry (34) currently results in identification of thousands of HLA bound peptides (35–38). HLA peptides were previously isolated from GBM cell lines (39) and fresh tumors, including GBM (4, 16). Similarly, the HLA peptidomes of tumors, such as melanoma (40, 41), renal cell carcinoma (42, 43), leukemia (44) and multiple myeloma (45) have been characterized. Further, several thousands of peptides have successfully been isolated from cancer patients' plasma-sHLA molecules (28, 30).

In this research, more than 35,000 plasma-sHLA and tumor-mHLA peptides from GBM patients were identified. These large HLA peptidomes included several hundred peptides, derived from both known TAAs and newly defined potential CTAs. We propose that these may serve as useful biomarkers and candidates for immunotherapeutics for GBM, as well as for other cancers.

EXPERIMENTAL PROCEDURES

Experimental Design and Statistical Rationale—One hundred six plasma samples were obtained from 52 different GBM patients, collected prior to tumor removal and at different time points after the surgery (supplemental Table S1). Additionally, control plasma of healthy donors ($n = 6$) and patients diagnosed with ankylosing spondylitis ($n = 30$), as an exemplar inflammatory process, were used for this study. In addition, tumor samples were obtained from 10 of the 52 GBM patients for mHLA peptidome analysis, as well as for tumor proteome analysis.

Patient Characterization—Peripheral blood (PB) and tumor tissues samples from GBM patients (University Hospital Heidelberg, Leiden University Medical Centre, Vall d'Hebron University Hospital, Université de Genève, Southampton Universities Hospital, Heidelberg University Medical Center, Herlev Hospital, and Rigshospitalet) and non-cancerous controls (Universitario Central de Asturias and Bnei-Zion Hospital). The HLA tissue typing of the GBM patients and noncancerous controls, healthy, and Ankylosing Spondylitis patients are listed only with gender, age, and stage of disease in supplemental Table S1. All human bio-specimens were obtained with informed consent, and after approval of the relevant ethics committees.

Plasma and Tumor Cells Collection—Peripheral blood samples were collected into EDTA tubes and cleared of the cells by centrifugation for 10 min at $1200 \times g$, at room temperature. The cleared plasma was stored frozen at -80°C until use for sHLA purification. Tumor samples were frozen in liquid nitrogen immediately after dissection and stored at -80°C until use, for both mHLA peptidome and proteome analyses of the same tissue extracts.

Affinity Purification of HLA Molecules—Membranal HLA class I molecules were purified from GBM tumor tissues by immunoaffinity from the cells after lysis with 0.25% sodium deoxycholate, 0.2 mM iodoacetamide, 1 mM EDTA, 1:200 Protease Inhibitors Mixture (Sigma), 1 mM PMSF, and 1% octyl-D-glucopyranoside (Sigma) in

¹ The abbreviations used are: GBM, glioblastoma; MHC, major histocompatibility complex; HLA, human leukocytes antigen; CTA, cancer/testis antigen; TSA, tumor specific antigens; TAA, tumor associated antigens; sHLA, soluble human leukocytes antigen; mHLA, membranal human leukocytes antigen; FDR, false discovery rate; BBB, blood brain barrier.

PBS at 4 °C for 1 h. The lysate was cleared by centrifugation for 45 min at 18,000 rpm, at 4 °C. HLA class I molecules from the cleared lysate or from the fresh human plasma were immunoaffinity purified using the W6/32 mAb bound to Amino-Link beads (Thermo-Fisher Scientific, Waltham, MA) as in (28, 46). The HLA molecules with their bound peptides were eluted from the affinity column with five column volumes of 1% TFA. The eluted HLA class I proteins, and the released peptides were loaded on disposable C18 micro-columns (Harvard Apparatus, Holliston, MA), and the peptides fraction was recovered with 30% acetonitrile in 0.1% TFA, whereas the protein fraction was recovered with 80% acetonitrile in 0.1% TFA, as in (46). The peptide fractions were dried using vacuum centrifugation, reconstituted in 100 μ l of 0.1% TFA, reloaded on C18 Stage-Tips (47), eluted with 80% acetonitrile, dried, and reconstituted with 0.1% formic acid for LC-MS/MS analysis.

Proteomics analysis was performed after trypsin digestion of the proteins in gel slices. Briefly, 40 μ g protein sample from each tissue was ran in 10% acrylamide gel, stained with Coomassie, and each lane of the gel was sliced into five slices. The proteins in the gel slices were reduced with 2.8 mM DTT at 60 °C for 30 min and carbamidomethylated with 8.8 mM iodoacetamide in 100 mM ammonium bicarbonate at room temperature for 30 min and digested overnight at 37 °C in 10% acetonitrile and 10 mM ammonium bicarbonate with modified trypsin (Promega) at a 1:10 (wt/wt) enzyme-to-substrate ratio. This was repeated for another 4 h with a similar amount of trypsin, followed by 150 min gradients of LC-MS/MS of the tryptic peptides from each gel slice.

Identification of the HLA and Tryptic Peptides—Both the HLA and tryptic peptides were resolved by capillary chromatography and electrospray tandem mass spectrometry using a Q-Exactive-Plus mass spectrometer fitted with a capillary Ultimate 3000 RSLC nano-capillary UHPLC (Thermo-Fisher Scientific). The reversed phase chromatographs were with about 30 cm long, 75-micron inner diameter capillary columns, home-packed with 3.5 μ m silica ReproSil-Pur C18-AQ resin (Dr. Maisch GmbH, Ammerbuch-Entringen, Germany) as in (48). HLA and tryptic peptides were eluted with a linear gradient of 5–28% of acetonitrile in 0.1% formic acid. For both the HLA peptides and for the in-gel digested fractions, the capillary HPLC gradients were run at flow rates of 0.15 μ l/minutes during 120 min. Data was acquired using a data-dependent “top 10” method, fragmenting the peptides by higher-energy collisional dissociation (HCD). Full scan MS spectra were acquired at a resolution of 70,000 at 200 m/z with a target value of 3×10^6 ions. MS/MS ions were accumulated to an AGC target value of 10^5 with a maximum injection time of 100 msec. For the HLA peptides with unassigned precursor ion charge states, or charge states of four and above, no fragmentation was performed. For tryptic peptides, the fragmentation was performed on charge states between 2 to 7. The peptide match option was set to Preferred. Normalized collision energy was set to 25% and MS/MS resolution was 17,500 at 200 m/z . Fragmented m/z values were dynamically excluded from further selection for 20 s.

Data Analysis—The MS data was analyzed by the MaxQuant computational proteomics platform (49) version 1.5.3.8 and searched with the Andromeda search engine (50). Peptide identifications were based on the human section of the UniProt database (<http://www.uniprot.org/>) of July 2015 containing 69,693 proteins from 91,459 entries, including one sequence of mutated EGFR-VIII. An additional FASTA file, containing 638 entries of mutated peptides (sequences in supplemental Table S10) was used during the same MaxQuant analysis (however, none of these was identified in the HLA peptidome analyses). Proteins were declared positive identification with at least two identified tryptic peptides per protein. Mass tolerance of 4.5 ppm for the precursor masses and 20 ppm for the fragments were allowed. Methionine oxidation was accepted as variable modification for both

tryptic and HLA peptides. Carbamidomethyl cysteine was accepted as a fixed modification for the proteomics data and as a variable modification for the HLA peptidome data. Methionine sulfoxide and *n*-acetylation were set as variable modifications for both the proteomics and HLA peptidomics analyses. Minimal peptide length was set to seven amino acids and a maximum of two miscleavages was allowed for tryptic peptides. The false discovery rate (FDR) was set to 0.01 for protein identifications, and 0.05 for the HLA peptides, because it resulted in identification of about twice as many true HLA ligands (see Discussion). The resulting identified protein tables were filtered to eliminate the identifications derived from the reverse database, as well as common contaminants.

Normalization of the HLA Peptidome and Proteome Data—The HLA peptidome of each of the LC-MS/MS runs was normalized according to its median. The medians were zeroed, and downregulated HLA peptides were defined as those changing by at least two folds in the plasma samples collected immediately before surgery, relative to samples collected at later clinic visits. The proteomics LC-MS/MS data was normalized using the LFQ and iBAQ tools of the MaxQuant software (49) and the statistical evaluation of the data was performed with Perseus (51).

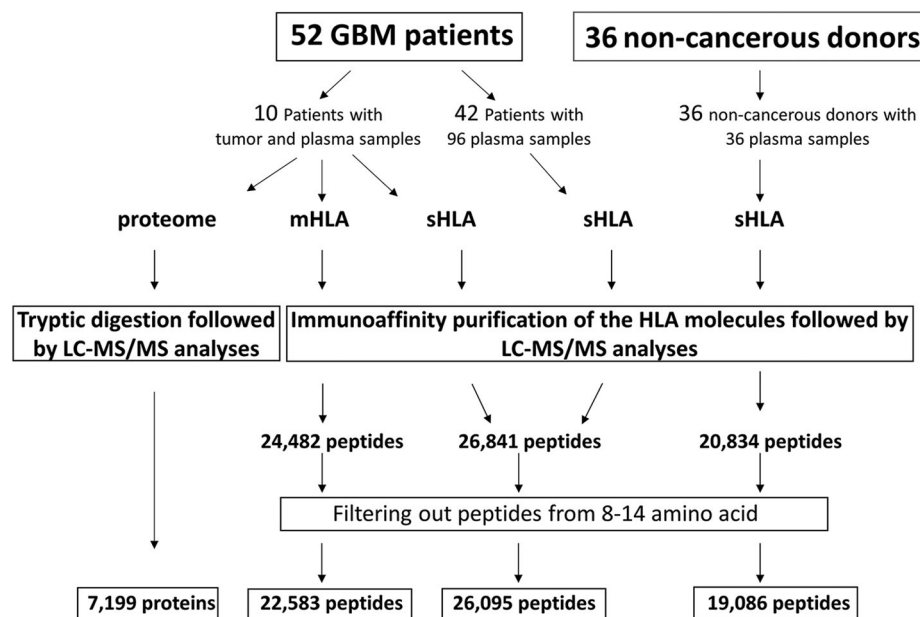
HLA Typing—DNA was extracted from blood of GBM patients using the DNeasy Blood & Tissue kit (Qiagen, Hilden, Germany) as per the kit instructions. The HLA typing was conducted with this DNA at the DKMS Life Science Lab (Germany). The samples were processed within high-throughput workflow using next generation sequencing. HLA typing was based on the regions covering the peptide-binding domains. Regions sequenced were Exons 2 and 3 for HLA-A, -B, -C, -DQB1, -DRB1, and -DPB1 (52).

Definition of the Tumor Associated Antigen (TAA) Group—Selection of TAAs was based on the cancer tumor database (CT gene database) (<http://www.cta.lncc.br/>) (53) comprised of 277 different TAAs (data accumulated between the years 2005–2009) and the Tumor T cell Antigen database (TANTIGEN) (<http://cvc.dfci.harvard.edu/tadb>), comprised of 259 different TAAs (54).

RESULTS

Large HLA Peptidomes Are Identified from Tumors and Plasma of GBM Patients—Tumor samples were obtained from 10 different patients undergoing surgery. These freshly frozen tumor tissues were used for analysis of their membranal HLA peptidomes and their tumor proteomes. In addition, 106 plasma samples from these 10 patients, and other 42 GBM patients (52 patients in total) were used for the analysis of their plasma-sHLA bound peptidomes. Additionally, control plasma of healthy donors ($n = 6$) and patients diagnosed with ankylosing spondylitis ($n = 30$), as an exemplar inflammatory process, were used for this study (Fig. 1, supplemental Table S1). The study was performed as part of the GAPVAC project (The European Glioma Actively Personalized Vaccine Consortium), which aimed to develop a personal immunotherapy treatment based on administration of synthetic copies of selected HLA peptides derived from tumor antigens to GBM patients (56). Here we describe the results of the plasma-soluble HLA (sHLA) peptidome analyses, the tumor membranal HLA (mHLA) peptidome and proteome analyses, performed in parallel with the GBM tumors. Overall, the tumor-mHLA peptidome analyses resulted in the identification of 22,583 different HLA peptides, derived from 7610 different source proteins, and the plasma-sHLA peptidome analyses resulted in the identification of 26,841 different HLA peptides,

Fig. 1. Overview of the sHLA and mHLA peptidome and proteome analyses schemes, and selection of likely HLA ligands.



derived from 8817 different source proteins (supplemental Table S2). The proteomics analysis of these 10 GBM tumor samples resulted in identification of 7199 different proteins (supplemental Table S3).

The GBM HLA Peptidomes Include Many Peptides from Multiple Tumor Antigens—Among the patients' plasma-sHLA and the tumor-mHLA peptidomes, 989 different HLA peptides derived from 179 known TAAs were identified. The reference set of TAAs was based on the cancer tumor database (CT gene database) (<http://www.cta.lncc.br/>) comprising 277 different TAAs (data accumulated between –2009) (53) and the Tumor T cell Antigen database (TANTIGEN) (<http://cvc.dfci.harvard.edu/tadb>) (54) with 259 different TAAs, for a total set of 496 TAAs.

Derived from these 496 TAAs, a total of 853 plasma-sHLA and 659 tumor-membranal HLA peptides were identified. Importantly, up to 63.3% of the plasma-sHLA peptides, derived from this TAA group, were also detected among the tumor-mHLA peptidomes of the different patients. These results indicate that the plasma-sHLA peptidome may indeed provide a useful source of tumor antigens for diagnosis and immunotherapeutics (selected examples in Table I, and the entire list in supplemental Table S4). 424 out of the 989 identified HLA peptides derived from TAAs were detected only in GBM plasma and tissue samples but not in any noncancerous donors' plasma samples, supporting their potential significance as authentic tumor antigens.

Among the TAAs expressed in the tumor cells, a subset of genes, normally expressed only in germline, embryonic and placenta cells, can be defined as CTAs, which are expressed at levels below a threshold of nine gcma units (expression units using background adjustment: GC content adjusted with Robust Multiarray Average, described in the BioGPS website)

in all normal and essential tissues. Such CTAs were further defined as those derived from genes whose transcripts are expressed at significantly higher levels in the tumors tissues (according to BioGPS (57, 58), see Discussion). Using this filtration, a list of 134 CTAs was established (supplemental Table S5). In the HLA peptidome analysis described here, 77 different HLA peptides, belonging to 32 CTAs, were identified. Of these, 36 HLA peptides were identified only in the GBM plasma and/or tumor samples (for example ACPP, supplemental Fig. S1A and the entire list in supplemental Table S2). In addition, HLA peptides derived from proteins expressed only in fetal tissues were observed in the GBM samples, and not at all, or in less than 5% of the control plasma samples. An example for such antigen is the transcription factor SOX11 gene (SOX11) that is expressed only in fetal brain, according to BioGPS (supplemental Fig. S1B). In this analysis, two of its derived HLA peptides, AHSASEQQL and NFSDLVFTY, were observed in the plasma-sHLA and tumor-mHLA of the GBM patients, and were not detected in any of the noncancerous blood donors' plasma-sHLA peptidomes (Table I). Moreover, HLA peptides derived from male tissues such as testis or prostate, and identified in women's plasma or GBM tissue, or HLA peptides derived from female tissues, such as placenta and ovaries, and identified in the men's plasma or GBM tissue, can serve as potential immunotherapeutics candidates (Table II). For example, six different HLA peptides originating from the ETV5 gene were identified in this analysis. This gene is expressed preferentially in the placenta. Moreover, the six ETV5 HLA peptides were identified mostly among the male GBM sHLA and mHLA samples relative to the noncancerous donors according to the BioGPS analysis, suggesting their relevance to serve as proper biomarkers for the disease.

TABLE I

List of HLA peptides derived from the TAA group, identified among the GBM sHLA and mHLA peptidomes

Gene name	Sequence	Count #		
		sHLA non-cancerous	sHLA GBM	mHLA GBM tissue
ETV5	FPDNQRPF	0	0	1
	GPAPAPHSL	0	10	1
	KVAGERYVY	0	2	3
	LPYAEFGAY	1	6	2
	MPGPPAHGF	1	0	3
SOX4	RPAMNYDKL	1	1	1
	KIMEQSPDM	3	8	4
	SAASASAAL	0	17	0
	SISNLVFTY	0	4	1
MDM2	GEISEKAKL	0	10	2
	YTMKEVLFY	0	0	1
DDR1	SEQETLVVR	2	1	1
	AVGDGPPRV	0	10	6
	FLAEDALNTV	1	4	7
	LPPPPQNSV	0	0	0
CSF1	YLQVDLQRL	1	6	2
	HSSGSLPLGELE	2	1	0
UBE2A	HTVDPGSAKQR	8	6	0
	REYEKRVSA	2	6	0
BST2	RLQEDPPAGV	1	15	6
	DASAEVERL	2	4	0
NPM1	AAAPQLLIV	0	8	0
	VPLIIFTI	0	16	1
	DENEHQLSL	0	4	1
	EGSPIKVTL	3	19	2
BCAP31	EITPPVVL	1	4	0
	FEITPPVVL	0	4	2
	HQLSLRTV	1	0	0
	MSPLRPQNYLFG	1	0	0
	NYEGSPIKVTL	0	0	0
	VEAEAMNY	1	5	1
	VEAKFINY	1	11	2
	YEGSPIKVTL	0	2	2
	AESASEAAKKY	3	8	1
	KLDVGNAAEV	1	11	3
CDKN2A	KYMEENDQL	1	3	0
	THVLEGAGNKL	0	5	1
	AAPGAPAAV	0	0	2
SART3	LPVDLAEL	0	6	2
	AAFTRALEY	1	10	1
	AEAPRLAEY	1	10	1
	ARLEKVHSL	0	1	4
PA2G4	AYIDFEMKI	2	10	0
	DHQVISVTF	0	5	1
	TVFVSNLPYSM	1	0	0
	DVAQGTQVTGR	1	2	0
COTL1	IAFPTSISV	2	16	1
	KEGEFVAQF	1	4	1
	TPIEGMLSH	0	8	2
	FVISDRKEL	4	10	3
AT1C	KTLTPISAAY	1	4	0
	NLYPFVKTV	0	3	5
GPNMB	AARLPDAA	0	3	0
	ALTSTLISV	0	7	4
	AYMREHNQL	3	10	1
	IENSPGNV	0	2	0
	IPTVCTII	0	0	1
	KGLSVFLNR	3	1	1
	VFLNRAKAVFF	1	1	0
	VLTSDSPAL	0	7	1
CDK4	ALTPVVVTL	0	7	5
	FGLARIYSY	1	8	2
SOX11	ISTVREVAL	3	4	1
	AHSASEQQL	0	2	1
	NFSDLVFTY	0	0	1

The Levels of Potential Biomarker Plasma-sHLA Peptides are Reduced Following Surgical Removal of the Tumors—The presence of the plasma-sHLA peptides derived from tumor antigens may serve as surrogate biomarkers for different cancers, including GBM. Peptides detected among the plasma-sHLA peptidome, derived from tumor antigens that are not normally expressed at elevated levels in normal tissues, should be reduced in their levels after removal or reduction of the tumor load by treatments (59) such as the surgery performed for the GBM patients studied here. Most of the plasma-sHLA peptides of the individual patients remain relatively stable before and after surgery, and even a few months later, because these are self-peptides of the healthy tissues (example in Fig. 2A). This facilitated focusing on the minority of HLA peptides that were reduced in their levels relative to the rest of the sHLA peptidome, to search among them for potential biomarkers. In this study, 94 different plasma samples were collected prior to, and following surgery of the same 34 GBM patients as part of the GAPVAC project. The sHLA molecules of the different plasma samples were affinity purified and their bound peptidomes were analyzed separately by LC-MS/MS. As many as 422 sHLA peptides, derived from 362 proteins, were down regulated by at least two folds in the plasma samples collected before and a few weeks after surgery, and remained low in the plasma samples collected at subsequent visits to the clinic. These HLA peptides were derived from genes/proteins from which no sHLA peptides were detected in any of the plasma samples of the noncancerous donors (supplemental Table S2, column B). Examples for such down-regulated peptides are displayed in Fig. 2A. Some of the downregulated peptides were derived from known cancer related genes, such as the HLA-A*32:01 peptide RVNPLVKSF of FRMD3, which is not expressed in any of the normal tissues of the body, according to BioGPS. The disappearance of sHLA peptides from the plasma samples after surgery can be because of real reduction in their amounts, or because of chance misidentifications caused by the shotgun LC-MS/MS approach used here (examples in Fig. 2A).

The Plasma-sHLA and Tumor-mHLA Peptidomes of the Same Patients Are Highly Correlated—The large HLA peptidomes of the patients' tumors-mHLA and plasma-sHLA enabled comparisons between these patients' peptidomes. The peptidome analyses of the ten patients, from which both types of samples were available, indicated overlaps of up to 75% of the identified sHLA peptides of the mHLA peptides of the individual patients (Fig. 2B). Multiple plasma samples of each patient were also considered as biological replica, and their LC-MS intensities were averaged. Many of the mHLA and the sHLA peptides of the individual patients were shared between both types of samples (supplemental Fig. S2A). In addition, the LC-MS signal intensities of these shared sHLA and mHLA peptides, of the individual patients, were significantly correlated with Pearson correlations between 0 and 0.56 (supplemental Fig. S2B). Most of these shared peptides

TABLE II
Examples for HLA peptides derived from typical male tissues and identified in women's plasma or GBM tumor tissue, and HLA peptides derived from typical female tissues and identified in men's plasma or GBM tumor tissue

Gene name	Sequence	CTA description	# of identifications					
			mHLA male	mHLA female	sHLA GBM male	sHLA GBM female	sHLA non-cancerous male	sHLA non-cancerous female
ETV5	FPDNQRPFL	expressed in placenta	0	1	0	0	0	0
	GPAPAPHS		0	1	4	6	0	0
	KVAGERYVY		3	0	2	0	0	0
	LPYAEGFAY		1	1	5	1	0	1
	MPGPPAHGF		2	1	0	0	1	0
	RPAMNYDKL		0	1	1	0	0	1
ACPP	SAHDTTVSGLQ	expressed only in prostate	0	0	0	0	0	0
	AHDTTVSGL		1	0	0	0	0	0
TTK	SPNSILKAA	expressed only in testis and fetal organs	0	0	0	2	0	0
SPAG17	DQEKEKEKEK	expressed only in testis	0	0	0	0	0	0
ODF2	KILDLETQL	expressed only in testis	3	1	2	6	1	2

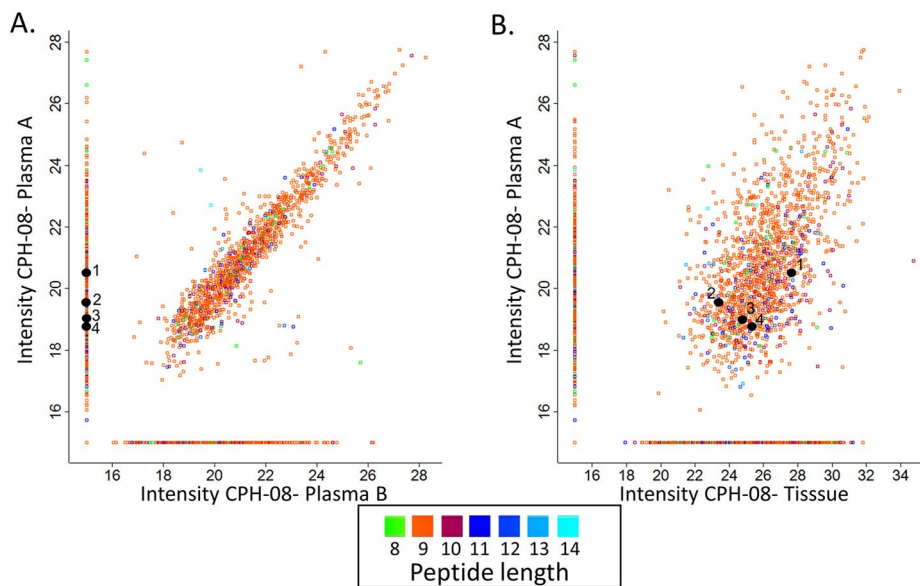


FIG. 2. Relative LC-MS signal intensities of the sHLA peptides identified in plasma A (before surgery) and plasma B (at least 4 weeks after surgery) of patient CPH-08. The group of HLA peptides detected only in plasma A are indicated on the vertical line and those observed only in plasma B are on the horizontal line with arbitrary numbers of 15 (A). Comparison between the plasma A sHLA peptidome and the tumor mHLA peptidome of patient CPH-08 (B). The LC-MS signals of the peptides are indicated on a Log2 scale. The numbered HLA peptides (1–4) were identified in plasma A and tissue but not in plasma B: 1) YERDEDNNL; 2) EHTEVLQLL; 3) NHALPLPGF; 4) EIIEKDTKY.

were true HLA ligands and received NetMHC scores below rank of 2%, according to the patient's HLA alleles. For example, in patient "11-002", as many as 1194 out of the 1324 (about 90%) of the sHLA peptides of plasma A (taken before surgery) and fitting this definition as HLA ligands, were shared with the tissue mHLA peptidome (supplemental Table S2). Importantly, many of the shared HLA peptides were derived from the TAA group (supplemental Table S4).

Similarity Between HLA Peptidomes of Individuals Sharing HLA Allotypes—As expected, both the tumor-mHLA and plasma-sHLA peptidomes of different blood donors were more similar when they shared HLA allotypes (examples in Fig. 3) and these patients shared more TAAa and CTAs peptides

(supplemental Table S6) even though only small fractions of the peptidomes were shared among most patients (Fig. 4, supplemental Table S7). The shared peptides identified in the plasma-sHLA peptidomes of people harboring very different HLA allotypes that belong to dissimilar HLA supertypes, are likely contaminating peptides, co-purifying with the HLA molecules during the affinity purification. Further, peptides that fit the sequence motifs of the HLA allotypes of the patients are more likely authentic HLA ligands of these allotypes (Fig. 3).

The HLA Peptidomes Do Not Correlate With the Proteomes of the Tumors—The protein repertoires and their levels (measured in iBAQ values) of the different tumors, were much better correlated than the HLA peptidomes of the different

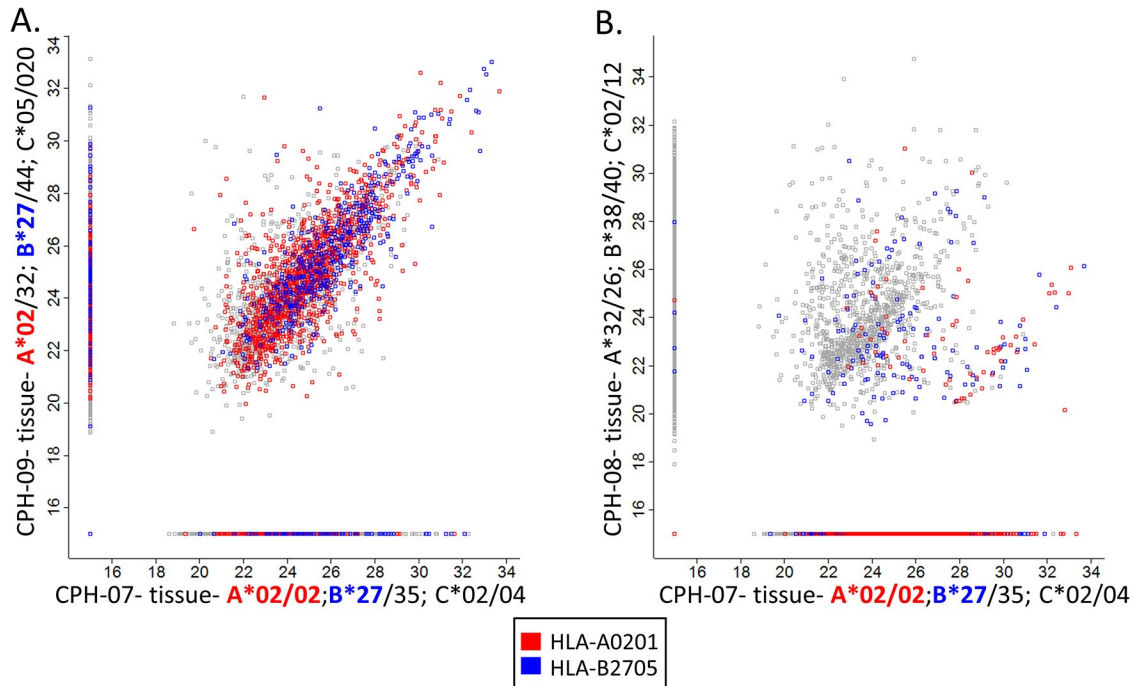


Fig. 3. **Comparisons of the mHLA peptidomes of three patients' tumors.** The HLA alleles of the patients are as following: CPH-07- A*02:01/02:01; B*35:01/27:05; C*02:02/04:01, CPH-09- A*02:01/32:01; B*27:05/44:02; C*05:01/02:02, CPH-08- A*32:01/26:01; B*38:01/40:02; C*02:02/12:03. Patients CPH-07 and CPH-09 share the same A*02:01 and B*27:05 HLA alleles (A). Patients CPH-08 and CPH-07 have completely different HLA alleles (B). The NAN values were replaced with arbitrary value of 15, the axes are on a Log2 scale and the color represents the HLA allele.

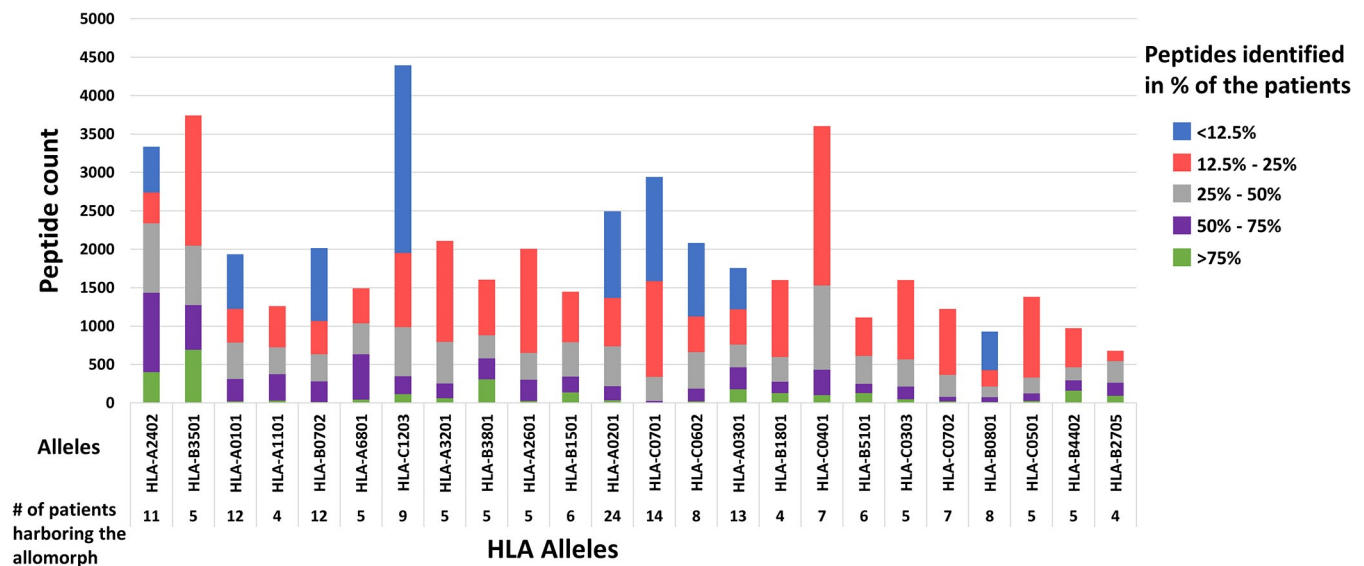


Fig. 4. **Numbers of HLA peptides shared between different patients harboring the same HLA allomorphs.** The bars represent the peptides counts that were detected in multiple mHLA and sHLA peptidomes of different GBM patients and the colors indicate the percentage of patients in which such peptides were detected.

tumors (Fig. 5, supplemental Fig. S3). As many as 4834 different proteins were identified in all ten tumors analyzed, out of the total number of 7199 proteins identified (supplemental Table S3). Importantly, 1652 different HLA peptides, derived from 947 of the proteins identified in all the tumor samples, were also detected in the tumor tissues mHLA and plasma

sHLA peptidomes. None of these peptides was derived from the source genes of the sHLA peptides that were detected in the noncancerous donors' plasma (supplemental Table S2, column E). Among these 1652 different peptides, a subset of 18 HLA peptides belong to the TAA group (Table III). Such genes, whose proteins products are expressed in the tumor

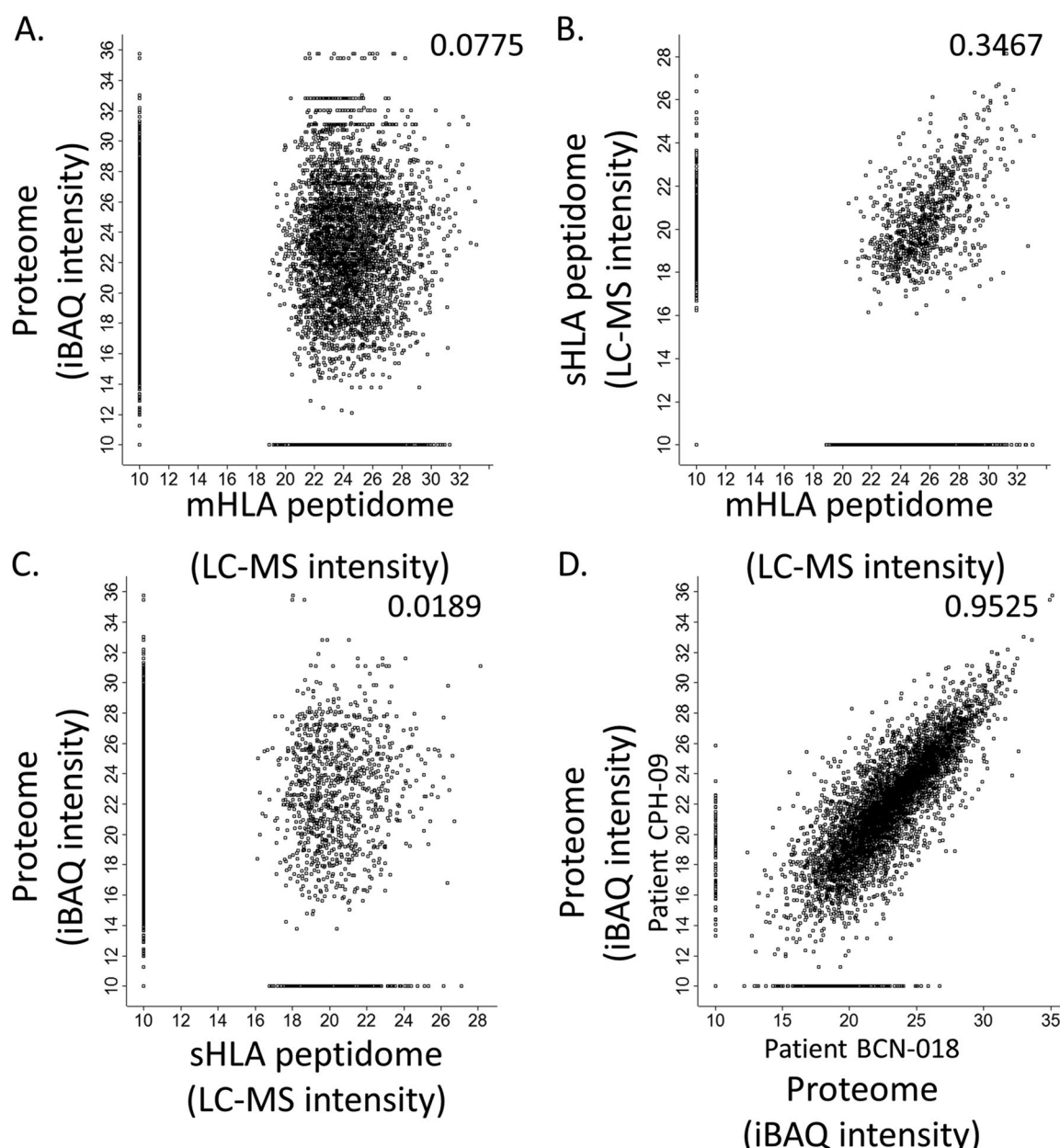


FIG. 5. Comparison between the tumor proteomes and HLA peptidomes: Comparison between the tumor proteome and mHLA peptidome (A), plasma sHLA and tumor mHLA peptidome (B), between the tumor proteome and plasma sHLA peptidome (C) of patient CPH-09, and comparison between the tumor proteomes of two different patients, CPH-09 and BCN-018 (D). Only gene products observed as both proteins and HLA peptides are displayed. Pearson correlations are indicated in each panel and the protein levels are calculated as iBAQ intensities on a Log_2 scales. Missing values were replaced with arbitrary value of 10.

tissues and their derived sHLA peptides appear in the plasma-sHLA peptidomes, may serve as both, potential biomarkers and candidates for immunotherapeutics.

Some HLA Allomorphs Present Larger Diversity of Peptides in Both Tumors and Plasma—The 52 different GBM samples and 36 different noncancerous samples have undergone a complete HLA typing analysis based on DNA sequencing (52). This analysis facilitated the use of the NetMHC platform to fit the HLA peptides sequences to their likely presenting HLA

allomorphs by selecting peptides with NetMHC rank equal or better than 2 (<http://www.cbs.dtu.dk/services/NetMHC/>). HLA allomorphs with a consensus sequence motif such as HLA-B*35:01 and HLA-A*24:02 presented larger numbers of peptides relative to allomorphs such as HLA-B*27:05 or HLA-B*40:01 (Fig. 4 and supplemental Table S8). This phenomenon was observed in both GBM and in the noncancerous donors' plasma samples, as well as the GBM tissues (Fig. 6, supplemental Fig. S4). The HLA-A*24:02 allele is common

TABLE III

List of the HLA peptides and their source proteins, belonging to the TAA group. Peptides and proteins were selected only if they were detected in all ten GBM tumors, and their derived HLA peptides were not detected in any of the non-cancerous donors' plasma-sHLA peptidomes

Gene names	Protein names	Sequence
ADAM17	Disintegrin and metalloproteinase domain-containing protein 17	THVETLLTF
CTNNA2	Catenin alpha-2	SEFKAMDSF
		SPVQALSEF
CTSH	Pro-cathepsin H	ESAIATGK
		TQDFMMYRT
ENAH	Protein enabled homolog	IYHHTGNNTF
		RIAEKGSTI
		SAMMHALEV
FMNL1	Formin-like protein 1	TGFHSDLHFLD
ITGB8	Integrin beta-8	ALMEQQHYV
		REKPEEIKM
PPIB	Peptidyl-prolyl cis-trans isomerase B	LLLPGPSAA
SCRN1	Secernin-1	AIIESDQEQGR
		AIIESDQEQGRKLR
SNRPD1	Small nuclear ribonucleoprotein Sm D1	HETVTIEL
SOX21;SOX14;SOX2	Transcription factor SOX-21;Transcription factor SOX-14;Transcription factor SOX-2	SEISKRLGAEW
TPM4	Tropomyosin alpha-4 chain	KEENVGLHQTLL
		VILEGELERA

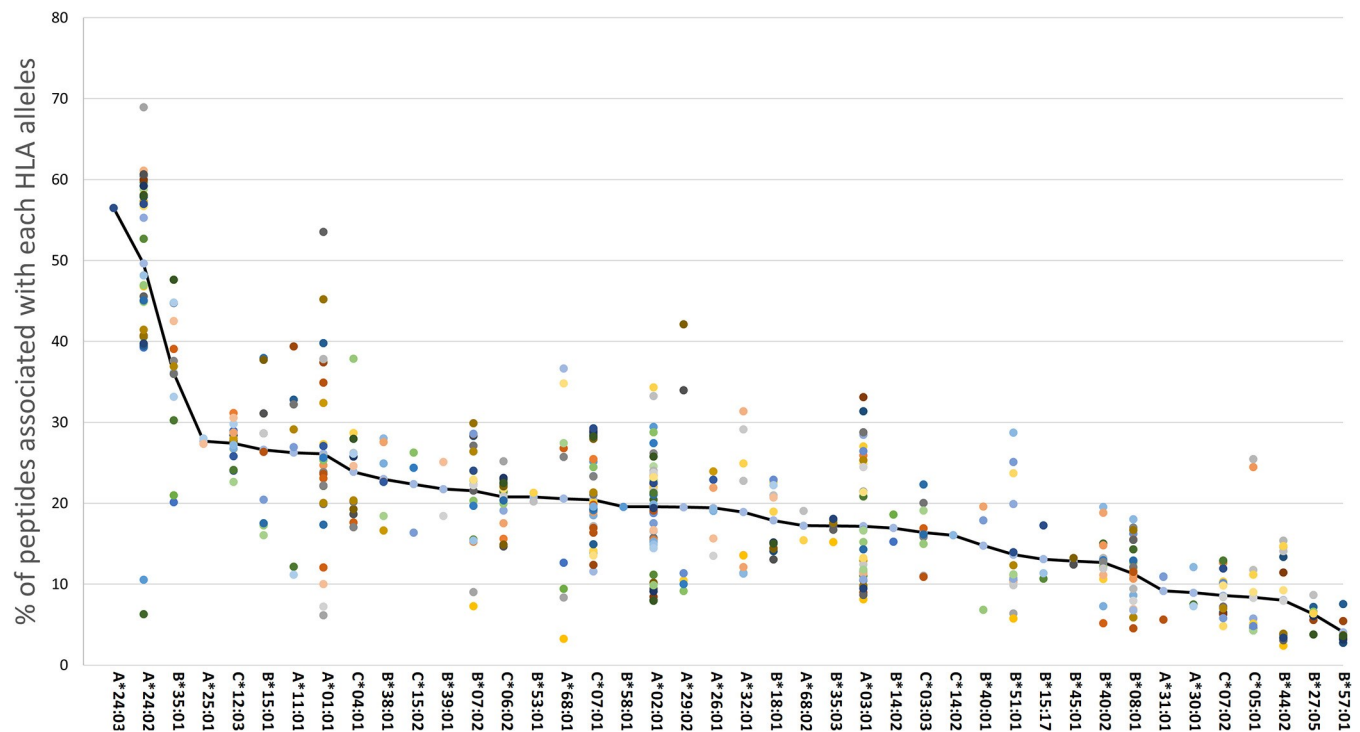


FIG. 6. **Fitness of HLA peptides to different HLA alleles presenting them.** Percentages of peptides discovered in each of the plasma sHLA peptidomes of the GBM patients that fit the sequence motifs of the HLA alleles of the individual patients, with rank below or equal 2% according to NetMHC. The colored dots represent different patients.

among glioma patients (60) and therefore HLA peptides presented by these alleles may become useful for treatment of larger groups of patients.

DISCUSSION

This study is an extensive immunopeptidome analysis of 106 GBM plasma samples, 10 GBM tumor tissues, and 36

control plasma samples. The analysis led to high confidence identification of 35,156 unique HLA peptides. It was suggested before that the plasma-sHLA peptidome represents the tumor-mHLA peptidome, because the tumors' cells shed larger amounts of sHLA molecules relative to the uninvolved tissues (28). Similarly, in this study, a significant overlap was observed between mHLA peptidomes of the tumor tissues

and matched plasma-sHLA peptidomes of each of the patients, demonstrating that the plasma-sHLA peptidome contains many peptides derived from the tumors. These correlations, between the repertoires and LC-MS intensities of the mHLA and sHLA peptidomes of the individual patients, were much larger than the correlations between the HLA peptidomes and the proteomes of the patients' tumors (supplemental Fig. S3). Low correlations in both repertoires and expression levels, between HLA peptidomes, proteomes, and transcriptomes, of the same cells, were suggested in previous publications (61, 62) but were found to be higher in others (63–66). Additionally, we demonstrate that the plasma-sHLA peptidomes are relatively stable, can be analyzed reproducibly, and represent the tumor-mHLA peptidomes. Comparative analysis of both mHLA and sHLA peptidomes may provide candidate peptides, potentially useful for immunotherapy and as biomarkers. Precision medicine based on large-scale body-fluid biomarkers may help to identify patients that are most likely to benefit from specific treatments, including, but not limited to immunotherapy (67–69). It is expected that different HLA peptides derived from the TAA and CTA genes will be presented among the patients' sHLA allomorphs' peptidomes. Thus, the search among the sHLA peptidomes of different people for disease biomarkers is not limited to specific peptides, but to the presence of different peptides derived from the selected TAA/CTA genes. A few of the 422 downregulated sHLA peptides observed in this study may serve as biomarkers for early detection of the disease or relapse. Indeed, potentially useful TAA can be selected based on different criteria. In our opinion, first and foremost are peptides that are derived from genes that not expressed in any healthy adult tissue, other than the immune privileged sites, such as the germline cells. Such HLA peptides are potentially useful for both immunotherapy and diagnosis. Plasma sHLA peptides that disappear after treatment or remission are potentially also useful for diagnosis, for early detection and for relapse. sHLA and mHLA peptides that appear in tumor tissues and not in the sHLA peptidome of the healthy donors are clearly more significant targets for further research because such peptides are more likely (but not necessarily) tumor specific. However, only a few of the downregulated peptides observed here are TAAs. Such selected biomarker peptides do not need to be derived from bona fide CTAs but need to be affected similarly in a large cohort of patients to become useful for clinical exploitation. One should consider that many peptides were not detected in this study in the plasma-sHLA peptidomes after surgery, also because of the nature of shotgun peptidomics methodology used here. Performing targeted LC-MS/MS analysis with the same or with different sets of samples is still required to exclude this possibility. The alternative use of data-independent analysis of HLA peptidome was already used for analyses of HLA peptidomes at higher reproducibility (70–73), and SRM were used to obtain more accurate presentation levels for the se-

lected peptides in multiple analyses (74, 75). Smaller numbers of HLA peptides were identified with the LC-MS/MS data collected in this study when the FDR was set to 0.01 instead of 0.05 in the Andromeda search (50), performed within the MaxQuant analysis tool (49). This is expected, because the MS/MS fragmentations of many peptides are suboptimal in the data-dependent (shotgun) analysis performed in this study. The use of data-independent analysis for the HLA peptidome analysis may help to solve some of the data loss incurred (71, 73, 76). Alternatively, one can increase the FDR to 0.05 and facilitate this way the discovery of candidates for vaccine or biomarkers HLA peptides that would have been lost with 0.01 FDR (77). Here as well, a significant fraction of the peptides that were lost by use of 0.01 FDR, instead of 0.05, are true ligands of the HLA allomorphs of the patient (supplemental Table S9), as defined by the similarity of sequence motifs to the peptides included in the 0.01 FDR (supplemental Fig. S5).

The blood-brain barrier (BBB) prevents entrance and exit of some cells and molecules (5, 6) to the brain and it is unknown if it allows passage of the circulating sHLA-peptide complexes. However, the BBB in GBM patients is partially broken by the local inflammatory conditions and differs in its tumor vessels morphology and in the hyper-permeability of its endothelial cells (5). The loss of functional integrity of the BBB allows passage of low molecular weight compounds, including different chemotherapeutics into the brain (78). Such changes in the BBB may also facilitate the release of sHLA molecules with their bound peptides into the circulation, thus allowing their detection in the plasma for further evaluations as tumor markers.

Ideally, HLA peptides useful to serve as immunotherapeutics or biomarkers should be derived from genes expressed at enough levels in the malignant tissues, but not at all in any of the other essential healthy tissues (18, 23). Fresh-frozen tumors can be used for exome, transcriptome, proteome and HLA peptidome analyses, allowing identification of neo-epitopes and tumor antigens (4, 16, 40–45, 65, 79). Selection of CTA candidates for immunotherapy can be based on high expression levels of their HLA peptides, mRNA and proteins in the tumors, and no expression in healthy tissues. Even though tumors and plasma are good sources of HLA peptides, healthy tissues are not normally available for analysis of their HLA peptidomes, whereas data about the gene expression levels is available in public databases. Very large and relatively accurate databases are publicly available, including data on gene and protein expression in many tissues, of numerous people. Examples include BioGPS (57, 58), TANTIGEN (54) and HPR (<https://www.proteinatlas.org/>) (80, 81). Although these databases are based on numerous studies, some discrepancies were observed between them in regard to different selected CTAs studied here. Using additional gene expression databases may alleviate some of these

concerns and provide candidate immunotherapeutics with lower risk of inducing adverse effects.

The preferred CTAs, providing potentially useful HLA peptides discovered in this study, were defined as those derived from genes whose transcripts are expressed at levels below nine gcrma units in all normal, essential tissues (according to BioGPS) and are expressed at significantly higher levels in the tumors tissues. Nine gcrma units of mRNA levels of expression were selected here as sufficiently low, because this is the highest measured mRNA expression level of many well-characterized CTAs in healthy (nontestis) adult tissues. Some of these CTAs, including CTAG1A (NY-ESO-1) and MAGE-A1 are well known TAAs, whose expression levels in healthy tissues are below 9 gcrma units. These CTAs were already used in multiple clinical studies without observable autoimmune reactions (19, 82–84) suggesting that genes expressed below these levels are possibly safe for clinical use.

The tumor-mHLA and plasma-sHLA peptidomes of different people can be compared while looking for HLA peptides shared between larger cohorts of individuals. HLAs with completely distinct binding motifs are not expected to bind and present shared peptide ligands. Importantly, significant similarities were observed in this study between the HLA peptidomes of different patients that share some of their HLA alleles (Fig. 3). In contrast, shared peptides, detected in the HLA peptidomes of different people who do not have any common HLA alleles or have HLA alleles belonging to different HLA supertypes (85) are more likely to be defined as contaminants, rather than true HLA ligands. For example, it is very unlikely that HLAs such as HLA-A*2, B*7 and B*27 will share any peptide ligands (85). On the other hand, shared tumor antigens that are detected in multiple patients that have similar HLA allotypes, are extremely important, because of their potential to become useful for treatment of multiple patients.

The HLA molecules of some HLA allomorphs, such as HLA A*24:02, are more abundant in the plasma of carriers of these alleles (32). In addition, some HLA allomorphs present more numerous peptides than others do (86). Indeed, here we observe similar phenomena at the HLA peptidome levels, some HLA allomorphs, such as HLA-A24, HLA-B35 and HLA-B51 present more diverse repertoires of peptides than others do, in both the tumors and the plasma. This implies that the use of plasma-sHLA peptidome analysis is probably more efficient for carriers of these alleles, because larger sHLA peptidomes can be recovered and identified from their plasma samples (Fig. 4 and Fig. 6). It may also mean that there is larger immune tolerance in the HLA-A*24 patients and, therefore, it may be more difficult to break the immune tolerance induced by circulating sHLA molecules, when attempting immunotherapy. Importantly, HLA-A*24 was claimed to be associated genetically with glioma (60). Therefore, the use of the sHLA peptidome analysis to search for disease biomark-

ers will need to be adjusted accordingly, to take into account the particular HLA allotypes of the individual patients.

In conclusion, the data described here suggests a useful method for selection of biomarkers and cancer immune-therapeutics for GBM, and provides large lists of such candidates. Such methodologies are likely useful for discovery of biomarkers and immune-therapeutics candidates for other cancers. Most importantly, the identification of numerous sHLA peptides derived from CTAs can represent a promising non-invasive strategy for the monitoring of patients' response and progression during standard treatment modalities.

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DATA AVAILABILITY

The mass spectrometry proteomics data of the tissue proteomes, the mHLA and the sHLA peptidomes have been deposited to the ProteomeXchange Consortium (55) (<http://proteomecentral.proteomexchange.org>) via the PRIDE partner repository with the data set identifier PXD008127.

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