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## HIV microarray for the mapping and characterization of HIV-specific antibody responses†

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We used the microarray technology to develop chips containing a comprehensive set of proteins and peptides covering the proteome of HIV-1 clade C, which is the HIV-1 subtype that causes the majority of infections worldwide. We demonstrate that the HIV microarray allows simultaneous, sensitive and specific detection of antibody responses for the major immunoglobulin classes (IgG, IgA, IgM, IgE) and subclasses (IgG<sub>1-4</sub>) with minute amounts of serum samples towards a large number of HIV antigens and peptides. Furthermore, we show that the HIV chip can be used for the monitoring of antibody responses during the course of the disease and during treatment. The HIV microarray should be useful to study antibody responses to multiple HIV antigens and epitopes in HIV-infected patients to explore pathomechanisms of the disease, for diagnosis and for monitoring of treatment and of vaccine trials.

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### Introduction

HIV infections represent a major health threat. At present more than 35 million individuals worldwide are infected.<sup>1</sup> More than 10% of the HIV-infected persons recorded in the United States were unaware of their infection.<sup>2</sup> Currently available diagnostic tests for HIV are based on the detection of viral materials such as RNA or proteins and on the detection of HIV-specific antibodies using first line tests with high sensitivity and second line procedures with high specificity.<sup>3</sup> The identification of new neutralizing antibodies which protect against infection and results obtained from the analysis of

antibody responses in HIV controllers and HIV vaccine trials indicate that a detailed investigation of antibody responses towards different viral antigens and epitopes may provide new information for the development of new immunological strategies for the treatment and prevention of HIV infections (e.g., antibody-based treatments, vaccines).<sup>4-8</sup> Evidence for the usefulness of multiplexed antibody analysis in the field of HIV comes from a recent study which analysed antibody responses in HIV-infected patients and subjects who participated in vaccine trials. A comprehensive set of small peptides derived from the virus envelope allowed the identification of antibody signatures that may be associated with protection.<sup>9</sup>

In fact, multiplex immune assays based on micro-arrayed antigens and epitopes are currently revolutionizing the analysis of pathomechanisms as well as the diagnosis of several immunological diseases such as allergy, autoimmunity, infectious diseases and cancer.<sup>10-15</sup> However, multiplex assays for the analysis of antibody responses against multiple HIV-proteins and peptides are not available. Therefore, the aim of this study was the development of an HIV microarray containing a large panel of HIV proteins and peptides for the mapping and characterization of HIV-specific antibody responses towards multiple viral antigens and epitopes with minimal amounts of sample and short assay-duration. For this purpose, we employed the microarray-chip technology which we originally had developed for the diagnosis of allergy (i.e., Immuno solid-phase allergen chip, ISAC).<sup>10</sup> We prepared a set of HIV proteins and peptides derived from HIV-1 clade C, because this is the HIV-1 subtype that causes the majority of infections worldwide (48%). HIV-1 clade C is also the

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subtype that predominates in Sub-Saharan Africa, where the highest rates of infection are reported (4.7% infected adults) (UNAIDS global report 2013).<sup>16</sup> We demonstrate that the HIV microarray allowed the measurement of isotypes and IgG subclasses against a comprehensive set of proteins and peptides covering the clade C proteome.

## Materials and methods

### Synthesis, purification and characterization of HIV peptides

Overlapping peptides covering the amino acid sequence of gp120, gp41, VIF, REV, VPR and VPU from HIV-1 subtype C (isolate ZA.04.04ZASK146, hiv.lanl.gov #AY772699) were produced by solid-phase synthesis. The peptides were synthesized with the 9-fluorenyl-methoxy-carbonyl (Fmoc)-method (CEM-Liberty, Matthews, NC, USA and Applied Biosystems, Carlsbad, CA, USA) on PEG-PS preloaded resins (Applied Biosystems, Carlsbad, CA, USA). After washing with dichloromethane, peptides were cleaved from the resins in 19 ml trifluoroacetic acid, 0.5 ml silane and 0.5 ml H<sub>2</sub>O and precipitated into pre-chilled *tert*-butylmethylether. Peptides were purified by reverse-phase HPLC in a 10–70% acetonitrile gradient using a Jupiter 4  $\mu$ m Proteo 90 Å, LC column (Phenomenex, Torrance, CA, USA) and an UltiMate 3000 Pump (Dionex, Sunnyvale, CA, USA) to a purity >90%. Their identities and molecular weights were verified by mass spectrometry (Microflex MALDI-TOF, Bruker, Billerica, MA, USA).

### Expression, purification and characterization of recombinant HIV proteins

Recombinant gp120 from HIV-1 subtype C was purchased from Sino Biological (Beijing, People's Republic of China). The ectodomain of gp41 from HIV-1 subtype C (isolate ZA.04.04ZASK146, hiv.lanl.gov #AY772699, amino acids 532–683 numbered accordingly to the HXB2 scheme) was expressed in a human T cell line (Jurkat E6 cell line) (Wollmann, unpublished data). Recombinant his-tagged structural (matrix, MA; capsid, CA and nucleocapsid, NC), functional (protease, PR; reverse transcriptase + RNaseH, RR and integrase, IN) and accessory proteins (NEF, TAT and VIF) were expressed in *E. coli*. The DNA sequences of MA, CA, NC, PR, NEF, TAT, VIF (HIV-1 clade C isolate ZA.04.04ZASK146, hiv.lanl.gov #AY772699) and of RR and IN (HIV-1 clade C isolate ET.86.ETH2220, hiv.lanl.gov #U46016) contained a 3' sequence coding for a C-terminal hexa-histidine tag. Sequences codon-optimized for bacterial expression were cloned into plasmid pET17b, between *EcoRI* and *NdeI* restriction sites (ATG:biosynthetics, Merzhausen, Germany). *E. coli* BL21(DE3) cells (Agilent Technologies, Santa Clara, CA, USA) were transformed and grown to an OD<sub>600</sub> = 0.4–0.6 in LB medium containing 100 mg l<sup>-1</sup> ampicillin. Isopropyl- $\beta$ -thiogalactopyranoside (0.5–1.0 mM) was used to induce protein expression and cells were harvested at time-points of maximal expression (*i.e.* PR: 30 min; MA, CA, RR, IN, NEF, TAT and VIF: 4 h; NC: ON). Recombinant proteins were purified by nickel-affinity chromatography under native

(MA, CA, NC) or denaturing (PR, IN, NEF, TAT) conditions (Qiagen, Hilden, Germany). VIF and RR were purified with an inclusion body preparation protocol.<sup>17</sup> Stepwise dialysis was used to remove the denaturing agent and to refold the recombinant proteins. The identity of the proteins was verified by SDS-PAGE followed by Coomassie Brilliant Blue staining. In addition, His-tagged proteins were blotted onto Whatman Protran nitrocellulose (GE Healthcare Bio-Sciences, Uppsala, Sweden) and detected with mouse  $\alpha$ -His IgG (0.2  $\mu$ g ml<sup>-1</sup>) (Dianova, Hamburg, Germany) followed by incubation with alkaline-phosphatase-labelled rabbit  $\alpha$ -mIgG (0.5  $\mu$ g ml<sup>-1</sup>) (BD, Franklin Lakes, NJ, USA). Mass spectrometry was performed to verify the molecular mass of recombinant proteins (Microflex MALDI-TOF, Bruker). The secondary structure of the proteins was measured by circular dichroism spectroscopy on a Jasco J-810 spectropolarimeter (Japan Spectroscopic, Tokyo, Japan) at a protein concentration of 0.1 mg ml<sup>-1</sup> in 10 mM sodium-phosphate.<sup>17</sup> The biochemical properties of the recombinant HIV proteins were calculated from their amino acid sequence with ProtParam (<http://web.expasy.org/protparam/>).

### Control proteins, labelling of detection antibodies and spotting of the HIV microarray

Human serum albumin (HSA) and bovine serum albumin (BSA) were purchased from Behring (King of Prussia, PA) and Sigma-Aldrich (St. Louis, MO), respectively. Recombinant VP1 of Rhinovirus 89 (VP1 89) was expressed as His-tagged protein.<sup>18</sup> Allergens and fluorescence-labelled BSA were from Phadia Austria GmbH (Part of Thermo Fisher Scientific Immuno-Diagnostics, Vienna, Austria). Detection system controls were: huIgG<sub>1</sub>, huIgG<sub>2</sub>, huIgG<sub>3</sub>, huIgG<sub>4</sub> (Sigma-Aldrich); huIgG, huIgA, huIgM (Jackson ImmunoResearch, West Grove, PA) and huIgE, isolated from a pool of plasma and sera from different patients by anti-IgE affinity chromatography.<sup>19</sup>

Anti-huIgG (Phadia-Thermo Fisher),  $\alpha$ -huIgG<sub>1</sub>,  $\alpha$ -huIgG<sub>2</sub> and  $\alpha$ -huIgA (Becton Dickinson, Franklin Lakes, NJ) and  $\alpha$ -HSA antibodies (Sigma-Aldrich) were labelled with DyLight 650 (Pierce, Thermo Fisher Scientific, Rockford, IL, USA).  $\alpha$ -huIgG<sub>3</sub> (Sigma-Aldrich),  $\alpha$ -huIgG<sub>4</sub>,  $\alpha$ -huIgM (Becton Dickinson) and  $\alpha$ -huIgE (Phadia-Thermo Fisher) were labelled with DyLight 550 (Pierce, Thermo Fisher Scientific).

Cleaning and coating of glass slides was performed as described by Harwanegg *et al.*,<sup>20</sup> In brief, microscopy glass slides were sonicated in acetone and then in a 1% Alconox solution (Sigma-Aldrich) to clean the surface. The glass surface was then made reactive for primary amine groups using a silylation procedure,<sup>21</sup> then the silane layer was heated, cooled and an amine-reactive complex organic polymer was added.<sup>20</sup> Customized spotting was performed by slow pin mode printing, each spot containing 50–200 fg of microarray component, corresponding to 1–5 attomol (Phadia Austria GmbH). For standard spotting, microarray components were used at a concentration of 0.5 mg ml<sup>-1</sup> in phosphate buffer pH 8.4 and spotted in triplicates. Optimization of spotting conditions was required for certain antigens and is described



in the ESI.† Sequential dilutions of antibody controls (0.125, 0.062, 0.031, 0.016, 0.008, 0.004 mg ml<sup>-1</sup>) were obtained by mixing the antibodies with the corresponding amount of BSA in order to obtain a final spotting concentration of 0.25 mg ml<sup>-1</sup> of total protein (component + BSA) in PBS pH 7.2. Two series of triplicates were spotted for gp120, gp41 and 120/15. Correct spotting of the negative control HSA was verified by detection with fluorescence labelled  $\alpha$ -HSA antibody.

### Serum samples

Sera from 15 African HIV-infected patients ( $n = 47$ ) were from the Asthma, Allergy and Immune Dysfunction Clinic, Harare, Zimbabwe. Fifteen European HIV-positive sera and 15 control sera (*i.e.* tested HIV-negative in conventional diagnostic tests at the date of microarray analyses) were obtained from the Department of Virology of the Medical University of Vienna. Sequential serum samples were obtained from the 15 African patients on highly active antiretroviral treatment (HAART) during 2009–2011. The HIV status of each of the sera was verified by confirmatory HIV Line-Immuno-Assay (InnoLIA, Innogenetics, Gent, Belgium). Analysis of the anonymized sera was approved by the ethics committee of the Medical University of Vienna (EK592/2010) and was performed according to standardized laboratory work procedures for infectious materials.<sup>22,23</sup>

### Microarray-based determination of antibody levels in human serum samples

To avoid assay-interference by lipid drops, serum samples were centrifuged for 1 min at 8000 g and the non-lipidic interphase was diluted in sample diluent (1:50 for detection of IgG, IgG<sub>1</sub>, IgA and IgM; 1:10 for IgG<sub>2</sub>, IgG<sub>3</sub>, IgG<sub>4</sub>; undiluted for IgE measurements). Microarrays were washed by stirring 5 min in washing buffer (Phadia Austria GmbH). After drying the slides by centrifugation (1000 g, 1 min, room temperature), 30  $\mu$ l sample were applied on each microarray and incubated for 2 hours at gentle rocking at room temperature. Microarrays were then rinsed and washed 5 min as described above. After centrifugation, they were incubated 30 min with 30  $\mu$ l of fluorescence-labelled antibodies (1  $\mu$ g ml<sup>-1</sup>), rocking at room temperature. Microarrays were again rinsed with washing buffer, washed 5 min by stirring in washing buffer and then washed 5 min in distilled water. After drying by centrifugation, the slides were scanned at 635 nm (DyLight 550-labelled antibodies) or at 532 nm (DyLight 650-labelled antibodies) using a confocal laser scanner (LuxScan-10K microarray scanner, CapitalBio, Beijing, People's Republic of China). For calibration and determination of background signals, a calibrator serum (*i.e.* a pool of HIV-positive sera, diluted 1:100) and sample diluent were included in each analysis run. IgG levels to control antigens (*i.e.*, allergens) listed in Table 3 were additionally tested for the calibrator serum by ImmunoCAP. A calibration curve was generated relating fluorescence intensities derived from scanning the microarrays with antibody levels measured by

ImmunoCAP. Due to the semi-quantitative character of antibody levels measured by microarray, results are given in ISAC standardized units (ISU).<sup>24</sup> The Phadia Microarray Image Analysis software was used to evaluate the measurements, to calculate the mean fluorescence intensities of triplicate analyses and to calibrate the results.

### Background signals, intra- and inter-batch variation and data analysis

Background reactivity of fluorescence-labelled  $\alpha$ -huIgG towards components spotted on the HIV microarray was determined by testing seven replicates of sample diluent alone. Variation among microarray measurements performed on the same day and on two consecutive days was analysed testing IgG reactivity of 4 and 3 replicates of the calibrator serum, respectively. Microarrays of the same lot were used in both cases. Mean coefficients of variation (CV = SD/mean) and signal-to-noise ratios (SNR = mean/SD) were calculated from ISU for component-specific reactivities. Serum titration experiments were performed testing HIV-positive and control samples at dilutions 1:10, 1:50, 1:100, 1:200 and 1:400. The cut-off for positive reactivity was set at 3 ISU, based on comparison with negative controls.

For data analysis, the background signal of each antigen was subtracted from the measured reactivity. The distribution of the reactivity of the detection antibodies towards spotted antibody controls was analysed with GraphPad Prism (La Jolla, CA, USA). For each HIV-derived component differences between IgG levels of HIV-positive samples and controls were analysed by Mann Whitney U tests; receiver operating characteristic (ROC) curves ( $x$ -axes: 1-specificity;  $y$ -axes: sensitivity) were generated and the respective area under the curve (AUC) values were calculated (GraphPad Prism). Median antigen-specific IgG levels measured in African and European HIV-positive sera were calculated (GraphPad Prism). Differences between IgG reactivities of the two populations were analysed by Mann Whitney U tests (IBM SPSS-Statistics, Version 20.0, IBM Corp, Armonk, NY, USA).

### Comparison of microarray- and ELISA-based determination of HIV-specific IgG levels

IgG levels of sera from 15 African HIV-positive, 15 European HIV-positive and 10 control subjects tested with the HIV microarray, were determined also by ELISA. Sera diluted 1:200 in PBS, 0.5% BSA, 0.05% v/v Tween 20 (PBST) were tested on plates coated overnight at 4 °C with 2  $\mu$ g ml<sup>-1</sup> of peptides/proteins in 100 mM sodium bicarbonate buffer pH 9.6, after blocking for 4 h at room temperature (2% BSA, PBST). After washing with PBST, bound antibodies were detected by 1 h incubation with HRP-labelled  $\alpha$ -huIgG (1:5000, 0.5% BSA, PBST). The colour reaction induced with 2,2'-azino-bis(3-ethylbenzothiazoline-6-sulfonic acid) di-ammonium salt was measured as optical density (OD<sub>405 nm</sub> - OD<sub>490 nm</sub>). Antibody levels measured with the HIV microarray (ISU) were plotted against levels measured by ELISA (OD) for the single peptides/proteins tested and correlation coefficients ( $R^2$ ) were calculated in Excel.



## Results

### Design of a chip containing a comprehensive set of micro-arrayed peptides and proteins of the HIV-1 clade C proteome

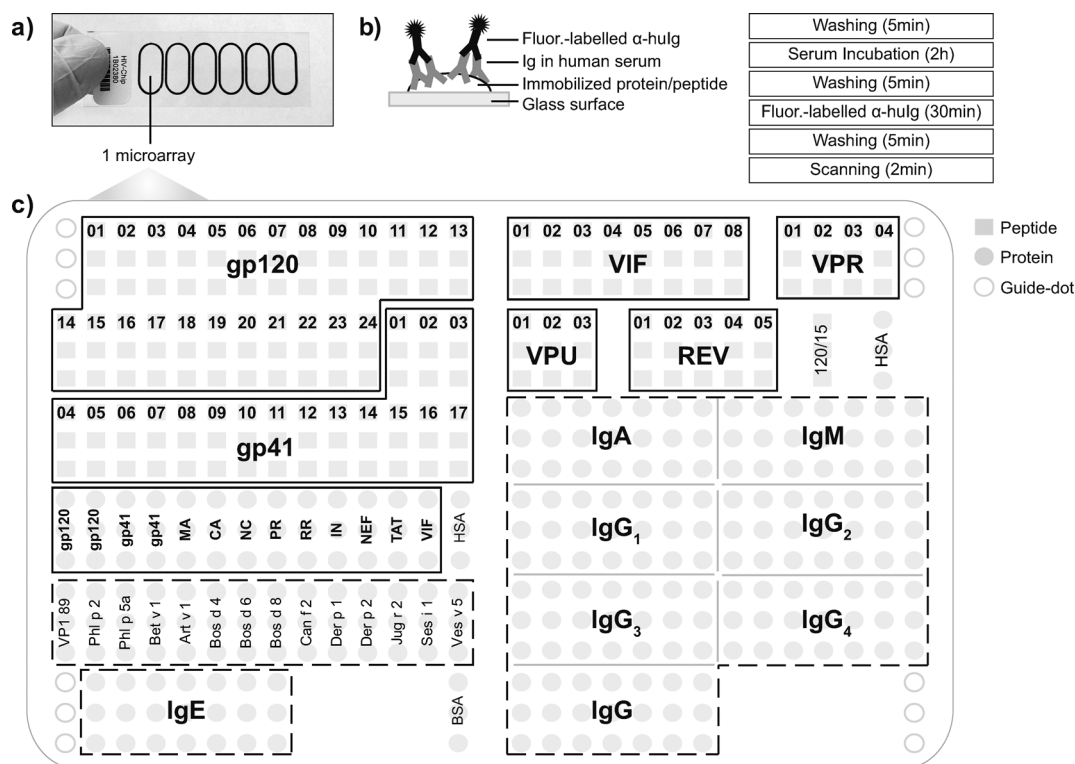
The HIV chips consist of glass slides containing six micro-arrays surrounded by a Teflon frame which allow the simultaneous application of six independent samples (Fig. 1a, b).<sup>24</sup> The Teflon frame was made in oval shape to allow incubation on a rocking plate so that serum samples can better access the areas in the outer parts of the arrays.

Each HIV microarray was designed to contain proteins and peptides from HIV-1 clade C and control components (*i.e.* antigens from other sources for which calibration sera were available such as VP1 from human rhinovirus, allergens or antigens which served as positive controls such as purified antibody isotopes, subclass dilutions) (Fig. 1c, Tables 1–3). Among HIV components, two panels of synthetic overlapping peptides from the envelope proteins gp120 and gp41 were included to map linear epitopes of envelope-specific antibodies (Table 1; Fig. 1c, left). Recombinant folded glycosylated envelope proteins were included together with folded structural, functional and accessory proteins to characterize HIV-specific conformation-dependent antibody responses (Table 2, Fig. 1c, left). Additionally, to map linear epitopes of

accessory proteins we included peptides derived from VIF, VPR, VPU and REV (Table 1; Fig. 1c, right).

Control components included: i) calibration components for which the amount of IgG antibodies contained in a calibrator serum had been determined by quantitative ImmunoCAP measurements (Table 3); ii) detection system controls, consisting of sequential dilutions of purified human antibody preparations (IgG, IgA, IgM, IgE) and purified monoclonal human IgG subclasses (IgG<sub>1-4</sub>); iii) positive controls (*e.g.* allergens showing IgE, IgG<sub>1</sub>, IgG<sub>4</sub>, IgG<sub>2</sub> reactivity, human rhinovirus-derived VP1 showing IgG<sub>1</sub>, IgG<sub>3</sub>, IgA, IgM reactivity with the calibrator serum) and negative controls (HSA, BSA); iv) fluorescence-labelled BSA molecules as “guide-dots” for software-based evaluation (Fig. 1c).

The identity and quality of each of the produced peptides and proteins were examined before spotting. Mass spectrometry analyses showed that the peptides had the correct molecular mass; SDS-PAGE followed by Coomassie Brilliant Blue staining and Western-blot confirmed the identity and purity of the recombinant proteins (*i.e.*, >95%). Furthermore, circular dichroism measurements showed that each of the recombinant HIV-1 clade C proteins was folded (Table 2). Thus, the current HIV microarray contained 147 components, of which 72 were derived from 14 different HIV proteins. All



**Fig. 1** HIV microarray. (a) Image of the HIV chip containing 6 microarrays. (b) Scheme and flowchart describing the procedure for the detection of HIV-specific antibodies on the microarray. (c) HIV-microarray layout depicting the position of peptide (squares, numbered), protein (filled circles) and guide-dot (open circles) triplicates. Peptides derived from the same HIV proteins (gp120, gp41, VIF, VPR, VPU, REV) and recombinant HIV proteins are boxed. Control proteins (*i.e.* antibody controls: IgA, IgM, IgG<sub>1-4</sub>, IgG and IgE at decreasing concentrations from left to right: 1st spot 0.250 mg ml<sup>-1</sup>, 2nd–7th spot 0.125, 0.062, 0.031, 0.016, 0.008, 0.004 mg ml<sup>-1</sup> obtained by mixing the antibody with BSA; control antigens: VP1 89 and allergens) are surrounded by broken lines.





Table 1 HIV-1 peptides spotted on the HIV microarray

Peptide	Amino acid sequence	Position <sup>a</sup>	No. of amino acids	Molecular weight (dalton)	Calculated isoelectric point <sup>b</sup>
120/01	RVRGILRNWPQWWIWGILGFWMIII	2–28	25	3210.9	12.3
120/02	WMIIICRGEENSWVTVYYGVPVWTE	24b–47	25	3031.5	4.5
120/03	PVWTEAKTTLFCASDAKAYEKEVHN	43–67	25	2839.2	5.5
120/04	KEVHNWVATHACVPTDPSQELVLE	63–87	25	2800.1	5.0
120/05	ELVLENTESFNMWENDMVDQMHEH	83–107	25	3055.3	3.8
120/06	QMHEDIIGLWDESLKPCVKLTPLCV	103–127	25	2868.4	4.7
120/07	TPLCVTLNCNTTSHNNSSPMTNC	123–157	25	2636.9	6.4
120/08	PMTNCSFNATTELDRKTQKVNALFY	153–177	25	2893.3	8.6
120/09	NALFYRSDIVPLEKNSSEYLINCNC	173–197	25	2916.3	4.7
120/10	LINCNTSTITQACPKVSFDPIPIHY	193–217	25	2776.2	6.7
120/11	IPIHYCAPAGYAILKCNCKTFNGTG	213–237	25	2667.1	8.8
120/12	FNGTGPCSNVSTVQCTHGIKPVVST	233–257	25	2533.9	8.1
120/13	PVVSTQLLNGSLAEGEIIIRSEN	253–277	25	2666.1	4.2
120/14	RSENLTDNAKTIVHLNKSVAIVCT	273–297	25	2740.2	8.2
120/15	AIVCTRPNNTTRKSIRIGPGQVFYT	293–320	25	2806.2	10.9
120/16	QVFYTNIEIIGNIRQAHCNISRELWN	315–339	25	3019.4	6.7
120/17	RELWNTLEQVKKLKEHFQNKTI	334–360	25	3154.6	9.4
120/18	NKTIEFQPPAGDLEVTTHSFNCRG	356–380	25	2719.0	5.4
120/19	FNCRGEFFYCNTSNLNFITASNASD	376–400	25	2836.1	4.7
120/20	SNASDANNNTITLPCIKQIINMWQ	396–428	25	2818.2	7.9
120/21	INMWQEVGRAMYAPPIAGNITCNSS	424–448	25	2724.1	6.0
120/22	TCNSSITGLLTRDGGNNNDTGNNN	444–465e	25	2565.7	4.2
120/23	TGNNNDTEIFRPGGNNMKDNWRSEL	465a–483	25	2823.0	4.8
120/24	WRSELYKYVVEIKPLGIAPTAKRRVVEREKR	479–511	33	4027.8	10.4
41/01	AVGLGAVLLGFLGTAGSTMGAASIT	512–536	25	2235.6	5.6
41/02	AASITLTVQARQLLSGIVQQSNLL	532–556	25	2653.1	9.8
41/03	QSNLLRAIEAQQHMLQLTVWGIKQL	552–576	25	2919.4	8.7
41/04	GKQLQARVLAIERYLKQDQLLGLW	572–596	25	2953.5	9.7
41/05	LLGLWGCSCGKLICTAVHWNSSWSN	592–616	25	2734.1	8.1
41/06	SSWNSKSDYIWNMTWMQWDREIN	612–636	25	3163.4	4.6
41/07	DREINNYTDIYTLLEESQSQQEKN	632–656	25	3044.2	4.1
41/08	QQEKNEKDLLALDSWNNLWNWFSIT	652–676	25	3093.4	4.3
41/09	WFSITKWLWYKIFIMIVGGLIGLR	672–696	25	3054.8	10.3
41/10	LIGLRIILGVLSIVKRVQGYSPS	692–716	25	2751.4	11.7
41/11	YSPLSFQTLPPNPRGPDRLRGIEEE	712–736	25	2869.2	5.1
41/12	GIEEGEGEQDKDRSIRLVSGFLALV	732–756	25	2718.0	4.4
41/13	FLALVWEDLRSLCLFSYHRLRDFIL	752–776	25	3126.7	6.7
41/14	RFILLAGRAAELLGRSSLRGLQGT	772–789	25	2671.1	11.5
41/15	GLQTGWQALKYGLSLVQYWGLELKK	787e–809	25	2880.4	9.4
41/16	LELKSAINLFDTTAIVVAEGTDRL	805–829	25	2718.1	4.8
41/17	GTDRLEGLQGIGRAIYNIPRRIRQGFEAALL	825–856	32	3568.1	10.6
VIF/01	MENRWQVLIVWQVDRMRIRTWNSLVKHHMY	1–30	30	3926.6	10.8
VIF/02	KHHMYISKRASRWVYRHYESRNPRISEV	26–55	30	3811.3	10.5
VIF/03	ISSEVHIPLGEARLVIKTYWGLHTGERDWQ	51–80	30	3491.9	6.0
VIF/04	ERDWQLGHGVSIEWRLRRYSTQVDPGLADQ	76–105	30	3568.9	5.5
VIF/05	GLADQLIHMHYFDCFADSIRKAILGQVVS	101–130	30	3319.8	6.0
VIF/06	GQVVSPPRCYQAGHNKVGSLQYLALTALIK	126–155	30	3230.7	9.1
VIF/07	TALIKPKRRKPLPSVRKLVEDRWNNPQKI	151–180	30	3579.3	11.6
VIF/08	NPQKIRDRRGNHTMNGH	176–192	17	2031.2	11.7
REV/01	AGRSQSDSEALLQAVRIKILYQSNPPPKP	2–31	30	3234.7	8.5
REV/02	LYQSNPPPKPEGTRQAQRNRRRRWRARQRQ	22–51	30	3788.2	12.3
REV/03	RRRWRARQRQIHVSERILSTCLGRPAEPV	42–71	30	3615.1	12.0
REV/04	TCLGRPAEPVPLQLPPIERLHIDRESSGT	62–91	30	3285.7	5.5
REV/05	HIDRESSGTSQTQSQGTTDRVASP	82–107	26	2705.8	5.4
VPR/01	EQPPEDQGPQREPYNEWALEILEELKQEA	2–31	30	3565.8	3.9
VPR/02	ILEELKQEA VRHFRPWLHNLGQYIYATYG	22–51	30	3643.1	6.9
VPR/03	LGQYIYATYGDITWTGVEALLRILQQLFIH	42–71	30	3497.0	5.3
VPR/04	RILQQLFIHFRIGCQHSRIGILRQRARRNGASRS	62–96	35	4158.9	12.5
VPU/01	SFLYASVDYRLGVALIHAL	2–16	20	2141.5	5.5
VPU/02	EYRKLRLQRKINKLIDRIRDREEDSGNESE	29–58	30	3760.1	8.5
VPU/03	REEDSGNESEGDIEELATMVDMGHLRLDDNNL	49–82	33	3717.9	3.9

<sup>a</sup> Position of the peptides in HIV-1 clade C proteins, numbered according to the HXB2 numbering scheme (www.hiv.lanl.gov). <sup>b</sup> Based on the amino acid sequence, calculated with ProtParam, ExPASy.



**Table 2** HIV-1 proteins spotted on the HIV microarray

Proteins	Calculated molecular weight <sup>a</sup> (kDa)	Calculated isoelectric point <sup>a</sup>	Secondary structure determined by CD	Expression system	Final solvent
gp120	54.8	8.2	α, β	Human cells	PBS, 5% trehalose, 5% mannitol pH 7.4
gp41	18.9	6.8	β (68%), α (11%)	Human cells	10 mM NaH <sub>2</sub> PO <sub>4</sub> , pH 4.7
MA	15.5	9.1	α (63%), β (10%)	<i>E. coli</i>	50 mM Tris, 200 mM NaCl, 1 mM β-met, 20% glycerol, pH 5.3
CA	26.5	6.6	α (57%), β (12%)	<i>E. coli</i>	10 mM Na-phosphate, 150 mM NaCl, 10% sucrose, pH 5.0
NC	7.2	10.2	β (34%), α (5%)	<i>E. coli</i>	50 mM Na-phosphate, 50 mM NaCl, 1 mM β-met, pH 5.5
PR	24.6	6.2	β (27%), α (14%)	<i>E. coli</i>	10 mM Na-phosphate, pH 4.7
RR	12.2	9.0	β (29%), α (12%)	<i>E. coli</i>	10 mM Na-phosphate, pH 4.7
IN	23.7	10.5	β (32%), α (12%)	<i>E. coli</i>	10 mM Na-phosphate, pH 4.7
NEF	11.7	8.7	β (42%), α (6%)	<i>E. coli</i>	20 mM Tris, 0.5 M NaCl, 15 mM β-met, pH 7.4
TAT	65.1	6.8	β (28%), α (17%)	<i>E. coli</i>	H <sub>2</sub> O
VIF	33.2	7.4	β (25%), α (19%)	<i>E. coli</i>	10 mM Na-phosphate, 150 mM NaCl, 20% glycerol, pH 6.0

<sup>a</sup> Based on the amino acid sequence, calculated with ProtParam, ExPASy. Abbreviations: kDa, kilo Dalton; CD, circular dichroism; α, alpha-helical structure, β, beta-sheet structure; *E. coli*, *Escherichia coli*; β-met, β-mercaptoethanol; Na-phosphate, sodium phosphate.

**Table 3** Allergens spotted on the HIV microarray

Microarray component	Allergen source	Recombinant/natural
Phl p 2	<i>Phleum pratense</i>	Timothy R
Phl p 5a	<i>Phleum pratense</i>	Timothy R
Bet v 1	<i>Betula verrucosa</i>	Birch N
Art v 1	<i>Artemisia vulgaris</i>	Mugwort N
Bos d 4	<i>Bos domesticus</i>	Cattle N
Bos d 6	<i>Bos domesticus</i>	Cattle N
Bos d 8	<i>Bos domesticus</i>	Cattle N
Can f 2	<i>Canis familiaris</i>	Dog R
Der p 1	<i>Dermatophagoides pteronyssinus</i>	House dust mite N
Der p 2	<i>Dermatophagoides pteronyssinus</i>	House dust mite R
Jug r 2	<i>Juglans regia</i>	Walnut N
Ses i 1	<i>Sesamum indicum</i>	Sesame N
Ves v 5	<i>Vespula vulgaris</i>	Wasp R

components were spotted in triplicates in order to obtain three independent determinations in each experiment (Fig. 1c).

### The HIV microarray allows sensitive and specific detection of antibodies towards a comprehensive set of proteins and peptides with minute serum volumes and short assay duration

Antibody detection on the chip involves a washing step followed by application of the serum sample, washing, detection of bound antibodies with fluorescence-labelled antibody conjugates, washing and scanning requiring less than 3 hours (Fig. 1b).

Fig. 2 exemplifies the specificity of the HIV microarray. A sample volume of 30 μl was sufficient to detect specific antibody responses against each of the micro-arrayed components. The following serum volumes were needed for detection of the antibody classes/subclasses: IgG (dilution 1:50; 0.6 μl), IgG<sub>1</sub> (1:50; 0.6 μl), IgG<sub>2</sub> (1:10; 3 μl), IgG<sub>3</sub> (1:10; 3 μl), IgG<sub>4</sub> (1:10; 3 μl) IgA (1:50; 0.6 μl), IgM (1:50; 0.6 μl), IgE (undiluted; 30 μl).

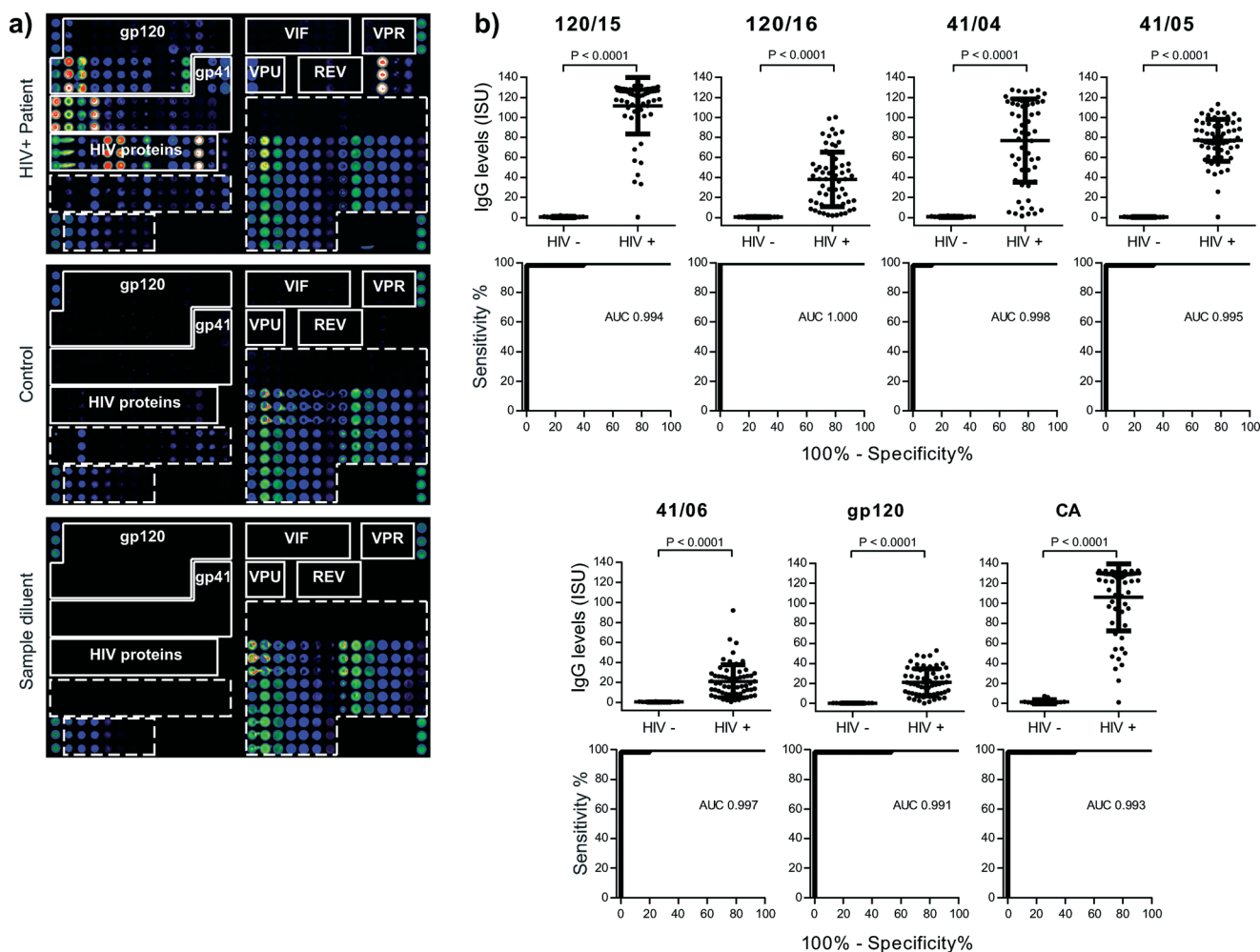
Thus, less than 50 μl of serum allowed determining the specificities of all antibody classes and subclasses to 72 HIV derived proteins and peptides in triplicate analyses.

On scan-images the bound antibodies could be clearly identified as dots of varying intensities and HIV-specific profiles could be used to discriminate HIV-positive from control samples (Fig. 2a, upper and mid panels). Tests performed with sample diluent alone showed specific binding of the IgG detection system towards the spotted antibody controls (IgG, IgG<sub>1-4</sub>, IgE-containing preparation) but no non-specific binding towards any of the other components (Fig. 2a, lower panel). To relate scanned fluorescence levels to amounts of antibody present in serum samples, we determined specific IgG levels of a calibrator serum with the HIV microarray and with quantitative ImmunoCAP (Fig. 3a). The results obtained were used to generate calibration curves and to convert fluorescence levels into ISAC standardized units (ISU). When the calibration curve could be approximated to a linear function, the amount of IgG (μg) per ml of serum could be calculated with the formula [ $\mu\text{g IgG per ml} = \text{ISU} \times \text{serum dilution factor}/1000$ ]: this was observed for values up to 6 ISU, which corresponds to 0.3 μg IgG per ml of serum (Fig. 3a).

Intra- and inter-assay replicates of calibrator serum were measured with high reproducibility, as characterized by mean coefficients of variation (CV) <1 ISU (*i.e.*, 0.36 and 0.38) and signal to noise ratios (SNR) >1 ISU (*i.e.*, 8.3 and 5.5). Intra- and inter-assay variation were even lower when analysed for IgG levels greater than 35 ISU (CV = 0.15, SNR = 12.4 and CV = 0.13, SNR = 33.6, respectively).

Background reactivity was measured by incubating arrays with sample diluent alone ( $n = 7$ ). The signals ranged between 0.0 and 0.14 ISU, confirming the absence of non-specific binding of the detection system. Assessment of the reactivity of anti-huIgG detection antibodies towards spotted antibody controls of different isotype and IgG subclasses





**Fig. 2** (a) Scan images obtained from a microarray tested for IgG reactivity with serum from a HIV-positive subject (upper panel), a control serum sample (middle panel) or sample diluent (lower panel). IgG-reactive vertical antigen triplicates are visualized with fluorescent-labelled antibodies. Increasing fluorescence intensities from blue to red/white correspond to the amount of bound IgG. The layout of spotted antigens is shown in Fig. 1c. (b) IgG levels of HIV-infected patients and controls are shown together with ROC curves for antigens with AUC > 0.99. Median IgG levels (horizontal lines), *P*-values of test significance and AUC values are indicated in each figure part.

showed specific and concentration-dependent binding of the fluorescence-labelled anti-huIgG antibody (Fig. 3b).

We then performed serum titration experiments with HIV-positive sera at dilutions of 1:10, 1:50, 1:100, 1:200, 1:400 that identified 1:50 as a suitable serum dilution (Fig. 3c). Thus, 0.6  $\mu$ l of serum were sufficient for the determination of IgG reactivities towards the panel of spotted components.

Next we tested 62 HIV-positive sera and 15 control sera, replicates of the calibrator serum ( $n = 2$ ) and sample diluent ( $n = 2$ ). Ranges and median IgG levels for each of the HIV components are shown in Table S1†. IgG levels to micro-arrayed HIV-derived peptides and proteins were significantly higher in HIV-positive samples than in controls for all components, except 120/01, 120/07 and VIF/08 (Table S1†). To estimate the use of microarrays for diagnostic purposes, we analysed sensitivity and specificity for each HIV-derived component with ROC curves (Table S1†). The highest areas under the curve (AUC) were measured for: 120/15, 120/16, 41/04, 41/05, 41/06, gp120 and CA (AUC > 0.99, Fig. 2b) and followed by 120/09, 120/17,

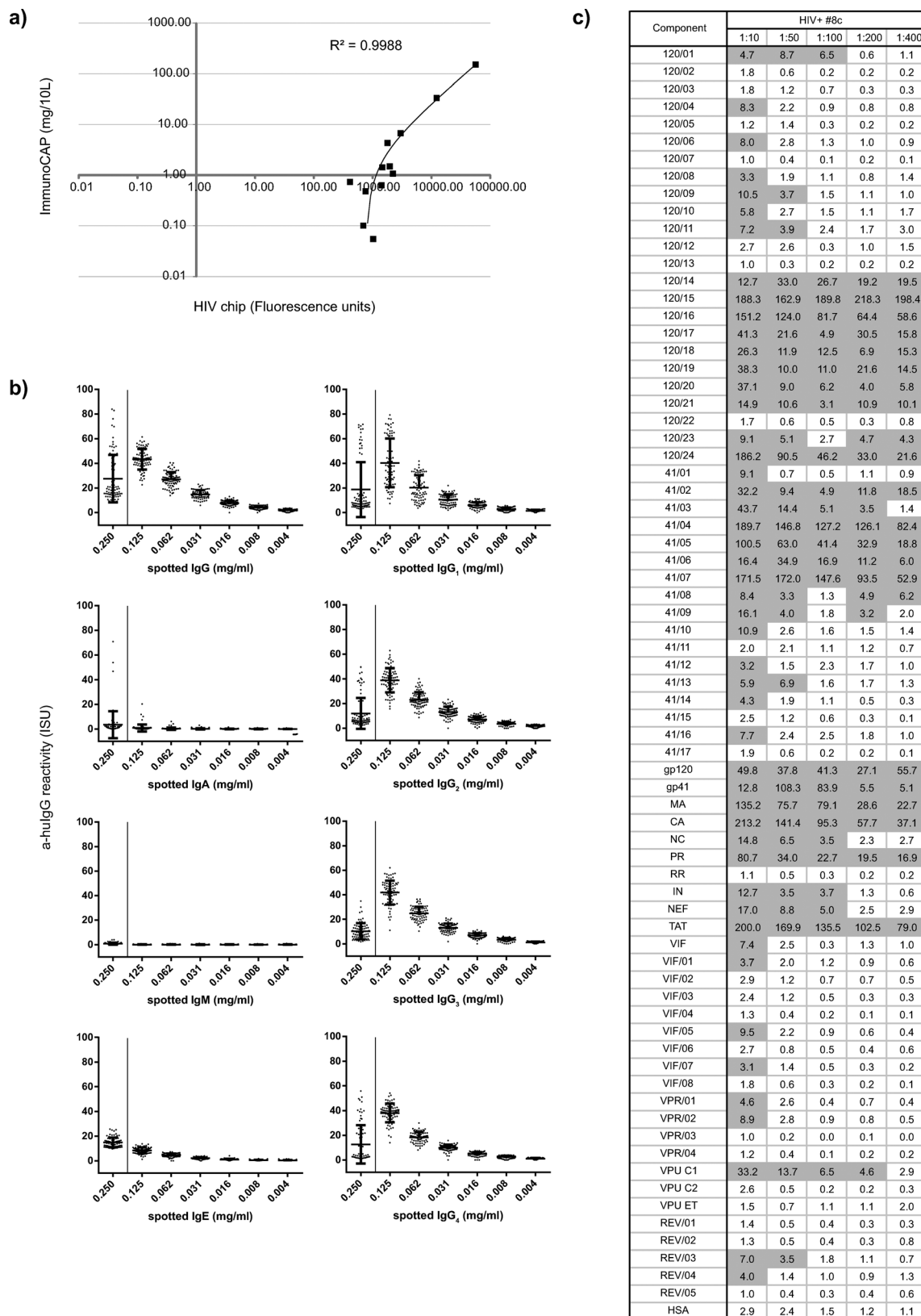
120/24, 41/07, 41/16, MA and IN ( $0.98 < \text{AUC} \leq 0.99$ ). Thus, determination of IgG levels towards these micro-arrayed HIV-derived components may be useful for diagnosis.

### Comparison of the HIV microarray with ELISA

For HIV components that had high median IgG levels and AUC > 0.98 we compared IgG levels determined by microarray with results obtained by ELISA measurements (Fig. 4). A positive correlation was found between the two assays for peptides 120/15, 120/24, 41/04, 41/05 and proteins gp120, MA and CA. Interestingly, IgG levels against two peptides 120/16 and 41/06 were detected only when immobilized on the microarray but not by ELISA and were specific for HIV-infected patients. The lack of IgG binding to the peptides by ELISA may be due to the fact that the peptides 120/16 and 41/06 did not bind to the ELISA plate.

IgG levels against gp120 were lower when detected on the microarray than by ELISA and were lower than envelope

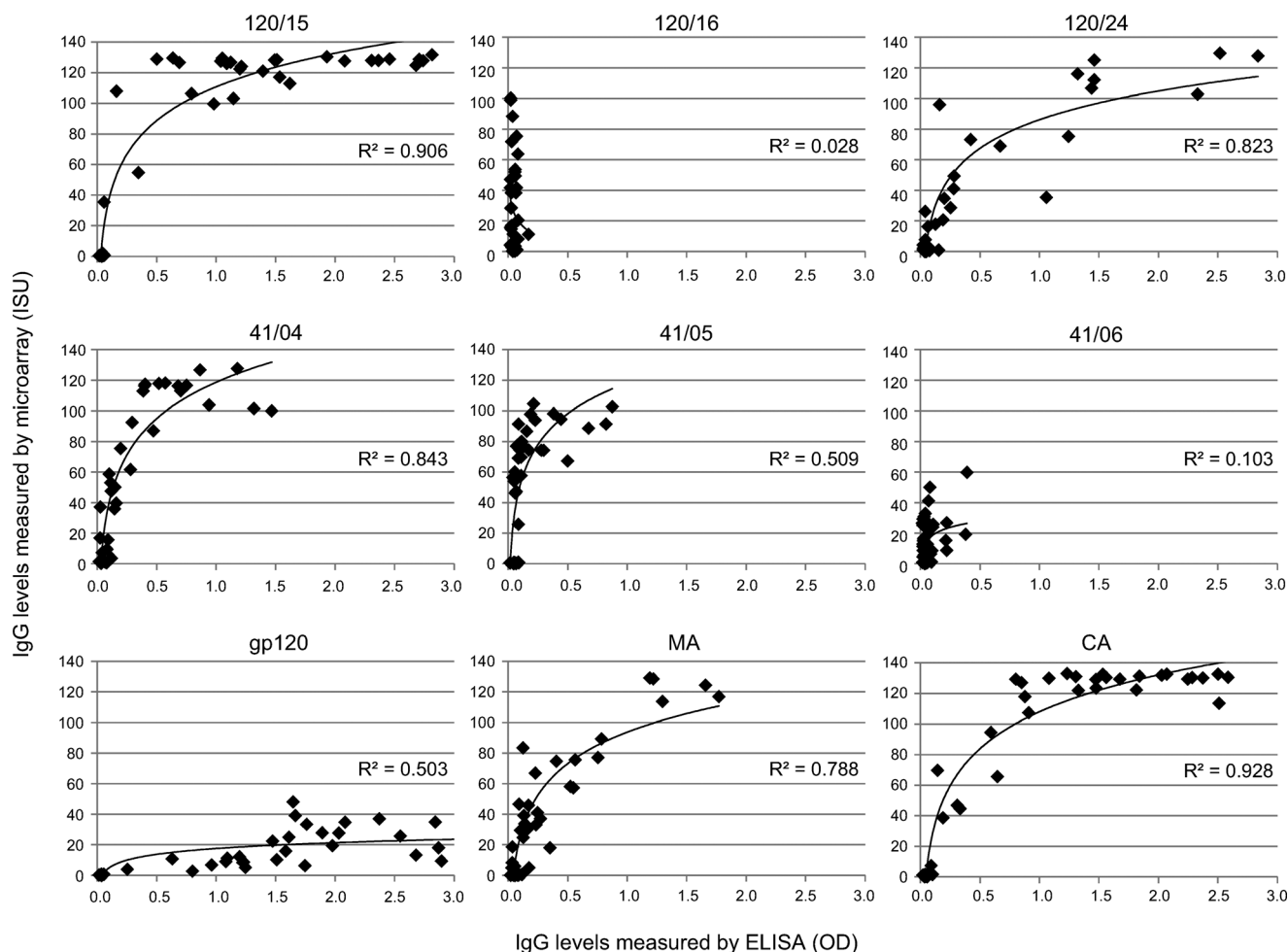




**Fig. 3** (a) Fluorescence levels determined by microarray (*x*-axes) for calibration components (*i.e.*, allergens listed in Table 3) are shown with corresponding IgG levels measured by ImmunoCAP (*y*-axes). (b) Reactivity of anti-human IgG detection antibody with spotted preparations of human antibody isotypes (IgG, IgA, IgM, IgE) and human monoclonal IgG subclasses (IgG<sub>1-4</sub>) measured by microarray (expressed as ISAC standardized units, ISU). Reactivity is shown for antibody controls spotted at 0.25 mg ml<sup>-1</sup> (*x*-axes, left) as well as mixed with BSA at decreasing concentrations (0.125, 0.062, 0.031, 0.016, 0.008, 0.004 mg ml<sup>-1</sup>, *x*-axes, right). (c) IgG levels determined by microarray (ISU) towards spotted components in serum titration analyses of an HIV-infected sample (#8c) at sequential serum dilutions (1:10–1:400). Positive reactivity is shown in grey.







**Fig. 4** Comparison of microarray- and ELISA-based determination of HIV-specific IgG. IgG levels measured by microarray, expressed in ISAC standardized units (ISU, y-axes) are plotted against IgG levels determined by ELISA as optical densities (OD, x-axes) for peptides/proteins with an AUC (area under the curve) >0.98 and high median IgG levels.

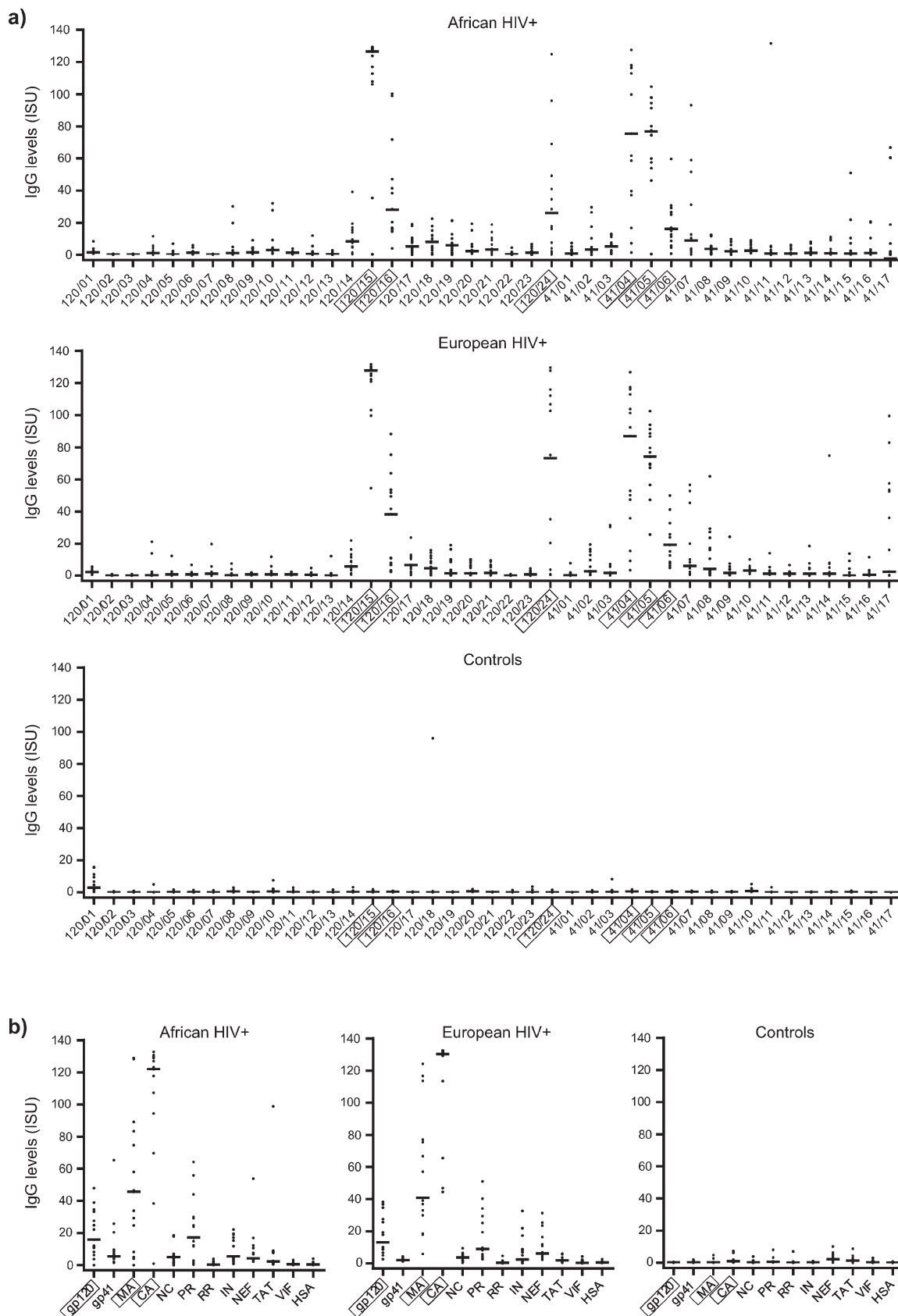
peptide-specific responses (Fig. 5). This could be due to the fact that proteins and peptides were immobilized at the same concentration. Thus, a smaller number of molecules was spotted in case of high molecular weight proteins such as the proteins gp120 and gp41, in comparison to low molecular weight peptides. Another possible explanation for the low reactivity compared to ELISA could be a lower binding of gp120 and gp41 to the chip surface. Finally, it is possible that the carbohydrate moieties on gp120 and gp41 blocked reactivity of peptide-specific antibodies or that these peptides represent cryptic epitopes which were not fully accessible on the intact glycosylated proteins used by us.

#### The HIV microarray allows mapping of IgG reactivity profiles towards a comprehensive set of HIV-proteins and peptides representing the proteome of given strains

Next, we tested the antibody recognition profiles of HIV-infected patients from an African region where clade C is predominant (*i.e.*, Zimbabwe)<sup>25</sup> and compared it with that of patients from a region where HIV-1 clade C is not endemic

(*i.e.*, Europe).<sup>26</sup> We found that clade C-derived envelope-derived peptides (Fig. 5a) and HIV proteins (Fig. 5b) were recognized by African and also European HIV-infected patients. The highest median IgG levels were observed in both populations towards gp120-derived peptides 120/15, 120/16 and 120/24, gp41-derived peptides 41/04, 41/05, 41/06 and gp120, MA, CA and PR proteins (Fig. 5a, b). Strong antibody reactivity was observed in both populations also towards 41/07 and 41/17 (Fig. 5a). Control subjects without HIV infections showed no binding to the micro-arrayed components except towards peptide 120/01, which covers the signal peptide of the protein, and occasional reactivity was found in single sera towards peptides 120/04, 120/10, 120/18, 41/03 and 41/10. IgG levels towards peptides derived from HIV accessory proteins were low in both African and European HIV-infected populations (Fig. S1a†). Common peptide epitopes recognized by African and European patients were VPU/01, VPU/02 and REV/03. Peptides VIF/04, VIF/05, VIF/07 showed IgG reactivity with single sera from control subjects. IgG responses towards control components to which most subjects are exposed (*i.e.*, allergens and VP1) were found in





**Fig. 5** Microarray-based mapping of HIV-specific IgG responses in African and European HIV-infected individuals and controls. IgG levels (y-axes, ISAC standardized units, ISU) to micro-arrayed gp120- and gp41-derived peptides (a) as well as to HIV proteins and HSA (b) are shown for African HIV-positive patients ( $n = 15$ ), European HIV-positive patients ( $n = 15$ ) and controls ( $n = 15$ ). Median IgG levels are indicated for each peptide/protein (horizontal lines). Peptides/proteins with an AUC (area under the curve)  $>0.98$  and high median IgG levels are boxed.



HIV-infected patients as well as in control individuals (Fig. S1b†).

### The HIV microarray can be used to measure HIV-specific antibody responses for various isotypes and IgG subclasses

We also evaluated if the HIV microarray could be used for detection of HIV-specific IgG subclasses (*i.e.* IgG<sub>1</sub>, IgG<sub>2</sub>, IgG<sub>3</sub> and IgG<sub>4</sub>) and for detection of specific IgA, IgM and IgE responses in sera from HIV-infected patients. Fig. 6 shows examples of scan-images of microarray-based antibody determinations of an HIV-positive and a control serum. In the HIV-positive serum sample shown, specific IgG and IgG<sub>1</sub> antibodies were found towards many of the HIV-derived components, whereas specific IgG<sub>3</sub>, IgA and IgM reactivity occurred only towards certain HIV-derived and allergen components (Fig. 6). In serum samples from other HIV-infected patients we detected also IgG<sub>2</sub> and IgG<sub>4</sub> subclass responses towards HIV antigens/peptides, whereas we found so far no IgE responses against HIV components (Gallerano *et al.*, unpublished data).

### The HIV microarray for monitoring the specificities and magnitudes of HIV-specific antibody responses during the course of disease and treatment

Next, we studied if the HIV microarray could be used to monitor HIV-specific IgG responses during the course of

disease and treatment. For this purpose we analysed IgG reactivity towards micro-arrayed peptides/proteins of 15 HIV-infected African patients from whom sera were available at different time-points (*i.e.*, 47 serum samples from different time points). In this set of experiments sera from 15 control serum samples and sample diluent alone were tested as negative controls. Interestingly, most of the HIV-infected patients preserved their antibody recognition patterns (*i.e.*, specificities, levels) over time (Fig. S2†). For single cases specific antibody reactivities were detected only at certain time-points (*e.g.*, Pat #8: gp41; Pat #9: 120/24, 41/11, REV/01; Pat #12: envelope peptides, MA, CA, PR). Fig. 7 shows two of these examples (patients #8, 9), which were analysed in relation to immunological, clinical and treatment parameters.

For patient #8 (Fig. 7a) four serum samples, taken over a period of 21 months, were analysed. Despite drops of the CD4 counts at the times when the second and third blood samples were taken, the IgG recognition profiles remained almost unchanged and there were no strong alterations regarding the levels of IgG responses towards the tested peptides and antigens. Only a few changes were noted. For example, IgG levels towards 120/24 and 41/02 decreased over time and 120/16-specific IgG levels went down in the second serum sample and increased in the third and fourth sample. gp41-specific IgG was only detectable in the

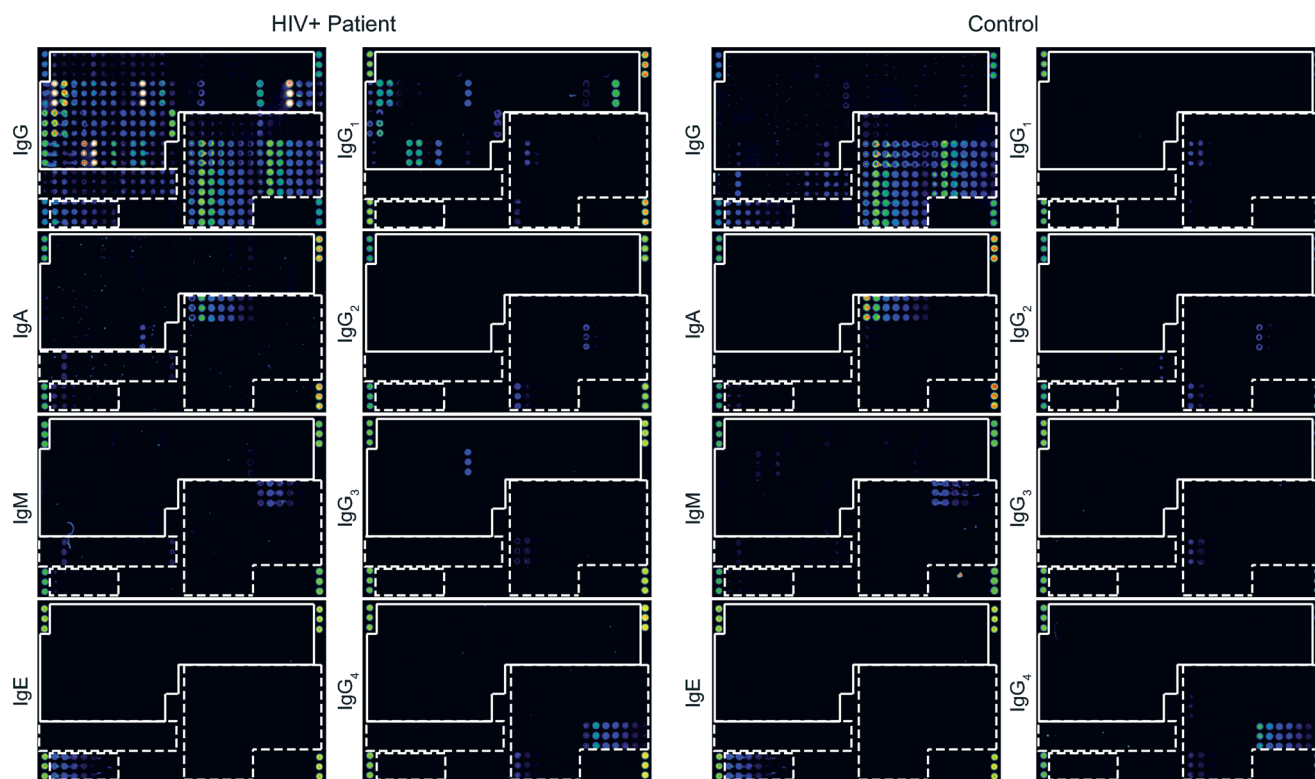
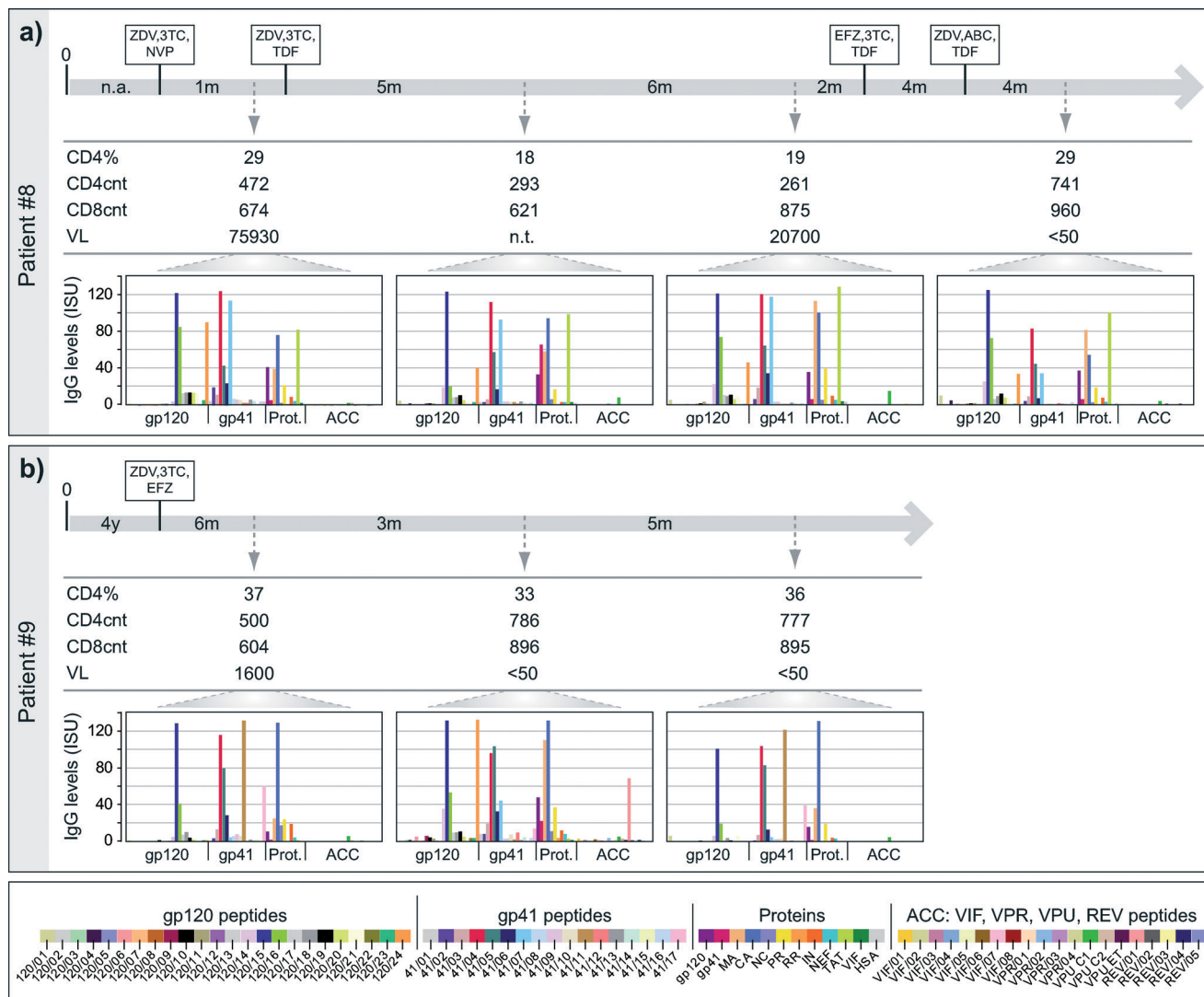


Fig. 6 IgG, IgA, IgM, IgE and IgG<sub>1-4</sub> reactivities to micro-arrayed antigens. Scan images obtained by testing serum from an HIV-positive subject (left panel) and control serum (right panel) for IgG, IgA, IgM, IgE and IgG<sub>1-4</sub> reactivity towards micro-arrayed HIV peptides/proteins and control proteins. Increasing fluorescence intensities from blue to red/white correspond to the amount of bound antibodies. The layout of spotted antigens is shown in Fig. 1c. HIV peptides and proteins are boxed and control antigens are surrounded by a broken line.





**Fig. 7** Time course of IgG reactivities to micro-arrayed HIV proteins and peptides in sequential serum samples from HIV-infected individuals. Serum samples obtained from two HIV-infected individuals, (a) patient #8 and (b) patient #9, were obtained at different time points (x-axes, with time intervals). Indicated are the time-points of infection (0), anti-retroviral treatment (HAART regimens boxed: Abacavir, ABC; Efavirenz, EFZ; Lamivudine, 3TC; Nevirapine, NVP; Tenofovir, TDF; Zidovudine, ZDV), time-points of serum collection (arrows), percentages of CD4 cells (CD4%, CD4cnt/CD45cnt  $\times 100$ ), CD4 and CD8 counts (CD4cnt and CD8cnt, cells  $\text{mm}^{-3}$ ), viral loads (VL, copies  $\text{ml}^{-1}$ , n.t., not tested) and IgG levels (y-axes: ISU) towards gp120 peptides, gp41 peptides, HIV proteins and accessory protein-derived peptides (bottom line: colour code).

third serum sample but not in the others. In patient #9 (Fig. 7b), who had a stable treatment response (CD4 counts  $>500$  cells  $\text{mm}^{-3}$  and VL decreasing  $<50$  copies  $\text{ml}^{-1}$ ), we also observed a rather conserved antibody recognition profile. Again, only a few changes were noted. For example, in the second serum sample IgG levels towards 41/11 and 41/17 decreased (60-fold and 4-fold, respectively) and high IgG levels were measured towards 120/14, 120/24, 41/07, gp120, MA and REV/01, although IgG was absent or low to these peptides/proteins at the other time-points.

The results thus demonstrate that HIV microarray allows detecting changes of epitope specificity in longitudinal assessments in given individuals.

## Discussion

We developed HIV chips containing six microarrays, on which a large panel of proteins and peptides of the HIV-1 clade C proteome were immobilized for the analysis of HIV-specific antibody responses. The miniaturized assay allowed the measurement of IgG, IgG subclass, IgM, IgA and IgE responses towards a broad panel of HIV peptides and proteins, with minimal amounts of spotted antigens (*i.e.*, 50–200 fg), minute volumes of serum (*i.e.*, 0.6  $\mu\text{l}$  for IgG, IgG<sub>1</sub>, IgA, IgM; 3.0  $\mu\text{l}$  for IgG<sub>2-4</sub>, 30  $\mu\text{l}$  for IgE measurements), consuming low amounts of detection antibodies (*i.e.*, 0.03  $\mu\text{g}$ ) and in short assay-duration (*i.e.*  $<3$  h). To test IgG reactivity to the same





72 HIV-derived components and a control component, immunoassays such as ELISA and Western-blot would require approximately 0.4  $\mu\text{g}$  of protein/peptide, 50–80  $\mu\text{l}$  of serum (considering a serum dilution 1:200) and 0.6–1.0  $\mu\text{g}$  detection antibody. With assay durations of 2–3 days for ELISA, only 20–80 tests per week could be performed manually by one person towards the same set of antigens in comparison to 600 tests per week that can be analysed by the microarray technology described here. Furthermore automated processing of a large number of chips is currently being developed.

The HIV microarray may well be used also for serological diagnosis of HIV infections but it must be born in mind, that currently available screening procedures that combine different antigen–antibody assays and nucleic acid based assays offer extremely high sensitivity and specificity.<sup>27–31</sup> Nucleic acid-based tests and tests focusing on the p24 antigen are particularly important for testing when antibody production is not yet detectable in patients shortly after infection and when the host's immune system is compromised.<sup>32,33</sup> Another focus in diagnostic testing is the discrimination of HIV-1 and HIV-2 infections<sup>33</sup> and the availability of tests that can be used for point of care testing and in countries under difficult conditions where laboratory facilities are lacking and costs are an important issue.<sup>34,35</sup>

Fast and comprehensive immunoassays such as the HIV microarray may therefore address another increasing need of the scientific HIV-research community. Multiplex tests will be useful for the screening of large numbers of samples in prevalence, population studies and vaccine trials. In fact it has been shown that multiplex assays deciphering immune antibody signatures towards a large panel of HIV antigens and epitopes may allow identifying protective immune responses.<sup>9,36</sup> One possibility to assess simultaneously antibody responses towards several antigens and epitopes as well as towards antigenic structures from different microorganisms is the use of various forms of microbeads containing different antigens.<sup>37–40</sup> However, with bead technology only a limited number of different antigens can be tested whereas the use of microarrays allows testing simultaneously for much larger numbers of antigens.

The HIV microarray developed by us contained 147 different components, which may be further increased by number. We noted that the micro-arrayed HIV-1 clade C peptides and proteins not only allowed mapping of HIV-specific antibody recognition profiles both in HIV-infected patients from an African region where clade C is endemic, but also in patients from Europe where clade B predominates. At present our micro-array comprised only clade C peptides and proteins but it may be considered to expand the repertoire of spotted components to include also antigens and peptides from other strains in order to test if we can identify strain-specific antibody signatures. At present the chip contains structurally folded recombinant HIV proteins as well as unfolded peptides and thus allows detection of antibody responses towards conformational as well as sequential epitopes. It may be also considered to expand the antigen repertoire regarding

carbohydrate epitopes and glycoproteins. We think that the HIV microarray is a suitable tool for the mapping of antibody responses towards HIV-derived peptides and also HIV recombinant proteins in large patient cohorts and trials and in populations from different geographic regions. The possibility to measure various isotypes and IgG-subclasses against a comprehensive set of HIV antigens and peptides in serum samples and other body fluids may also provide new information for the development of new therapeutic strategies. For example, the assessment of different Ig isotypes and IgG subclasses may be important because they are relevant for the effector functions of these antibodies and thus in virus defense (*e.g.*, complement activation, ADCC, *etc.*). Several studies indicate that certain isotypes/subclass responses may be associated with infection control or bad prognosis. For example, protective effects were suggested for gp120-specific IgG<sub>3</sub> in the RV144 vaccine trial<sup>41</sup> or for gp41-specific IgG<sub>2</sub> antibody responses regarding persistence of long-term non-progression.<sup>42</sup> HIV-specific IgA responses are found in sera and mucous secretions of HIV-infected patients (*e.g.* genital secretions, saliva) and in breast milk of infected mothers.<sup>43,44</sup> Micro-arrayed HIV components may therefore be interesting to study the fine specificities of these responses and to relate them to clinical findings.

First results from our study indicate that the HIV microarray can be used to detect changes of epitope specificity in longitudinal assessments in given individuals. Interestingly, we found changes in antibody recognition profiles even when plasma viral loads were undetectable. It is thus possible that the presence of HIV cellular reservoirs in anatomical compartments other than blood boosts antibody production and/or that long-lived plasma cells continuously secrete HIV-specific antibodies.<sup>45–48</sup> Unfortunately detailed PCR data from the investigated subjects were not available for a comparison with the antibody signatures. However, any direct comparison of nucleic acid-based tests and tests measuring specific antibodies must be considered difficult because nucleic acid-based tests will measure already the presence of a microorganism whereas antibody-based tests reflect the host immune response against the microorganism.

In summary we believe that the HIV chip will be valuable for the analysis of isotype and subclass responses towards a comprehensive panel of HIV components and thus may be useful for gaining new insights into HIV-specific immune responses, for diagnosis and monitoring of treatment strategies.

## Conclusion

The HIV chip offers a miniaturized platform containing a comprehensive set of antigens and peptides covering the HIV proteome for the assessment of the specificity and magnitude of HIV-specific IgG, IgG-subclass, IgA, IgM and IgE responses. We showed that the HIV microarray allows diagnosis as well as monitoring of HIV-specific antibody responses during the natural course of infection and treatment. The HIV



multiplexed immunoassay will be particularly useful for the mining of complex protein and peptide-specific antibody signatures in populations, cohorts and vaccine trials.

## Conflict of interest

DG, ES and RV are authors on a patent application regarding the use of clade C peptides/proteins for diagnosis of HIV infections. RV serves as a consultant for Biomay AG, Vienna, Austria, ThermoFisher, Uppsala, Sweden and Fresenius Medical Care, Bad Homburg, Germany.

## Abbreviations

$\alpha$ -huIg	anti-human Ig
AUC	area under the curve
BSA	bovine serum albumin
CA	capsid
ELISA	enzyme-linked immunosorbent assay
gp120	glycoprotein 120
gp41	glycoprotein 41
huIg	human immunoglobulin
HAART	highly-active anti-retroviral treatment
HSA	human serum albumin
IN	integrase
ISAC	immuno solid-phase allergen chip
ISU	ISAC standardized unit
MA	matrix
NC	nucleocapsid
NEF	negative factor
OD	optical density
PBS	phosphate buffered saline
PR	protease
REV	regulator of virion expression
ROC curve	receiver operating characteristic curve
RR	reverse transcriptase + RNaseH
SD	standard deviation
TAT	trans-activator of transcription
VIF	virus infectivity factor
VPR	viral protein R
VPU	viral protein U

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