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Human B-cell ontogeny in humanized NOD/SCID γ_c^{null} mice generates a diverse yet auto/poly- and HIV-1 reactive antibody repertoire

Hong Chang^{1,2}, Subhabrata Biswas¹, Aimee St. Clair Tallarico, Phuong Thi Nguyen Sarkis, Shusheng Geng³, Madhura M. Panditrao, Quan Zhu, and Wayne A. Marasco

Department of Cancer Immunology and AIDS, Dana-Farber Cancer Institute and Department of Medicine, Harvard Medical School, Boston, MA 02215

Abstract

Characterization of the human antibody (Ab) repertoire in mouse models of the human immune system is essential to establish their relevance in translational studies. Single human B-cells were sorted from bone marrow and periphery of humanized NOD/SCID γ_c^{null} mice at 8–10 months post-engraftment with human cord blood-derived CD34⁺ stem cells. Human immunoglobulin variable heavy (V_H) and kappa (V_K) genes were amplified, cognate V_H - V_K gene-pairs assembled as single-chain variable fragment-Fc antibodies (scFvFc) and functional studies performed. Although overall distribution of V_H genes approximated the normal human Ab repertoire, analysis of the V_H -third complementarity determining regions (H-CDR3) in the mature B-cell subset demonstrated an increase in length and positive charges suggesting autoimmune characteristics. Additionally, >70% of V_K sequences utilized V_K4-1 , a germline gene associated with autoimmunity. The mature B-cell subset-derived scFvFc displayed the highest frequency of autoreactivity and polyspecificity, suggesting defects in checkpoint control mechanisms. Furthermore, these scFvFc demonstrated binding to recombinant HIV envelope corroborating previous observations of poly/autoreactivity in anti-HIVgp140 antibodies. These data lend support to the hypothesis that anti-HIV BnAbs may be derived from auto/polyspecific Abs that escaped immune elimination and that the hNSG mouse could provide a new experimental platform for studying the origin of anti-HIV neutralizing Ab responses.

Keywords

humanized mouse; single B-cell; antibody repertoire; autoreactive; checkpoint control

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CORRESPONDENCE: Wayne A. Marasco, M.D., Ph.D., Department of Cancer Immunology and AIDS, Dana-Farber Cancer Institute JFB824, Boston, MA 02215; Phone: (617) 632 2153; Fax: (617) 632 3889; wayne_marasco@dfci.harvard.edu.

¹These authors contributed equally to this work

²PRESENT ADDRESS: Department of Biochemistry, School of Basic Medicine, Hebei Medical University, Shijiazhuang, China 050091

³PRESENT ADDRESS: Tian Guang Shi Biotechnology Co. LTD, Beijing, China 101111

Conflict of interest

The authors declare no conflict of interest.

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Introduction

Advancement in high-throughput screening techniques has led to the recent discovery of several highly potent broadly neutralizing antibodies (BnAbs) against HIV¹⁻⁴ and Influenza A⁵ recovered from peripheral blood (PB) derived B-cells of infected individuals; however, the occurrence of these BnAbs is extremely rare. This has spurred a renewed interest in 'rational vaccine design' where it may be feasible to analyze the patient's antibodyome in order to obtain insight into the ontogeny of the BnAbs.⁶ This information may then be used to design candidate vaccines in order to improve BnAb responses.⁴

Given the cost and ethical constraints of using human subjects for investigative vaccine studies, there is a growing need for a predictive and surrogate system to study human Ab evolution at the single-cell level. "Humanized" mouse models are increasingly being used to study human immunity, developmental and disease processes.^{7, 8} Newer mouse models deficient in the expression of the interleukin-2 receptor (IL2R) γ -chain (γ^{null}), including NOD/SCID γ^{null} (NSG), BALB/c-Rag2^{-/-} γ^{null} and H2^d- Rag2^{-/-} γ^{null} mice support the development of a multi-lineage human hemato-lymphoid system following transplantation with fetal or adult hematopoietic stem cells (HSC). Additionally, these engrafted γ^{null} mice exhibit normal life spans, unlike previous models, thus enabling long-term studies.⁹ In spite of these favorable advances, the adaptive Ab responses of these animals are weak with barely detectable secondary responses including class switching and affinity maturation.⁷ Growth factor supplementation with human BLYS¹⁰ and T cell-cytokines¹¹ in order to support growth and differentiation of the transplanted cells has resulted in only marginal improvement. Treatment of these mice with human cytokines and other costimulatory/growth factors delivered by a variety of techniques are being actively investigated to further improve human immune system development.¹²

Clonal diversity and immune tolerance are two major cornerstones of an effective Ab response that must also be considered in the evaluation of these mice as a relevant platform system to study human Ab responses. Several studies have evaluated immune repertoire complexity in hNSG mice by TCR CDR3 spectratyping,⁹ BCR H-CDR3 immunoscope analysis¹³ and multiplex PCR of V-J rearrangements of TCR β and H-CDR3¹⁴ and have concluded that both repertoires show levels of diversity comparable to those of humans. In addition, there have been two reports that analyzed the diversity of the IG repertoire with a focus on only the V_H4 family in NOD/SCID and NOD/SCID/ β 2m^{null} mice.^{15, 16} However, a systematic study in which the diversity of the human B-cell repertoire is analyzed via genetic and functional analysis of the variable (V), diversity (D) and joining (J) gene segments of the IG heavy and light chain genes has not been performed in hNSG mice. Analysis of immune tolerance in hNSG mice by the evaluation of the physiologic checkpoint control mechanisms that are normally operative during B-cell development^{17, 18} has also not been reported (see Mouquet et al¹⁷ for schema).

In the present study, analysis of V_H and V_K gene arrangements in hNSG-derived single human B-cells sorted at different developmental stages was performed. Nucleotide and amino acid sequence analysis of the heavy chain genes indicated the presence of a diverse antibody repertoire; however, characterization of H-CDR3 regions and a specific restriction

in the V_{κ} repertoire suggested an autoreactive potential. This was further confirmed by functional studies where scFvFc clones from single B-cells were found to exhibit binding to self-antigens. Intriguingly, many autoreactive clones also displayed affinity for HIV-1 envelope protein gp140 (HIV-1gp140). These data lend support to the contemporary hypothesis that anti-HIV BnAbs may be derived from auto/polyspecific Abs that escaped immune elimination.^{19, 20} Thus, the defects in immune tolerance in these hNSG mice may provide a unique model to study the development of anti-HIV BnAbs where ancestral origins of antibodies can be determined and the existing HIV-reactive Ab clones can be modulated via experimental infection or immunization.

Results

B-cell development in long-term engrafted hNSG mice

Significant numbers (>10% of total lymphocytes) of human CD45⁺ (hCD45⁺) cells could be detected in the PB of hNSG mice between 1–2 months post HSC injection; however, total human CD3⁺ T cells remained low (<5% of total hCD45⁺ cells) during that time. At 6 months post HSC injection, the hCD45⁺ frequency in the PB of the hNSG mice was 43.87 ± 10.11% (n = 8) of total lymphocytes, of which 41.49 ± 22.75% stained for hCD3 (T cells) and 53.28% ± 21.5% for hCD19 (B-cells). Of the total CD19⁺ human B-cells, 53.64 ± 12.69% expressed the CD5 surface antigen, which was higher than the reported level of circulating CD19⁺CD5⁺ cells (~15%) found in humans.²¹ Profiles of the different CD19⁺ B-cell subpopulations in the hNSG mice at >6 months post HSC transplantation at the time of single B-cell sorting are shown in Figure 1. B-cells at all major stages of the developmental pathway (early immature, immature, new emigrant and mature naïve) were detected, but CD27⁺ B-cells, associated with a memory phenotype²² were notably absent. Overall, the data demonstrate that the human B-cell developmental pathway is recapitulated in the hNSG mice; however the memory B-cell compartment is not or only sparsely populated.

Analysis of human V_H sequences

Table 1 summarizes the total data set analyzed in the present study. Single B-cells from different B-cell subpopulations were sorted, and IG gene sequences were obtained as described in Methods. In total, 257 and 264 sequences unique for human immunoglobulin (IG) heavy and light chains, respectively, were obtained and further analyzed. Intriguingly, in the first round PCR reactions to amplify kappa (V_{κ}) and lambda (V_{λ}) light chain genes, only those sequences that corresponded to kappa light chain gene segments were recovered. PCR mediated primer bias was ruled out by using the same primer mix to amplify heavy and light chain genes from single-sorted mature B-cells from a normal human donor. From a total of 192 single-sorted human PB B-cells, 137 V_H , 44 V_{κ} and 33 V_{λ} unique and productive gene segments could be amplified (Supplementary Figure 1). The reason(s) for this observed exclusivity of the V_{κ} genes is currently under investigation and may include factors, such as V_{λ} transcript instability, greater V_{λ} variability and V_{κ} gene rearrangement bias in the hNSG mice.

Individual IG sequences were aligned with the International Immunogenetics Information System (IMGT) database, and the overall germline usage was determined. Human V_H gene

segments are divided into 7 families, named V_H1–7 with 51 known functional genes and >40% of the genes representing the V_H3 family (IMGT). V_H sequences from all except V_H2 and V_H6 families were represented in the hNSG mice, and the sequence usage from different V_H gene families across the B-cell subpopulations is shown in Figure 2, upper panel. In comparison to the human V_H repertoire found in normal adults,^{23, 24} gene segments corresponding to the V_H3 family were most frequently utilized followed by V_H4 in most of the B-cell subpopulations. An exception was observed in new emigrant B-cells where the V_H3 usage (17.4%) was marginally lower than that of V_H4 (22.5%). Additionally, V_H7 gene segments were detected at a frequency higher than normally found (~2% in the human V_H repertoire, IMGT), particularly in the early immature stage of B-cell development where it was ~12% of the total V_H sequences (Supplementary Table 1). No other significant differences were found in the distribution of the V_H gene families across the different B-cell subsets. These observations suggest that the V_H repertoire in the hNSG mouse is quite diverse with 36 out of 51 known functional genes found at least once in the various subpopulations (Supplementary Table 1). Among individual gene segments, V_H4–34, V_H3–30 and V_H1–2 were utilized most often. V_H4–34 was most frequently observed in the repertoire of all the B-cell subsets, except for the immature subpopulation where V_H3–30 had the highest occurrence.

Analysis of V_H rearrangement and H-CDR3 composition

The complexity of the IG repertoire is enhanced by gene rearrangement of V_H with D_H and J_H segments and V_{κ/λ} with J_{κ/λ} segment for the light chain. Therefore, the utilization of the D_H and J_H gene segments across the different B-cell subpopulations was analyzed next (Figure 2, middle and lower panels, respectively). The D_H genes are classified into 7 families, D_H1–D_H7. As reported for humans,²⁵ the usage of the D_H3 family in the hNSG mice was significantly more frequent than the other families (p<0.005, Exact Fisher-Freeman-Halton Test). An increase in the usage of D_H1 and D_H2 segments along the CD5⁻ B-cell developmental pathway was observed (early immature<immature<new emigrant<mature). There were differences between the mature CD5⁻ and CD5⁺ B-cells in terms of D_H gene segment usage, particularly with regards to the utilization of D_H2 (lower in CD5⁺ B-cells), D_H3 (higher in CD5⁺ B-cells) and D_H5 gene segments (absent in CD5⁺ B-cells). Rearrangement of D_H gene segments may occur either through inversion or deletion processes which determines the D_H reading frame utilized. An analysis of the usage of the D_H reading frame (RF) indicated a significantly higher utilization of RF2, found on an average of 51% of all the sequences across all B-cell subsets in the hNSG mice (p<0.005, Fisher's Exact Test, data not shown). This draws a parallel to that observed for normal human D_H gene rearrangements where deletion was found to be the favored process along with a lower frequency of the use of RF3.²⁵ The distribution of J_H genes clearly showed a higher bias for J_H4 usage (p<0.005, Fishers' Exact Test). No significant difference was observed between the mature B-cell subsets (p>0.005, Fisher's Exact Test). However, distribution of J_H6 usage compared to all other J_H segments (except J_H3) was found to be significantly higher in the mature B-cell subsets (CD5⁺ and CD5⁻) compared to the BM-derived B-cells (early immature and immature B-cells) (p<0.005, Fisher's Exact Test).

The H-CDR3 region is considered to be an important determinant that imparts antibody specificity as well as affinity. The overall composition of H-CDR3 regions in terms of length, positive charge content, N-nucleotide insertions and exonuclease activity was analyzed (Figure 3). Extremely long (>20 amino acids) H-CDR3 regions were more frequent in the CD5⁻ mature B-cell subset, which may be explained by the higher utilization of the J_H6 segment by this group of cells that results in longer CDR3.²⁵ Additionally, this same B-cell subset carried more cells with 2+ and 3+ positively charged residues in the H-CDR3 compared to the CD5⁺ B subset. While increases in length and positive charges in the H-CDR3 region are expected in the IG repertoire from BM-derived immature B-cell populations, these are less frequently observed when the B-cells mature and migrate to the periphery. Thus, the mature B-cell subset in the hNSG mice with longer and increased positive charges in their H-CDR3 regions is a major and previously unrecognized difference from their counterpart in normal humans. No significant difference in N-insertions and J_H exonuclease activity between the B-cell subpopulations was observed (p<0.005, Fishers' Exact Test) (Figure 3b). A bias in Tyrosine (Y) and Glycine (G) composition in the H-CDR3 regions (40–50% of total residues) that is observed in normal humans²⁶ was recapitulated in the hNSG mice (Figure 3a).

Presence of hypermutated V_H sequences

Hypermutation in the V_H sequences was analyzed by comparing against the closest germline sequences, and the B-cell subpopulations were classified as those with identical or highly homologous (99–100% homology), moderately mutated (94–99%) and highly mutated (<94%) sequences as described earlier for analysis of human IG V_H repertoire²⁴ (Table 2). Accumulation of hypermutated V_H sequences (<99%) was significantly lower in the mature B-cell subsets in the periphery compared to the earlier stages in development in the bone marrow (p<0.005, Fishers' Exact Test). In contrast to an earlier report,²⁴ where IgM⁺CD5⁺ human B cells were found to accumulate significantly higher number of mutations compared to the IgM⁺CD5⁻ B cells, we did not observe a significant difference between the CD5⁻ and CD5⁺ mature B-cell subsets (p<0.005, Fishers' Exact Test). Moreover, the V_H repertoire in the peripheral B cells were notably devoid of highly mutated sequences, e.g., displaying <94% homology to germline sequences. Analysis of the mutations in the Framework Regions (FR) and H-CDR3 both at the nucleotide and amino acid residue levels revealed the occurrence of both silent as well as replacement mutations (Table 2).

Potential V_H replacement in hNSG mice

The cryptic recombination signal sequence (cRSS, TACTGTG)²⁷ at the 3'-end of rearranged V_H genes allowed the detection of potential V_H replacement products. The cRSS is the binding motif for the recombinase activating gene (RAG) family of enzymes that mediate V_H replacement, a process in which a secondary arrangement of an upstream V_H sequence occurs in an already formed V_H-D_H-J_H gene.²⁷ Remarkably, 13 potential V_H replacements in the 257 V_H sequences were detected (Table 3). This secondary rearrangement process was not confined within the early stages of B-cell development and did not only involve V_H4 genes as reported earlier.^{28, 29} In order to confirm the validity of the sequences scored as the products of V_H replacement, a similar scan was performed on the D_H-J_H junction of the V_H sequences. One major criteria of such replacement products, as shown in Table 3, is

that they often encode highly charged amino acid residues (R/D/E).³⁰ No such sequences were identified in the D_H-J_H region of the V_H genes.

Analysis of human V_κ sequences and cognate V_H-V_κ pairs

Figure 4a presents an analysis of the V_κ sequences recovered from the different human B-cell subsets obtained in the hNSG mice. The human κ locus contains 44 functional gene segments divided into 5 gene families: V_κ1–V_κ5 (IMGT). Only a single member is known for each of the V_κ4 and V_κ5 families, V_κ4-1 and V_κ5-2 respectively (IMGT). At least 24 unique V_κ gene segments, representing 4 out of 5 V_κ gene families in the B-cell subpopulations, were detected with a distinct bias towards the utilization of V_κ4-1 (Supplementary Table 2). Notably, the V_κ4-1 gene segment is located most proximally in the human κ locus on chromosome 2; however, the significance of this location with regards to the overutilization of this gene in the hNSG mice is presently unclear. V_κ5 family genes remained undetected in the hNSG mice, which may be attributed to the limited sample size. An analysis of the J_κ sequences demonstrated the utilization of all known J_κ segments in the human IG repertoire (J_κ1–J_κ5) with an overrepresentation of the J_κ4 gene segments across all B-cell subpopulations (Figure 4a).

V_H-V_κ pairs derived from single B-cells were interrogated for the frequency of pairing of individual heavy and light chains across different cell subsets (Figure 4b), and a complete list of the pairings is shown in Supplementary Figure 2. Given that each V_H-V_κ pair represents a unique Ab, the profile of heavy and light chain pairing across the different B-cell subsets can provide information on antibody evolution, clonality, editing and other physiological processes that shape the IG repertoire. Interestingly, as represented in Figure 4b, a significant decrease in V_H-pairing with V_κ4-1 in mature B-cells (CD5⁺ and CD5⁻) was found when compared to BM-derived B-cell subsets (p<0.005, Fishers' Exact Test). Given that the usage of V_κ4-1 has been associated with autoreactivity and anti-DNA autoantibodies,³¹ the decreasing frequency of V_κ4-1 usage with increasing B-cell maturation suggests that checkpoint control mechanism(s), which are associated with peripheral tolerance, are at least partially intact.

Immunoglobulin levels and auto/polyreactive antibodies in hNSG mice

Total human IgM and IgG concentrations in hNSG mouse serum ranged between 15–50 and 20–90 µg/ml, respectively, which were ~10–50 and ~100–200 fold lower than normal human serum IgM and IgG levels. By comparison, in newborns the antibody levels approximate that found in adults, albeit they increase with time, plateauing at ~1 year for IgM and ~5–6 years for IgG.³² It has also shown that in human bone marrow transplants, serum IG levels return within normal levels by day 90 post transplant.³³

The physiological presence of auto/polyreactive antibodies was next tested by analyzing the binding of serum IgM and IgG antibodies by ELISA to a panel of antigens, including cardiolipin, insulin, ssDNA, dsDNA and LPS, and the overall reactivity was compared to normal human sera. As shown in Figure 5, the hNSG mice sera showed significantly higher binding levels over that of normal human sera to all antigens tested, and this auto/polyreactivity was almost entirely restricted to the IgM fraction (p<0.005, Mann Whitney

Test). An increased level of binding to LPS by both normal human IgG and IgM may be attributed to 'natural antibodies' present in the serum.³⁴

To investigate this auto/polyreactivity in more detail, additional studies were performed on recombinant Abs recovered from single B-cells. As shown in Table 1, 176/219 cognate V_H-V_K pairings representing all different human B-cell subsets were successfully assembled into scFvs (80% efficiency), expressed as scFvFc and purified for functional studies. To determine the autoreactivity of these Abs, HEp-2 (human epidermoid cancer cell) ELISA which is a standard clinical assay for detection of anti-nuclear antibodies (ANAs) was performed. As shown in Figure 6a, autoreactive ANAs were detected in all B-cell subsets. Circa 50% of the scFvFc from all B-cell subsets were scored as autoreactive, with the exception of the CD5⁻ mature B-cell subset where autoreactivity was observed in a significantly higher percentage of the Abs (80.7%) compared to the other B-cell subsets (p<0.005, Mann Whitney Test). The reactivity of all 176 scFvFc against each antigen is shown in Figure 6b. There was a significant increase in the median reactivity levels of antigen binding in the mature B-cell subsets when compared to that observed for the B-cells at earlier stages in development (p<0.005, Mann Whitney Test). No significant difference was observed between the CD5⁺ and CD5⁻ mature B-cell subsets. All polyreactive Abs also scored positive in the ANA assay. Additionally, scFvFc demonstrating high level reactivity to one antigen also bound strongly to other antigens. Thus, the results of both the serologic and single B-cell derived binding studies provide evidence for the retention of a substantial pool of auto/polyreactive B-cell clones centrally in the bone marrow and particularly in the periphery, suggesting that the first two checkpoint control steps (between early immature to immature and new emigrant to mature) involved in clearance of autoreactive clones are impaired.¹⁷ ELISA binding studies using a comparable mouse cell line to test scFvFc autoreactivity to mouse ANA antigens, may provide additional information on autoreactivity given the development of these human B-cells in the murine host.

A total of 22 scFvFc were scored based on their absolute MSD values (>3 fold background as obtained from reactivity of buffer alone to antigen coated wells) as low, medium, and highly polyreactive based on the relative reactivity of 4E10 to the same antigens (<20%, 20–60%, and 60% of 4E10 scFvFc reactivity, respectively). Of these 22, 2 were from the early immature B-cell subset, 1 from immature, 2 from new emigrant, 13 from the CD5⁻ mature B-cells and 4 from CD5⁺ B-cells. To identify any common features between the 22 polyreactive scFvFc in terms of sequence composition, the respective V_H and V_K families and H-CDR3 length and positive charge composition were compared (Supplementary Table 3); however, except the fact that the majority of the light chain sequences corresponded to V_K 4-1, no other obvious similarities was observed.

Polyreactivity extends to HIV-1gp140 binding

Previous studies have reported that several anti-HIV-1 envelope directed BnAbs including 2F5 and 4E10 show promiscuous binding to multiple auto-antigens, which raises the hypothesis that their rare appearance may be due to escape from immune tolerance.^{35, 36} A more recent study demonstrated that 75% of all anti-HIV antibodies isolated from infected individuals showed polyreactivity against ssDNA, dsDNA, cardiolipin, LPS and keyhole

limpet hemocyanin by ELISA.¹⁹ Given the auto/polyreactive binding profiles above, both serum and the hNSG-derived scFvFc were tested for reactivity to HIV-1 envelope. As shown in Figure 7a, sera samples from hNSG mice displayed significantly higher reactivity to HIV gp140 compared to human serum controls with most of the reactivity restricted to the IgM fraction. Given that the IgG content in normal human serum is 100–200 folds higher than that in hNSG sera, the apparently higher reactivity of normal human IgG is most likely a concentration effect rather than antigen specificity. Furthermore, in a similar pattern to that described earlier, the scFvFc derived from the mature B-cells displayed significantly higher binding to HIV-1 envelope protein compared to the scFvFc derived from B-cells earlier in developmental pathway (Figure 7b).

Kinetic analysis of two random scFvFc displaying high (N1M10-1, isolated from CD5⁺ B-cell subset) and medium reactivity (NM10-1, isolated from CD5⁻ mature B-cells) for binding to HIVgp140 trimer protein was performed using Biolayer Interferometry, and k_{on} , k_{off} and K_D values of the scFvFc (b12 and 4E10 IgG used as positive controls) were measured. The sensorgrams of each antibody-binding (which were immobilized on Protein A sensors) to a series of different concentrations of HIV envelope protein in solution are shown in Figure 7c. The association rates (k_{on}) of the scFvFc were comparable to that of b12 and 4E10 [k_{on} (1/Ms) values were 4.69×10^6 and 5.54×10^5 for N1M10-1 and NM10-1, respectively, whereas it was 2.89×10^5 and 2.55×10^5 for b12 and 4E10 respectively]; however, the scFvFc dissociated from the target protein at a faster rate [k_{off} (1/s) values were 2.68×10^{-2} and 2.40×10^{-2} for N1M10-1 and NM10-1, respectively, whereas it was 4.19×10^{-5} and 1.42×10^{-4} for b12 and 4E10 respectively]. Intriguingly, the equilibrium dissociation constant (K_D , in M) values of the scFvFc obtained from naïve hNSG mice closely approximated those of the HIV BnAbs, with K_D values of 5.72×10^{-9} for both N1M10-1 and NM10-1, and 1.45×10^{-10} and 5.56×10^{-10} for b12 and 4E10 respectively. Although these Abs displayed substantial affinity to HIV envelope protein, they did not show neutralizing activity to HIV-1 or compete with target binding of b12 or 4E10 (data not shown) most likely due to their fast dissociation rates.

Auto/polyreactivity is still present in “humanized” BLT and GTL mice

To address the question of whether T cells affect the antibody repertoire as suggested earlier,³⁷ similar studies were performed with humanized BLT and GTL mice. Although both the BLT³⁸ as well as the GTL mice display a rapid development of human T cells (human T:B cell levels in these mice approximate that found in normal adults) as a result of co-engraftment of human thymic tissue in addition to CD34⁺ stem cells, preimmune serum antibody levels of human IgM and IgG were not improved over that observed for the hNSG mice. Following similar methods as described for the hNSG mice, single human B-cell sorting of mature splenic B-cells (CD19⁺IgM⁺CD10⁻), Ab gene recovery and scFvFc assembly were performed from a single BLT and GTL mice 16–20 weeks post tissue transplantation. A total of 59 and 37 scFvFc was assembled from mature human B-cells isolated from the spleen of a BLT and GTL mouse, respectively. The gene usage profile for heavy and light chain variable regions was comparable to that observed for the hNSG mice, including the lack of somatic hypermutations in the V_H sequences, sole recovery of V_K genes and an overutilization of V_K4-1 (data not shown). A majority of the scFvFc, ~75%

and ~78%, from the BLT and GTL mice respectively, also displayed auto- and polyreactivity as shown in Supplementary Figure 3. These results suggest that the polyreactive IG repertoire is not exclusive to the hNSG model, and it is likely that other “humanized” mouse models may share common deficiencies in peripheral tolerance mechanisms.

Discussion

In this study, we performed an in depth analyses and characterization of the human antibody repertoire in the humanized NSG mouse model. The hNSG mice supported the development of a multilineage human hematopoietic system following transplantation with human hematopoietic stem cells, and at 8–10 months post-transplantation, phenotypic characterization indicated a normal B-cell developmental pathway with immature B-cells residing predominantly in the bone marrow and the mature cells in the periphery. However, several deviations were noted, some of which have been reported earlier, e.g., slow development of human T cells⁹ and a higher than normal frequency of CD19⁺CD5⁺ B-cells in the periphery.^{38, 39} Additionally, the current work provides evidence for the first time that the human antibody repertoire that develops in the hNSG and other humanized mouse models is largely auto- and polyreactive.

The human V_H diversity in the hNSG mice appears to closely approximate to that of the normal adult IG repertoire in terms of germline gene usage, but with an unexpectedly high level of positively charged, long H-CDR3s in the mature CD5⁻ population that displayed increased auto/polyreactivity. There was also an abnormal high contribution to the rearranged V_H gene pool from the CD5⁺ B cells in the periphery (Figure 6A). It has been previously shown that increased numbers of CD5⁺ B cells are found in the cord blood which display auto/polyreactivity.^{40, 41} It is doubtful that the source of human stem cells induces the development of CD5⁺ B cells in the humanized mice, since CD5⁺ B cells have also been shown to develop in the hNSG mice when engrafted with stem cells derived from the adult bone marrow or mobilized peripheral blood.³⁹ In early life, a restricted V_H repertoire resulting from limited IG diversity has been observed which coincides with the observed autoreactivity in the B cell clones isolated from fetal liver and cord blood.^{42, 43} In addition, in fetal liver B cells there is skewing in the distribution of V_H genes which are chromosomally located closer to the D_H-J_H.C_H locus.⁴⁴ Recently, next-generation sequencing (NGS) of expressed antibody repertoires from human cord blood cells have shown comparable frequencies of V_H germline gene usage to those present in adult IgM repertoire except for the high frequency of V_H1–2 germline gene that was preferentially expressed in the cord blood cells. As expected lower degree of somatic mutation in the CDR and framework regions was observed in the cord blood cells.⁴⁵ While our genetic analyses show both similarities and differences compared to the different V_H repertoires discussed above, the auto/polyreactivity of the hNSG antibodies set their functional maturity at an early stage.

The observations of a large pool of auto/polyreactive antibodies in the humanized mice made in this study raise two inter-related questions; 1, why are the autoreactive B cell clones retained in the periphery; and 2, are the auto/polyreactive antibodies in the humanized mice

akin to normal circulating ‘natural antibodies’ found in both men and mice.⁴⁶ During normal course of B-cell development, a majority of the self-reactive B-cell clones are eliminated by checkpoint control mechanisms as characterized by Wardemann and colleagues.¹⁸ A ‘central’ checkpoint occurs in the BM during the development of immature B-cells from the previous early immature stage and a ‘peripheral’ checkpoint between the new emigrant and the mature phase of B-cell development. A third checkpoint has also been predicted that eliminates residual autoantibodies from the IgM⁺ memory B-cells.⁴⁷ These checkpoint controls result in a significant reduction in H-CDR3 length and positive charge content from B clones isolated at the immature and mature naïve stages when compared to the corresponding early immature and new emigrant B-cell stages. Our data support that the peripheral checkpoint control mechanism in the hNSG mice may be impaired given the significant increase in the number of mature B-cells carrying long and highly charged H-CDR3 regions. The factors regulating these checkpoint control steps have not been identified, and human B-cell ontogeny in a xenogeneic environment adds to the complexity. Autoreactive B-cells are thought to be selected for elimination by their reactivity to self-antigens by deletion and receptor editing. In the hNSG BM, human B-cell development conceivably occurs by selection against both donor (human) as well as self (mouse) antigens and additionally, the distribution of these antigens may be varied in different compartments (e.g., bone marrow vs. periphery). This may result in improperly ‘educated’ human B-cells, particularly in the periphery where self-antigens predominate and the peripheral checkpoint control mechanism(s), designed to eliminate autoreactive clones, could be more impaired. Despite the apparent autoreactivity in the periphery, these mice did not display any apparent signs of autoimmunity, i.e. symptoms of graft-versus-host disease (GvHD), e.g., anorexia, wasting, alopecia, etc.

We verified to a great extent, that the predominance of V_κ sequences, particularly V_κ 4-1 gene segment, as the exclusive light chain partner in all assembled antibodies was not a PCR induced bias. The fact that cloned and serum antibodies alike displayed autoreactivity and polyspecific responses to multiple antigens provide a strong correlation with V_κ 4-1 prevalence. These results also illustrate an apparent defect or insufficiency in one of the major mechanisms of silencing self-reactive B-cells; light chain receptor editing which is mostly mediated by V_λ genes.⁴⁸ Polyreactive antibodies have also been detected in the IgG⁺ memory B-cell pools from normal human subjects, and these have been described as a by-product of extensive somatic hypermutation occurring during antigen-induced B-cell differentiation.⁴⁷ This observation is, however, irrelevant to the current study, as neither a memory phenotype nor a high level of hypermutation was observed in all the B-cell clones studied.

Natural antibodies secreted by B cells are predominantly of the IgM subclass, but are also represented by IgG and IgA isotypes. These antibodies have been characterized extensively in mice⁴⁹ and are present during early human life.⁵⁰ A distinct B cell subset, called B-1 cells, characterized by surface expression of the CD5 marker has been found to be the major producer of natural antibodies in mice. In addition to displaying low affinity interactions with self-antigens, these antibodies have also been found to play a major role in early protective responses against pathogens.³⁴ Further studies remain to be performed to understand the nature of the auto/polyreactive human antibodies that develop in the hNSG

mice under pathogen-free conditions and whether the CD5⁺ human B cells represent the murine equivalent.

A number of anti-HIV envelope BnAbs have shown characteristics of auto/polyreactive antibodies and thus, it has been proposed that the rare emergence of such antibodies in the vaccinated or infected host may be due to physiologic tolerance mechanisms that are operative during B cell development.⁵¹ Recently, it was also shown that HIV envelope reactive antibodies isolated from ‘elite controllers’ of infection were polyreactive and that heterologation to gp120 and self-antigen can increase the apparent affinity of the antibodies for HIV.¹⁹ The preponderance of peripheral auto/polyreactive antibodies in naïve and germ-free hNSG mice that also show binding to HIV envelope proteins may provide a model in which the generation and origins of protective antibody responses against HIV can be investigated. Indeed, since many of these antibodies already display binding to the HIV envelope, their potency could be further enhanced by experimental infection or immunization. Therefore, the hNSG and other similar mouse models may provide a novel experimental system in which the breakdown in physiologic tolerance mechanisms can be exploited to further investigate the role that auto/polyreactivity may play in the development and evolution of HIV BnAbs. This knowledge will be of critical importance to current and future HIV/AIDS vaccine efforts.

Materials and methods

Construction of humanized mice

Male NSG mice, 6 wk of age (Jackson Laboratory, Bar Harbor, ME), were housed under BSL-2 conditions at the Animal Research Facility (ARF), Dana-Farber Cancer Institute (DFCI, Boston, MA). Mice received autoclaved food and Baytril (fluoroquinolone) -treated water. CD34⁺ HSC were isolated from human umbilical cord blood, obtained from Brigham and Women’s Hospital (BWH, Boston, MA), using immunomagnetic column purification techniques (MiniMACS) per the manufacturer’s protocol (Miltenyi, Auburn, CA). At ~8 wk of age, mice were sublethally irradiated with 325 cGy (Gammacell-40, Best Theratronics, Ottawa, Canada) and then injected with 2.5×10^5 HSC resuspended in 200 μ l of phosphate buffered saline (PBS) via the tail vein. NOD/SCID-thy/liv (also called BLT) and NSG-thy/liv (GTL) mice were also constructed as previously described.³⁸ Briefly, 6–8 week old NOD/SCID or NSG female mice were ‘humanized’ following implantation of human fetal liver and thymus tissues under the kidney capsule along with an intravenous delivery of fetal liver-derived autologous CD34⁺ stem cells. All fetal tissues (17–20 weeks of gestational age) were obtained from Advanced Bioscience Resources, Alameda, CA. Tissues were also screened for the presence of HIV-1 and 2 and Hepatitis B virus and determined to be negative.

Mice were bled monthly via the mandibular route, and engraftment levels were determined by flow cytometry analysis of human CD45⁺ cells in the peripheral blood. All studies involving human tissues were approved by the Institutional Review Boards at both DFCI and BWH.

Flow cytometry

Analysis of human immune reconstitution following CD34⁺ HSC delivery was performed by flow cytometry (supplemental methods). Cell isolation techniques and criteria for single cell sorting from different human B-cell subsets were adopted from a previous report⁵² with modifications as described in Supplementary Methods. Single cell sorting from NSG mice was performed on early immature and immature B-cell subsets from the bone marrow (BM) and new emigrant/transitional and mature B-cell subsets (both CD5⁻ and CD5⁺) from the peripheral blood (PB).

Single cell RT-PCR and analysis of Ig heavy and light chain gene segments

RT-PCR reactions were performed as described earlier (Supplementary Methods).^{18, 52}

Expression and purification of scFvFc

Cognate variable heavy and light chain gene segments amplified from single B-cells were assembled following cloning into a human IgG1-Fc expressing vector, transiently transfected in 293FT cells (Invitrogen). The expressed single chain antibody fragments (scFvFc) were purified as described in Supplementary Methods.

Detection of self (auto)-reactive and polyspecific antibodies

QUANTA Lite™ ANA (anti-nuclear antibody) ELISA (INOVA Diagnostics, San Diego, CA) was used to test self-reactivity of the antibodies. 4E10 scFvFc and sera from patients and healthy individuals (manufacturer provided) were used as the positive or negative controls in this assay. Purified antibodies were tested at 50 µg/ml, and reactive samples were further confirmed at 25 µg/ml. The cutoff at OD450 for positive reactivity was calculated at 80% of the absorbance reading obtained for the manufacturer provided low positive control.

The electrochemiluminescent (ECL) based MSD platform (Meso Scale Discovery, Gaithersburg, MD) was used to evaluate the auto- and polyreactivity of hNSG-derived scFvFc. Human single- and double stranded DNA (ssDNA and dsDNA), recombinant insulin, cardiolipin, lipopolysaccharide (LPS, Sigma, St. Louis, MO) at 10 µg/ml and recombinant HIV-1 gp140-trimer from YU2 strain³⁸ at 5 µg/ml were used as polyreactive antigens in this assay. Each well in MSD 384-well High Bind MULTI-ARRAY® plates was coated with 5 µl of polyreactive antigens in PBS (cardiolipin in 20% alcohol) at 4°C overnight. The plates were blocked with 35 µl of 10% FBS/PBS for 1 h with shaking at 800 rpm on the microplate shaker. After washing with 35 µl PBST, purified scFvFc were added at 5 µg/ml, and the plates were incubated at room temperature for 2 h with shaking. 4E10 scFvFc was used as a positive control. Goat anti-human IgG-Fc Ab with SULFO-Tag (15 µl at 1 µg/ml in 1% FBS/PBS) was used as the detection antibody, and the plates were read on the MSD Sector Imager 2400 according to the manufacturer's protocol. Routine ELISA (Supplementary Methods) was also performed to analyze auto/polyreactivity in hNSG and normal human serum samples. 4E10 IgG (NIH, AIDS Research and Reference Reagent Program) was used as a positive control to estimate the level of autoreactivity.

Affinity measurements by Biolayer Interferometry (BLI)

Kinetic analysis of the hNSG-derived scFvFc_s, b12-IgG and 4E10 IgG was performed by measuring binding affinity to recombinant HIVgp140 trimeric protein. Analysis was performed by BLI using the Octet Red instrument (Forte Bio, Menlo Park, CA) (Supplementary Methods).

Statistical analysis

To analyze differences in distribution of various gene segments across different B-cell subsets, non-parametric statistical methods were used since the data were not normally distributed. Differences were assessed for statistical significance by the Fisher-Freeman-Halton test. Monte Carlo estimation (10 000 samples) was used to approximate the exact tests. Significant differences in the H-CDR3 length and N1 and N2 insertion across B-cell subsets were determined by fitting one-way analysis of variance using SAS PROC GLM (SAS, Cary, NC). Differences in the utilization of individual gene segments were analyzed by one sample binomial test. Differences in utilization of particular gene segments between mature and immature B-cell subsets were analyzed by using a two-sample test for equality of proportions with continuity correction. Mann-Whitney U tests (two-tailed) (Graph Pad, La Jolla, CA) were performed to determine statistically significant differences between median values of each ELISA data set. P values < 0.005 were considered significant.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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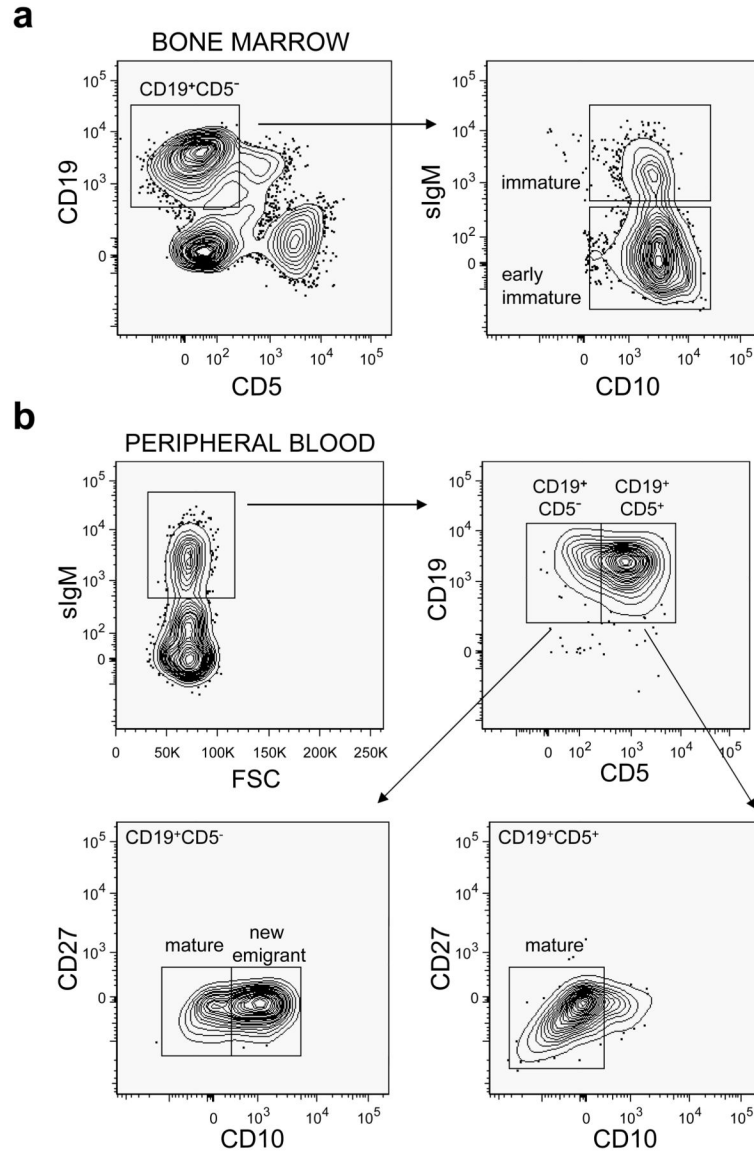


Figure 1. Sorting criteria of human B cell subsets from humanized NOD/SCID γ_c^{null} (hNSG) mice. (a) Single cell suspension was prepared from the bone marrow of a representative single hNSG mouse and cells in the lymphoid gate (not shown) were further gated for CD19⁺CD5⁻ expression (left). Cells were then plotted for surface IgM (sIgM) and CD10 expression (right), and single B cells from the early immature (sIgM⁻ CD10⁺) and immature (sIgM⁺CD10⁺) subsets were sorted. (b) PBMCs were prepared from 3 hNSG mice, all transplanted with the same donor and were gated for lymphoid population (not shown). These lymphocytes were gated on IgM⁺ cells (middle left) and further characterized as CD19⁺CD5⁻ or CD19⁺CD5⁺. Sorting was performed in the CD19⁺CD5⁻ gate to obtain single B cells from the new emigrant (CD10⁺CD27⁻) and mature (CD10⁺CD27⁻) B cell subsets (lower left). Similarly the CD5⁺CD19⁺ gated cells were single sorted to obtain CD5⁺ mature B cells (CD10⁺CD27⁻) (lower right). Single cell sortings were performed between

8–10 months post engraftment of the hNSG mice. All antibodies used were specific for human cell surface markers.

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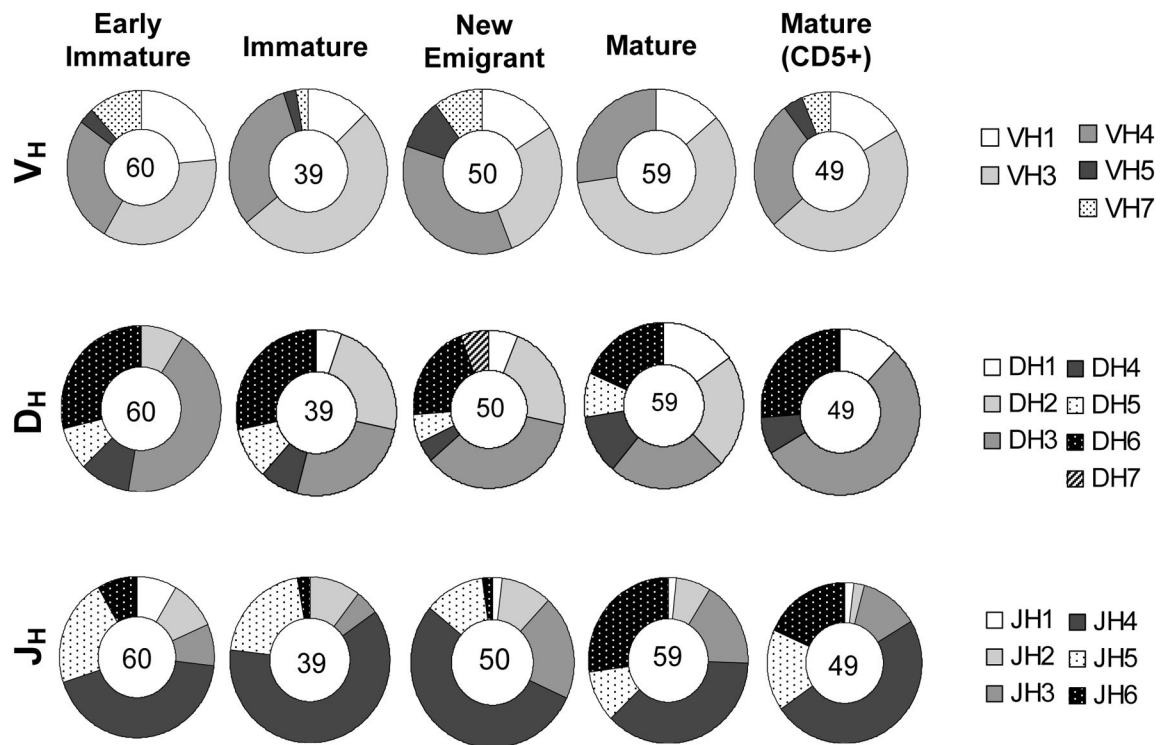
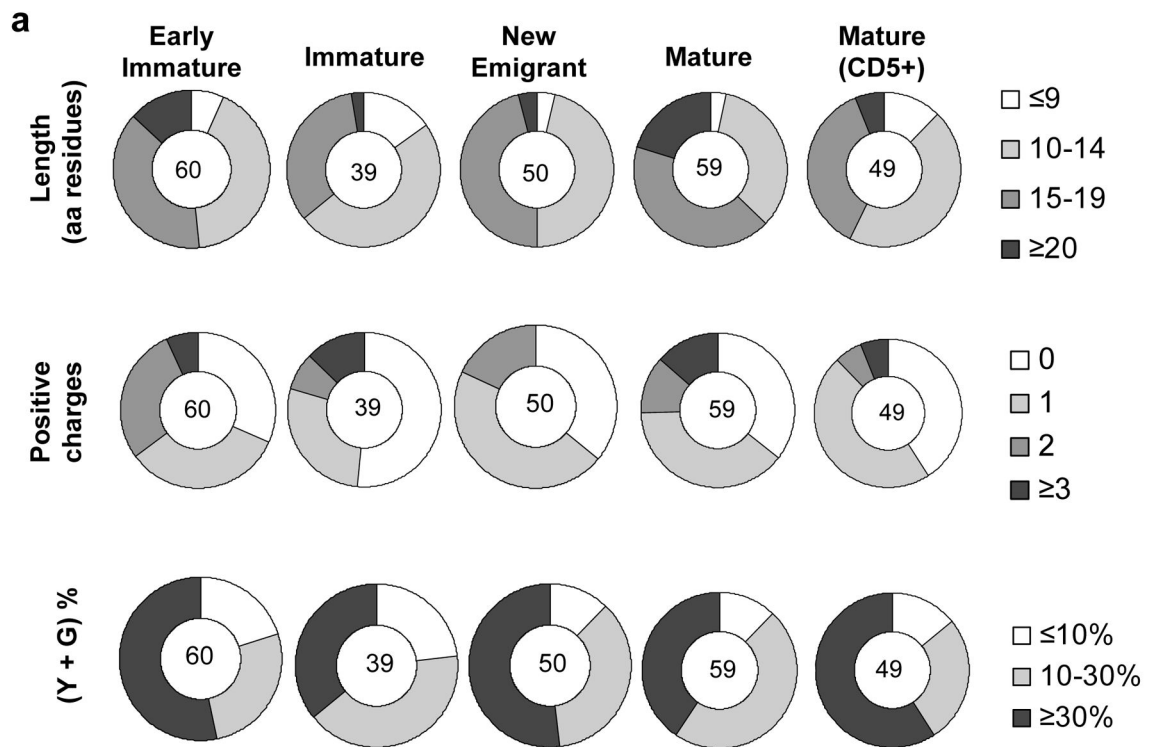


Figure 2.

Analysis of the human immunoglobulin V_H repertoire in hNSG mice. The frequency of utilization of the different V_H gene families (upper panel), D_H (middle) and J_H (lower) gene families across the different B cell subpopulations (indicated above pie charts) is shown. The number in the center of each pie chart indicates the total sample size for the particular B cell subset. Any significant difference in the overall usage of specific gene families in each B cell populations was assessed by Fisher's test. Usage of V_H3 family was significantly higher in all cell subsets ($p < 0.005$) except the new emigrant B cells. Sequence identities and categorizations into different V_H gene families were performed using the IMGT database.



b

N-Nucleotide insertions (bp)		Early Immature	Immature	New Emigrant	Mature	Mature (CD5+)
N1 (V-D)		7.8	6.4	7.6	7.3	5.7
N2 (D-J)		6.7	5.4	4.4	8.0	4.7
JH exonuclease activity (bp)						
Median		3.28	3.33	3.08	3.98	4.29
Range		0-13	0-13	0-9	0-14	0-13

Figure 3.

Composition of the third complementarity region in the human immunoglobulin heavy chain (H-CDR3) regions. (a) The pie charts indicate length (amino acid residues) (upper panel), total positive charge content (middle) and percentage composition of total Y and G residues in each H-CDR3 region analyzed (lower). The different B-cell subpopulations from which the samples were derived are indicated above each pie chart. The number in the center of each pie chart indicates the total sample size for the particular B cell subset. (b) Average N-nucleotide insertions and J_H exonuclease activity across the different B cell subsets are listed. All analyses were performed using the IMGT database.

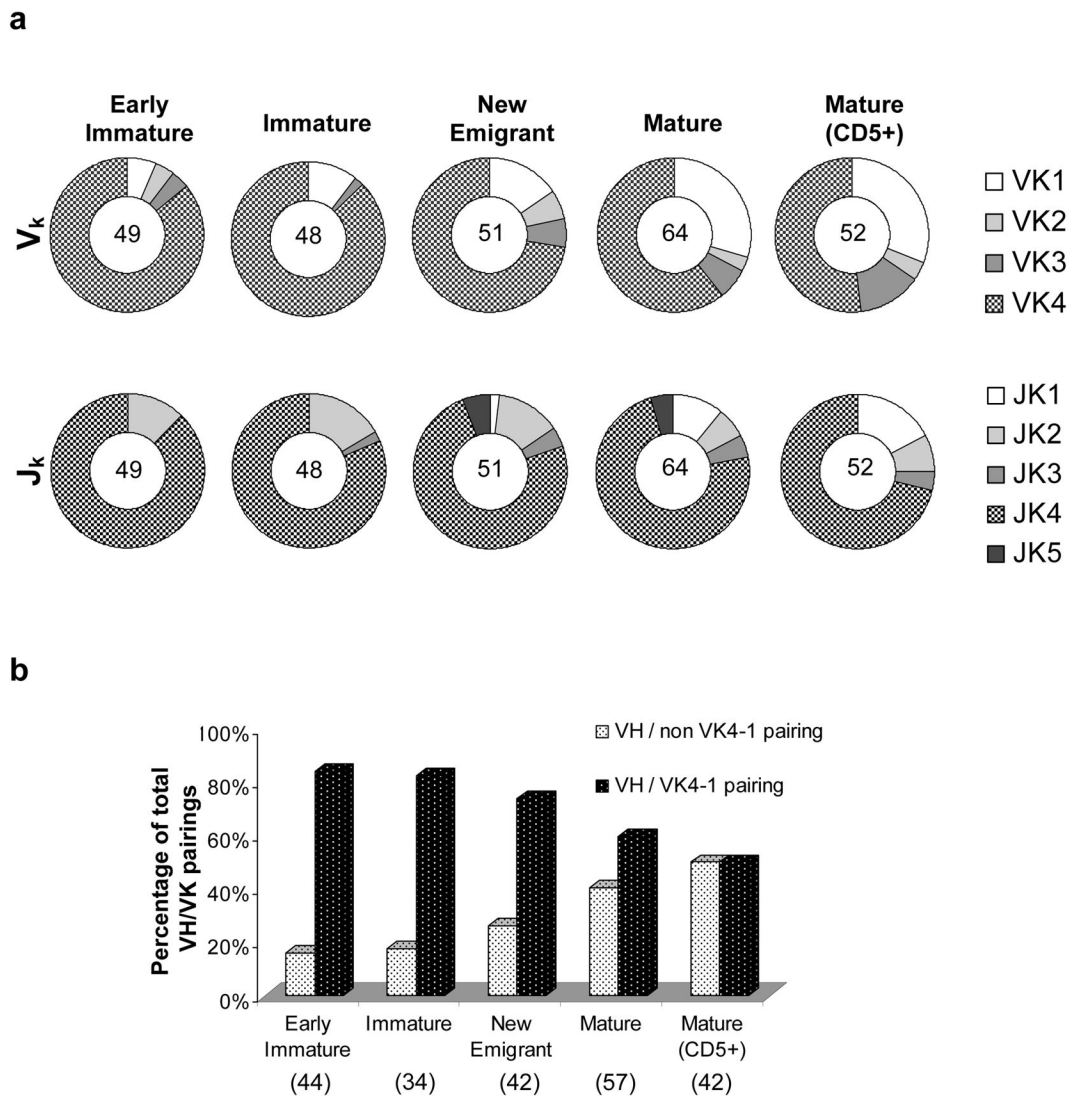


Figure 4. Analysis of the V_κ repertoire and V_H-V_κ pairing in human B cells from hNSG mice. (a) The frequency of utilization of the different V_κ (upper panel) and J_κ (lower) gene families across the different B cell subpopulations as indicated above each pie chart is shown. The number in the center of each pie chart indicates the total sample size for the particular B cell subset. (b) V_H-V_κ pairs were classified into two groups – V_H/non V_κ4-1 pairings and V_H/V_κ4-1 pairings, their percentages over total number of V_H-V_κ pairing in a particular B cell subset were calculated and plotted as shown. A significant decrease in the incidence of V_H/V_κ4-1 pairings (or conversely, an increase in V_H/non V_κ4-1 pairings) was observed in the mature B cell subsets (CD5⁻ and CD5⁺ subsets, combined) in comparison to the bone marrow derived B cell (early immature and immature subsets, combined) (Fisher's Test, p<0.005).

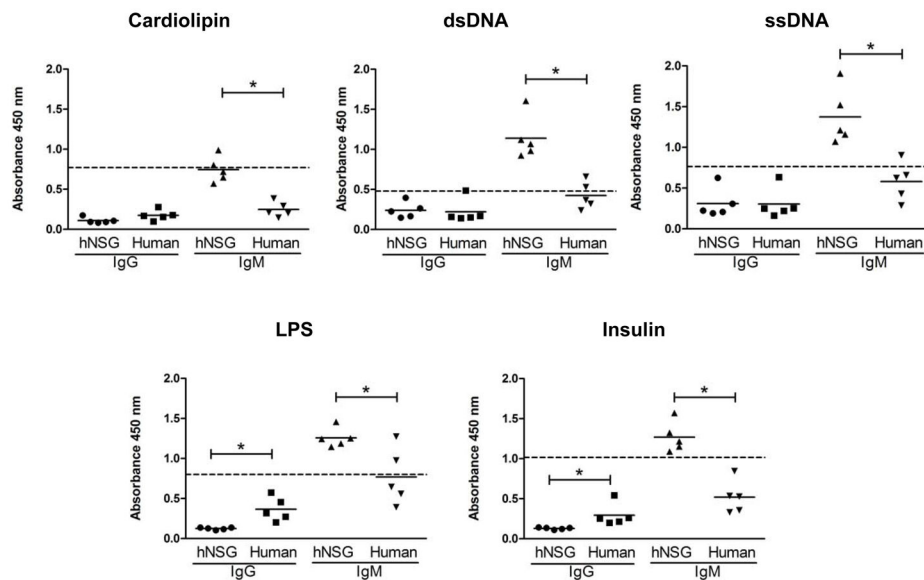


Figure 5.

Polyreactivity in the sera of hNSG mice. Sera from 5 hNSG mice at 10 months post engraftment and 5 normal human subjects were tested by ELISA for binding to different antigens as listed above each graph. Antigens were coated on 96 well plates, and each serum sample was tested in two sets of duplicates, each set scored for human IgM or IgG (x-axis) mediated antigen binding using the appropriate detection antibody, i.e. anti-human IgM and anti-human IgG, respectively both conjugated to HRP. All serum samples were used at 1:100 dilution. Total human IgM and IgG content were 10–50 and 100–200 fold higher respectively, in the human sera compared to the hNSG sera. Sera from unengrafted hNSG mice showed no reactivity (not shown). Each dot represents an individual hNSG mouse or human subject as indicated on the x-axis, and a scatter plot along with the mean for each group is shown. The horizontal dashed line represent the level of reactivity by 4E10 IgG (5 μ g/ml) used as a positive control. Significant differences in reactivity between data sets as indicated (* $p < 0.005$) were calculated using the Mann-Whitney unpaired t-test.

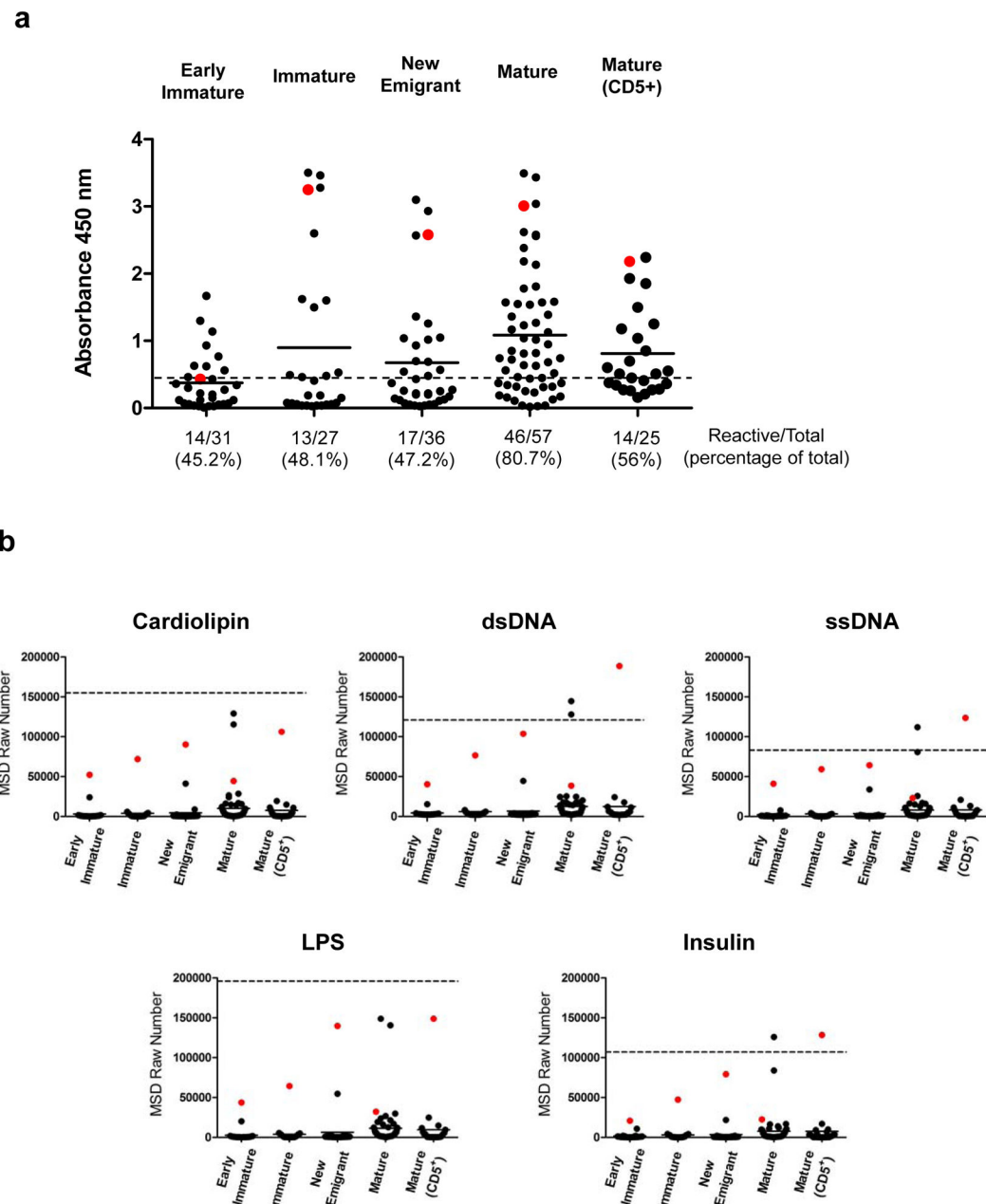
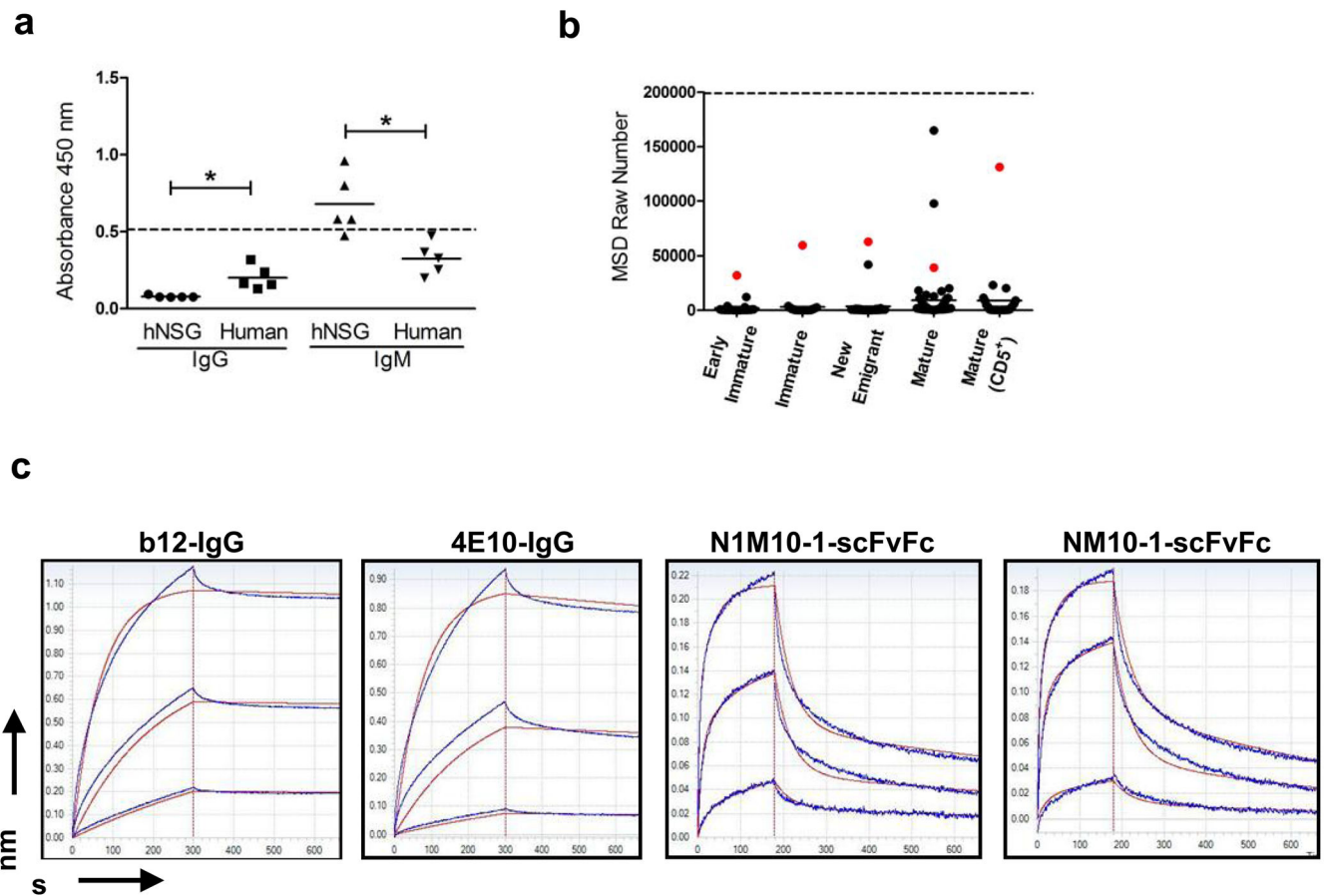


Figure 6.

Auto and polyreactivity analysis of the hNSG derived scFvFc. (a) Hep-2 ELISA was performed with scFvFc assembled and expressed from each B cell subset and that were purified from 293FT cell culture supernatants by Protein A chromatography. ELISA was performed using commercially available QUANTA Lite™ ANA ELISA plates. The assay was performed twice once with 50 µg/ml and repeated with 25 µg/ml of the scFvFc. The horizontal dashed line represents the low positive control (manufacturer provided) cut-off and the numbers below each scatter plot represent the total number and percentage of the scFvFc those were scored as reactive in the Hep-2 assay. Each dot represents a single scFvFc, and the horizontal bar represents mean reactivity for each cell subset. A single

scFvFc determined to be polyreactive (see Figure 6b) in each B cell subset has been color coded (red dot) for correlation between autoreactivity and polyspecificity. Autoreactivity was significantly higher in the mature B cell (median reactivity in Mature vs. Early immature/Immature/New emigrant and Mature CD5⁺ vs. Early immature/Immature/New emigrant, $p < 0.005$). (b) Polyreactivity of the scFvFc's isolated from the different B cell subsets (indicated on the x-axis) was tested using the MSD platform. 384-well plates were coated with the different antigens as listed above each plot. scFvFc's were tested at a concentration of 5 $\mu\text{g/ml}$, and all samples were tested in duplicates. The mean value in MSD raw numbers (y-axis) as obtained in the MSD Sector Imager was plotted. 4E10 scFvFc was used as positive control represented as the horizontal dashed line in each plot and readings obtained with PBS binding to antigen coated wells was considered as background. Each dot represents the binding of individual scFvFc's, and the horizontal bar represents mean reactivity for each cell subset. A single scFvFc in each B cell subset was color coded (red dot) for correlation of polyspecificity across different antigens. Polyspecificity was significantly higher in the mature B cell subsets (median reactivity in Mature vs. Early immature/Immature/New emigrant and Mature CD5⁺ vs. Early immature/Immature/New emigrant, $p < 0.005$). Differences in median reactivity between the scFvFc's of each B cell subset were calculated using the Mann-Whitney unpaired t-test.

**Figure 7.**

Binding of hNSG Abs to recombinant HIV-1gp140. (a) Sera from 5 hNSG mice and 5 normal human subjects were tested by ELISA for binding to recombinant HIV-1gp140 coated at a concentration of 5 $\mu\text{g}/\text{ml}$ per well. Each serum sample was tested in two sets of duplicates, each set scored for human IgM or IgG (x-axis) mediated antigen as described previously (see Figure 5). All serum samples were used at 1:100 dilution. The horizontal dashed line represent the level of reactivity by 4E10 IgG (5 $\mu\text{g}/\text{ml}$) used as a positive control. Significant differences in reactivity between data sets as indicated (* $p < 0.005$) was calculated using the Mann-Whitney unpaired t-test. (b) HIV-1gp140 reactivity of the scFvFc's isolated from the different B cell subsets (indicated on the x-axis) was tested using the MSD platform. HIV envelope protein was coated at a concentration of 5 $\mu\text{g}/\text{ml}$ per well. All samples were tested in duplicates, and the mean value in MSD raw numbers (y-axis) as obtained in the MSD Sector Imager was plotted. 4E10 scFvFc was used as positive control (represented as the horizontal dashed line), and readings obtained with PBS binding to antigen coated wells was considered as background. Each dot represents the binding of an individual scFvFc's, and the horizontal bar represents mean reactivity for each cell subset. The same scFvFc's in each B cell subset has been color coded (red dot) as in Figure 6. Reactivity to HIV-1gp140 was significantly higher in the mature B cell subsets (median reactivity in Mature vs. Early immature/Immature/New emigrant and Mature CD5⁺ vs. Early immature/Immature/New emigrant, $p < 0.005$). Differences in median reactivity between the

scFvFc of each B cell subset were calculated using the Mann-Whitney unpaired t-test. (c) Affinity measurements of the hNSG derived scFvFc to HIV-1gp140 were obtained using Biolayer interferometry. Representative sensorgrams of two HIV BnABs (b12-IgG and 4E10-IgG) and two randomly selected polyreactive scFvFc (NM10-1 and N1M10-1, color coded in Figure 6) binding to different concentrations of recombinant HIVgp140 trimer are shown. Antibodies were loaded on Protein A sensors which were dipped into recombinant HIVgp140 in solution to determine association followed by dissociation in PBS. Traces were obtained for antibody binding to a range of HIVgp140 trimer concentrations. Dissociation was performed for up to 1800 s, but steady state was attained by 600 s which is shown here. Binding responses to <10 nM HIV envelope protein were negative in case of the scFvFc; therefore only the binding traces obtained for 50 nM and 16.6 nM and 5.6 nM of HIVgp140 are shown. Experiments were repeated at least three times with different lots of proteins. Affinity measurement values for each antibody are described in Results.

Numeric representation of the human IG repertoire analyzed from humanized NSG mice

Table 1

Source	Human CD5 ⁻ /CD5 ⁺ B cell subsets ¹	Number of cells sorted	Number of unique IG sequences		Number of scFv-Fc	
			VH	Vκ cognate VH-Vκ pair ²		
Bone marrow	Early Immature	72	60	49	44	31
	Immature	72	39	48	34	27
	New Emigrant	96	50	51	42	36
Peripheral blood	Mature	96	59	64	57	57
	Mature (CD5 ⁺)	72	49	52	42	25
Total		408	257	264	219	176

¹ except the 'Mature (CD5⁺)' B cells that were sorted following gating on a CD19⁺CD5⁺ cell population, all the other human B cell subsets were sorted on a CD19⁺CD5⁻ gate (see Figure 1)

² inclusive of numbers indicated for individual VH and Vκ sequences

Table 2

Distribution of hypermutated V_H genes in hNSG mice

B cell subpopulations	Homology to germline sequence											
	100–99%				99–94%				94%			
	nt/aa mutations ²		nt/aa mutations		nt/aa mutations		nt/aa mutations		nt/aa mutations		nt/aa mutations	
# (%) Sequences ¹	FR	CDR	# (%) Sequences	FR	CDR	# (%) Sequences	FR	CDR	# (%) Sequences	FR	CDR	
Early Immature	36 (60%)	0.46/0.32	0.07/0.05	20 (33%)	1.77/1.14	1.18/0.79	4 (7%)	0.44/0.23	0.53/0.30	0.44/0.23	0.53/0.30	
Immature	26 (67%)	0.68/0.38	0.16/0.08	8 (21%)	0.97/0.57	0.51/0.32	5 (13%)	1.46/0.86	1.14/0.54	1.46/0.86	1.14/0.54	
New Emigrant	33 (66%)	0.30/0.22	0.18/0.08	17 (34%)	1.56/1.14	1.14/0.72	0 (0%)	-	-	-	-	
Mature	50 (85%)	0.60/0.38	0.17/0.10	9 (15%)	0.36/0.24	0.14/0.05	0 (0%)	-	-	-	-	
Mature (CD5 ⁺)	41 (84%)	0.38/0.23	1.18/0.79	8 (16%)	0.68/0.49	0.47/0.30	0 (0%)	-	-	-	-	

¹Total number and corresponding percentage (in parentheses) of total sequences per B cell subset²Total number of nucleotide/amino acid changes in the Framework region (FR) and H-CDR3 (CDR) regions per V_H sequence

Table 3

Potential V_H replacement analysis of the human IG sequences from various B cell subsets¹

Sequence	V _H Donor Family	V _H Sequence	V-D (N1)	D _H	D _H	J _H	R _{Fam}
<i>Early Immature</i>							
NE14-1	4-59	TACTGTGcgagag	cttcct tagata atagtgaacaccc (SLD) NSEHP	tattgtggtggtgactgctatt	2-21	2	1-45
NE31	3-13	TACTGTGcaagag	gacc caaaa atataga (GT) ND	tggtacggtgactacgtcgg	4-17	4	3-43
<i>Immature</i>							
NI47-1	4-61	TACTGTGcgagag	cttcct tagata atagtgaacaccc (SLD) NSEHP	tattgtggtggtgactgctatt	2-21	2	1-45
<i>New Emigrant</i>							
NN37-2	3-11	TACTGTGcgagaa	tctagaag (PLE)	gggggag	3-16	4	3-72
NN70	4-39	TACTGTGcgag	cacgga caaggggggoga (TQ) QGR	atgatatagtggttat	3-22	4	2-26
NN80	4-34	TACTGTGcgagag	cttcct tagata atagtgaacaccc (SLD) NSEHP	tattgtggtggtgactgctatt	2-21	2	3-43
<i>Mature</i>							
NM10-1	4-31	TACTGTGcgagaga	tacagg ccactcaagccgctagtcc (HG) SRAASP	gggtatagcagcagctggtac	6-13	3	2-26
NM48-4	3-23	TACTGTGcgaaaag	ttcaatc cgat tggccagcattcccggcg (GCP) QIBRRY	atggaccgggg	3-10	3	2-26
NM54-1	1-2	TACTGTGcgagag	ggg ggcag t-c (VGS) I	attactatgatagtagtggtta	3-22	4	1-58
NM7-2	3-33	TACTGTGcgagaga	tttcag gcaga ggttg (FRQ) L	gtggtggtaccag	6-19	5	1-58
<i>Mature (CD5⁺)</i>							
N1M26	4-34	TACTGTGcgaga	cggaactcaaa agaaaa (RNS) KKN	attactatggttcggggag	3-10	6	4-28
N1M43-1	3-30	TACTGTGcgaaaag	ct caaaa (GK) I	ggtcgggggagtattata	3-10	3	3-23 4-28
N1M51-1	7-4-1	TACTGTGcgagag	tacacg ta (VTS)		ND	4	2-5

¹ Nucleotide sequence nomenclature used in the table is from IMGT. The listed sequences start from the first nucleotide of the cryptic recombination signal sequence (indicated in upper case), through the V-D junction into the D_H region. The V_H donor family, D_H family, J_H family and Replacement (Recipient) family (R_{Fam}) designations are listed as well. The bold and underlined nucleotides and amino acids in the V-D (N1) region match the 3'-end of the V_H germline of the replacement family and the charged amino acids are shown in red. V_H replacement sequences represent 5% of the total number of Abs analyzed. Italics in the V_H donor column represent gene segments that are downstream to the recipient and thus correspond to a potential non-conventional V_H replacement process.