Title
Dynamics of HSPC repopulation in nonhuman primates revealed by a decade-long clonal-tracking study

Permalink
https://escholarship.org/uc/item/67z6b70v

Journal
Cell Stem Cell, 14(4)

ISSN
1934-5909

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Publication Date
2014-04-03

DOI
10.1016/j.stem.2013.12.012

Peer reviewed
SUMMARY

In mice, clonal tracking of hematopoietic stem cells (HSCs) has revealed variations in repopulation characteristics. However, it is unclear whether similar properties apply in primates. Here, we examined this issue through tracking of thousands of hematopoietic stem and progenitor cells (HSPCs) in rhesus macaques for up to 12 years. Approximately half of the clones analyzed contributed to long-term repopulation (over 3–10 years), arising in sequential groups and likely representing self-renewing HSCs. The remainder contributed primarily for the first year. The long-lived clones could be further subdivided into functional groups contributing primarily to myeloid, lymphoid, or both myeloid and lymphoid lineages. Over time, the 4%–10% of clones with robust dual lineage contribution predominated in repopulation. HSPCs expressing a CCR5 shRNA transgene behaved similarly to controls. Our study therefore documents HSPC behavior in a clinically relevant model over a long time frame and provides a substantial system-level data set that is a reference point for future work.

INTRODUCTION

Blood is a highly regenerative, multifunctional tissue and a relatively small population of self-renewing hematopoietic stem cells (HSCs) can regenerate all blood cell types for the lifetime of the host (Kondo et al., 2003). Transplant of HSCs exploits these properties for treatment of various human diseases and as a means to gene-modify hematopoietic cells for genetic therapies (Kohn and Candotti, 2009; Weissman, 2000). Repopulation following transplant occurs through the combined engraftment, growth, and differentiation of a large number of hematopoietic stem and progenitor cells (HSPCs) including self-renewing HSCs as well as more differentiated and lineage committed progenitor cells; yet, the individual and cumulative behavior of the HSPCs at the system level is not well understood. Murine models have provided valuable insights into the regenerative potentials of HSCs, having the benefit of established assays for purification of HSCs from among the diverse HSPCs in the bone marrow (Purton and Scadden, 2007). Historically, HSCs have been presumed to be biologically homogeneous and possess an unlimited self-renewal potential. More recently, however, single-cell studies have reported significant cell-to-cell variations in repopulation kinetics, life span, and sensitivity to extracellular stimuli (Benveniste et al., 2010; Copley et al., 2012; Jordan and Lemischka, 1990; McKenzie et al., 2006; Morita et al., 2010; Müller-Sieburg et al., 2002; Osawa et al., 1996; Pina et al., 2012; Smith et al., 1991). Interestingly, even the multipotent behavior of HSCs has recently been shown to vary, with some HSCs being biased toward either myeloid or lymphoid lineages (Copley et al., 2012; Lu et al., 2011; Müller-Sieburg et al., 2012). Unlike the extensive polyclonal repopulation seen in primates, murine studies demonstrated the behaviors of a single or a few repopulating clones per mouse (Copley et al., 2012; Jordan and Lemischka, 1990; Lemischka et al., 1986; Müller-Sieburg et al., 2012; Smith et al., 1991). Some recent studies have investigated oligoclonal repopulation (dozens of clones) in mice (Cornils et al., 2012; Gerrits et al., 2010; Naik et al., 2013; Verovskaya et al., 2013); nevertheless, insights gained by translating these studies into an understanding of human repopulation are limited because of the far greater complexity of polyclonal reconstitution as well as the greater demands placed on stem cells within the larger and longer-lived...
human system. Thus, despite the significant advances in our understanding of HSC biology from murine models, the functional properties of primate HSPCs following transplant remain poorly characterized.

All current transplant protocols for humans and nonhuman primate models utilize partially purified CD34+ cell populations, of which only a small proportion represent true HSCs. Because purification of primate HSCs is not as yet technically feasible, the primary method utilized to track repopulation in transplant for human diseases has been through the use of integrating retroviral vectors that “mark” individual HSPCs through differential semirandom genomic integration sites. In early clinical studies, the use of gamma-retroviral vectors skewed repopulation due to insertional mutagenesis that often resulted in malignant transformation (Hacein-Bey-Abina et al., 2010; Nienhuis et al., 2006; Stein et al., 2010). More recent clinical trials using lentiviral vectors have tracked repopulation without obvious genotoxic effects, revealing long-term repopulation by thousands of gene-engineered cells (Aiuti et al., 2013; Biffi et al., 2013; Cartier et al., 2009; Cavazzana-Calvo et al., 2010). These studies showed that the early phase of the hematopoietic reconstitution in patients (1.5–2 years) was achieved by a large number of low-frequency clones, many of which transiently contributed to the blood. However, the long-term behavior patterns of the vast number of HSPCs working together to reconstitute the complex polyclonal hematopoietic system remain unclear. It is also important to note that human clinical studies may not be the best setting to study normal HSPC repopulation because for ethical reasons all such studies are in the context of a diseased hematopoietic system and the introduction of disease-corrective transgenes may provide favorable repopulation by the gene-corrected HSPCs.

The nonhuman primate autologous HSPC transplant model is arguably the animal model most closely resembling humans; its treatment conditions—including the use of CD34+ cells and mobilization and conditioning regimens—all parallel those of human (Heim and Dunbar, 2000; Shi et al., 2002). We previously demonstrated long-term stable hematopoietic reconstitution in rhesus macaques after transplantation of autologous CD34+ HSPCs transduced with lentiviral vectors (An et al., 2001, 2007; Sander et al., 2006). These animals maintained stable long-term marking for as long as 12 years without any genotoxic effects resulting from transplant. The behaviors of HSPC clones at the system-level were characterized using a deep-sequencing-based clonal tracking assay (Kim et al., 2010), which can quantify a large number of individual clones in parallel by vector integration site (VIS) sequence enumeration. We observed that thousands of HSPCs engrafted, and they maintained long-term hematopoiesis at a constant level through sequential expansions of different groups of clones. High-throughput, long-term clonal tracking in our test animals revealed high variability in HSCs for their lineage outputs, including myeloid-biased, lymphoid-biased, and balanced subtypes.

**RESULTS**

**Long-Term Stable Marking after CD34+ HSPC Transplantation**

The subjects of this study were four rhesus macaques transplanted with autologous CD34+ HSPCs from mobilized peripheral blood (mPB) after a myeloablative dosage of total body irradiation for periods of 4 to 12 years (An et al., 2001, 2007; Sander et al., 2006). Long-term repopulation of HSPCs in two animals (95E132 and 2RC003) was analyzed after transplant with CD34+ cells marked with lentiviral vectors (human immunodeficiency virus [HIV]-based, self-inactivating [SIN-HIV] vectors expressing EGFP) (An et al., 2001; Sander et al., 2006). As a model of genetic therapy for HIV-1 disease, two other animals (RQ3570 and RQ5427) were transplanted with CD34+ HSPCs engineered with simian immunodeficiency virus (SIV)-based vectors expressing both EGFP and shRNA against the HIV-1 coreceptor, C-C chemokine receptor type 5 (CCR5) (An et al., 2007). The potency and safety of the shRNA (sh1005), selected from hundreds of other candidates, was previously demonstrated both in vivo and in vitro (Shimizu et al., 2009, 2010). Animal 2RC003 was cotransplanted with CD34+ HSPCs transduced with SIN-HIV vectors expressing the mutant herpes simplex virus-1 thymidine kinase (HSV1-sr39tk) gene for position emission tomography (Sander et al., 2006). These HSV1-sr39tk-expressing cells repopulated at 0.01%–0.2% of total blood for the first year and were removed by ganciclovir IV treatment at 23 months posttransplant. In all animals, the lymphocyte recovery was slower than that of the granulocytes (grans) or monocytes. After an initial phase of fluctuation for about 6–12 months, all tested blood lineages showed stable marking over the long-term (Figure 1). To date, all animals remain healthy, with no sign of malignant transformation.

Total peripheral blood cells (PBCs) for 95E132 and fractionated grans and peripheral blood mononuclear cells (PBMCs) for animals 2RC003, RQ5427, and RQ3570 were serially collected over the entire analysis period so we could study the longitudinal behavior patterns of repopulating HSPC clones in these animals (Table 1). Different blood cell types (CD4+, CD8+, CD14+, CD18+, and CD20+ cells and mPB CD34+ HSPCs) were also isolated near the experiment end point to study lineage-output potentials of long-term engrafted clones. All sorted cell samples showed at least 98.7% purity except for CD4+ cells from RQ5427 and 2RC003 (94.1% purity) (Table S1 available online).

**Clonal Tracking Assay**

The repopulation of individual HSPCs was tracked by the distinguishing VIS of each vector-marked clone. The frequency of each clone was quantitatively analyzed by large-scale VIS sequencing and bioinformatics analysis of VIS sequences. We optimized the assay conditions to enable quantitative VIS sequencing, including the amount of input DNA, the intensity of sequencing, data analysis standards to restrict detection biases, and the conditions for concurrent analysis of the 5′ and the 3′ vector-host junctions (Kim et al., 2010). Briefly, the genomic loci and the relative frequencies of individual vector integrants were determined by a Taqal-mediated, bidirectional VIS assay (see Supplemental Experimental Procedures for details). During the assay, the Taqal restriction enzyme generated different lengths of PCR amplicons, each harboring a distinct VIS sequence (Figure S1 available online). Due to inefficient sequencing of PCR amplicons of >500 bp with our experimental conditions, only the vector integrants that generated ≤500 bp VIS PCR amplicons at either the left or right vector-host
Long-Term Clonal Tracking of Primate HSPCs

Cell Stem Cell

Figure 1. Longitudinal EGFP Marking in Four Rhesus Macaques

The levels of EGFP expression in granulocytes (grans; green), monocytes (Monos; brown), lymphocytes (Lymphs; red), red blood cells (RBC; purple), and platelets (PLT; blue) were assayed by flow cytometry, based on the forward scatter versus side scatter plot. PLTs were also gated on CD41+ cells. Data represent longitudinal follow-up for 38–145 months posttransplant, depending on the test animal. Data gathered during the ganciclovir IV treatment (see arrow at 23 months posttransplant) and within the 2 week period after leukapheresis have been excluded. See Table S1 for more details.

Long-Term Reconstitution by Thousands of HSPC Clones

Variable levels of polyclonal repopulation were observed in each animal. VIS analysis revealed a range of 30 to 1,338 unique vector integrants, each representing progeny derived from a single vector-marked HSPC per test sample, totaling 846–5,758 per animal (Table 1 and Table S2). The total number of unique vector integrants recovered from each animal was proportional to the number of vector-marked HSPCs that were infused at the beginning of the experiment (Figure 2A). We previously estimated that approximately 40.3% of vectors were QVI, based on a simulation with random vector integration sites (Kim et al., 2010). In our data sets, 61%–83% of the total recovered vector integrants were QVIs (Table 1). The discrepancy between the computational prediction and the assay results may be due to an assay limitation to detect some vectors remotely located from the TaqI restriction sites in the genome. Further biostatistical analysis demonstrating optimization of VIS analysis is in the Supplemental Information (see Figure S2B). Individual QVI clones in each animal showed highly variable detection frequencies. While thousands of different clones presented, only a small number of clones dominated the blood repopulation over time. For example, the 5% most frequent QVI clones contributed on average 49%–72% of the total QVI repopulation (up to 60%–88% over time) in each animal with the most prevalent clone showing an average relative detection frequency of 0.0322–0.1028 (based on an average across all time points) (Figure 2B). In contrast, the majority of QVI clones (43%–71%) were infrequently detected with an average frequency of <0.0002, contributing to <7% of total QVI repopulation over time in all animals except RQ3570. In animal RQ3570, the QVI clones with <0.0002 frequency contributed to <23% of total repopulation. In this animal, the mean and the standard deviation of the relative frequencies among individual clones were lower than those of other animals (see Figure 2B), due most likely to a relatively short follow-up time (3 years) and also to the greater presence of clones (four to seven times as many) in this animal.

Long-Term, Stable Hematopoietic Reconstitution Was Maintained through Sequential Expansion of Different HSPC Clones

In test animals, the levels of total vector-marked cells in the blood were maintained at relatively constant levels for years following the first 6–12 months posttransplant. To determine how the large number of HSPC clones maintained hematopoietic homeostasis, we analyzed the patterns of clonal repopulation over time by clustering QVIs based on their frequency profiles in serially collected PBCs (in animal 95E132) or grans/PBMCs (in animals 2RC003, RQ5427, and RQ3570) (Figure 3A). Only QVIs with an average relative frequency of ≥0.0002 (calculated across all time points) were included for cluster analysis because they provided reliable signals of clonal behavior over time. These QVIs were often detected at more than three time points, thereby enabling a traceable trajectory of their longitudinal behaviors (see Figure S2D for details). Low-frequency clones (<0.0002 relative frequency) were excluded from the analysis because their signals were expected to be noisier and inconsistently available across time points due possibly to uneven PCR amplification of low-frequency clones during the assay process. QVI clones with an average frequency of ≥0.0002 were clustered into 5–8 groups, depending on cell types and animals, using the WGCNA package (Langfelder and Horvath, 2008) (Figure 3A and Figure S3A). Except for the smallest cluster among the data sets (cluster-1 with nine clones in the 2RC003 PBMC data set), all clusters were clearly separable, with a significance level (Zsummary.qual) of >10 (Langfelder and Horvath, 2008). The replication kinetics of clusters-1 to -8 demonstrate sequential expansions of different clones, peaking at different time points. Cluster-1 corresponds to the first wave of clones repopulating at the earliest time point, followed by cluster-2 and continuing through the final cluster (cluster-5 or -6 for RQ3570 and cluster-7 or -8 for all other animals) (Figure 3A). Fluctuations in clonal frequencies were higher early after transplant (2–9 months)—when the cluster-1 and -2 clones were the primary source of blood repopulation—than at later time points (Figure S3B). Most of the cluster-1 and -2 clones lasted for only months whereas the clones that emerged later expanded and decayed over years. These results indicate that repopulation occurs in
at least two kinetically distinct phases: (1) an initial phase lasting about 1 year where clones rapidly appear and disappear and (2) a long-term phase where more stable clones gradually emerged and predominated the HSPC repopulation. Furthermore, even the most stable clones appear to have a finite life span. Sequential declines of hundreds of clones are compensated for by expansion of others, maintaining a relatively stable level of marked cells over the ensuing years in each animal (Figure S3C).

### Segregation of Long-Term and Short-Term Repopulating Clones

Clones persisting for many years likely represent repopulation by the progeny of HSCs. To specifically study the long-term repopulating clones, we segregated QVI clones that were detected near the experimental end point (long-term QVIs [LT-QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter). We determined the lineage output of LT-QVI clones by fractionating cells near the experimental end point (long-term QVIs [LT-QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter) in mature blood lineages (at 36, 64, 70, or 116 months from animals RQ3570, RQ5427, 2RC003, and QVIs], hereafter).

### Lineage Disparate Subtypes of Long-Term Repopulating Clones

We determined the lineage output of LT-QVI clones by fractionating cells near the experimental end point into subpopulations of myeloid (CD14+ and CD18+) and lymphoid (CD4+, CD8+, and CD20+) cells. Individual LT-QVIs showed highly variable detection frequencies in different cell types. To obtain reliable patterns of clonal lineage potentials, LT-QVIs with an average detection frequency of <0.0002 (average of myeloid cells [CD14+ and CD18+]). T cells [CD4+ and CD8+], and B cells [CD20+] were excluded from the analysis. The results revealed three clearly distinguishable subtypes among the LT-QVIs that differed in the proportion of lineage output to myeloid or lymphoid cells (Figures 4A–4D and Figures S5A and S5B). Myeloid-biased (My-bi), lymphoid-biased (Ly-bi), and balanced (Bal) LT-QVIs were defined based on the myeloid (GM) and...
lymphoid (BT) ratios of $\geq 3:1$, $\leq 1:3$, and between 3:1 and 1:3, respectively. These showed a high quality of separation for all data sets ($|Z_{summary.qual}| > 10$ using the WGCNA R package; Langfelder and Horvath, 2008), except within animal 2RC003, which had the two smallest clusters (with only 30 and 38 clones) that did not achieve significance. In animals 95E132, RQ5427, and RQ3570, the proportions of the My-bi, Bal, and Ly-bi clones among the analyzed LT-QVIs were $25\%$/$31\%$, $23\%$/$33\%$, and $40\%$/$46\%$, respectively. In animal 2RC003, the My-bi proportion was much greater ($59\%$) than the others ($18\%$ and $23\%$ for Bal and Ly-bi, respectively), possibly because it was affected by ganciclovir IV treatment at 23 months posttransplant. These results provide a demonstration of highly heterogeneous lineage output behaviors of long-term polyclonal repopulation in primates.

**Dynamics of My-bi, Ly-bi, and Bal Clone Repopulation over Time**

We observed that LT-QVIs became the dominant source of hematopoietic reconstitution after rapid but transient reconstitution by a large number of ST-QVIs that declined primarily within the first year (see Figure 3B). Here, we tracked the clones constituting three LT-QVI subtypes over time to understand how these subtype clones contributed to long-term hematopoietic reconstitution. Interestingly, a small number of Bal type LT-QVI clones, constituting only 4%–10% of total recovered QVIs, expanded extensively during the first 10–20 months and, after which time, became the predominant source of long-term hematopoietic reconstitution, ultimately contributing up to 71% (95E132), 42%–52% (2RC003), 61%–67% (RQ5427), and 25%–32% (RQ3570) of total QVI repopulation (Figure 4E). The My-bi LT-QVIs, constituting 6%–16% of total QVIs, also increased gradually but maintained relatively lower levels than the Bal type. In contrast to the My-bi and Bal clones, which increased consistently over time, the Ly-bi LT-QVIs (6.2%–12.8% of total QVIs) expanded for the first 10–20 months and then gradually declined. The greatest proportion of repopulation is ultimately contributed by the Bal type clones, which contribute relatively equally to myeloid and lymphoid repopulation.

We further examined the utilization of these clones over time in blood cells fractionated for granulocytes and peripheral blood mononuclear cells (PBMCs). Serial analysis of fractionated granulocytes and PBMCs in animals 2RC003, RQ5427, and RQ3570 showed that the lineage-biased or -balanced patterns of these clones started to be apparent at about 10 months posttransplant and sustained until the end point (Figure 4E and see Figure S6A for 10 most frequent clones).

**Correlation of Clonal Behavior with Cluster Analysis**

Through clonal tracking analysis we were able to define four functionally distinct repopulating subpopulations arising from the transplanted CD34+ HSPC pool, including short-term repopulating clones and three subtypes (My-bi, Bal, and Ly-bi) of long-term repopulating clones. We also determined their relative contributions to hematopoietic reconstitution over time (see Figure 5A for a summary). In order to investigate the more detailed dynamics of the clones constituting these subgroups, they were compared with the kinetics clusters defined in Figure 3A. As expected, the ST-QVIs were highly enriched in cluster-1,
representing clones that appeared first and disappear shortly after, except serial PBMCs in 2RC003 (Figure 5B). Conversely, LT-QVIs were found primarily in clusters appearing later. As expected from the increasing patterns over time, My-bi and Bal type QVIs were enriched in the clusters that appeared the latest (cluster-7 or -8 for 95E132, 2RC003, and RQ5427, and cluster-5 or -6 for RQ3570), whereas Ly-bi clones did not show such a pattern (Figure 5B). The results indicate that within each given lineage subtype, individual clones rise, peak, and fall at different times.

**The Relative Proportions of My-bi, Bal, and Ly-bi Clones in the CD34+ HSPC Pool**

Because some mature cells persist for years, the relative proportions of My-bi, Bal, and Ly-bi clones assessed in mature cells may not reflect those in stem cells or progenitors. The relative frequencies of these long-term QVI clones in mature lineages were compared near the experimental endpoints for each animal with those in the underlying HSPC pool (37, 63, 71, or 117 months). mPB CD34+ HSPCs were collected within 1 month of the blood used for lineage fractionation (Table 1). For most animals the

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**Figure 3. Repopulation Kinetics of HSPC Clones**

(A) Sequential expansions of HSPC clones over time. Major clonal kinetics patterns in serial PBCs (in animal 95E132) and PBMCs and grans (in animals 2RC003, RQ5427, and RQ3570) were derived using average linkage hierarchical clustering based on clonal frequency profiles over time. The relative frequencies of QVI clones at different time points (shown in months at the top of each chart) appear in a white-black-yellow color scheme, with white denoting zero frequency and yellow the highest frequency (see the numbers at the bottom of each chart). QVIs were grouped into 5–8 clusters (denoted by different colored arrows), depending on animal and cell type, using the WGCNA package (Langfelder and Horvath, 2008). All clusters were clearly separable, with a significance level ($|Z_{summary.qual}| > 10$ (Langfelder and Horvath, 2008), except for cluster-1 in the 2RC003 PBMC data set (only nine QVI clones). Results showed that different groups of QVI clones, beginning with cluster-1, expanded sequentially over time. See Figure S3 for more details.

(B) Repopulation of long-term and short-term QVI Clones. Long-term QVI clones (LT-QVIs) and short-term QVI clones (ST-QVIs) were segregated based on the presence of these clones in blood lineages isolated at 116–117 months (95E132), 70–71 months (2RC003), 63–64 months (RQ5427), or 36–37 months (RQ3570) posttransplant. The relative contributions (y axis) of LT-QVIs (orange) and ST-QVIs (black) in serial PBCs (95E132) or grans/PBMCs (2RC003, RQ5427, and RQ3570) are shown over time. See Figure S4 for more details.
majority of the Bal and My-bi clones were present in the HSPC pool (57%–96% of the Bal type and 38%–80% of My-bi clones were shared with the HSPC pool) (Figure 5C and also see Figures 4A–4D). In contrast, only 17%–50% of Ly-bi clones were common to the HSPC pool. When My-bi, Bal, and Ly-bi LT-QVI clones that were present in the CD34+ HSPCs were tracked over time, we found that the unique lineage-biased or -balanced patterns of these clones became apparent at around 1 year posttransplant and were maintained over time (Figure S6B). The majority of Ly-bi clones occurred in both grans and PBMCs at earlier time points but persisted primarily in PBMCs until the end point. We also observed that the clonal composition between the HSPCs and the lymphoid cells (CD4+, CD8+, and CD20+ cells) was generally less correlated than the clonal compositions between the HSPCs and CD18+ grans (Figure 5D), as is expected because the relatively short life span of grans would reflect the immediate output of precursor cell differentiation. These results indicate that the long-term HSPC population is relatively enriched for the Bal and My-bi clones. A smaller proportion of Ly-bi LT-QVIs were maintained above the detection level in the long-term engrafted HSPC pool. This result may reflect the longer life span of lymphoid cells and consequently a decreased requirement for Ly-bi clones with age.

**VIS Analysis Indicates Benign Integration Bias without Selection for Any Particular Cell Type**

As seen in previous gene-therapy clinical studies using murine leukemia virus (MLV) vectors, insertional mutagenesis may cause aberrant clonal proliferation or skewed differentiation. While low genotoxic effects of lentivirus vectors have been demonstrated in recent studies by VIS analysis for up to 2 years in humans (Aiuti et al., 2013; Biffi et al., 2013; Cartier et al., 2009; Cavazzana-Calvo et al., 2010), careful monitoring of the long-term effects by lentivirus vectors is required because gene-modified HSCs will be maintained for a lifetime. Genomic VIS patterns of repopulating cells in our test animals largely reflect those of freshly transduced CD34+ HSPCs (acute infection) (Figure S7A). We also found several local VIS hot spots common in all test animals but undetected in acute infection of mPB CD34+ cells (Figure 6). These hot spots were characterized using a Bayesian change-point (BCP) model that defines VIS hot spots relatively independently of data set size (Presson et al., 2011). Lentivirus vectors showed distinctive VIS hot spot patterns that differed from oncogenic MLV vectors. Clinical gene-therapy studies with MLV vectors have revealed hot spots consisting of densely clustered VISs in a small genomic region, typically near the transcription start site of a gene, that often differed across individuals (Deichmann et al., 2007; Presson et al., 2011; Wang et al., 2010). In contrast, the lentivirus vector hot spots in our test animals consisted of VISs spread across multicentric, megabase-scale regions (Figure S7B). These hot spots were conserved in all time points and cell types without any notable changes in size or intensity (Figure S7C). Similar hot spot patterns were also observed at homologous regions in humans (Presson et al., 2011) and in mice (Biffi et al., 2011). Through analysis of PBMC, grans, and PBC data sets, we found that VIS hot spots were not correlated with highly expanded clones (data not shown). When the genes hosting VISs in My-bi, Bal, Ly-bi, or ST-QVI data sets were compared with each other, no notable functional gene group was associated with any specific cell type (Figures S7D and S7E). Thus, these results demonstrate an apparently benign integration bias not associated with any particular HSC subtype.

**DISCUSSION**

To date, our understanding of the primate hematopoietic system lags far behind the level desired for current and future use of HSPC-based therapies (Cavazzana-Calvo et al., 2011; Copley et al., 2012; Doulatov et al., 2012). Despite the significant advances in HSPC-based therapies, the phenotypic properties of primate HSCs during hematopoietic reconstitution after transplant remain unclear. We provide here a detailed system-level description of the decade-long properties of a large number of heterogeneous clones following nonhuman primate CD34+ cell transplant and address how long-term and stable hematopoietic reconstitution over 4 to 12 years was achieved through the combined contributions of thousands of clones. Remarkably, despite waves of HSPC utilization lasting for periods of months or years, total marking remains relatively constant for over 10 years. And, despite lineage-disparate behaviors of HSCPs, normal repopulation of blood cell lineages is maintained, initially through the contribution of short-lived HSC clones, but eventually the majority of repopulation is determined by a few stable clones having a relatively balanced multilineage output. The results suggest tremendous diversity as well as intricate control of HSPC repopulation.

Classically, HSCs were thought to be largely biologically homogeneous in regards to self-renewal and differentiation potential. Recent single-cell transplant studies in mice introduced new complexity into the regulation of the hematopoietic system, showing significant heterogeneity in longevity, self-renewal, and differentiation potentials among the most primitive HSCs (Copley et al., 2012; Muller-Sieburg et al., 2012). Unlike the murine transplant studies using purified HSCs, primate transplant occurs through the combined and competitive repopulation by a vastly greater number of HSPC clones over a longer period of time. The majority of repopulating clones in our test animals showed a kinetic pattern indicating finite life spans of engrafted HSCPs. These clones made their peak contribution at different time points and could be grouped into several clusters based on their repopulation kinetics, showing sequential expansions of different groups of clones over months for the earliest clones and over years for those clones appearing later after repopulation (Figure 3A). Of the clones detected, approximately equal numbers are long-term (LT-QVI) and short-term (ST-QVI) clones. Although the assay end point given to each animal to distinguish long-term and short-term clones ranged from 3 to 10 years, this variation in end points did not significantly affect our findings regarding the relative frequencies of these two populations insofar as the clonal kinetics in all animals became more stable after 1–2 years. We estimated approximately 0.01% of transplanted CD34+ cells (1 in 1–2 million mobilized PBMCs) were capable of long-term repopulation above the detection limit. Our estimate is comparable to an estimate of the human HSC frequency (1 in 1–6 million cells in cord blood, adult bone marrow, or mPB) by limiting dilution analysis using immunodeficient mouse xenotransplant models (Doulatov et al., 2012; Wang et al., 1997). An exquisite balance exists between the expansions of newly arising clones and the declines...
of preceding ones, ensuring stable hematopoietic homeostasis of transplanted clones in the blood over years. The clones detected early after transplant (within the first 7 to 9 months) were of large number, were mostly low-frequency, and persisted transiently, whereas the clones detected later were kinetically more stable, expanding slowly and often intensively over years before subsiding and being replaced by new groups of clones. This sequential pattern of polyclonal repopulation suggests that stable long-term hematopoietic reconstitution is achieved by expansions of HSPCs with different longevities. Differences in the life span of HSCs have also been observed in mice where transplantation of single HSCs exhibited wide differences in longevity (Benveniste et al., 2010; Sieburg et al., 2011). We hypothesize that the large number of early clones represent more differentiated progenitors that repopulate into the blood relatively quickly, making only brief contributions but ensuring adequate hematopoiesis and consequently survival of the host during this critical early period after myeloablation. More detailed insights into the clonal behaviors during this period are provided in a related nonhuman primate study by Wu et al. (2014), published in this issue of Cell Stem Cell, showing transient repopulation of unilineage clones occurring primarily at 1–2 months followed by more stable repopulation of multilineage clones. Our long-term analysis data show that those HSCs at a more primitive stage take more time to make their peak contribution to the blood and persist for much longer but still finite times. Many clones remain undetected until many years after transplant before arising to replace decaying clones. The replacement of subsiding clones with new groups of HSCs in order to maintain a constant level of marked hematopoietic output further suggests that there exist host environmental cues that instruct new clones to emerge as old clones decay.

Long-term clones exhibit highly variable lineage output. Many clones were relatively balanced in terms of their contributions to myeloid and lymphoid lineages (balanced subtype), but others showed a biased differentiation pattern favoring either myeloid output (My-bi) or lymphoid output (Ly-bi). Among the Ly-bi clones, some were highly skewed toward either T or B cells. Over the long term, a small number of the Bal subtype clones (4%–10%) expanded extensively and predominantly contributed to the total blood repopulation (25%–71%). While the detailed lineage cell-sorting fractionations were performed at a single late time point, we demonstrated in three animals that within blood fractionated at multiple time points for grans, representing myeloid cells, and PBMCs, representing 80%–90% lymphoid cells, the numbers of QVI clones appear in parentheses. In subpanels (ii) and (iii), the ternary diagrams show the lineage output potentials of each clone (indicated by the positions of the circles relative to the points of the triangle). The relative frequencies of clones (indicated by the size of the circles) in mature lineages (ii) are compared to those of identical clones in mPB CD34+ cells (iii). The dominance of the Bal subtype was also suggested in a recent murine study (Lu et al., 2011); however, this study was performed at only a single time point, thus lacking longitudinal HSC behavior data. Thus, our results provide evidence for the coexistence of phenotypically lineage disparate clones contributing to hematopoiesis in primates. The dominance of Bal type HSC clones is to be teleologically expected in order to maintain uniform blood lineage production over time.

The Ly-bi and My-bi clones are present in each of the animals over the duration of our study although they contribute less than Bal clones to total repopulation. The analysis of these clones in granulocyte and PBMC fractions suggests conservation of these lineage properties over time, although more detailed longitudinal cell fractionation will be required for confirmation. In murine studies, the clones restricted for myeloid output (Ly-bi) correlate with short longevity; self-renewal activity of this type of cell was exhausted within 6–7 months (Dykstra et al., 2007). In our studies, Ly-bi clones were maintained over the long term; however, each animal showed a notable decrease in the number of Ly-bi clones within the CD34+ progenitor cell population at the assay end point (3–10 years posttransplant), suggesting potential quiescence or exhaustion of some previously activated Ly-bi progenitors. By contrast, balanced and My-bi long-term clones were relatively well conserved in the CD34+ cells (see Figures 4A–4D and Figure 5C). Since mature lymphoid cells have a far greater life span than myeloid cells, the reduced frequency of Ly-bi clones in CD34+ cells may reflect a reduced requirement for Ly-bi HSCs over time. The role of the lineage-biased clones is unknown, but it is interesting to speculate that they may be instructed by environmental factors to contribute to certain blood lineages in times of particular need or stress.

Previous clinical trials with gamma-retroviral vectors have been associated with cancers (Hacein-Bey-Abina et al., 2010; Nienhuis et al., 2006; Stein et al., 2010); hence, the safety and efficacy of therapeutic vectors have been a concern for future HSPC-based genetic therapies. Despite safer clonal repopulation patterns found in recent human gene therapy studies with lentiviral vectors followed for 1.5–2 years by VIS analysis (Aiuti et al., 2013; Biffi et al., 2013; Cartier et al., 2009; Cavazzana-Calvo et al., 2010), potentially life-long persistence of engineered stem cells warrants more careful monitoring of repopulating clones for a much longer period. We demonstrate safe and stable hematopoietic reconstitution in nonhuman primates over 4–12 years (equivalent to 10–30 years in humans, based on the average life span of the two species) without evidence of clones with obvious aberrant behavior. Gene modification of HSPCs with an shRNA directed to CCR5, an essential coreceptor for CCR5...
R5-tropic strains of HIV-1 (An et al., 2007), and targeting the same region of CCR5 as a candidate anti-HIV gene therapeutic for human clinical trials (http://www.clinicaltrials.gov; protocol NCT01734850) showed no notable effects on HSPC longevity or lineage-output behaviors. Thus, our long-term analysis can serve as a frame of reference for monitoring the safe use of HSPCs and lentiviral delivery in the clinic, including HIV gene therapy and other therapies using HSPCs.

Figure 5. Summary Diagrams Showing the Relative Proportions of HSPC Subpopulations and Their Contributions to Blood Repopulation over Time

(A) Summary diagrams. (Ai) shows the relative proportions of long-term repopulating cells within the transplanted CD34+ cell pool. (Ai) shows the contributions of short-term clones (ST-Clones) and the three subtypes of long-term clones over time.

(B) These charts show variable repopulation kinetics among the clones within each subpopulation. The numbers of My-bi, Bal, Ly-bi, and ST-QVI clones in the various kinetics clusters are indicated by bar graphs. The clusters and color codes for each cluster are the same as those in Figure 3A.

(C) The bar graphs show the proportions of My-bi, Bal, and Ly-bi clones present in the CD34+ HSPC pool near the end point (denoted by red bars) relative to the total detected in the mature lineages (denoted by black bars) (see Figure 4Ai–4Di) for individual clones in mature lineages and CD34+ cells. The long-term HSPC population is relatively enriched for the Bal and the My-bi clones.

(D) Comparison of clonal compositions across cell types. The relative frequencies of individual clones are compared between cell types. The Pearson correlations coefficient value ($r$) is indicated by a rainbow color scheme.
EXPERIMENTAL PROCEDURES

Animal Care and Peripheral Blood Cell Isolation

Four rhesus macaques were maintained in accordance with federal guidelines and the policies of the Veterinary Research Program of the National Institutes of Health according to the protocols approved by the Animal Care and Use Committee of the National Heart, Lung, and Blood Institute. Procedures for peripheral blood cell isolation can be found in the Supplemental Information.

VIS Analysis

VISs in repopulating cell samples were analyzed by TaqαI-mediated, bidirectional VIS sequencing and bioinformatics analysis, as described previously (Kim et al., 2010) with modifications for animals 2RC003, RQ3570, and RQ5427 (see Supplemental Information for details).

Kinetics Clusters

Major clonal kinetics patterns were derived using the WGCNA R package v.1.20 (Langfelder and Horvath, 2008), Clusters were defined by using thresholds that were specific to each data set, where each threshold was selected to yield clusters with visually distinct kinetics profiles. The significance of the clusters was quantified using the modulePreservation function in the WGCNA package (Langfelder and Horvath, 2008) as described above (see Supplemental Information for details).

Lineage Analysis

Analogous to previous studies (Dykstra et al., 2007; Lu et al., 2011), the relative contributions of individual LT-QVIs toward grans/monocytes (GM: CD14 and CD18), T cells (CD4 and CD8), and B cells (CD20) were determined with ternary plots and cluster analysis using the WGCNA R package v.1.20 (see Supplemental Information for details).

Genomic Vector Integration Site Hot spots

The BCP model (Presson et al., 2011) was used to define hot spots (see Supplemental Experimental Procedures for details). The correlation of hot spots among different HSPC subtypes was tested based on a Fisher’s exact test for hot bin overlap (see Supplemental Information for details).

SUPPLEMENTAL INFORMATION

Supplemental Information for this article includes Supplemental Experimental Procedures, seven figures, and five tables and can be found with this article online at http://dx.doi.org/10.1016/j.stem.2013.12.012.

ACKNOWLEDGMENTS

We thank the animal support and veterinary staff of both 5 Research Court and the Division of Veterinary Resources for maintaining these nonhuman primates for all these years; Keyvan Keyvanfar of the Hematology Branch and Ann Williams of the Core Flow Cytometry Laboratory, NHLBI, for performing cell sorting; and the UCLA Genotyping and Sequencing Core for performing DNA sequencing. This work was supported in part by the intramural program of the NIH, Hematology Branch, National Heart, Lung and Blood Institute; NIH grants AI055281-06A2 (I.S.Y.C.), CA68859 (S.A.C.), P01 HL073104 (G.M.C.), and K99 HL116234 (S.K.); the California Institute of Regenerative Medicine (S.K.), and the California Institute of Regenerative Medicine (S.K.).
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