3’ UTR-truncated HMGA2 overexpression induces non-malignant in vivo expansion of hematopoietic stem cells in non-human primates

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Vector-mediated mutagenesis remains a major safety concern for many gene therapy clinical protocols. Indeed, lentiviral-based gene therapy treatments of hematologic disease can result in oligoclonal blood reconstitution in the transduced cell graft. Specifically, clonal expansion of hematopoietic stem cells (HSCs) highly expressing HMGA2, a chromatin architectural factor found in many human cancers, is reported in patients undergoing gene therapy for hematologic diseases, raising concerns about the safety of these integrations. Here, we show for the first time in vivo multilineage and monoclonal expansion of non-human primate HSCs expressing a 3’ UTR-truncated version of HMGA2 without evidence of any hematologic malignancy >7 years post-transplantation, which is significantly longer than most non-human gene therapy pre-clinical studies. This expansion is accompanied by an increase in HSC survival, cell cycle activation of downstream progenitors, and changes in gene expression led by the upregulation of IGF2BP2, a mRNA binding regulator of survival and proliferation. Thus, we conclude that prolonged ectopic expression of HMGA2 in hematopoietic progenitors is not sufficient to drive hematologic malignancy and is not an acute safety concern in lentiviral-based gene therapy clinical protocols.

INTRODUCTION
Hematopoietic stem cells (HSCs) are the target cell for gene therapy of many hematologic and non-hematologic diseases because of their lifelong self-renewal and differentiation potential, which ensures long-term clinical benefits.1,4 Lentiviral vectors represent the favored approach for ex vivo gene transfer into HSCs and have been used routinely in clinical gene therapy protocols.7 However, the safety of lentiviral genomic integration remains a concern.7 Indeed, hematopoietic clonal expansion has been observed in several clinical protocols employing lentiviral vectors. However, the clinical significance of these expansions and their propensity to convert to hematologic malignancy remains unclear.7

Disruption of HMGA2 via vector integration has been a safety concern in gene therapy for years. Hmga2 encodes an architectural transcription factor whose chromosomal rearrangement or truncation results in tumors in patients and transgenic mice.10–15 Upregulation of a 3’ UTR-truncated HMGA2 caused by intronic integration of a lentiviral vector resulted in HSC clonal expansion in a beta-thalassemia gene therapy trial.16–18 Numerous clones containing intronic vector integrations in HMGA2 have also been seen in gamma-retroviral and lentiviral SCID-X1 gene therapy.19 Moreover, cases of clonal hematopoiesis have been linked to overexpression of proteins functionally downstream of HMGA2.20–22 These findings are not surprising given that ectopic Hmga2 can expand murine HSCs and knockdown or overexpression of HMGA2 perturbs the repopulating and differentiation potential of human HSCs.21–23 However, a single corrected and markedly expanded clone persisted without malignant transformation in an early ADA-SCID gene therapy trial,24 suggesting that clonal expansion does not irrevocably progress to malignancy. Thus, the clinical prognosis of gene therapy patients with HMGA2-associated hematopoietic clonal expansions is unclear.

Here, we provide evidence that persistent expression of a 3’ UTR-truncated HMGA2 cDNA, which causes a massive overexpression of this protein, leads to the competitive oligoclonal expansion of long-term HSCs in transplanted pigtailed macaques (M. nemestrina) without perturbed cellular differentiation or evidence of malignancy >7 years post-transplantation.

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RESULTS

3' UTR-truncated HMGA2 yields multiclonal CD34+ cell expansion in non-human primates

HMGA2 is an architectural non-histone chromatin protein that regulates gene expression by altering chromatin structure.25–28 HMGA2 contains seven let-7-micro RNA (miRNA) binding sites in the 3' UTR, which negatively regulates HMGA2 translation.29 To overexpress HMGA2, we amplified the protein coding region devoid of let-7 binding sites (HMGA2D7) from human HEK293T cells by reverse transcription-PCR. The PCR product was subcloned into the Cl20c lentiviral vector so that expression of HMGA2D7 would be driven by a murine stem cell virus (MSCV) promoter, which has good activity in HSCs (Figure 1A). An internal ribosome entry site (IRES)-GFP expression cassette was also included to allow for tracking of HMGA2D7-expressing cells by flow cytometry. A control vector was constructed in which mCherry cDNA is also driven by the MSCV promoter (Figure 1A). An internal ribosome entry site (IRES)-GFP expression cassette was also included to allow for tracking of HMGA2D7-expressing cells by flow cytometry. A control vector was constructed in which mCherry cDNA is also driven by the MSCV promoter (Figure 1A). Both vectors were produced transiently in HEK293T cells with a titer of 2 × 10⁶ tu/mL and 5 × 10⁶ tu/mL after concentration, respectively. HEK293T, HeLa, and 3T3 cells transduced with the HMGA2D7 vector expressed the expected 19 kDa HMGA2 protein (Figure 1B). Pigtailed macaque CD34+ cells transduced with the HMGA2D7 vector also expressed HMGA2 (Figure 1D). GFP expression was comparable to HMGA2 (data not shown). Thus, the lentiviral HMGA2 vector can be produced with sufficient titer to transduce CD34+ cells from pigtailed macaque.

To test whether HMGA2 expression expands HSCs in large animals, we isolated CD34+ cells from granulocyte-colony stimulating factor (GCSF)-mobilized bone marrow (BM) of pigtailed macaques. Cells were transduced with a lentiviral vector containing HMGA2D7-GFP or the control lentiviral vector (containing mCherry only). A mix of transduced cells by the two different vectors was transplanted into the same animal after lethal irradiation (Figure 2A). Two animals (A10W016 and A10W027) were transplanted. The transduction efficiency in the total graft for animal A10W016 was 42.9% for GFP and 19.1% for mCherry, and that for animal A10W027 was 6% for GFP and 7.9% for mCherry. See Table S1 for transplant details. Blood and platelets were supplied as needed until absolute neutrophil counts (ANCs) reached 500/mL and platelet counts reached 50,000/mL (Figure S1A). Six days post-transplant, peripheral blood (PB) chimerism was 4%–6% for GFP and mCherry in both animals, despite higher contribution of GFP-marked cells in A10W016's graft (Figure 2Bi). White blood cell (WBC) chimerism decreased thereafter, and by 3 months post-transplant, PB chimerism was 2.9% GFP+ and 1.1% mCherry+ for animal A10W027 (Figure 2Bi), suggesting equivalent and low HSCs and progenitor cell transduction with both vectors. Thereafter, GFP chimerism slowly but progressively increased, while mCherry chimerism decreased further in both animals (Figure 2Bi). By day 200, the GFP marking in WBCs reached 8.9% and 4.5% in A10W016 and A10W027, respectively, while the mCherry marking decreased to 0.9% and 1.5%, respectively (Figure 2Bi).
HMGA2 overexpression.

Differentiation of transduced cells. GFP+ and mCherry+ cells were sorted Vector integration site (VIS) analysis over time and in multiple was further reected by the presence of GFP label in BM cells, where the most primitive hematopoietic cells displayed 60% and 80% GFP chimerism (HSC: CD34+CD38−CD90+CD45RA−) for animals A10W016 and A10W027, respectively, at the latest time point of 74 months (Figure 2C). BM hematopoietic progenitors (multipotent progenitors [MPPs]: CD34+CD38−CD90−CD45RA− and multi-lymphoid progenitors [MLPs]: CD34+CD38−CD90−CD45RA−) also show increased GFP marking (Figure 2C). HMGA2 protein was detectable in sorted GFP+ WBCs but not GFP− WBCs (Figure S1C).

Vector integration site (VIS) analysis over time and in multiple mature lineages allows tracing of cellular clonal dynamics and the differentiation of transduced cells. GFP+ and mCherry− cells were sorted from PB and the genomic DNA subjected to VIS analysis. The VIS in the mCherry− subpopulation in A10W027 was not analyzed because of limited cell number. More than 15 different clones contributing to the blood were detected in all the lineages of both animals (Figure 2D; Table S2), indicating expansion of multipotent HSCs. HSC expansion could be due to dysregulation of multiple genes and/or independent of gene regulation. Affymetrix array analysis of RNA from BM CD34+GFP+ and CD34+GFP− cells isolated 22–26 weeks post-transplant revealed 39 transcripts upregulated and three transcripts downregulated in the CD34+GFP+ subpopulation compared to the CD34−GFP− subpopulation in both animals (Table S3). As expected, HMGA2 was the highest differentially expressed gene in both animals. The only other gene upregulated by >2-fold was IGF2BP2 (Figure 4A; Table S3), which is also differentially expressed in HSCs and progenitors compared to differentiated cells in humans and mice (Figure S2). qRT-PCR analysis showed that the p16INK4a was not altered in animal A10W016 and was slightly reduced (67% of CD34+GFP+ cells) in animal A10W027 (Figure 4B).

Complete blood counts, including total WBC and red blood cell (RBC) counts and PB lineage distribution, were in the normal range (Table S4), demonstrating lack of any detectable hematopoietic abnormality. BM smears also were normal (Figure S3A). Despite IGF2BP overexpression being linked to changes in hemoglobin expression patterns, hemoglobin was also normal in recipients of HMGA2−ΔL7-expressing cells (Figure S3B).

DISCUSSION

HMGA2 high expression has been linked to expansion of single corrected clones that persisted without malignant transformation for years in multiple gene therapy trial.40–43 These observations suggest that clonal expansion does not irrevocably progress to malignancy. In contrast, there are concerns that HMGA2 overexpression may increase the risk of expanding clones with pre-existing oncogenic mutations, as in clonal hematopoiesis of undetermined potential.35 Indeed, HMGA2 is considered a putative oncogen.10–13 Chromosomal translocations with breakpoints in HMGA2 that eliminate the 3’ UTR let-7 miRNA binding sites cause multiple tumor types.36–38 High expression of HMGA2 in cancers of epithelial origin associates with increased aggressiveness39,40 and is seen in some leukemias.13 Chromosomal translocation of HMGA2 has also been seen in two cases of paroxysmal nocturnal hemoglobinuria.41 Six distinct transgenic mouse models have been made with different promoters driving expression of HMGA2.12 Interestingly, these mice never develop hematologic malignancies, even after three rounds of serial BM transplantation.34 A key difference between these transgenic mice and our study is that advantage of HMGA2ΔL7-expressing cells over time in transplanted pigtailed macaques.

3’ UTR-truncated HMGA2 expression perturbs the gene expression profile of primate CD34+ cells without malignancy

HMGA2 alters transcription of multiple genes.30–33 Our observed HSC expansion could be due to dysregulation of multiple genes and/or independent of gene regulation. 3’ UTR-truncated HMGA2 expression perturbs the gene expression profile of primate CD34+ cells without malignancy. 3’ UTR-truncated HMGA2 expression perturbs the gene expression profile of primate CD34+ cells without malignancy.
overexpression of HMGA2 started in the fetus and could not be downregulated later during development because of a pan-cellular promoter. Additionally, IGF2BP overexpression is also implicated in the malignant transformation of human CD34+ cells.\(^5\) However, animals transplanted here with HMGA2ΔL7-expressing cells displayed a normal hematologic profile. HMGA2 directly regulates IGF2BP2 during myogenesis\(^46\) and embryonic development. IGF2BP2 upregulation has also been associated with the regulation of stem cells during tissue regeneration.\(^47\) IGF2BP proteins have been ascribed an anti-apoptotic role in multiple cell contexts,\(^48,50\) and IGF2BP is known to boost proliferation of several tissues.\(^51–53\)

Thus, upregulated IGF2BP2 may explain our observed increased survival and proliferation of hematopoietic stem and progenitor cells (HSPCs). Alternatively, HMGA2 binds the DNA replication fork in murine embryonic stem cells and protects against nucleolytic fork collapse, which can cause double strand DNA breaks and trigger apoptosis, a differentiation block, or senescence.\(^54\) Thus, it is possible that HMGA2 stabilizes replication forks in HSC, thereby reducing apoptosis while also enhancing self-renewal, resulting in the slow but progressive competitive expansion of GFP+ HSCs in transplanted animals. Future studies will be needed to test these potential molecular mechanisms. HMGA2 expression also suppresses the CDKN2a locus in neural stem cells in young mice.\(^54\)

In sum, here we show for the first time in non-human primates that long-term HSCs dramatically expand in vivo when 3’ UTR-truncated HMGA2 (HMGA2ΔL7) is overexpressed. This expansion was slow but progressive, taking ∼2 years for cells to expand from 1%–3% to 40%–60%. Once at those levels, expansion stabilized for >7 years, reaching 60%–80% in the most primitive HSPCs. Importantly, expansion was oligoclonal, as multiple GFP-marked clones were observed. Further, expanded HSCs maintained multilineage differentiation potential as evidenced by the progressive increase of GFP+ cells and the presence of a multiple identical VIS in all PB lineages. Finally, expansion was accompanied by increased HSPC survival and enhanced proliferation of downstream progenitors, which likely contributed to the competitive advantage of HMGA2ΔL7-expressing cells.

Importantly, the two transplanted pigtailed macaques in our study showed no evidence of hematologic malignancies >7 years post-transplant. Thus, even in this “worse-case scenario” where HMGA2 is massively overexpressed and no longer subject to typical post-transcriptional regulation, HMGA2 overexpression was not oncogenic in non-human primate HSCs. Recently, one case of acute myeloid leukemia (AML) and one case of myelodysplastic syndrome (MDS) have been reported in two participants with sickle cell disease who were treated with lentiviral vector gene therapy; it is currently unclear whether these malignancies are related to vector integration.\(^55\) Yet our study, with truncated HMGA2 overexpression driven by a strong viral promoter in a lentiviral vector and no detected hematopoietic malignancy, suggests that massive HMGA2 overexpression in the context of lentiviral integration is not sufficient to drive malignancy. Although these clones should be monitored in the gene therapy setting, our data suggest that they are benign and unlikely to progress to malignancy.
MATERIALS AND METHODS

Animal usage
Healthy juvenile pigtailed macaques (M. nemestrina) weighing between 3 and 8 kg were housed at the St. Jude Children’s Research Hospital Animal Research Center (Memphis, TN, USA) and at the Keeling Center for Comparative Medicine and Research at MD Anderson Cancer Center (Houston, TX, USA). All studies and procedures were reviewed and approved by the St. Jude Children’s Research Hospital Institutional Animal Care and Use Committee.

Cell culture
HEK293T, HeLa, and NIH 3T3 cells were obtained from the American Type Culture Collection (Manassas, VA, USA). Cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) with 10% fetal bovine serum, 2 mM l-glutamine (Invitrogen, Waltham, MA, USA), 50 U/mL penicillin, and 50 mg/mL streptomycin (Invitrogen, Waltham, MA, USA), unless otherwise specified, and maintained at 37°C and 5% CO2.

Lentiviral vector design, production, and titration
Human HMGA2DL7 (HMGA2 lacking the Let-7 target sites) was amplified from HEK293T cDNA with primers designed to exclude the 3' UTR Let-7 target sites (forward: TGA<em>G</em>GAATTCCAGCGAGTGAAGGACGCGG; reverse: GCGAGCAATCGATCTAGTCTTCGCG). The forward primer contains an EcoRI restriction site (underlined), and the reverse primer contains a ClaI restriction site (underlined). The HMGA2DL7 amplicon (359 bp) was subcloned by using the designed EcoRI and ClaI restriction sites to create pCL20c-MSCV-HMGA2DL7-IRE5-GFP. Lentiviral vector was produced with HEK293T cells, using calcium phosphate precipitation to cotransfect the four plasmids: the CL20 vector plasmid (HMGA2DL7 or mCherry), pCAG-VSVG, pCAG4-RTR2, and pCAGkGP1R. The following day, cells were washed once in PBS and viral collection media. Stemline supplemented with 1% human serum albumin (HSA) and 2 mM l-alanyl-l-glutamine (Cellgro, Herndon, VA, USA) was added. Vector-containing supernatant was collected 48 h post-transient transfection. Human HMGA2 has been used previously in murine cells and shown to be functional.56 Also, human and macaque HMGA2 proteins are nearly identical (only 7/147 amino acids differ, while 55/147 amino acids differ compared with murine HMGA2 protein).

To determine vector titer, 2 × 10^5 HeLa cells were transduced in 6-well dishes with 6 μg/mL polybrene and various volumes of vector in a final volume of 2 mL/well. The following day, 5 mL of medium was added to each well. Cells were analyzed for GFP or mCherry expression 4–6 days later with flow cytometry.

Western blot
Cells (1–2 × 10^6) were collected, washed with PBS, resuspended in 50 mL of PBS with protease inhibitor (Halt) and then mixed with 50 mL of 2× Laemml buffer with 5% β-mercaptoethanol. Samples were boiled for 10 min and then loaded onto 4–12% NuPage gels. Antibodies were against HMGA2 (Abcam, Cambridge, MA, USA) and Actin (SCBT, Dallas, TX, USA).

Flow cytometry
Whole blood or BM samples were treated with BD Pharm Lyse (BD Biosciences, San Jose, CA, USA) for 5 min at room temperature and then spun for 10 min at 300 g at 4°C. Leukocytes were resuspended in magnetic cell sorting (MACS) buffer and stained with indicated antibodies against cell surface markers at 4°C for 20–40 min. Table S5 details all antibodies used in this article. Samples were analyzed with an LSR Fortessa (BD Biosciences, San Jose, CA, USA). Intracellular HMGA2 staining was achieved with the Foxp3/Transcription Factor Staining Buffer Set (eBioscience, San Diego, CA, USA) according to manufacturer’s protocol and HMGA2 antibody (Cell Signaling Technology, Danvers, MA, USA).

CD34+ cell enrichment
GCSF-mobilized BM was harvested from two pigtailed macaques. BM was diluted 10-fold with RBC lysis buffer (BD Biosciences, San Jose, CA, USA) and incubated at room temperature for 5 min prior to centrifugation at 300 × g for 10 min at 4°C. Cells were resuspended in MACS buffer (Miltenyi Biotec, Carlsbad, CA, USA) and passed through a 70 mm cell strainer. Cells were counted and resuspended.
at $2 \times 10^8$ cells/mL in MACS buffer. Cells were incubated with CD34-PE antibody for a total of 45 min on ice with gentle shaking. Antibody-stained cells were then washed, resuspended at $2 \times 10^8$ cells/mL, and incubated with a combination of anti-PE microbeads (Miltenyi Biotec, Carlsbad, CA, USA) for 30 min on ice with gentle shaking. Cells were washed, resuspended at $2 \times 10^8$ cells/mL, and enriched with a LD column (Miltenyi Biotec, Carlsbad, CA, USA). CD34+ cells were eluted and analyzed by flow cytometry for purity.

**CD34**+ cell culture and transduction

CD34+ cells were cultured in X-VIVO 10 medium (Lonza, Portsmouth, NH, USA) supplemented with 1% HSA and 100 ng/mL human stem cell factor (hSCF), human thrombopoietin (hTPO), and human Fms-like tyrosine kinase 3 ligand (hFLT3-L) (CellGenix, Portsmouth, NH, USA). Cells were cultured on tissue culture dishes coated with RetroNectin (TaKaRa Bio, Shiga, Japan). Cells were cultured for one night prior to the first overnight vector exposure in the presence of 4 mg/mL protamine sulfate. After the first vector exposure, cells were washed and cultured in the previously described X-VIVO 10 medium for 8–12 h before a second overnight vector exposure. The multiplicity of infection (MOI) of transduction was between 50 and 100. The next day, cells were washed in fresh medium and resuspended in PlasmaLyte-148 (Baxter Healthcare, Cleveland, MS, USA) with 2% HSA (Sigma-Aldrich, St. Louis, MO, USA) and passed through a 70-mm filter prior to infusion into autologous recipients irradiated with 950 cGy.

**Vector insertion site determination**

The method for vector insertion site determination has been detailed elsewhere.

Genomic DNA from sorted cell populations was isolated with DNeasy Blood & Tissue Kits (QIAGEN, Germantown, MD, USA). A control sample with 19 known and confirmed integration sites was spiked into each sample at 2% of the final amount. DNA samples were subjected to random shearing with a E210 sonicator (Covaris, Woburn, MA, USA) (200 cycles/burst, Intensity 3, 65 s). After shearing, the DNA ends were repaired with NEBNext End Repair Module, and then a poly(A) tail was added with the NEBNext dA-Tailing Module (New England Biolabs, Ipswich, MA, USA). Then, the NEBNext Adaptor for Illumina was added, followed by a round of linear amplification with a biotinylated primer. The amplified product was then enriched using streptavidin Dynal M-270 beads (Thermo Fisher Scientific, Waltham, MA, USA) and amplified further with a final nest PCR. Samples were pooled and run on an E-gel and DNA ranging from ~250 to 800 bp was extracted with a DNA Gel Extraction Kit (QIAGEN, Germantown, MD, USA) and analyzed with Mi-Seq (Illumina, San Diego, CA, USA). 100–1,000 ng of genomic DNA isolated from sorted subpopulations of PB was used for VIS analysis using a quantitative PCR.

**Microarray analysis**

Total RNA was extracted from sorted GFP+CD34+ or GFP-CD34+ BM cells and subjected to Cynomolgus/Rhesus Gene 1.0 ST Array (Thermo Fisher Scientific, Waltham, MA, USA) with established protocols.
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