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WHIM syndrome immunodeficiency is caused by autosomal dominant gain-of-function mutations in chemokine receptor CXCR4. Patient WHIM-09 was spontaneously cured by chromothriptic deletion of one copy of 164 genes, including the CXCR4WHIM allele, presumably in a single hematopoietic stem cell (HSC) that repopulated HSCs and the myeloid lineage. Testing the specific contribution of CXCR4 hemizygosity to her cure, we previously demonstrated enhanced engraftment of Cxcr4+/o HSCs after transplantation in WHIM (Cxcr4+/w) model mice, but the potency was not quantitated. We now report graded-dose competitive transplantation experiments using lethally irradiated Cxcr4+/+ recipients in which mixed BM cells containing ~5 Cxcr4+/o HSCs and a 100-fold excess of Cxcr4+/w HSCs achieved durable 50% Cxcr4+/o myeloid and B cell chimerism in blood and ~20% Cxcr4+/o HSC chimerism in BM. In Cxcr4+/o/Cxcr4+/w parabiotic mice, we observed 80-100% Cxcr4+/o myeloid and lymphoid chimerism in the blood and 15% Cxcr4+/o HSC chimerism in BM from the Cxcr4+/w parabiont, which was durable after separation from the Cxcr4+/o parabiont. Thus, CXCR4 haploinsufficiency likely significantly contributed to the selective repopulation of HSCs and the myeloid lineage from a single chromothriptic HSC in WHIM-09. Moreover, the results suggest that WHIM allele silencing of patient HSCs is a viable gene therapy strategy.

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\textbf{Short title:} Development of gene therapy for WHIM syndrome

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ABSTRACT

WHIM syndrome immunodeficiency is caused by autosomal dominant gain-of-function mutations in chemokine receptor CXCR4. Patient WHIM-09 was spontaneously cured by chromothriptic deletion of one copy of 164 genes, including the CXCR4\textsuperscript{WHIM} allele, presumably in a single hematopoietic stem cell (HSC) that repopulated HSCs and the myeloid lineage. Testing the specific contribution of CXCR4 hemizygosity to her cure, we previously demonstrated enhanced engraftment of Cxcr4\textsuperscript{+/-} HSCs after transplantation in WHIM (Cxcr4\textsuperscript{+/-}) model mice, but the potency was not quantitated. We now report graded-dose competitive transplantation experiments using lethally irradiated Cxcr4\textsuperscript{+/-} recipients in which mixed BM cells containing ~5 Cxcr4\textsuperscript{+/-} HSCs and a 100-fold excess of Cxcr4\textsuperscript{+/-} HSCs achieved durable 50% Cxcr4\textsuperscript{+/-} myeloid and B cell chimerism in blood and ~20% Cxcr4\textsuperscript{+/-} HSC chimerism in BM. In Cxcr4\textsuperscript{+/-}/Cxcr4\textsuperscript{+/-} parabiotic mice, we observed 80-100% Cxcr4\textsuperscript{+/-} myeloid and lymphoid chimerism in the blood and 15% Cxcr4\textsuperscript{+/-} HSC chimerism in BM from the Cxcr4\textsuperscript{+/-} parabiont, which was durable after separation from the Cxcr4\textsuperscript{+/-} parabiont. Thus, CXCR4 haploinsufficiency likely significantly contributed to the selective repopulation of HSCs and the myeloid lineage from a single chromothriptic HSC in WHIM-09. Moreover, the results suggest that WHIM allele silencing of patient HSCs is a viable gene therapy strategy.
INTRODUCTION

Warts, Hypogammaglobulinemia, Infections, Myelokathexis (WHIM) syndrome is a rare, combined primary immunodeficiency disease caused in almost all cases by autosomal dominant, gain-of-function mutations in the gene encoding chemokine receptor CXCR4 (1, 2). In response to its principal agonist CXCL12, CXCR4 signaling normally mediates homing, retention, and multilineage differentiation of hematopoietic stem and progenitor cells (HSPCs) in bone marrow (BM) niches by regulating their migration, survival, and quiescence (3-6). CXCR4 signaling also normally regulates migration and distribution of most types of mature leukocytes. WHIM mutations, which block normal arrestin-dependent CXCR4 desensitization and downregulation processes, distort normal leukocyte trafficking and distribution resulting in neutropenia, and even panleukopenia in many patients, despite normal hematopoietic development (7-9). Neutropenia is typically severe and occurs despite BM myeloid hyperplasia due to BM retention of fully mature and degenerating neutrophils (10-13). This striking hematopathologic picture, which is almost pathognomonic of WHIM syndrome, is called myelokathexis, a Greek neologism meaning ‘marrow retention’ and the ‘M’ in the acronym WHIM (14).

We previously reported that patient WHIM-09 was cured of WHIM syndrome when the disease allele $\text{CXCR4}^{R334X}$ and 163 other genes were spontaneously deleted on one copy of chromosome 2 by chromothripsis presumably in a single HSC (15). Chromothripsis refers to random chromosome shattering resulting in all-at-once complex chromosomal rearrangements and gene deletions and duplications (16). It occurs in a small percentage of cancers and has been reported in a few patients with congenital neurodevelopmental disorders (17, 18). WHIM-09 is the only patient known to have had a beneficial outcome from chromothripsis (15). Fortuitously, the original chromothriptic HSC in WHIM-09 acquired a selective advantage and repopulated both
the HSC pool and the entire myeloid lineage. The absolute neutrophil count (ANC) and absolute monocyte count (AMC) rose to new baselines ~1.5-2 fold above normal limits where they have remained stable for >20 years associated with both spontaneous clearance of warts and cessation of recurrent acute infections. In contrast to the myeloid lineage, the chromothriptic HSCs were unable to repopulate the patient’s lymphoid lineage, so that she became a hematopoietic chimera of WHIM lymphoid cells and chromothriptic non-WHIM myeloid cells hemizygous (genotype +/o) for CXCR4 and the other 163 deleted genes. This implies that the WHIM allele drives major clinical phenotypes of the disorder in a myeloid-dependent manner.

Competitive bone marrow and HSC transplantation of 50:50 mixtures of donor cells from all paired combinations of Cxcr4+/o, Cxcr4+/+ and Cxcr4+/w (‘w’ denotes knockin of the human WHIM allele S338X) mice into lethally-irradiated congenic Cxcr4+/+ recipient mice suggested that deletion of the WHIM allele per se might be necessary and sufficient to account for WHIM-09’s cure, independently of the other 163 deleted genes (15). In particular, the results of the Cxcr4+/o/Cxcr4+/w HSC competition showed >95% Cxcr4+/o myeloid and B cell chimerism in the blood and ~80% Cxcr4+/o HSPC and mature leukocyte chimerism in the bone marrow, associated with enhanced proliferation of Cxcr4+/o HSCs in vivo. Consistent with this, CXCR4 signaling had previously been shown to suppress HSC proliferation (5), suggesting a cure mechanism for WHIM-09 involving a competitive proliferative advantage in HSCs conferred by CXCR4 haploinsufficiency.

We also reported a competitive advantage for Cxcr4+/o HSCs over Cxcr4+/+ and Cxcr4+/w HSCs for hematopoietic reconstitution after transplantation into unconditioned congenic Cxcr4+/w recipient mice (19). In this case, competitive transplantation of 50 million Cxcr4+/o total bone marrow cells (~5000 HSCs (20)) into unconditioned WHIM recipient mouse resulted in ~70%
durable Cxcr4+/o donor CD11b+ myeloid cell chimerism in the blood despite achieving only 6% Cxcr4+/o donor HSC chimerism in the bone marrow.

Our previous studies did not define the magnitude of the selective advantage conferred by CXCR4 haploinsufficiency on HSC engraftment efficiency, which is important for considerations of gene therapy strategy in the disease. Moreover, a major difference between transplantation experiments in mice and the spontaneous genetic cure that occurred in WHIM-09, other than the additional 163 genes deleted in WHIM-09, is the manipulation and transplantation of HSCs in the former but not the latter. Here we address both these issues by experiments in: 1) lethally-irradiated recipient mice competitively transplanted with graded ratios of Cxcr4+/o and Cxcr4+/w donor BM cells, and 2) parabiotic mice, in which HSC manipulation and transplantation are obviated.
RESULTS

*Cxcr4*+/o donor BM cells have a strong selective advantage over *Cxcr4*+/w donor BM cells for reconstituting myeloid and B cells in the blood after transplantation of lethally irradiated mice

To better understand the role played by CXCR4 haploinsufficiency in the cure of patient WHIM-09, we performed competitive BM transplantation experiments under increasingly stringent conditions, first by transplanting 5 million donor BM cells with different ratios of *Cxcr4*+/o and *Cxcr4*+/w cells into lethally irradiated congenic *Cxcr4*+/+ recipient mice. Our hypothesis was that if CXCR4 haploinsufficiency per se in the original chromothriptic HSC was necessary and sufficient to cure her, then in competitive *Cxcr4*+/o and *Cxcr4*+/w BM cell transplantation experiments in lethally irradiated mice we might observe a threshold dose of *Cxcr4*+/o donor BM cells for full reconstitution, but not a graded *Cxcr4*+/o donor BM cell dose-response. Donor and recipient mouse hematopoietic cells were differentially marked by CD45 polymorphisms for quantitation of reconstitution by flow cytometry over time (CD45.1 marked *Cxcr4*+/+, CD45.2 marked *Cxcr4*+/w and CD45.1/CD45.2 marked *Cxcr4*+/o). We have previously reported for these donors that the HSC frequencies in total bone marrow are equivalent and that transplantation of purified HSCs gives equivalent results to transplantation of total bone marrow cells when the input cell ratio is 50:50 (15).

Confirming our previous report, transplanting an ~50:50 mixture of the two BM donors resulted in durable ~100% *Cxcr4*+/o donor-derived myeloid and B cell chimerism in the blood (15, 19) (Figure 1A-C). In contrast, as we have previously reported, the post-transplantation donor-derived T cell ratio in recipient blood was similar to the input value (Figure 1B) (15,19). The reason for T cell exceptionalism in this transplantation model is not clear and will require additional focused research. Full *Cxcr4*+/o donor-derived myeloid and B cell chimerism in the
blood was also observed even when only 25% of the transplanted BM cells were from the Cxcr4⁺/⁻ donor (N.B. this represents ~125 HSCs, since ~0.01% of whole bone marrow cells are HSC (20)) (Figure 1B). However, a clear Cxcr4⁺/⁻ donor BM cell dose-response became apparent at lower ratios. Remarkably, at the lowest ratio tested, 1% input Cxcr4⁺/⁻ donor BM cells (~5 HSCs) still resulted in over 50% Cxcr4⁺/⁻ donor-derived myeloid and B cell chimerism in the blood (Figure 1B), which was sufficient to stably improve circulating myeloid and B cell levels in the transplanted animals to the normal range. In contrast, and similar to our previous studies, the absolute numbers of Cxcr4⁺/⁺ donor-derived myeloid cells and B cells in the blood after competitive transplantation remained below even the leukopenic norm for unmanipulated Cxcr4⁺/⁺ mice.

When the mice were sacrificed 278 days post-transplantation (a date chosen arbitrarily once durability of chimerism was established), we found that the frequencies of Cxcr4⁺/⁻ donor-derived myeloid cells and B cells also significantly increased in the bone marrow compared with the input donor cell frequencies across all input donor cell ratios tested, although to a lesser extent than in peripheral blood (Figure 1C). Consistent with this, the frequencies of Cxcr4⁺/⁻ donor-derived HSCs and HPCs (hematopoietic progenitor cells) also significantly increased in the bone marrow compared with the input frequencies across all input donor cell ratios tested (Figure 1D). Thus, the predominance of Cxcr4⁺/⁻ over Cxcr4⁺/⁺ donor-derived leukocytes in the blood may be the result of both superior engraftment of Cxcr4⁺/⁻ HSCs and superior egress of mature Cxcr4⁺/⁻ leukocytes. Nevertheless, the dose-response results do not entirely explain WHIM-09’s cure since there is a limit to the ability of transplanted and engrafted Cxcr4⁺/⁻ HSCs to fully reconstitute hematopoiesis when placed in competition with Cxcr4⁺/⁺ HSCs in this conditioned transplantation system.
Cxcr4+/- leukocyte chimerism predominates in both parabionts of Cxcr4+/w: Cxcr4+/o parabiotic mice

We next used parabiotic pairing of unconditioned Cxcr4+/w and Cxcr4+/o mice to test whether the process of harvesting Cxcr4+/o BM cells for transplantation, which did not happen in WHIM-09, might affect HSC engraftment efficiency. As with the transplantation model, in the parabiosis model Cxcr4 was the only one of the 164 genes deleted in WHIM-09 that was tested for the effects of haploinsufficiency on engraftment and chimerism. In particular, Cxcr4+/o mice were surgically joined with age- and sex-matched congenic CD45 isoform-marked Cxcr4+/w mice (designated as parabiotic pair Cxcr4+/o/Cxcr4+/w) to model the HSC competition that occurred in WHIM-09 but limited to a competition between Cxcr4 genotypes (Figure 2A). As a control, we generated parabiotic pairs comprised of Cxcr4+/+ mice joined to congenic CD45 isoform-marked Cxcr4+/+ mice (designated as parabiotic pair Cxcr4+/+ /Cxcr4+/+) (Figure 2A). This control allows assessment of the background pattern of leukocyte distribution in peripheral blood and BM established by parabiosis per se. After 5 months of parabiosis (the connected phase), 3 of the 5 pairs of parabiotic mice from both the experimental and control groups were terminated for BM analysis without first being separated. To test whether chimeric HSCs were stably engrafted, we surgically separated the remaining three parabiotic pairs (two in the Cxcr4+/o/Cxcr4+/w group and one in the Cxcr4+/+ /Cxcr4+/+ group) 5 months after parabiotic joining and then followed the separated parabionts for an additional 5 months (the separated phase) when they were arbitrarily sacrificed for BM analysis (see Figure 2A for experimental design).

After connecting the control Cxcr4+/+/Cxcr4+/+ parabiotic pair, peripheral blood chimerism was established quickly at ~30% for myeloid cells and ~45% for both B and T cells and was sustained at these levels across the entire time course of the connected phase with no significant
differences observed between the parabionts of each parabiotic pair (Figure 2B, upper panels). The reason the frequencies are not 50%, as would be predicted from simple mixing of the circulations, and differ between different subsets may be explained by the limited circulatory connection in the connected skin, which is made up of only small blood vessels across a limited area of contact, and the different rates of production of different leukocyte subsets (21). At the end of the connected phase, we found no difference between chimerism levels measured in blood and BM for either B cells or T cells (~45%), whereas chimerism for myeloid cells was only ~5% in BM compared to ~30% in blood (Figure 2C, upper panels). HSC chimerism in BM from the Cxcr4+/+ Cxcr4+/+ parabiotic pairs was even lower at ~3% without significant differences for all subsets in both parabionts (Figure 2E). After separation, mature leukocyte chimerism in the peripheral blood of each Cxcr4+/+ parabiont fell for each subset to a new stable level. In particular, Cxcr4+/+ myeloid chimerism fell quickly from ~30% to ~5%, whereas B and T cell chimerism both fell much more gradually over 5 months from 45% to ~20% (Figure 2B, upper panels). The different rates of reduction of chimerism between myeloid and lymphoid cells may be explained by their different half-lives. At the end of the separated phase, BM HSC and HPC chimerism was 3% for both parabionts (Figure 2E), the same as at the end of the connected phase. Although we only tested one +/+ : +/+ parabiotic pair in the separated phase, the results were similar to previous reports for this phase in wild type parabiotic mice (22, 23).

Compared to this control parabiosis hematopoietic chimerism pattern, chimerism levels in the Cxcr4+/o Cxcr4+/w parabiotic pairs were very different. In particular, Cxcr4+/o myeloid cells, B cells and T cells all rapidly reconstituted in the Cxcr4+/w parabiont after establishment of parabiosis, peaking at ~day 20 post-parabiotic connection at ~60-70% for myeloid cells and ~80-90% for B and T cells. These levels were maintained with limited excursion for 5 months, when
the animals were sacrificed or separated. Conversely, $Cxcr4^{+/w}$ chimerism levels were very low (<20% for each subset) in the $Cxcr4^{+/o}$ parabionts (Figure 2B, lower panels).

At the end of the connected phase, $Cxcr4^{+/o}$ HSC and HPC chimerism was much higher in BM from the $Cxcr4^{+/w}$ parabiont than was $Cxcr4^{+/w}$ HSC and HPC chimerism in BM from the $Cxcr4^{+/o}$ parabiont (15-20% versus 1-4%, respectively) (Figure 2E). Mature $Cxcr4^{+/o}$ myeloid and lymphoid cell chimerism in the $Cxcr4^{+/w}$ parabionts was much lower in BM than in blood, with myeloid chimerism in BM only ~15% and lymphoid chimerism in BM only 35% and 25% for B and T cells, respectively (Figure 2C, lower panels). $Cxcr4^{+/w}$ chimerism in BM from the $Cxcr4^{+/o}$ parabionts was similar to the blood with extremely low (<5%) levels of myeloid chimerism (Figure 2C, lower panels). BM $Cxcr4^{+/w}$ B cell and T cell chimerism was 40% and 60% for the $Cxcr4^{+/o}$ parabionts, respectively, the same as for BM $Cxcr4^{+/o}$ B cell chimerism in the $Cxcr4^{+/w}$ parabionts and more than double the level for BM $Cxcr4^{+/o}$ T cell chimerism in the $Cxcr4^{+/w}$ parabionts (Figure 2C, lower panels).

As with the control $Cxcr4^{+/+}/Cxcr4^{+/+}$ parabiotic pair, separation of both remaining $Cxcr4^{+/w}/Cxcr4^{+/o}$ pairs resulted in a decline in mature $Cxcr4^{+/w}$ leukocyte chimerism in the peripheral blood of the $Cxcr4^{+/o}$ parabionts to a new stable level with myeloid and B cell chimerism declining from ~5% to zero and T cell chimerism from ~20% to ~10% at the end of the separated phase, 5 months post parabiosis (Figure 2B, lower panels). However, $Cxcr4^{+/o}$ chimerism in the $Cxcr4^{+/w}$ parabionts was still maintained at the relatively high levels of ~40% myeloid and 60% lymphoid chimerism after the initial decline (Figure 2B, lower panels).

The pattern of BM HSC and HPC chimerism at the end of the separated phase was similar to the pattern at the end of the connected phase with ~15% $Cxcr4^{+/o}$ chimerism in the $Cxcr4^{+/w}$ parabiont and ~3% $Cxcr4^{+/w}$ chimerism in the $Cxcr4^{+/o}$ parabiont (Figure 2E). The BM myeloid
cell chimerism pattern was also similar in the two phases (Figure 2D, lower panels). The BM lymphoid chimerism pattern at the end of the separated phase was slightly different from the pattern at the end of the connected phase with \( Cxcr4^{+/o} \) chimerism in the \( Cxcr4^{+/w} \) parabiont \(-10\) and \(45\)% for B and T cells, respectively, and \( Cxcr4^{+/w} \) chimerism in the \( Cxcr4^{+/o} \) parabiont \(-35\) and \(50\)% for B and T cells, respectively (Figure 2D, lower panels).

Thus, compared to the isogenic \( Cxcr4^{+/+}/Cxcr4^{+/+} \) control parabiotic mice, in which there is no difference in chimerism levels between each parabiont in any leukocyte subset in either bone marrow or blood and in either the connected or separated phase, chimerism levels in paired parabionts are strongly skewed by \( Cxcr4^{+/w}/Cxcr4^{+/o} \) competition in parabiotic mice in all subsets analyzed in both compartments and both phases. Owing to the stress placed on mice by parabiotic life, we limited the analysis to these two pairs which gave results consistent with the results of \( Cxcr4^{+/w}/Cxcr4^{+/o} \) competitive transplantation experiments in both irradiated and unconditioned congenic recipients and with published reports for isogenic \( Cxcr4^{+/+}/Cxcr4^{+/+} \) parabiotic mice. The likelihood that the other two isogenic control pairings \((Cxcr4^{+/o}/Cxcr4^{+/o}, Cxcr4^{+/w}/Cxcr4^{+/w})\) would give different skewed results is very low.
DISCUSSION
The present study demonstrates that Cxcr4 haploinsufficiency enhances hematopoietic reconstitution in the setting of both bone marrow transplantation and parabiosis in mice and suggests that it may have been a contributory genetic mechanism, but possibly not the sole genetic mechanism, underlying the remarkable cure of WHIM patient WHIM-09. The results confirm and extend our previous results testing the effects of Cxcr4 haploinsufficiency on HSC engraftment in mice using less stringent conditions that were limited to transplantation (15, 19). Most importantly, the results indicate that even low-level engraftment of Cxcr4 haploinsufficient HSCs in bone marrow is sufficient to correct leukopenia in WHIM mice, which has important implications for gene therapy strategies in the disease.

Precise delineation of the hematologic cure mechanism for WHIM-09 requires a minimum of two elements: a mechanism for silencing the hyperactive WHIM allele in an HSC and a mechanism for replacing WHIM hematopoiesis with non-leukemic clonal expansion of the original chromothriptic HSC. The chromothriptic event provides a unifying genetic mechanism for both elements. Chromothriptic deletion on one copy of chromosome 2 is a straightforward and obvious explanation for WHIM allele silencing in HSCs and the myeloid lineage. Moreover, the myeloid selectivity of the chromothriptic event not only reveals that CXCR4 haploinsufficiency in the myeloid lineage is a viable genetic state but also that myeloid expression of the WHIM mutation may be a dominant factor driving the main clinical manifestations of the disease. However, identifying the gene(s) that contributed to clonal expansion of the original chromothriptic HSC is not straightforward since all combinations of the 164 genes deleted in the original chromothriptic HSC must be considered as potential candidates, a puzzle that cannot be solved in the patient. Defining the relevant gene(s) is of particular importance for strategies to cure WHIM syndrome but also has general importance because this information may provide a target
for development of interventions to facilitate HSC engraftment after any transplantation application, with the goal of limiting or even eliminating the toxicity of current conditioning methods.

The challenge to identifying the relevant engraftment-enhancing gene(s) lies not only in the large number of genetic possibilities but also in the fact that the chromothriptic event occurred in vivo presumably in an HSC niche, a setting that we cannot precisely mimic experimentally. Nevertheless, any translational engraftment-enhancing method based on the underlying selective mechanism in WHIM-09 would involve transplantation, so we focused initially on mouse models of transplantation. We tested the most parsimonious hypothesis, that deletion of the WHIM allele by itself is sufficient to explain the selective advantage of the chromothriptic HSC and the complete myeloid conversion in WHIM-09. This hypothesis is consistent with the known anti-proliferative effect of CXCR4 signaling in HSCs (5), and indeed we previously demonstrated that Cxcr4+/o HSCs are hyperproliferative in vivo relative to Cxcr4+/w HSCs when competitively transplanted into lethally irradiated mice (19). Our previously published (15, 19) and current results show that both in lethally irradiated wild type recipient mice and non-conditioned unmanipulated Cxcr4+/w WHIM model recipient mice, Cxcr4 haploinsufficiency dramatically enhances hematopoietic reconstitution of the B cell and myeloid lineages. Remarkably, in irradiated recipients as few as 1% Cxcr4+/o donor HSCs (~5 HSC transferred per mouse) could successfully compete against 99% Cxcr4+/w donor HSCs to establish >20% long-term Cxcr4+/o HSC and HPC chimerism in BM and >50% stable Cxcr4+/o myeloid and B cell chimerism in the blood. As expected, as the ratio of donor Cxcr4+/o bone marrow cells increases, the proportion of Cxcr4+/o hematopoiesis increases. Since the Cxcr4+/w genotype favors skewing of mature hematopoietic cells to the bone marrow (the key feature of myelokathexis) and the Cxcr4+/o does
not, the total number of cells in the circulation for each subset should decline as the donor Cxcr4+/o : Cxcr4+/o dose ratio increases, which is what we observed. While the potency of Cxcr4 haploinsufficiency revealed by these new graded dose-response studies is impressive, the results in transplanted lethally irradiated recipient mice clearly indicate that a single Cxcr4+/o HSC is unable to compete with a great excess of Cxcr4+/w HSCs to fully repopulate both the HSC pool and the entire myeloid lineage, which is what happened in WHIM-09 (15). Previously, we performed Cxcr4+/o BM transplantation in completely unconditioned Cxcr4+/w recipient mice, an experiment in which the donor Cxcr4+/o HSCs, but not the recipient mice and recipient Cxcr4+/w HSCs, were manipulated ex vivo, and demonstrated that Cxcr4+/o BM donor cells (~5000 donor Cxcr4+/o HSCs) could establish ~70% donor myeloid and B cell chimerism in the blood (19). This was enough chimerism to correct leukopenia in Cxcr4+/w mice but also could not repopulate the entire recipient Cxcr4+/w myeloid lineage, suggesting that the Cxcr4+/o HSCs transplanted in this experiment are less potent than the chromothriptic HSC in WHIM-09.

To obviate potential artifacts of the transplantation process on engraftment, we surgically created parabiotic Cxcr4+/o/Cxcr4+/w mouse pairs. In this parabiotic system, hematopoietic cells including HSCs and HPCs in the peripheral blood could migrate to each parabiont, and some of the HSCs and HPCs in the peripheral blood could home to BM of the opposite parabiont resulting in HSC chimerism without any in vitro manipulation. The results showed durable 80-100% Cxcr4+/o myeloid and B and T cell chimerism in the blood and 15% Cxcr4+/o HSC chimerism in BM of the Cxcr4+/w parabiont, which was maintained for 5 months after separation from the Cxcr4+/o parabiont when the experiment was terminated. This was much higher chimerism than that observed in the Cxcr4+/o parabiont partner and the wild-type parabiotic control pairs. However, this experiment also indicated that Cxcr4-haploinsufficiency alone is not sufficient to
replace the entire population of Cxcr4+/w HSCs and entire Cxcr4+/w myeloid lineage. A possible difference between this experiment and the chromothriptic HSC in WHIM-09, other than the additional 163 genes deleted in WHIM-09, could be related to different environmental conditions for the Cxcr4+/o HSCs in the BM of the Cxcr4+/w parabiont which must have transited through the blood from the Cxcr4+/o parabiont, relative to the chromothriptic HSC in WHIM-09, which may have been resident in the BM. In this regard, it has been reported that HSCs in the circulation are in general less totipotent than those in the BM (22, 24, 25).

We previously suggested that hematopoietic reconstitution may be enhanced by Cxcr4 haploinsufficiency in donor HSCs because of two mechanisms: enhanced proliferation of HSCs and enhanced release of mature leukocytes from BM. The former mechanism is most evident in the lethally irradiated recipient model, where there is maximum availability of stem cell niches and a massive excess of Cxcr4+/o HSCs over Cxcr4+/w HSCs in BM at equilibrium, overcoming a BM homing advantage conferred by the WHIM allele early on. In contrast, in unconditioned Cxcr4+/w recipient mice and parabiotic mice, niche availability is minimal, which may explain why the frequencies of Cxcr4+/o and Cxcr4+/w HSCs in BM at equilibrium were much less skewed in the parabiosis experiments and not different in the unconditioned Cxcr4+/w recipients. Thus, in these two models, Cxcr4+/o enhancement of hematopoietic reconstitution in the blood appears to be driven by enhanced release of Cxcr4+/o mature leukocytes from BM, probably supported by enhanced proliferation of engrafted Cxcr4+/o HSCs. In the two transplantation models, a large number of donor HSCs were transplanted all at once, whereas in parabiosis, the number of the donor HSCs is limited by the frequency of HSCs in the blood (about 1 in every 100,000 nucleated blood cells (26)) and the transplantation process continues until the two parabionts are separated. Thus, in this model Cxcr4+/o enhancement of hematopoietic reconstitution in the blood may be
driven more by enhanced mature leukocyte release from BM with a contribution from enhanced HSC engraftment in BM. Taken together, our results suggest that the increased proliferative potential of Cxcr4+/o HSCs over Cxcr4+/w HSCs may only become manifest when stem cell niches are present in great excess over available donor HSCs or when it is needed for an immune response. In contrast, the foundational chromothriptic HSC in WHIM-09 was able to replace the entire WHIM HSC population.

In conclusion, our study of WHIM-09 and efforts to model her cure in mice suggest that other factors besides chromothriptic deletion of the WHIM allele may have played a significant role in conversion of the myeloid lineage. Future work will be needed to prioritize the 163 other genes for analysis of effects of haploinsufficiency on hematopoietic reconstitution activity.

Although complete Cxcr4 deficiency is embryonic lethal in mice (27), and CXCR4 gain-of-function mutations cause WHIM syndrome and are associated with worse outcomes and poorer responses to therapy in patients with Waldenstrom’s Macroglobulinemia (28), the only phenotype we have identified to date that defines Cxcr4 hemizygosity as a haploinsufficient state is enhanced hematopoietic reconstitution activity in the context of transplantation. Importantly, our combined results in all three competitive transplantation models show that even low-level Cxcr4+/o HSC and HPC engraftment is sufficient to correct leukopenia in WHIM model mice, suggesting that to correct leukopenia and cure WHIM patients only low-level engraftment of donor CXCR4 haploinsufficient HSCs may be necessary. Further, our parabiosis experiments indicated that Cxcr4+/o HSCs from the circulation also have enhanced hematopoietic reconstitution activity in blood after homing to the BM of the WHIM recipient, suggesting the feasibility of gene therapy for WHIM syndrome by deletion of the CXCR4 disease allele in HSCs collected from the blood of WHIM patients. Efforts to test this concept using gene editing are currently underway in our
laboratory. More broadly, the results also imply that CXCR4 knockdown may be a useful general genetic adjuvant strategy to enhance engraftment of donor HSCs and spare patients from genotoxic conditioning in the setting of syngeneic and possibly allogeneic transplantation.
METHODS

Mice
Mouse strains Cxcr4+/o on CD45.1 or CD45.2 or a heterozygous CD45.1/CD45.2 background, and Cxcr4+/w on CD45.2 or a heterozygous CD45.1/CD45.2 background have been previously described (15, 19). All the wildtype mice used were the litter-mates of Cxcr4+/o or Cxcr4+/w mice. The mice were kept in a specific pathogen free facility at NIH and were 6-8 weeks old at the time of transplantation.

Transplantation Experiments
Total BM cells (from a single donor strain or as a mixture of cells from two donor strains) were transferred via tail vein injection into recipient mice that had undergone lethal irradiation (900 rads) 8 hours prior to transplantation. The donor ratio for mixed BM cells in each competitive transplantation was determined by flow cytometric analysis of CD45 markers. In each experiment, donors and recipients were sex-matched. Results were similar in both male and female mice. Irradiated mice were given water containing neomycin for four weeks post irradiation to reduce mortality from infection.

Parabiosis
Each pair of female mice was cohoused before surgery for 2 wks beginning at 6 weeks of age. Pairs were surgically connected as previously described (23, 29). The mice were anesthetized to full muscle relaxation with xylazine and ketamine HCl (1.67 mg per 10 g of body weight) by intraperitoneal injection. After shaving the corresponding lateral aspects of each mouse, matching skin incisions were made, starting 0.5 cm distal to the elbow to 0.5 cm below (distal to) the knee joint, and the subcutaneous fascia was bluntly dissected to create about 0.5 to 1.0 cm of free skin. The olecranon and knee joints were connected by a single 2-0 silk suture and tie, and the skin of the two animals was connected with a continuous absorbable 5-0 Vicryl suture material. The
parabiotic mice were treated with Trimethoprim/Sulfamethoxazole-containing drinking water for at least 10 days post-surgery to prevent bacterial infections and given ibuprofen-containing drinking water for 48 hours to limit post-surgical pain. In some experiments, parabiotic mice were surgically separated by a reversal of the above procedure.

To track leukocytes, each parabiont in each parabiotic pair has a different CD45 background (CD45.2 or CD45.1/2). To exclude possible effects of CD45 background on the parabiosis results, we switched the CD45.2 and CD45.1/2 background in \( Cxcr4^{+/o} \), \( Cxcr4^{+/w} \) and \( Cxcr4^{+/+} \) mice in a separate experiment, which showed similar results. Therefore, we combined the experimental results in Figure 2.

**Blood cell counts**

Blood was collected from mandibular veins of recipient mice using EDTA as anticoagulant (Becton Dickinson, Franklin Lakes, NJ). Total leukocyte counts were measured with a Cellometer Auto 2000 Cell Viability Counter (Nexcelom Bioscience, Lawrence, MA). Absolute leukocyte subset counts, and frequencies were quantitated by flow cytometry.

**Flow cytometry analysis**

For leukocyte subset analysis of the cells from blood and BM, 100 µl of mouse blood was collected, and incubated for 10 min with 2 µl of Fc block (BioLegend, San Diego, CA) prior to incubation with specific antibodies on ice for 30 min. The monoclonal Abs used were CD45.1-PECy7 and CD45.2-eFlour450 (catalog #25-0453-82 and #48-0454-82, eBioscience, San Diego, CA), and Ly6G-APCCy7, CD11b-PerCP-Cy5.5, CD19-FITC, and CD3-APC (catalog #127624, #101228, #115506, and #100312, Biolegend). Erythrocytes were lysed with 2 ml ACK lysis buffer (Quality Biologicals, Inc., Gaithersburg, MD) for 2 min at room temperature before centrifugation. Cells were then washed twice with FACS buffer, fixed with 1% paraformaldehyde, stored on ice,
and analyzed using an LSRII FACS cytometer (BD Biosciences, San Jose, CA) and FlowJo software (TreeStar Inc., Ashland, OR). A similar procedure was followed for staining BM cells isolated from femurs and tibiae. For HSC and HPC analysis, BM cells were isolated from femurs and tibiae. 8x10^6 cells in 80 µl of FACS buffer were incubated for 20 min with biotinylated anti-CD34 mAb (clone RAM from eBioscience), washed once and resuspended in 70 µl of FACS buffer. Then, the cells were stained with Abs Lin-FITC (FITC-conjugated antibodies against lineage markers, including B220, CD3, CD4, CD8, CD11b, Gr1, and Ter119), Sca1-APC-Cy7, c-Kit-APC, Flt3-PE-Cy5, IL7ra-Violet 605, and streptavidin-PE (catalog #133302, #108126, #105812, #135312, #135025, and #405203, Biolegend). The HSCs and HPCs were defined as: long-term HSC (LT-HSC: Lin^−^Sca1^+^c-Kit^+^CD34^−^Flt3^−^), short-term HSC (ST-HSC: Lin^−^Sca1^+^c-Kit^+^CD34^+^Flt3^+^), common lymphoid progenitors (CLP: IL7ra^+^ Lin^−^ Sca1^low^ c-Kit^low^), multipotent progenitors (MPP: Lin^−^Sca1^+^c-Kit^+^CD34^+^Flt3^+^), and LSK (Lin^−^Sca1^+^c-Kit^+^).

Statistical analysis

Data are expressed as means ± SEM. Two-tailed student’s t test for single comparisons and 2-way ANOVA for multiple comparisons were used. P less than 0.05 was considered significant.

Study approval

All animal experiments were performed using an NIAID Animal Care and Use Committee-approved protocol in approved and certified facilities.
AUTHOR CONTRIBUTIONS

Experimental design was provided by J.G., D.H.M. and P.M.M. Generation of experimental data was provided by J.G., A.A., A.P., K.B., E.Y., M.S., A.Y., and Q.L. with supervision and analysis by J.G. and P.M.M. J.G. and P.M.M. were principally responsible for writing the manuscript.
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Figure 1. Cxcr4+/o donor BM cells have a strong selective advantage over Cxcr4+/w donor BM cells for reconstituting myeloid and B cells in the blood of lethally irradiated mice. (A) Experimental design. Cxcr4 genotypes and congenic CD45 isoforms are indicated above and below the mouse cartoons, respectively. A total of 5 million total BM cells were transplanted at different donor cell ratios, indicated as the % Cxcr4+/o input cells to the right of panels B, C and D. (B) Time course of hematopoietic reconstitution. Each graph corresponds to the immunophenotypically defined blood cell subset indicated at the top of the column in which it is
found. Top row of graphs: $C_{xcr4}^{+/o}$ donor-derived blood cell chimerism, presented as the percentage (mean ± SEM) of the total cell number for each subset. Bottom row of graphs: total absolute blood counts for each leukocyte subset, presented as mean ± SEM. In the bottom row of graphs, solid lines and dashed lines indicate average values of blood counts for each subset in unmanipulated control $C_{xcr4}^{+/+}$ (n = 58) and $C_{xcr4}^{+/w}$ (n = 38) mice in our colony. (C and D) Chimerism in BM versus blood for $C_{xcr4}^{+/o}$ donor-derived mature leukocytes (C) and HSCs and HPCs in BM (D). The blood data are the same as at day 256 for the top graphs in panel B, the day when the experiment was terminated for BM analysis. For all conditions, n was at least 5 mice per group. SEM was less than 5% of the mean at all time points lacking visible error bars. The experiment was repeated once with a similar pattern. In panel B, p-values are color-coded to refer to the corresponding color-coded $C_{xcr4}^{+/o}$ input % result compared to the 1% $C_{xcr4}^{+/o}$ input cell result. In panels C and D, * p<0.05; ** p<0.01; *** p<0.005. BM, bone marrow; LSK, lineage-Sca-1⁺c-Kit⁺ bone marrow cells; LT-and ST-HSC, long-term and short-term hematopoietic stem cells; MPP, multipotential progenitor cells; CLP, common lymphoid precursor cells. Single comparisons, student’s t test; multiple comparisons, 2-way ANOVA.
Figure 2. Durable low level Cxcr4+/0 HSC chimerism in BM supports a high level of blood leukocyte chimerism in Cxcr4+/w mice. (A) Experimental design. Genotype code: +/0: Cxcr4+/0, +/w: Cxcr4+/w, and +/-: Cxcr4+/+. To track leukocytes, each parabiont has a different congenic CD45 background from its partner (CD45.2 or CD45.1/2). (B-D) Mature leukocyte chimerism analysis in blood and BM. Leukocyte subsets are defined by immunophenotype at the top of each column of panels. Symbols for the genotypes of each parabiont are defined at the right of each row. Upper row of graphs in each panel: data from Cxcr4+/+ /Cxcr4+/+ parabiosis; lower row of graphs in each panel: data from Cxcr4+/0 /Cxcr4+/w parabiosis. (B) Time course in blood. Vertical dashed lines indicate the time when the parabiotic mice were separated. (C and D) BM chimerism 5 months after parabiotic surgery, the time of separation of the parabionts (C) or 5 months after separation of the parabionts (D). Blood chimerism at the same two time points is reproduced in panels C and D from panel B for direct comparison. (E) HSC and HPC analysis in BM. The time of BM analysis is indicated at the top, genotypes of parabiotic pairs are indicated at the bottom, and the symbols for each parabiont genotype are defined at the upper left within each panel. LSK, lineage-Sca-1+c-Kit+ bone marrow cells; LT-and ST-HSC, long-term and short-term hematopoietic stem cells; MPP, multipotential progenitor cells; CLP, common lymphoid precursor cells. (B-E) Data are the summary of all parabiotic pairs created and are presented as the mean ± SEM % of partner cells in each parabiont for each subset. In the connected phase, a total of 5 Cxcr4+/0:Cxcr4+/w pairs and 4 Cxcr4+/+:Cxcr4+/+ pairs were studied. Of these, three pairs were terminated in each group for BM analysis at the end of the connected phase, leaving two Cxcr4+/0:Cxcr4+/w pairs and one Cxcr4+/+:Cxcr4+/+ pair for study in the separated phase. Cxcr4+/0:Cxcr4+/w data are combined from parabiotic pairs that were differentially marked with CD45 isoforms both possible ways (i.e. Cxcr4+/0 CD45.1/2 : Cxcr4+/w
CD45.2; and Cxcr4+/o CD45.2 : Cxcr4+/w CD45.1/.2), since the results were the same indicating no effect of CD45 polymorphism. * p<0.05; ** p<0.01; *** p<0.005. Single comparisons, student’s t test; multiple comparisons, 2-way ANOVA.