NUP98-HOXA10hd fusion protein sustains multi-lineage haematopoiesis of lineage-committed progenitors in transplant setting

Yong Dong1,2,3,4 | Kaitao Wang1 | Qitong Weng1,2,3 | Tongjie Wang1,2 | Peiqing Zhou1,2,3 | Xiaofei Liu1,2 | Yang Geng1,2 | Lijuan Liu1,2 | Hongling Wu1,2 | Jinyong Wang1,2,3,5,6 | Juan Du1,2,3,5,6

1CAS Key Laboratory of Regenerative Biology, Joint School of Life Sciences, Guangzhou Institutes of Biomedicine and Health, Chinese Academy of Sciences, Guangzhou Medical University, Guangzhou, China
2Guangdong Provincial Key Laboratory of Stem Cell and Regenerative Medicine, Guangzhou Institutes of Biomedicine and Health, Guangzhou, China
3University of Chinese Academy of Sciences, Beijing, China
4Institute of Blood Transfusion, Chinese Academy of Medical Sciences & Peking Union Medical College (CAMS & PUMC), Chengdu, China
5Guangzhou Regenerative Medicine and Health Guangdong Laboratory, Guangzhou, China
6Institute for Stem Cell and Regeneration, Chinese Academy of Sciences, Beijing, China

Abstract

Objectives: Exploring approaches of extending the haematopoiesis time window of MPPs and lineage-committed progenitors might produce promising therapeutic effects. NUP98-HOXA10hd (NA) fusion protein can expand long-term haematopoietic stem cells (HSCs) and promote engraftment competitiveness without causing obvious oncogenesis. Our objectives were to investigate the roles of NA fusion protein in MPP and downstream lineage-committed progenitor context.

Material and Methods: 300 sorted MPPs (Lin−CD48−c-kit+Sca1+CD135−CD150+) were mixed with 5 × 10^5 total BM helper/competitor cells and injected into irradiated recipients. For secondary transplantation, 5 × 10^6 total BM cells from primary recipient mice were injected into lethally irradiated recipients. NA-MPP recipient mice were sacrificed for flow cytometric analysis of bone marrow progenitors at indicated time points. Sorted MPPs and myeloid progenitors were used for RNA-seq library preparation.

Results: We showed that NA-expressing MPPs achieved significantly longer multi-lineage haematopoiesis (>44-week) than natural MPPs (20-week). NA upregulated essential genes regulating long-term haematopoiesis, cell cycle, epigenetic regulation and responses to stress in MPPs. These molecular traits are associated with the earlier appearance of a Sca1+c-kit+ myeloid progenitor population, and more abundant...
1 | INTRODUCTION

Allogeneic stem cell transplantation is widely used for blood disorders, such as leukaemia, myeloproliferative diseases, anaemia. However, lack of timely available HLA matched-donor hindered the application of allogeneic transplantation in clinical applications. Cord blood banks hold the potential to serve as the cell source for transplantation, while the insufficient haematopoietic stem cell number in one unit of cold blood become the main obstacle for the application of cord blood. One solution is to expand haematopoietic stem cell in vitro prior to transplantation. Great efforts have been made to achieve haematopoietic stem cell (HSC) expansion, including the use of small molecules, such as SR1, UM171, UM729, as well as the introduction of intrinsic regulators into HSC-enriched stem cell population, HOXB4 for example. However, the clinical efficacy of these approaches remains unknown and need further investigations. Another possible solution is to enhance self-renewal potential of progenitors or generate abundant progenitor cells to reconstitute haematopoiesis. If this progenitor population could successfully sustain multi-lineage haematopoiesis for a prolonged time period, infusion of enhanced progenitor cells regularly could be an alternative option when allogenic donor is unavailable. Until now, limited but encouraging success has been achieved in this aspect. Ectopic expression of transcription factor Sox17 via retroviruses has been demonstrated to increase the self-renewal potential of multipotent progenitors (MPPs) and therefore conferred on MPPs the potential for long-term multi-lineage reconstitution. Another study has successfully conferred long-term repopulating ability on MPPs with a single miRNA, miR-125a. The enforced expression of miR-125a endowed MPPs with enhanced self-renewal potential, resulting in robust long-term multi-lineage repopulation.

NUP98-HOXA10hd fusion gene (NA) has been shown to be potent for haematopoietic stem cells to expand, survive under stress and promote engraftment competitiveness. However, whether the ectopic expression of NA in MPPs and lineage-committed progenitors could confer long-term multi-lineage haematopoiesis remains unknown. In this study, we explored the roles of NA fusion protein in MPPs and their downstream lineage-committed progenitor context. We used the MPPs sorted from NA compound mice for transplantation assays, which enable the stable overexpression of Nup98-Hoxa10 fusion gene from Rosa26 locus instead of retroviral transduction to avoid risk of insertional mutagenesis. Here, we found that ectopic expression of NA fusion protein in MPPs conferred long-term multi-lineage haematopoiesis in recipient mice, offering promising means to involve MPPs to augment cell source in clinical transplantation settings.

2 | MATERIAL AND METHODS

2.1 | Mice

NA^LSL/+ mice were generated by targeting a mouse ES line (C57BL/6 line, CD45.2^+, Beijing Biocytogen Co., Ltd.) through homologous recombination as previously described. C57BL/6 (CD45.2, CD45.1) and Vav-Cre strain (CD45.2) mice were purchased from the Jackson Laboratory. NA^LSL/+ mice were subsequently bred with Vav-Cre mice to generate NA^LSL/+; Vav-Cre (NA) mice. Mice were housed in the SPF grade animal facility of the Guangzhou Institute of Biomedicine and Health, Chinese Academy of Science (GIBH, CAS, China). All animal experiments were approved by the Institutional Animal Care and Use Committee of Guangzhou Institutes of Biomedicine and Health (IACUC-GIBH).

2.2 | Transplantation

Adult C57BL/6 recipient mice (CD45.2^+, 8-10 weeks old) were irradiated with 2 doses of 4.75 Gy (RS 2000, Rad Source) for a 4-hour interval. For MPP transplantation assay, 300 sorted MPP cells (Lin^−CD48^−c-kit^Sca1^CD135^−CD150^) were mixed with 5 × 10^5 total BM helper/competitor cells or Sca1^− BM helper cells and subsequently injected into the retro-orbital vein of the irradiated recipients. The transplanted mice were maintained on trimethoprim-sulphamethoxazole-treated water for 2 weeks. After transplantation, peripheral blood was obtained from the retro-orbital vein regularly for flow cytometric analysis. For secondary transplantation, 5 × 10^6 total BM cells from primary recipient mice were injected into the retro-orbital vein of the lethally irradiated recipients.

2.3 | Flow cytometry analysis

Antibodies for haematopoietic lineage and haematopoietic progenitors or stem cells analysis: CD45.2(104), CD2 (RM2-5), CD3e (145-2C11), CD4 (RM4-5), CD8a (53-6.7), Ter119 (TER-119), Mac1...
Genes expression and differential expression genes (DEGs) analysis was performed by RSEM and DESeq2. GO enrichment analysis was performed with clusterProfiler package. Gene set enrichment analysis (GSEA) was performed as described. Heatmaps were plotted using gplots (heatmap.2) and ggplot2 package.

2.5 | Statistical analysis

Statistical analysis was performed with SPSS (SPSS v.23, IBM Corp.). Normal distribution of data was tested with SPSS applying Shapiro-Wilk normality test. The data were represented as mean ± SEM. Two-tailed independent Student’s t tests were performed for comparison of two groups of data. *P values <.05 were considered statistically significant (**P < .01 and ***P < .001).

3 | RESULTS

3.1 | NA-overexpressing MPPs sustain long-term multi-lineage haematopoiesis in primary mice

We have previously established NA-overexpression mouse model and found that ectopic expression of NA confers engraftment competitiveness of LT-HSCs in competitive transplantation assay. We further investigated the cell-context roles of NA in MPPs. We directly sorted 300 MPP (Lin−c-kit− Sca1−) cells (Figure S1) from bone marrow nucleated cells of NA mice, and transplanted them together with 5 × 10^5 total bone marrow helper/competitor cells into lethally irradiated recipient (Figure 1A). Recipient mice were bled regularly to monitor donor contribution and donor-derived lineages. Strikingly, 300 NA MPPs continuously contributed to multi-lineage reconstitution over 44 weeks. Donor contribution started at 41% at 4-week time point, increased gradually and persisted around 60% from week-16 to week-36 and decreased to 40% at 44-week time point. In contrast, donor contribution of WT MPP recipient mice peaked at 35% at 4-week time point, dramatically decreased and became undetectable from 20-week time point (Figure 1B). NA MPPs contributed to myeloid, B and T cell lineages at all time points (Figure 1C-E). Expectedly, non–NA-derived multi-lineage haematopoiesis was observed in recipient mice (Figure S2). Interestingly, when 300 NA MPPs were transplanted with 5 × 10^5 Sca− bone marrow helper cells, we also observed donor-derived long-term multi-lineage reconstitution in recipient mice and the donor contribution started from 83% at week-5 and decreased gradually to 61% at week-40. Moreover, NA MPPs contributed to all lineages including myeloid, B and T cells (Figure S3). Taken together, our data demonstrated that NA-overexpressing MPPs repopulated long-term multi-lineage haematopoiesis both with and without of WT HSC/MPP competitors in primary mice.

2.4 | RNA-Seq and data analysis

MPPs were sorted separately into 200 µL DPBS-BSA buffer (0.5% BSA) using 1.5 mL EP tube. Sequencing libraries of MPPs were generated as previously described for HSC. Briefly, cDNA of sorted 1000 MPP aliquots was generated, amplified and used for sequencing library preparation with illumina Nextera XT DNA Sample Preparation Kit (FC-131-1096). For MP sequencing, 1 × 10^5 myeloid progenitor cells (Lin−c-kit− Sca1−) were sorted as one sample from bone marrow nucleated cells of respective recipient mice. Donor-derived NA MPP cells (Tdtomato+ Sca1−) were sorted from NA MPP recipient mice 4 months post-transplantation, while WT MPP cells (CD45.2+) were obtained from control group received WT MPP transplantation at the same time point. Following RNA extraction with RNeasy micro kit (QIAGEN), total RNA of each sorted myeloid progenitor sample was used for sequencing library preparation with illumina Trueseq RNA Sample Preparation Kit (RS-122-2001). All libraries were sequenced by illumina sequencer NextSeq 500 (illumina). The fastq raw data files were generated using illumina bcl2fastq software and uploaded to Gene Expression Omnibus public database (GSE146781). Alignment, normalization of genes expression and differential expression genes (DEGs) analysis were performed by RSEM and DESeq2. GO enrichment analysis was performed with clusterProfiler package. Gene set enrichment analysis (GSEA) was performed as described. Heatmaps were plotted using gplots (heatmap.2) and ggplot2 package.
RNA-Seq analysis. Differential gene expression analysis identified 2347 differential expression genes (DEGs) in NA MPPs compared with WT MPPs (Table S1). GO enrichment analysis of upregulated DEGs (>2 fold) using the NA MPPs showed that these DEGs were largely involved in the regulation of cell cycle, epigenetic regulation, responses to stress, RNA splicing and regulation of signaling pathways (Figure 2A). Further gene set enrichment analysis (GSEA) showed NA regulated genes were enriched for gene sets of long-term haematopoiesis and cell cycle (Figure 2B). Overexpression of Hoxa9 has been reported to expand HSC compartment and results in lympho-myeloid long-term repopulation; meanwhile, Hoxa9 deficiency impairs the multi-lineage repopulating ability of HSC. Interestingly, NA regulated genes were enriched for Hoxa9 targets and targets_of_Hoxa9_and_Meis1 (Figure 2C). Collectively, our data imply that NA endows MPPs with the capability to support long-term haematopoiesis through regulating gene involving in the
regulation of long-term haematopoiesis, cell cycle regulation, epigenetic regulation and response to stress.

3.3 | NA-MPPs efficiently generated abundant lineage-committed progenitors

To investigate whether NA-MPPs produce abundant lineage-committed progenitors to sustain long-term haematopoiesis, we investigated the bone marrow progenitor components in primary MPP recipient mice. Firstly, we transplanted 300 sorted MPPs and analysed the bone marrow myeloid progenitor compartment at day 16, 22, 28 post-transplantation. Not only we observed donor-derived MPs at an earlier time point (day-16), but also at higher ratios compared with control (Figure 3A). As expected, NA-MPPs produced more myeloid progenitor cells and eventually output much more mature white blood cells which contributed to higher number of bone marrow cellularity at day 16, 22, 28 after MPP transplantation (Figure 3B). Since NA-MPPs achieved long-term multi-lineage haematopoiesis, we further analysed lineage-committed progenitor population including common lymphoid progenitors and myeloid progenitors at week-20. Flow cytometric analysis of bone donor-derived progenitors showed that both percentage and absolute number of common lymphoid progenitors and myeloid progenitors were significantly higher compared with control (Figure 3C,D). To investigate the possibility that NA might induce dedifferentiation of MPP back to LT-HSCs, we analysed the LSK compartment in recipient mice at week-20. Interestingly, no detectable donor-derived LSK population persisted at week-20 (Figure 3E). To further test whether NA-MPPs possess long-term capability of haematopoiesis, we transplanted 5 million total BM cells from primary NA-MPP recipient mice into lethally irradiated secondary recipient mice and bled these mice to monitor NA-MPP-derived multi-lineage haematopoiesis. Interestingly, donor contribution in PB of secondary recipients was 19% at 4 weeks post-transplantation, decreased gradually to 6% at week-20 (Figure 3F). We sacrificed these secondary recipient mice at week-20 to examine the NA MPP-derived donor contribution. Flow cytometric analysis of bone marrow nucleated cells of these secondary recipients revealed less than 0.4% donor contribution (Figure 3G). Taken together, our results suggest that NA MPPs achieve long-term haematopoiesis via generating abundant lineage-committed progenitors rather than acquiring self-renewal capability.

3.4 | NA MPs upregulate genes involving in regulation of phosphorylation and myeloid differentiation

Donor-derived progenitor cells supported haematopoiesis up to 44 weeks while there was no stem cell-enriched LSK population, indicating that these progenitor cells gained new function features. To gain molecular insights of NA-expressing lineage-committed progenitor cells, we investigated the transcriptome patterns of myeloid progenitor cells by RNA-seq analysis. Differential gene expression analysis identified 3051 differential expression genes (DEGs) in NA MPs compared with WT control (Table S2). GO enrichment analysis of upregulated DEGs (>2 fold) using the NA MPs showed that these DEGs were largely involved in the regulation
of haematopoiesis, homoeostasis, phosphorylation in addition to myeloid cell differentiation (Figure 4A). Notably, further gene set enrichment analysis (GSEA) for MPs also identified enrichment of gene sets of Hoxa9 targets as well as other Hox family targets (Figure 4B). Together, these data suggest NA MPs sustain haematopoiesis through actively upregulating genes involving in haematopoiesis, homoeostasis, phosphorylation and myeloid cell differentiation-related pathways.
However, development of myeloproliferative disease/leukaemia in secondary and tertiary recipients was observed. Self-renewal of MPPs by retroviral ectopic expression of miR125a. The term ectopic expression of Sox17 eventually led to leukemogenesis. In addition, the NA MPP primary recipient mice as well as the secondary mice remained leukaemia free, demonstrating no obvious oncogenesis of our approach. This is consistent with the previous report that the restriction of NUP98-fusion to the homeodomain of Hoxa10 blunts leukemogenic potential of Nup98-Hoxa10.17

In our experiment, overexpression of NA in MPPs upregulated genes regulating long-term haematopoiesis, cell cycle, epigenetic regulation, response to stress and RNA splicing. These molecular signatures could contribute to the capability to sustain long-term multi-lineage haematopoiesis of NA MPPs. We have validated the expression of representative genes of leading edge genes for gene sets of cell cycle regulation and targets of Hoxa9 and Meis1 by real-time PCR (Figure S4). Importantly, further investigation is needed to determine whether changes at transcriptome level ultimately confer alterations of cellular characteristics of NA MPPs, which may contribute to their capacity to sustain long-term multi-lineage haematopoiesis. We have examined the cell cycle status of NA MPPs and the results showed NA MPPs was more quiescent compared to control (Figure S5). Further study of other alteration of cellular characteristics of NA MPPs, such as proliferation rate, apoptosis and cell division, would provide more clues on the mechanism how overexpression of NA endows MPPs with the ability to sustain long-term multi-lineage haematopoiesis.

(A) Up-regulated genes in MPs (NA vs WT)

(B) Hoxa9 Target analysis

FIGURE 4 Transcriptome profiling of NA MPs. A, Gene ontology enrichment analysis of upregulated genes in NA MPs. MPs were sorted from primary recipient mice to construct RNA-seq libraries. Genes with at least 2-fold change over WT MPs were included for GO enrichment analysis. B, GSEA analysis of gene sets of Hoxa9 targets, targets_of_Hoxa9_and_meis1, immortalized_by_Hoxa9_and_Meis1, Hoxa5 targets, Hoxa10 targets and myeloid differentiation in transcriptome of NA MPs compared with WT MPs. Gene sets were obtained from the data set of c2.all.v5.2 symbols of the GSEA website (https://gsea-msigdb.org). Selected pathways with significant changes are shown (FDR < 0.25, P < .05).
We found the complete absence of LSK population at 20 weeks post-transplantation in primary mice. Yet, the NA MPP recipient mice could sustain haematopoiesis for another 24 weeks as well as support a low but constant donor-contributed multi-lineage haematopoiesis in secondary recipients. This could be induced by abundant number of lineage-committed progenitors generated by NA MPPs. In order to generate enough lineage-committed progenitors to sustain haematopoiesis, certain number of input NA MPPs are needed. This is supported by our parallel transplantation experiment with 100 NA MPPs and 300 NA MPPs. Transplantation of one hundred NA MPPs was insufficient to maintain steady high level of donor-contributed haematopoiesis within 20 weeks post-transplantation while 300 MPPs achieved steady high level of donor-contributed multi-lineage reconstitution (Figure S6). In addition, NA may also alter the proliferation and survival of lineage-committed. Long-lived progenitor cells have been shown to sustain steady-state haematopoiesis.18 It is possible that NA-overexpressing lineage-committed progenitor cells become long-lived and sustain long-term multi-lineage constitution when stem cell containing LSK compartment exhausts. Transplantation of NA MPPs combined with lineage-tracing strategy will illustrate whether long-lived committed progenitor cells existed and persisted in primary and secondary transplantation.

Notably, Hoxa9 targets appeared to be common gene sets enriched for GSEA analysis both in NA MPPs and MPs, and it was among the 23 overlapped differentially expressed TFs between MPPs and MPs (Figure S7), indicating it may be the key target of NA across various cellular contexts. Overexpression of Hoxa9 expands HSC compartment and results in lympho-myeloid long-term repopulation,15 while the loss of Hoxa9 impairs the multi-lineage repopulating ability of HSC.16 implying Hoxa9 could also be an important player as the downstream target of NA in our case. Transplantation assay with NA-Hoxa9- MPPs will reveal the dependency of NA on Hoxa9 to exert its role to endow MPPs with the capacity to sustain long-term multi-lineage haematopoiesis. Following up screening and functional validation of NA targets in MPPs and MPs could open up the opportunity searching for regulators conferring long-term multi-lineage haematopoiesis capacity on progenitor cells.

In summary, our data show that NA-overexpressing MPPs can support long-term multi-lineage haematopoiesis in primary mice through generating abundant lineage-committed progenitors, which will benefit the application of MPPs in the absence of LT-HSCs to improve clinical outcomes in transplantation settings.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS

YD, KW, QW, TW, PZ, XL, YG, LL and HW performed the experiments and analysed data. JD and JW conceived and supervised the study, wrote the manuscript and approved the final manuscript.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding author upon request. All RNA-Seq data are in the GEO database with accession code GSE146781.

ORCID

Jinyong Wang https://orcid.org/0000-0002-7218-0659
Juan Du https://orcid.org/0000-0003-4701-3573

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

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