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# Motixafortide and G-CSF to mobilize hematopoietic stem cells for autologous transplantation in multiple myeloma: a randomized phase 3 trial

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Autologous hematopoietic stem cell transplantation (ASCT) improves survival in multiple myeloma (MM). However, many individuals are unable to collect optimal CD34<sup>+</sup> hematopoietic stem and progenitor cell (HSPC) numbers with granulocyte colony-stimulating factor (G-CSF) mobilization. Motixafortide is a novel cyclic-peptide CXCR4 inhibitor with extended in vivo activity. The GENESIS trial was a prospective, phase 3, double-blind, placebo-controlled, multicenter study with the objective of assessing the superiority of motixafortide + G-CSF over placebo + G-CSF to mobilize HSPCs for ASCT in MM. The primary endpoint was the proportion of patients collecting  $\ge 6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> within two apheresis procedures; the secondary endpoint was to achieve this goal in one apheresis. A total of 122 adult patients with MM undergoing ASCT were enrolled at 18 sites across five countries and randomized (2:1) to motixafortide + G-CSF or placebo + G-CSF for HSPC mobilization. Motixafortide + G-CSF enabled 92.5% to successfully meet the primary endpoint versus 26.2% with placebo + G-CSF (odds ratio (OR) 53.3, 95% confidence interval (CI) 14.12-201.33, P < 0.0001). Motixafortide + G-CSF also enabled 88.8% to meet the secondary endpoint versus 9.5% with placebo + G-CSF (OR 118.0, 95% CI 25.36-549.35, P < 0.0001). Motixafortide + G-CSF was safe and well tolerated, with the most common treatment-emergent adverse events observed being transient, grade 1/2 injection site reactions (pain, 50%; erythema, 27.5%; pruritis, 21.3%). In conclusion, motixafortide + G-CSF mobilized significantly greater CD34<sup>+</sup> HSPC numbers within two apheresis procedures versus placebo + G-CSF while preferentially mobilizing increased numbers of immunophenotypically and transcriptionally primitive HSPCs. Trial Registration: ClinicalTrials.gov, NCT03246529

Multiple myeloma (MM) is the second most common hematologic malignancy<sup>1</sup>, historically associated with median overall survival (OS) of 24–30 months. However, the development of high-dose chemotherapy and autologous stem cell transplantation (ASCT), immunomodulatory drugs

(IMiDs), proteasome inhibitors (PIs), anti-CD38 monoclonal antibodies (mAbs) and other novel therapies has greatly expanded therapeutic options for newly diagnosed MM. Currently, median OS exceeds 45–82 months with ASCT playing a central role in the treatment paradigm for  $MM^{2,3}$ .

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**Fig. 1** | **GENESIS trial enrollment. a**, A total of 162 patients signed informed consent (IC). Screen failures were due to patients not meeting study eligibility criteria; 124 patients began G-CSF mobilization, two elected to withdraw consent before randomization, leaving a total of 122 patients who were randomized (2:1) to either motixafortide + G-CSF or placebo + G-CSF and were included in the ITT analysis. In the motixafortide + G-CSF arm, one patient did not perform apheresis due to an adverse event unrelated to study drug. In the placebo + G-CSF arm, one patient elected not to undergo apheresis due to personal concerns regarding the COVID-19 pandemic. Both these patients were included as mobilization failures in the ITT analysis (that is, did not meet the primary endpoint) but were not remobilized on study. In total, 98.8% (79 of 80) of patients in the

motixafortide + G-CSF arm and 97.6% (41 of 42) of patients in the placebo + G-CSF arm received all study-related mobilization injections and underwent apheresis on protocol without any events of treatment arm crossover. **b**, Patients were enrolled across 18 centers and five countries, with the majority treated in the United States. **c**, Enrollment numbers are presented by individual study center, grouped by geographic region (United States and Europe) and mobilization regimen (motixafortide + G-CSF or placebo + G-CSF), with the proportion of patients at each site meeting the primary endpoint (collection of  $\geq 6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> within two apheresis days) shown in red and the proportion not meeting the primary endpoint shown in black.

Autologous stem cell transplantation in MM has been shown to improve event-free survival and OS compared with conventional chemotherapy alone in previously untreated standard-risk MM<sup>4,5</sup>. However, the effectiveness of ASCT relies, in part, on the ability to collect sufficient hematopoietic stem and progenitor cells (HSPCs), typically from peripheral blood (PB). A minimum of  $\ge 2 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> are necessary, while infusion of optimal numbers of  $\geq 5-6 \times 10^{6} \text{ CD34}^{+}$  cells kg<sup>-1</sup> is associated with improved engraftment, disease-free survival and OS compared with lower transplant doses<sup>6-8</sup>. Clinically, CD34 expression remains the most common immunophenotypic cell surface marker defining human HSPCs. However, multicolor fluorescence-activated cell sorting (FACS) and single-cell RNA sequencing (scRNA-seq) have illustrated the heterogeneous nature of CD34<sup>+</sup> HSPCs, identifying immunophenotypically and transcriptionally distinct CD34<sup>+</sup> subsets ranging from primitive hematopoietic stem cells (HSCs) capable of long-term self-renewal and multilineage potential to relatively differentiated, lineage-committed progenitors<sup>9,10</sup>.

Granulocyte colony-stimulating factor (G-CSF) is widely considered the standard agent for PB HSPC mobilization. Nevertheless, despite the use of G-CSF in mobilization of HSPCs to the PB and after multiple days of apheresis, 40-50% of patients with MM remain unable to collect optimal numbers of HSPCs for ASCT<sup>11,12</sup>. The addition of chemotherapy to G-CSF may incrementally increase mobilization, but also prolongs HSPC mobilization with multiple apheresis days and chemotherapy-related toxicities<sup>13,14</sup>. Meanwhile, protracted mobilization substantially increases the financial and logistical burden to both patients and the healthcare system<sup>14,15</sup>.

The interaction between CXCL12 and its receptor, CXCR4, is critically involved in the retention of HSPCs within the bone marrow and blockade of CXCR4 mobilizes HSPCs to PB<sup>16</sup>. Previous studies have shown the low-affinity (inhibitory constant (Ki), 652 nM), short-acting CXCR4i, plerixafor + G-CSF enhanced mobilization of CD34<sup>+</sup> HSPCs to PB compared with G-CSF<sup>12,17</sup>. However, despite up to eight injections of G-CSF, four injections of plerixafor and four apheresis procedures, 15–35% of patients remained unable to mobilize optimal HSPC numbers<sup>12,18</sup>. Preclinical and clinical data suggest that CXCR4 expression on CD34<sup>+</sup> HSPC subsets is variable, with relatively lower CXCR4 expression on primitive CD34<sup>+</sup> HSCs and multipotent progenitors (MPPs) compared with higher expression on certain lineage-committed CD34<sup>+</sup> progenitors<sup>19</sup>. These studies suggest that optimization of CXCR4 blockade may increase CD34<sup>+</sup> HSPC mobilization and mobilize differential HSPC subsets<sup>19</sup>.

Motixafortide (BL-8040) is a selective cyclic-peptide inhibitor of CXCR4 with high affinity (Ki, 0.32 nM), long receptor occupancy and extended clinical activity (>48 h)<sup>20-22</sup>. In a phase 1, two-part study (NCT02073019), motixafortide administered to healthy subjects was safe, well tolerated and led to a sustained five- to seven-fold increase in PB CD34<sup>+</sup> HSPCs, enabling collection of a median of 11.2 × 10<sup>6</sup> CD34<sup>+</sup> cells kg<sup>-1</sup> in one leukapheresis. In another, single-arm, open-label, dose-escalation study (NCT01010880), motixafortide administered before ASCT to patients with MM undergoing standard HSPC mobilization was safe, well tolerated and led to significant increases in PB CD34<sup>+</sup> HSPCs.

Therefore, this phase 3 study (GENESIS) was designed to compare the safety and efficacy of motixafortide + G-CSF versus placebo + G-CSF in patients with MM undergoing HSPC mobilization before ASCT (NCT03246529). In addition, immunophenotypic and transcriptional profiling was performed via multicolor FACS and scRNA-seq of CD34<sup>+</sup> HSPCs mobilized on the GENESIS trial, as well as a contemporaneous, prospectively enrolled cohort of demographically similar patients with MM mobilized with plerixafor + G-CSF (protocol no. 201103349) and three cohorts of healthy, allogeneic HSPC donors (allo-donors) undergoing single-agent mobilization with motixafortide, plerixafor or G-CSF alone (NCT02639559, NCT00241358, protocol no. 201106261, respectively).

## Results

**Patient demographics were comparable across study cohorts** From 22 January 2018 to 30 October 2020, a total of 122 patients from 18 sites in five countries were enrolled and randomized 2:1 to receive either motixafortide + G-CSF (80 patients) or placebo + G-CSF (42 patients) for HSPC mobilization (Fig. 1 and Extended Data Fig. 1a). Demographics between the two treatment arms of the GENESIS trial were similar (Table 1). In addition, the cohort of contemporaneous patients with MM enrolled prospectively and mobilized with plerixafor + G-CSF shared demographics comparable to those of patients on the GENESIS trial (Extended Data Fig. 1b and Extended Data Table 1).

**Motixafortide** + G-CSF rapidly mobilized high numbers of HSPCs Enumeration of CD34<sup>+</sup> HSPCs in the apheresis product was performed by both local and central laboratories. According to the prespecified protocol, all clinical decisions were made based on local laboratory results, including the determination that the patient met the collection goal for the primary endpoint and determining the number of CD34<sup>+</sup> cells kg<sup>-1</sup> infused for ASCT. Local and central laboratory assessments were included in the prespecified statistical analysis plan, with statistically significant results observed via both assessments favoring the increased effectiveness of motixafortide + G-CSF compared with placebo + G-CSF (Extended Data Fig. 2).

Mobilization with motixafortide + G-CSF resulted in 92.5% of patients meeting the primary endpoint of collecting  $\ge 6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> within two apheresis days versus 26.2% with placebo + G-CSF, by local laboratory assessment using an intention-to-treat (ITT) analysis (OR 53.3, 95% CI 14.12–201.33, P < 0.0001) (Fig. 2a). Furthermore, 88.8% of patients mobilized with motixafortide + G-CSF met the key prespecified secondary endpoint of collecting  $\ge 6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> in one apheresis day versus 9.5% with placebo + G-CSF (OR 118.0, 95% CI 25.36–549.35, P < 0.0001).

The baseline level of PB CD34<sup>+</sup> HSPCs before G-CSF administration was similar between the motixafortide + G-CSF and placebo + G-CSF cohorts, at 1.5 and 1.7 CD34<sup>+</sup> cells  $\mu$ l<sup>-1</sup>, respectively (Fig. 2b). Following four doses of G-CSF and before motixafortide or placebo administration, PB CD34<sup>+</sup> counts remained similar between the motixafortide + G-CSF and placebo + G-CSF cohorts, at 15.8 and 12.0 CD34<sup>+</sup> cells  $\mu$ l<sup>-1</sup>, respectively. However, following administration of either motixafortide or placebo but before apheresis, the median number of PB CD34<sup>+</sup> HSPCs in the motixafortide + G-CSF cohort (*n* = 74) was 116.0 versus 19.0 cells  $\mu$ l<sup>-1</sup> in the placebo + G-CSF cohort (*n* = 40) (*P* < 0.001), with PB CD34<sup>+</sup> cells  $\mu$ l<sup>-1</sup> correlating well with apheresis yields.

Meanwhile, 96.3% of patients mobilized with motixafortide + G-CSF collected  $\ge 2 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> within one apheresis

	Motixafortide + G-CSF (n=80)	Placebo + G-CSF (n=42)	
Median age (±s.d.), years	63.5 (9.4)	62.0 (9.6)	
Male sex, n (%)	55 (68.8)	24 (57.1)	
Race/ethnicity, n (%)			
African	9 (11.3)	2 (4.8)	
Asian	2 (2.5)	0(0)	
Caucasian	65 (81.3)	40 (95.2)	
Hispanic/Latino	1 (1.3)	0(0)	
Other/NOS	3 (3.8)	0(0)	
Median time from diagnosis to consent, months	4.0	4.5	
IMWG response at screening, n (%)			
sCR	4 (5.0)	2 (4.8)	
CR	12 (15.0)	7 (16.7)	
VGPR	33 (41.3)	23 (54.8)	
PR	31 (38.8)	10 (23.8)	
Median no. of induction cycles (±s.d.)	4 (0.9)	4 (0.8)	
Lenalidomide-containing induction, <i>n</i> (%)	57 (71.3)	28 (66.7)	
Anti-CD38 antibody-containing induction, n (%)	0(0)	1 (2.4)	
Previous radiotherapy, n (%)	9 (11.3)	4 (9.5)	

Similar demographics were observed for patients treated with motixafortide+G-CSF (n=80) compared with placebo+G-CSF (n=42). IWMG, International Myeloma Working Group. NOS, not otherwise specified; sCR, stringent complete remission;VGPR, very good partial remission.

day versus 64.3% with placebo + G-CSF (OR 18.9, 95% CI 4.47-80.04, P < 0.0001). The median number of CD34<sup>+</sup> HSPCs mobilized in one apheresis day with motixafortide + G-CSF was  $10.8 \times 10^6$  versus  $2.25 \times 10^{6}$  cells kg<sup>-1</sup> with placebo + G-CSF. The total number of CD34<sup>+</sup> HSPCs infused for ASCT was determined independently by each investigator according to local practice, with a median of  $<6 \times 10^6$  cells kg<sup>-1</sup> infused for ASCT in both arms (minimum  $2 \times 10^{6}$  CD34<sup>+</sup> cells kg<sup>-1</sup> required according to the protocol). Median time to neutrophil engraftment (TNE) was 12 days in both arms (hazard ratio (HR) not estimable, P = 0.9554). Median time to platelet engraftment (TPE) was 18 days with motixafortide + G-CSF and 17 days with placebo + G-CSF (HR 0.95, 95% CI 0.2–5.7, P = 0.9554). Graft durability at day 100 post ASCT was 92.2% in the motixafortide + G-CSF arm and 91.9% in the placebo + G-CSF arm (OR 1.04, 95% CI 0.2-4.5, P = 0.96). Progression-free survival (PFS) and overall survival (OS) were comparable between the two cohorts and consistent with contemporary outcomes (Extended Data Table 2).

#### $Motix a fortide + G\text{-}CSF \, was \, safe \, and \, well \, tolerated$

Treatment-emergent AEs (TEAEs) were defined as starting with the first dose of G-CSF through 30 days following the last apheresis procedure or the first dose of conditioning chemotherapy, whichever occurred first. Overall, TEAEs were reported in 93.8% (grade 3, 27.5%) of patients with motixafortide + G-CSF versus 83.3% (grade 3, 4.8%) with placebo + G-CSF (Table 2). The most common TEAEs related to motixafortide were transient, grade 1/2 local injection site reactions and systemic reactions. Local injection site reactions most commonly included pain (50%), erythema (27.5%) and pruritis (21.3%). Systemic reactions most commonly included flushing (32.5%), pruritis (33.8%), urticaria (12.5%) and erythema (12.5%). Meanwhile, bone pain was



**Fig. 2** | **GENESIS trial efficacy. a, b**, Patients with MM were mobilized with either motixafortide + G-CSF or placebo + G-CSF with the goal of collecting  $\ge 6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup>. **a**, The proportion of patients meeting the primary endpoint of collecting to goal within two apheresis days and the key secondary endpoint of collecting to goal in one apheresis day are shown, along with collection rates after three and four apheresis procedures. No patients in the motixafortide + G-CSF arm underwent a fourth apheresis on protocol. The primary endpoint and prespecified secondary efficacy endpoint were analyzed using the CMH test and are presented as OR with two-sided 95% Cls and *P* values. **b**, The numbers of PB CD34<sup>+</sup> cells  $\mu$ l<sup>-1</sup> in each cohort are presented by mobilization day with standard

box and whisker plots, where the measure of center is the median value for PB CD34<sup>+</sup> cells  $\mu$ l<sup>-1</sup> with the exact median value of CD34<sup>+</sup> cells  $\mu$ l<sup>-1</sup> noted below the *x* axis by day and cohort; the box represents the interquartile range and lines represent the minimum/maximum (±1.5 × interquartile range). Each patient contributed a single biologically independent sample examined over one independent experiment per sample/patient. Total sample counts associated with motixafortide + G-CSF were: day 0 (*n* = 80), day 4 (*n* = 80), day 5 (*n* = 80), day 6 (*n* = 11), day 7 (*n* = 6) and day 8 (*n* = 0). Total sample counts associated with placebo + G-CSF were: day 0 (*n* = 39), day 4 (*n* = 40), day 5 (*n* = 39), day 6 (*n* = 32), day 7 (*n* = 27) and day 8 (*n* = 17).

commonly observed in the placebo + G-CSF arm (31.0%). No grade 4 TEAEs or deaths occurred during the mobilization period of the study.

**Motixafortide + G-CSF reduced healthcare resource utilization** As a prespecified analysis, healthcare resource utilization was assessed comparing motixafortide + G-CSF with placebo + G-CSF. Patients mobilized with motixafortide + G-CSF received 5.26 G-CSF injections per patient versus 8.12 in the placebo + G-CSF cohort (P < 0.0001). Additionally, patients mobilized with motixafortide + G-CSF required an average of 1.23 apheresis procedures per patient to mobilize optimal CD34<sup>+</sup> HSPCs versus 3.24 with placebo + G-CSF (P < 0.0001). No patients mobilized with motixafortide + G-CSF underwent a fourth day of apheresis for primary mobilization and only 1% (n = 1) required remobilization, whereas 63.5% of patients mobilized with placebo + G-CSF required a fourth day of apheresis and 23.8% (n = 10) required remobilization with plerixafor + G-CSF.

# Motixafortide + G-CSF mobilized high numbers of primitive HSCs

Extended immunophenotyping via multicolor FACS of CD34<sup>+</sup> HSPCs from the day 1 apheresis product of patients (n = 51) mobilized with placebo + G-CSF (n = 13), plerixafor + G-CSF (n = 14) and motixafortide + G-CSF (n = 24) as a prespecified correlative analysis demonstrated nine distinct CD34<sup>+</sup> HSPC subsets, ranging from primitive HSCs to lineage-committed progenitors (Fig. 3a and Extended Data Fig. 3). Compared with placebo + G-CSF, motixafortide + G-CSF significantly increased percentages of common lymphoid progenitors (CLPs), natural killer cell precursors (NKPs) and basophil precursors (BPs) (Fig. 3b). When compared with plerixafor + G-CSF, motixafortide + G-CSF significantly increased percentages of multipotent progenitors and common myeloid progenitors (MPPs/CMPs), NKPs and BPs, with fewer lymphomyeloid primed progenitors (LMPPs/CLPs) (Fig. 3b). Quantitation of absolute numbers of HSPC subset yields demonstrated significantly increased quantities of eight out of nine HSPC subsets in the motixafortide + G-CSF products versus placebo + G-CSF, with 10.5-fold higher absolute numbers of primitive HSCs (Fig. 3c). Compared with plerixafor + G-CSF, motixafortide + G-CSF significantly increased numbers of MPPs/CMPs, CLPs and BPs (Fig. 3c). Taken together, these data suggest that motixafortide induced pan-mobilization of multiple HSPC subsets capable of broad multilineage hematopoietic reconstitution, with notable increases in the absolute number of primitive HSCs and MPPs/CMPs.

CD34<sup>+</sup> HSPCs from apheresis were split and stained with two different antibodies to CD184 (CXCR4), clones 12G5 and 1D9. The 12G5 antibody recognizes an epitope involving the first and second extracellular domains of CXCR4 and competes with motixafortide and plerixafor for CXCR4 binding<sup>23,24</sup>. In contrast, the 1D9 antibody binds the N terminus of CXCR4 and is unaffected by motixafortide or plerixafor bound to CXCR4 (ref. 24). Binding of 1D9 to CD34<sup>+</sup> HSPCs was similar among all three arms (P = 0.45 - 0.75). In contrast, both the percentage of 12G5<sup>+</sup> cells and total 12G5 antibody-binding capacity were significantly lower in the motixafortide + G-CSF cohort versus placebo + G-CSF (P < 0.0001) and plerixafor + G-CSF (P < 0.0001) for all HSPC subsets, consistent with extended CXCR4 occupancy by motixafortide (Extended Data Fig. 4). Despite extended CXCR4 occupancy by motixafortide, there was no impact on rehoming of HSPCs to the bone marrow with comparable engraftment kinetics and graft durability (Extended Data Table 2). These findings are consistent with previously reported data, and may be due to inhibition by motixafortide of CXCL12-induced CXCR4 internalization in a dose-dependent manner, leading to extended CXCR4 half-life on the cell surface and ultimately net upregulation of CXCR4 expression<sup>20,25</sup>.

#### HSPC number and subsets infused impact engraftment

The number of CD34<sup>+</sup> cells kg<sup>-1</sup> infused for ASCT was determined independently by each investigator according to local practice, with a median of  $<6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> infused in both cohorts and similar TPE and TNE (Extended Data Table 2).

However, a post hoc pooled analysis of all patients (n = 114) revealed an inverse correlation between increasing levels of CD34<sup>+</sup> cells kg<sup>-1</sup> infused and TPE (R = -0.30, P = 0.00097; Extended Data Fig. 5a). Sensitivity analyses determined that infusion of  $\ge 7 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> was associated with faster median TPE of 14 versus 18 days with  $<7 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> (HR 0.57, 95% CI 0.34–0.94, P = 0.0276). A dose–response relationship was also observed, including (1) TPE of 11 days with  $\ge 8 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> infused versus 18 days with  $<8 \times 10^6$ CD34<sup>+</sup> cells kg<sup>-1</sup> (HR 0.28, 95% CI 0.15–0.53, P = 0.0001) and (2) TPE of 10 days with  $\ge 9 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> infused versus 18 days with  $<9 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> (HR 0.28, 95% CI 0.14–0.57, P = 0.0004). TNE was not impacted by total CD34<sup>+</sup> cells kg<sup>-1</sup> infused (R = -0.13, P = 0.15) (Extended Data Fig. 5b).

#### Table 2 | GENESIS trial safety and toxicity

Motixafortio	de+G-CSF	Placebo+	G-CSF
Any grade	Grade 3	Any grade	Grade 3
93.8 (75 of 80)	27.5 (22 of 80)	83.3 (35 of 42)	4.8 (2 of 42)
reactions, % (n)			
50 (40 of 80)	6.3 (5 of 80)	4.8 (2 of 42)	0
27.5 (22 of 80)	0	0	0
21.3 (17 of 80)	0	0	0
reactions, % (n)			
32.5 (26 of 80)	7.5 (6 of 80)	0	0
33.8 (27 of 80)	11.3 (9 of 80)	0	0
12.5 (10 of 80)	1.3 (1 of 80)	0	0
12.5 (10 of 80)	0	0	0
17.5 (14 of 80)	0	31.0 (13 of 42)	0
17.5 (14 of 80)	0	14.3 (6 of 42)	0
13.8 (11 of 80)	0	11.9 (5 of 42)	0
13.8 (11 of 80)	0	11.9 (5 of 42)	0
11.3 (9 of 80)	0	14.3 (6 of 42)	0
	Motixafortic   Any grade   93.8 (75 of 80)   reactions, % (n)   50 (40 of 80)   27.5 (22 of 80)   21.3 (17 of 80)   reactions, % (n)   32.5 (26 of 80)   33.8 (27 of 80)   12.5 (10 of 80)   12.5 (10 of 80)   17.5 (14 of 80)   13.8 (11 of 80)   13.8 (11 of 80)   13.3 (9 of 80)	MotixafortiJe + G-CSF   Any grade Grade 3   Jas (75 of 80) Z7.5 (22 of 80)   reactions, % (n) Z7.5 (22 of 80)   50 (40 of 80) 6.3 (5 of 80)   27.5 (22 of 80) 0   21.3 (17 of 80) 0   gas (27 of 80) 7.5 (6 of 80)   32.5 (26 of 80) 7.5 (6 of 80)   33.8 (27 of 80) 11.3 (9 of 80)   12.5 (10 of 80) 1.3 (1 of 80)   12.5 (10 of 80) 0   17.5 (14 of 80) 0   13.8 (11 of 80) 0   13.8 (11 of 80) 0   13.3 (9 of 80) 0	MotixafortiJet + G-CSF Placebet + Grade 3   Any grade Grade 3 Any grade   93.8 (75 of 80) 27.5 (22 of 80) 83.3 (35 of 42)   reactions, % (n) : .   50 (40 of 80) 6.3 (5 of 80) 4.8 (2 of 42)   27.5 (22 of 80) 0 0   21.3 (17 of 80) 0 0   21.3 (17 of 80) 0 0   32.5 (26 of 80) 7.5 (6 of 80) 0   32.5 (26 of 80) 11.3 (9 of 80) 0   32.5 (26 of 80) 13.3 (1 of 80) 0   32.5 (10 of 80) 13.3 (1 of 80) 0   12.5 (10 of 80) 0 0   12.5 (10 of 80) 0 0   17.5 (14 of 80) 0 10.13 of 42)   17.5 (14 of 80) 0 14.3 (6 of 42)   13.8 (11 of 80) 0 11.9 (5 of 42)   13.8 (11 of 80) 0 14.3 (6 of 42)

All TEAEs occurring at a frequency of >10% in either motixafortide+G-CSF or placebo+G-CSF during the period from the first dose of G-CSF through 30 days following the last apheresis procedure or the first dose of conditioning chemotherapy, whichever occurred first. No grade 4 TEAEs or deaths events occurred during this period.

In addition, a post hoc pooled analysis of 37 patients (motixafortide + G-CSF, n = 24; placebo + G-CSF, n = 13) with extended CD34<sup>+</sup> immunophenotyping via multicolor FACS revealed that infusion of higher numbers of combined CD34<sup>+</sup> HSC, MPP, CMP and granulocyte monocyte progenitor (GMP) subsets was associated with more rapid TPE (R = -0.49, P = 0.0025) (Extended Data Fig. 5c). Also, infusion of higher numbers of GMPs alone was associated with more rapid TPE (R = -0.57, P = 0.00029) (Extended Data Fig. 5d). Sensitivity analyses determined that infusion of higher numbers (>75th percentile) of GMPs was specifically associated with more rapid TPE of 13 versus 19 days with lower numbers of GMPs (P = 0.0116). TNE was not impacted by specific CD34<sup>+</sup> HSPC subsets infused (all P > 0.05).

#### Motixafortide mobilized transcriptionally primitive HSPCs

Single-cell transcriptional profiling was performed via scRNA-seq on CD34<sup>+</sup> HSPCs from the day 1 apheresis products of patients with MM (n = 12) mobilized with placebo + G-CSF (n = 4), plerixafor + G-CSF (n = 4) and motixafortide + G-CSF (n = 4), along with CD34<sup>+</sup> HSPCs from the apheresis product of healthy allo-donors (n = 6) mobilized with G-CSF alone (n = 2), plerixafor alone (n = 2) and motixafortide alone (n = 2). When compared with the MM cohorts, allo-donor patients were generally younger (median age of MM cohort, 62–68 years versus allo-donor cohort, 55 years) and lacked a history of MM or recent chemotherapy exposure. Otherwise, the cohorts were demographically similar.

A total of 144,982 purified CD34<sup>+</sup> HSPCs were sequenced across all 18 samples (range 2,086–20,062 cells per patient, average 2,767 genes per cell). Uniform manifold approximation and projection (UMAP) clustering was performed, with cell identity determined by cross-referencing of gene expression profiles with previously published datasets and genes reported as reliable markers of lineage commitment<sup>9,26-29</sup>. A total of 20 transcriptionally distinct CD34<sup>+</sup> HSPC subsets were identified, including (1) six transcriptionally unique HSC subclusters (HSC1–6), (2) a large multilineage progenitor (MLP) cluster and (3) ten lineage-biased lymphoid (CLP\_Ly1/2), myeloid (GMP), monocytic/dendritic (MDP1/2), megakaryocytic/erythrocytic (MEP, MKP and ERP), histiocytic (HIST) and eosinophil/basophil/mast cell (Eo\_B\_Mast) progenitors (Fig. 4a and Extended Data Fig. 6). Transcriptional trajectory analysis over pseudotime predicted the HSC1 population as the most transcriptionally primitive HSC population, differentiating into HSC2–6 as well as MLP. Subsequent lineage-biased populations appear to differentiate from HSC2–6 and MLP (Fig. 4b).

The proportion of each transcriptionally defined HSPC cluster in healthy allo-donors (mobilized with G-CSF, plerixafor or motixafortide alone) and in patients with MM (mobilized with placebo + G-CSF, plerixafor + G-CSF or motixafortide + G-CSF) was evaluated (Extended Data Fig. 7). Notably, healthy allo-donors receiving motixafortide alone mobilized a higher proportion of transcriptionally primitive HSC1 cells (25.73%) relative to both plerixafor (1.35%) and G-CSF (1.63%). In addition, allo-donors receiving CXCR4i mobilization with either motixafortide or plerixafor mobilized a higher proportion of MDP1 and MDP2 cells relative to those mobilized with G-CSF alone (MDP1, G-CSF 1.46% versus CXCR4i 9.78% and MDP2, G-CSF 0.31% versus CXCR4i 1.98%). By contrast, within the MM cohorts similar proportions of HSC2-6, CLPs and GMPs were mobilized across the three mobilization regimens. However, the large proportion of MLPs mobilized by all regimens in the allo-donor cohorts (33.56%) was notably lower in patients with MM mobilized with placebo + G-CSF (3.66%). However, this large population of MLPs was preserved when either motixafortide or plerixafor was added to G-CSF mobilization in patients with MM (26.18%), suggesting that MLPs may be uniquely reduced in the G-CSF mobilized HSPC graft of patients with MM and that the addition of CXCR4i with either motixafortide or plerixafor to G-CSF may preserve this population within the HSPC graft in those patients. Taken as a whole, these data suggest that the mobilization regimen used in both allo-donors and patients with MM has an important impact on the proportion of various HSPC subsets mobilized. In addition, the presence of underlying MM, recent exposure to MM-targeting therapies, advanced age or a combination of factors may further impact mobilization of particular HSPC subsets.

Differential gene expression (DEG) analysis across all subsets identified unique transcriptional profiles for each CD34<sup>+</sup> HSPC subset, with specific differences based on mobilization regimen (motixafortide versus plerixafor) and donor type (allo-donor versus patient with MM). Within the HSC1-6 and MLP populations in healthy allo-donors, the HSC1 population was preferentially mobilized at higher numbers by motixafortide compared with either plerixafor or G-CSF (Fig. 4c,d). When comparing motixafortide with G-CSF or plerixafor-mobilized HSCs in the allo-donor cohort, we found EGR1, JUNB, NR4A1, IER2 and RPS26 to be notably upregulated, which has been associated with enhanced quiescence and self-renewal (Fig. 4e,f)<sup>30-32</sup>. By plotting the expression of these markers relative to single-cell populations, most of the expression of these genes is driven by HSC1 and HSC5. Moreover, these genes have markedly higher expression in HSCs mobilized by the long-acting CXCR4i motixafortide relative to either the short-acting CXCR4i plerixafor or G-CSF alone. In addition, KAT7 (HBO1), BMI1, PBX1 and MEIS1 and a related network of genes, which have been associated with HSC maintenance, quiescence and self-renewal, were highly expressed in the HSC5 population mobilized by motixafortide relative to plerixafor or G-CSF<sup>28</sup>. Similarly, HSC2–6 cells mobilized in allo-donors with motixafortide, but not plerixafor or G-CSF, expressed increased levels of *JUNB* and *NR4A1*. By contrast, gene expression profiles of the HSC1–6 and MLP populations were more homogenous within the MM cohorts, potentially reflecting the impact of recent MM-targeting induction therapies on the bone marrow niche, as well as increased age and/or a pre-existing diagnosis of MM. Nevertheless, motixafortide + G-CSF preferentially mobilized a higher proportion of the transcriptionally primitive HSC1 population (31.42%) relative to either plerixafor + G-CSF (17.36%) or placebo + G-CSF (22.77%) in patients with MM.

Gene set enrichment analysis of upregulated genes in the primitive HSC1 population relative to other HSC/MLP populations indicated increased EGF- and TNF- $\alpha$ /NF $\kappa$ B-related signaling, which have been associated with hematopoietic regeneration and/or regenerative potential. Meanwhile, HSC5 expressed upregulated HBO1-related genes associated with self-renewal and quiescence relative to other HSC/MLP populations. To further evaluate EGF, TNF- $\alpha$ /NF $\kappa$ B and HBO1 pathway signaling within each HSC1-6 and MLP subset across patient/treatment/disease characteristics (allo-donor versus MM) and mobilization regimen, pathway expression scores were generated for each pathway (Extended Data Fig. 8 and Methods). These data demonstrated numerically higher average levels of EGF-, TNF- $\alpha$ /NF $\kappa$ B- and HBO1-related gene expression with the use of the long-acting CXCR4i motixafortide as compared with short-acting CXCR4i with plerixafor or G-CSF alone, within most HSC1-6 and MLP subsets in healthy allo-donors, whereas the differences in gene expression scores for EGF-, TNF-α/NFκB- and HBO1-related genes in HSC1-6 and MLP subsets in the MM cohorts were relatively similar across mobilization regimens. In summary, extended CXCR4i with motixafortide mobilizes HSC populations, with upregulated gene expression profiles associated with enhanced self-renewal, quiescence and regeneration in healthy allo-donors.

## Discussion

The effectiveness of ASCT relies, in part, on the ability to collect an adequate number of HSPCs. The 'ideal' HSPC mobilization regimen would be rapid, robust, reliable, well tolerated and capable of safely mobilizing an optimal number of CD34<sup>+</sup> HSPCs in nearly 100% of patients in one apheresis day. Although G-CSF remains the most widely used mobilization agent, 40-50% of patients remain unable to collect an optimal number of HSPCs with G-CSF alone despite multiple injections and up to four apheresis days<sup>11,12</sup>. In addition, 15-35% of patients remain unable to collect optimal numbers of cells despite up to eight injections of G-CSF, four injections of plerixafor and four apheresis days<sup>12,18</sup>. Therefore, the ideal mobilization regimen remains elusive. In addition, progressive improvements in transplant-related care have enabled ASCT to be safely performed in patients with MM  $\geq$ 65 years of age, increasing from 11% of ASCTs in 2000 to 36% of ASCTs in 2019 (ref. 33). Similarly, advances in induction therapy have established three-drug (IMiDs, PIs and glucocorticoids) and four-drug induction regimens (IMiD, PI, glucocorticoids and anti-CD38 mAbs) as the standard of care for newly diagnosed transplant-eligible patients with MM<sup>34,35</sup>. However, increased age, exposure to lenalidomide and four-drug induction therapy are all associated with impaired HSPC mobilization, further emphasizing the unmet need for more effective HSPC mobilization regimens<sup>36-38</sup>.

**Fig. 3** | **Immunophenotyping with percentages and quantitation of CD34**<sup>+</sup> **HSPC subsets. a**-**c**, CD34<sup>+</sup> HSPCs from day 1 apheresis products, collected following treatment of patients with MM with G-CSF plus either placebo, plerixafor or motixafortide, were purified by immunomagnetic selection and evaluated by multicolor FACS. Each patient (*n* = 51) contributed a single biologically independent sample examined over one independent experiment per sample/patient. **a**, *t*-Distributed stochastic neighbor-embedding (t-SNE) projection of merged flow cytometry file showing nine HSPC subsets based on defined cell surface markers. **b**, Percentage of CD34<sup>+</sup> cells within each HSPC subset is shown for patients with MM treated with G-CSF plus either placebo (n = 13), plerixafor (n = 14) or motixafortide (n = 24). **c**, Absolute numbers (Abs. no.) of each HSPC subset are shown for patients with MM treated with G-CSF plus either placebo (n = 12), plerixafor (n = 12) or motixafortide (n = 24). **b**, **c**, Data presented as mean  $\pm$  s.d. Mean HSPC subset yields were compared using ANOVA followed by a post hoc Tukey–Kramer test for pairwise comparisons among groups. Exact two-sided *P* values for significant differences are listed.

In this international, phase 3, multicenter, randomized, double-blind, placebo-controlled trial, patients were representative of the typical MM population undergoing ASCT in the current era, with a median age of 63 years and 70% of patients receiving lenalidomide-containing induction therapy. Despite the presence of these risk factors for poor HSPC mobilization, 92.5% of patients treated with one injection of motixafortide + G-CSF collected  $\geq 6 \times 10^6$ CD34<sup>+</sup> cells kg<sup>-1</sup> within two apheresis days versus 26.2% mobilized with



placebo + G-CSF (P < 0.0001). In addition, 88.8% of patients mobilized with motixafortide + G-CSF collected  $\ge 6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> in one apheresis day versus 9.5% with placebo + G-CSF (P < 0.0001), with a median of 10.8 × 10<sup>6</sup> CD34<sup>+</sup> cells kg<sup>-1</sup> collected in one apheresis with motixafortide + G-CSF. By comparison, in a contemporary cohort of demographically similar patients with MM prospectively enrolled in parallel to the GENESIS trial and mobilized with plerixafor + G-CSF, 50.0% of patients collected optimal numbers of CD34<sup>+</sup> cells kg<sup>-1</sup> in one apheresis (median, 5.47 × 10<sup>6</sup> CD34<sup>+</sup> cells kg<sup>-1</sup>). Meanwhile, motixafortide + G-CSF was well tolerated with rapid and durable engraftment kinetics. Motixafortide + G-CSF usage and the number of apheresis days required to collect the optimal number of HSPCs.

Randomized controlled trials (RCTs) comparing the efficacy of G-CSF alone, chemotherapy + G-CSF, plerixafor + G-CSF and motixafortide + G-CSF to mobilize optimal HSPC numbers (≥6 × 10<sup>6</sup> CD34<sup>+</sup> cells kg<sup>-1</sup>) in the current era of MM therapy are lacking. In addition, comparisons between the present study and previous trials should be undertaken with caution, acknowledging the limitations of cross-trial comparisons. The largest previous RCT comparing plerixafor + G-CSF with placebo + G-CSF in patients with MM was published in 2009, with a median patient age of 58 years and only 5.9% of patients receiving lenalidomide before HSPC mobilization. In the context of relatively younger patients with minimal lenalidomide exposure compared with current practice, 17.3% of patients in the placebo + G-CSF arm and 54.2% in the plerixafor + G-CSF arm mobilized to goal in one apheresis<sup>12</sup>. Other contemporary studies commonly utilize pre-apheresis PB CD34<sup>+</sup> counts to screen for poor mobilizers, and add pre-emptive plerixafor mobilization in patients predicted to fail mobilization with G-CSF alone, making direct comparisons difficult<sup>39,40</sup>. Meanwhile, other studies targeted lower mobilization goals ( $\geq 2-3 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup>), limited premobilization lenalidomide exposure, used alternative growth factors or allowed more than four apheresis days to collect to goal, again making direct comparisons difficult<sup>13,41,42</sup>. Nevertheless, most RCTs and nonrandomized interventional trials in patients with MM since the development of CXCR4is for HSPC mobilization have reported that currently available regimens frequently yield suboptimal CD34<sup>+</sup> numbers despite multiple injections and multiple days of apheresis<sup>12,13,39,40</sup>. Thus it appears, based on these data, that a single injection of motixafortide added to G-CSF substantially improves on currently approved mobilization regimens in terms of the rapidity, robustness and reliability of HSPC mobilization for ASCT in patients newly diagnosed with MM following modern induction therapy in the current era.

There is also an increasing appreciation of the immunophenotypic and transcriptional heterogeneity of CD34<sup>+</sup> HSPCs, ranging from primitive HSCs capable of long-term self-renewal and multilineage potential down to differentiated, lineage-committed progenitors<sup>19</sup>. Moreover, the impact of extended CXCR4i with motixafortide relative to short-acting CXCR4i with plerixafor on graft composition remains incompletely understood. Previous studies evaluating PB HSPCs mobilized by plerixafor compared with those mobilized by G-CSF or collected from bone marrow in allo-donors demonstrated that plerixafor increased mobilization of strongly CXCR4<sup>+</sup>, lineage-committed plasmacytoid dendritic cell progenitors (pDCPs) comprising nearly 25% of the HSPC graft<sup>19</sup>. We observed that motixafortide mobilized a similar proportion of pDCPs compared with plerixafor; but also mobilized a significantly higher number of combined HSCs, MPPs and CMPs compared with plerixafor. This unique impact of extended CXCR4i with motixafortide relative to short-acting CXCR4i with plerixafor may be due to the observation that these more primitive HSPC subsets are less strongly CXCR4<sup>+</sup> at baseline, and thus are preferentially mobilized at higher numbers by extended CXCR4i with motixafortide. Also of note, scRNA-seq suggests that extended CXCR4i with motixafortide in healthy allo-donors mobilizes transcriptionally unique subsets of HSCs (HSC1 and HSC5; Fig. 4), which exhibit upregulated transcriptional programing associated with enhanced self-renewal, regeneration and quiescence (EGR1, JUNB, BTG2, NR4A1, MYB, IER2, EGR3, BMI1, PBX1, MEIS1 and KAT7 (HBO1)). Recent work by Desterke et al. revealed that quiescence markers (EGR1, JUNB, BTG2 and NR4A1) and other genes (MYB and IER2) were upregulated in long-lived HSCs<sup>30</sup>. The self-renewal transcription factor EGR3, which suppresses cell cycle progression, is upregulated in motixafortide-treated clusters and is also highly expressed in primitive HSCs during leukemic transformation<sup>31,32</sup>. Shepard et al. observed that HSCs in JAK2 mutant myeloproliferative neoplasms harbored defective self-renewal properties, while robust self-renewal capacity and HSC repopulation was noted when BMI1, PBX1 or MEIS1 were overexpressed within mutant myeloproliferative neoplasms<sup>43,44</sup>. Additionally, Yang et al. observed that the histone lysine acetyltransferase KAT7 (HBO1) and a related network of genes is necessary for HSC maintenance and self-renewal<sup>28</sup>.

The clinical implications of the observed immunophenotypic and transcriptional heterogeneity within CD34<sup>+</sup> HSPC subsets mobilized with various regimens remain unclear. This study was not designed to evaluate how differences in the total number of CD34<sup>+</sup> cells infused, or the number of specific HSPC subsets infused, might impact clinical outcomes. In addition, given that extended immunophenotyping was performed on only a subset of patients mobilized at Washington University in St. Louis (n = 37), comparative analyses between cohorts are probably underpowered. Nevertheless, in a post hoc pooled analysis, a significant association was observed between increasing total number of CD34<sup>+</sup> cells infused and faster TPE. These data suggest that infusion of higher numbers of CD34<sup>+</sup> cells ( $\geq 7 \times 10^{6}$  CD34<sup>+</sup> cells kg<sup>-1</sup>) may result in faster TPE. This observation is supported by previous publications suggesting that infusion of  $>6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> is associated with improved long-term platelet recovery<sup>45</sup>. In addition, we observed that patients infused with the upper quartile (>75th percentile) of GMPs alone had significantly faster TPE (13 versus 19 days). These data suggest that the number of GMPs within the HSPC graft specifically contributed to platelet engraftment kinetics, which may indicate that the CD34<sup>+</sup> HSPC subset immunophenotypically defined in this study as GMPs (CD45RA<sup>+</sup>/<sup>-</sup>CD123loCD38<sup>+</sup>/<sup>-</sup>CD10<sup>-</sup>) contains a population of

**Fig. 4** | **Single-cell transcriptional profiling and trajectory mapping of CD34**<sup>+</sup> **HSPCs with differentially expressed gene subanalysis of HSC1-6 and MLP populations. a**, UMAP plot of annotated single-cell clusters across entire cohort. Cells colored by abbreviations of each transcriptional annotation as defined. b, UMAP plot of single-cell clusters with Monocle3 trajectory mapping overlayed and colored by pseudotime. Overlayed trajectory indicates transitions between distinct transcriptional states. Pseudotime is a measure of progress along the overlayed trajectory, with the white spot labeled '1' indicating the earliest state on the transcriptional path. c, UMAP of early progenitor populations (HSC1-6, MLP) across each cohort and mobilization regimen, with each cell colored by cell-type annotation and separated by mobilization regimen and cohort (M, multiple myeloma; H, healthy allo-donor). d, Average cell-type proportions across each sample within each treatment group and cohort. Spot size indicates relative average expression, with the exact value overlayed. **e**, Highlighted genes found to be differentially expressed between early progenitor populations in the healthy allo-donor cohort across the listed mobilization regimens. Each column represents a different DEG analysis between the two groups, and values for each DEG listed show the average log<sub>2</sub>(fold change (FC)) in the first cohort relative to the second cohort in each column. Each element in heatmap is colored by average log<sub>2</sub>(FC), with red and blue denoting increased and decreased expression, respectively. **f**, Expression of a subset of DEGs and genes (from the literature) across stem cell and early progenitor clusters. Each spot is colored by average expression, its size indicating total number of cells expressing the gene of interest in the mobilization regimen and cohort labeled.

megakaryocytic progenitors capable of rapidly reconstituting platelet engraftment. Based on our gating strategy, this observation is consistent with a recent report demonstrating that unipotent megakaryocyte progenitors lie within the CD34<sup>+</sup> CD38<sup>+</sup> CD45RA<sup>-</sup> population<sup>46</sup>. In conclusion, the upfront use of a single injection of motixafortide added to G-CSF resulted in rapid, robust and reliable mobilization of optimal numbers of CD34<sup>+</sup> HSPCs in patients with MM undergoing ASCT. Moreover, extended CXCR4i with motixafortide preferentially



mobilized increased numbers of immunophenotypically and transcriptionally primitive HSCs. Future studies may consider similar regimens for HSPC-based, gene-edited platforms where the optimal HSPC collection goal is typically much higher  $(10-15 \times 10^6 \text{ CD34}^+$ cells kg<sup>-1</sup>) than that of ASCT (5–6 × 10<sup>6</sup> CD34<sup>+</sup> cells kg<sup>-1</sup>), given losses of HSPC-viability-associated gene/base editing<sup>47,48</sup>. In addition, regimens mobilizing higher proportions of primitive HSCs may be particularly advantageous for HSPC-based, gene-edited therapies given that the long-term effectiveness of such therapies relies on the successful ability of modified HSPCs to establish stable, long-term engraftment.

# **Online content**

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41591-023-02273-z.

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# Methods

#### **GENESIS trial design**

All patients treated on the GENESIS trial (Clinicaltrials.gov: NCT03246529) provided written informed consent before enrollment. Institutional Review Board (IRB) approval of the study protocol, investigator brochure, amendments, informed consent and any other documents provided to the subject was performed by the following: Washington University IRB; University of Miami IRB; Mayo Clinic IRB; University of Oregon Oregon Health & Science University IRB; University of Kansas Medical Center IRB; University of Maryland IRB; Western Cooperative Group IRB; The Loyola University Chicago Health Sciences Campus IRB; UCLA IRB; University of Utah IRB; University of Cincinnati IRB; The Weill Cornell Medical College IRB; UT M.D. Anderson Cancer Center IRB: Ethics Committee Catania 1 Azienda Ospedaliero-Universitaria 'Policlinico-Vittorio Emanuele' Catania; Ethics Committee South Reggio Calabria Division Grande Ospedale Metropolitano 'Bianchi-Melacrino- Morelli'; University of Cologne Ethics Committee; University of Debrecen, Clinical Center, Clinic of Internal Medicine, Hematology Scientific Council for Health - Ethical Committee for Clinical Pharmacology, Central Hospital of Southern Pest, National Institute of Hematology and Infectious Diseases Scientific Council for Health - Ethical Committee for Clinical Pharmacology. Please see Reporting Summary for additional details for each IRB. The study was conducted in accordance with the Guideline for Good Clinical Practice ICH E6(R2)-ICH Harmonized Guideline Integrated Addendum to ICH E6 (R1) (International Conference on Harmonization of Technical Requirements for the Registration of Pharmaceuticals for Human Use), Step 5, 14 June 2017; the Declaration of Helsinki: Seoul, 2008; the US Code of Federal Regulations (Title 21, CFR Part 11, 50, 54, 56 and 312) and/or EU Directives; and/or local country regulations and guidelines.

The GENESIS trial included a preplanned, lead-in, single-arm, open-label period previously reported<sup>49,50</sup>, demonstrating that motixafortide + G-CSF was safe, well tolerated and resulted in 82% (9 of 11) of patients with MM mobilizing  $\ge 6 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> within two apheresis days. The preplanned review of these Part 1 data led the independent data monitoring committee (DMC) to recommend transitioning to Part 2 of the study. In the prospective, phase 3, double-blind, placebo-controlled, multicenter Part 2 of the study, 122 patients undergoing ASCT were enrolled at 18 sites across five countries and were randomized 2:1 to receive either motixafortide + G-CSF or placebo + G-CSF for HSPC mobilization before ASCT for MM. Key eligibility criteria included: patients 18-78 years of age with a confirmed diagnosis of MM; an Eastern Cooperative Oncology Group performance status of 0-1; and adequate organ function undergoing first ASCT in first or second complete response (CR) or partial response (PR) (according to IMWG Response Criteria). Sex and/or gender were recorded for demographic purposes, as self-reported by the participant. Patients were excluded if they had undergone prior HCT or failed previous HSPC collection attempts (see full inclusion and exclusion criteria in supplemental GENESIS trial protocol). Randomization of eligible patients was performed using an interactive web response system and was conducted in permuted blocks with stratification by response status (CR versus PR) and baseline platelet count ( $<200 \times 10^9 l^{-1}$  or  $\ge 200 \times 10^9 l^{-1}$ ). Randomization was performed by the clinical research coordinator, who was not involved in direct patient care. All patients, investigators and providers/staff were blinded to treatment assignment. All patients received G-CSF (10 mcg kg<sup>-1</sup>) on days 1–5 (and 6–8, if needed). Patients received either motixafortide (1.25 mg kg<sup>-1</sup>, subcutaneous injection) or placebo on day 4 (and 6, if needed). Apheresis began on day 5 (four blood volumes), with the primary and secondary endpoints of collect $ing \ge 6 \times 10^6 CD34^+$  cells kg<sup>-1</sup> in up to two or one apheresis days, respectively. Apheresis continued on days 6-8 if needed. Total CD34<sup>+</sup> cells kg<sup>-1</sup> were analyzed locally to determine whether patients met the primary endpoint, and all samples were subsequently sent for assessment by a central laboratory. Patients who did not collect  $\ge 2 \times 10^{6}$  CD34<sup>+</sup> cells kg<sup>-1</sup>

by day 8 proceeded to rescue mobilization. To assess the impact of each mobilization regimen on PB CD34<sup>+</sup> cells and the relationship to collection of CD34<sup>+</sup> HSPCs via apheresis, peripheral blood CD34<sup>+</sup> HSPC counts were assessed using both local and central laboratories at the following time points: day 0 (baseline), before first dose of G-CSF; day 4, before first dose of motixafortide/placebo but after four doses of G-CSF; and day 5 (apheresis day 1), after first administration of motixafortide or placebo and after five administrations of G-CSF. The number of CD34<sup>+</sup> cells infused for ASCT was determined independently by each investigator according to local practice; however, a minimum of  $\geq 2 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> was required. Although initial power calculations called for 177 patients to be enrolled in Part 2 of the study, a preplanned interim analysis after 122 patients were enrolled was performed by an independent unblinded statistician and these results were communicated only to the independent DMC for review, leading the DMC to recommend halting the study due to statistically significant efficacy favoring the motixafortide + G-CSF mobilized cohort (prespecified threshold  $P \le 0.0108678$ ).

#### Correlative study design

All patients who participated in correlative studies provided written informed consent before enrollment on their respective protocols. GENESIS trial patients were enrolled and mobilized as previously described. In addition, a demographically similar, contemporaneous cohort of patients (n = 14) undergoing mobilization with plerixafor + G-CSF (regardless of PB CD34<sup>+</sup> cell count preapheresis) for ASCT for MM were prospectively enrolled on a parallel tissue-banking protocol (no. 201103349). All patients with MM received G-CSF (10 mcg kg<sup>-1</sup>) on days 1-5 (and 6-8, if needed). Patients then received either motixafortide (1.25 mg kg<sup>-1</sup>) or placebo on day 4 (and 6, if needed) via subcutaneous injection, or plerixafor (0.24 mg kg<sup>-1</sup>) on day 4 (and 5-7, if needed) via subcutaneous injection. Apheresis began on day 5 (and 6-8, if needed). Three healthy allo-donor cohorts underwent single-agent mobilization with either (1) G-CSF alone (10 mcg kg<sup>-1</sup>) on days 1-5 followed by apheresis beginning on day 5 (protocol no. 201106261); (2) plerixafor alone (0.24 mg kg<sup>-1</sup>) on day 1 followed by apheresis within 4 h of plerixafor administration (NCT00241358); or (3) motixafortide alone (1.25 mg kg<sup>-1</sup>) on day 1 followed by apheresis within 3 h of motixafortide administration (NCT02639559). Study samples were obtained from the apheresis product on the first day of apheresis in all patients, and CD34<sup>+</sup> cells were analyzed as detailed below for correlative studies.

#### Post hoc engraftment kinetics

Although the number of CD34<sup>+</sup> cells kg<sup>-1</sup> infused for ASCT on the GENESIS trial was determined independently by each investigator according to local practice, a minimum of  $\ge 2 \times 10^6$  CD34<sup>+</sup> cells kg<sup>-1</sup> was required. A post hoc pooled analysis was performed using Pearson correlation to evaluate TPE (platelet count  $\ge 20 \times 10^9$  l<sup>-1</sup> without platelet transfusion ×7 days) and TNE (absolute neutrophil count  $\ge 0.5 \times 10^9$  l<sup>-1</sup> ×3 days) based on the total number of CD34<sup>+</sup> cells kg<sup>-1</sup> infused without regard to mobilization regimen, as well as the total number of specific CD34<sup>+</sup> HSPC subsets infused. CD34<sup>+</sup> HSPC immunophenotyping was performed via multicolor FACS, as detailed below.

#### **Multicolor FACS**

Peripheral blood CD34<sup>+</sup> HSPC counts were assessed using a Stem Cell Enumeration Kit (BD Biosciences, no. 344563) at both local and central laboratories. For correlative studies, CD34<sup>+</sup> HSPCs from n = 51patients (placebo + G-CSF, n = 13; plerixafor + G-CSF, n = 14; motixafortide + G-CSF, n = 24) were purified from apheresis product collected on the first day of apheresis via CD34<sup>+</sup> immunomagnetic selection using an AutoMACS device (Miltenyi Biotech). CD34<sup>+</sup> HSPCs were washed in PBS and stained for 15 min at room temperature with a LIVE/DEAD Fixable Aqua Dead Cell Stain kit (Invitrogen). Cells were then washed in PBS supplemented with 0.5% bovine serum albumin and 2 mMEDTA

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and incubated for 10 min at room temperature with human Fc Block and Brilliant Stain Buffer (BD Biosciences). Samples were then incubated for 30 min at room temperature with pretitrated saturating dilutions of the following fluorochrome-labeled antibodies (clone, source designated, catalog no, and dilution in parentheses): CD45-BUV395 (HI30, BD Biosciences, no. 563792, 1:60); CD123-BUV737 (7G3, Biosciences, no.741769,1:120); CD49f-BV421 (GoH3, BioLegend, no.313624, 1:300); CD14-BV650 (M5E2, BioLegend, no. 301836, 1:120); CD45RA-BV785 (HI100, BioLegend, no. 304140, 1:120); CD34-VioBright515 (REA1164, Miltenyi, 1:120); CD10-PECF594 (HI10a, Biosciences, no. 562396, 1:484); CD38-PE-Cv7 (HIT2, BioLegend, no. 303516, 1:120); CD90-APC (5E10, BioLegend, no. 328114, 1:60); CD303-APCVIO770 (REA693, Miltenyi Biotech, no. 130-120-517, 1:150); CD184-PE (1D9, BD, no. 551510, 1:242); and CD184-PE (12G5, Biosciences, no. 555974, 1:150). Fluorescence minus-one controls were used to assess background fluorescence intensity and set gates for negative populations. After washing twice, samples were analyzed on a ZE5 (Bio-Rad) flow cytometer. Single-stain compensation controls were obtained using UltraComp eBeads (Thermo Fisher Scientific) and data were analyzed using FCS Express (DeNovo Software). The antibody-binding capacity per cell of the different CD34<sup>+</sup> HSPC subsets was determined for CD184 clones 12G5 and 1D9 using saturating concentrations of antibody and the Quantum Simply Cellular (QSC, Bangs Laboratories) system for fluorescence quantitation according to the manufacturers recommendations.

#### Single-cell library preparation and sequencing

CD34<sup>+</sup> HSPCs from n = 12 patients with MM (placebo + G-CSF, n = 4; plerixafor + G-CSF, n = 4; motixafortide + G-CSF, n = 4), and from n = 6healthy allogeneic HSPC donors (G-CSF, n = 2; plerixafor, n = 2; motixafortide, n = 2), were purified from apheresis products collected on the first day of apheresis via CD34 immunomagnetic selection. Transcriptional profiling was performed by 10×, 5' scRNA sequencing. For sample preparation on the 10X Genomics platform, the Chromium Next GEM Single Cell 5' Kit v.2, 16 rxns (no. PN-1000263), Chromium Next GEM Chip K Single Cell Kit, 48 rxns (no. PN-1000286) and Dual Index Kit TT Set A, 96 rxns (no. PN-1000215) were used. The concentration of each library was accurately determined through quantitative PCR utilizing the KAPA library Quantification Kit according to the manufacturer's protocol (KAPA Biosystems/Roche), to produce cluster counts appropriate for the Illumina NovaSeq6000 instrument. Normalized libraries were sequenced on a NovaSeq6000 S4 Flow Cell using XP workflow and  $a151 \times 10 \times 10 \times 151$  sequencing recipe according to the manufacturer's protocol. A median sequencing depth of 50,000 reads per cell was targeted for each gene expression library.

#### scRNA-seq data preprocessing

For each sample we obtained the unfiltered feature-barcode matrix by passing demultiplexed FASTQs to Cell Ranger v.6.0.1 'count' command using default parameters and the prebuilt GRCh38 genome reference (no. GRCh38-2020-A), and Chemistry flag (Single Cell 5' PE) for scRNA (data files available via the gene expression omnibus (accession no. GSE223972). Seurat 4.1.0 was used for all subsequent analyses. Cells were further filtered to maintain only those cells with <20% human mitochondrial DNA content and a minimum of at least 200 and maximum of 20,000 genes expressed. We constructed a Seurat object using the unfiltered feature-barcode matrix for each sample. Each sample was scaled and normalized using Seurat's 'SCTransform' function to correct for batch effects (with parameters 'vars.to.regress = c('nCount\_RNA', 'percent.mito')', variable.features *n* = 2,000). Any merged analysis or subsequent subsetting of cells/samples underwent the same scaling and normalization method. Cells were clustered using the original Louvain algorithm, and the top30 principal component analysis dimensions via functions 'FindNeighbors' and 'FindClusters' (with parameters: resolution = 0.5). The resulting merged and normalized matrix was used for subsequent analysis.

#### scRNA-seq cell-type annotation

Cell types were assigned to each cluster by manually reviewing the expression of a comprehensive set of marker genes derived from several publications  $^{9,26-29}$ .

#### scRNA-seq Monocle trajectory analysis

Monocle3 (https://cole-trapnell-lab.github.io/monocle3/) was used for pseudotime analysis. Analysis was completed following the standard tutorial for construction of single-cell trajectories (https:// cole-trapnell-lab.github.io/monocle3/docs/trajectories/).

#### scRNA-seq DEG analysis

For cluster-level differential expression we used the 'FindMarkers' or 'FindAllMarkers' Seurat function as appropriate, with a minimum percentage of 0.3 (parameter min.pct = 0.3). The resulting DEGs were then filtered for adjusted P < 0.05 and sorted by FC. All differential expression analyses were carried out using the 'SCT' assay.

#### **Gene expression scores**

Gene expression scores for each cell were annotated using the AddModuleScore function in Seurat. Gene sets were extracted from www.gsea-mdsigdb.org. The TNFA\_NFKB\_Score gene set was derived from the HALLMARK\_TNFA\_SIGNALING\_VIA\_NFKB pathway. EGF\_Score was derived from NAGASHIMA\_EGF\_SIGNALING\_UP and MIT\_EGF\_RESPONSE\_40\_HELA. HBO1\_score was derived from Yang et al<sup>28</sup>, including the following genes: *KAT7, MPL, TEK, GFI1B, EGR1, TAL1, GATA2, ERG, PBX1, MEIS1, HOXA9* and *GATA1*. For all gene sets, the percentage of cells expressing each gene in the set was calculated for each treatment group. Any genes with <10% expression across a given treatment group were filtered out and not used in the final gene score annotation.

#### Healthcare resource utilization

As a prespecified analysis, healthcare resource utilization items were collected alongside the GENESIS trial in each treatment arm, including (1) the number of motixafortide and G-CSF doses, (2) the number of apheresis procedures used in primary mobilization and (3) the proportion of patients requiring rescue mobilization due to inadequate primary mobilization.

#### **Statistics**

The primary endpoint was the proportion of patients mobilizing  $\geq 6 \times 10^{6} \text{ CD34}^{+}$  cells kg<sup>-1</sup> within up to two apheresis sessions in preparation for auto-HCT following G-CSF and a single administration of BL-8040/placebo. Prespecified secondary endpoints included the proportion of patients mobilizing  $\geq 6 \times 10^6$  CD34+ cells kg<sup>-1</sup> in one apheresis, the proportion of patients mobilizing  $\geq 2 \times 10^{6} \text{ CD34}^{+}$  cells kg<sup>-1</sup> in one apheresis, TPE, TNE, graft durability, OS and PFS. The primary endpoint, prespecified secondary efficacy endpoints and graft durability were analyzed using the Cochran-Mantel-Haenszel (CMH) test with calculation of OR, two-sided 95% Cl and P values. Time to engraftment, PFS and OS secondary endpoints were analyzed using Cox's proportional hazards model, and time-to-event analyses (Kaplan-Meier method) with calculation of HR, two-sided 95% CI and P values. Analyses of all prespecified primary and secondary endpoints were performed on an ITT basis unless otherwise stated. P < 0.05 was considered statistically significant unless otherwise stated. Pearson correlation was performed to assess associations between total CD34<sup>+</sup> cells kg<sup>-1</sup> infused and specific CD34<sup>+</sup> subset cells kg<sup>-1</sup> infused and TPE/TNE. Sensitivity analyses were performed using time-to-event analyses (Kaplan-Meier method) to determine thresholds for both total CD34<sup>+</sup> HSPCs infused and specific HSPC subsets infused above which TPE was significantly faster. Mean HSPC subset yields in apheresis were compared using analysis of variance, followed by post hoc Tukey-Kramer test for pairwise comparisons among groups. Two-sided statistical analyses were used.

### **Reporting summary**

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

# Data availability

All data and/or supporting documents related to the paper that Nature reviewers and/or editors will request for the purposes of evaluating this paper and verifying its contents will be provided through Egnyte system, at any time. Data will also be available to researchers and/or scientists in alignment with the International Committee of Medical Journal Editors' policy on clinical data sharing. Specifically, the authors will provide access to individual deidentified participant-level data that underlie the data presented in this paper, including data dictionaries, the study protocol and other relevant information, to any researcher who provides a methodologically sound proposal for academic purposes beginning 6 months and ending 5 years after article publication. These data can be requested via email to iritg@biolinerx.com, and will be made available to requesting parties through the Egnyte system following approval. This request and availability mechanism for accessing the clinical dataset will similarly apply to requests for the 'minimum dataset' necessary to interpret, verify and extend the research in the article. All requests for clinical data will be reviewed by the sponsor (BioLineRx) to verify whether the request is subject to any intellectual property or confidentiality obligations. The gene count matrices for scRNA-seq are available via Gene Expression Omnibus (accession no. GSE223972). References (GRCh38 genome reference, namely refdata-gex-GRCh38-2020-A) used for single-cell analysis of human genomes are available from public sources: https:// support.10xgenomics.com/single-cell-gene-expression/software/ release-notes/build.

# **Code availability**

Code relevant for single-cell analysis can be found on GitHub (https:// github.com/reykajayasinghe/WashU\_Genesis\_Study). Additionally, software and code used for the analyses presented in this study are enumerated within the Reporting Summary section on software and code and detailed here: Commercially available R, SPSS, SAS and Xcel statistical and graphics software packages were used for data analysis and graphics; Monocle3 (https://cole-trapnell-lab.github. io/monocle3/) was used for pseudotime analysis. Analysis was completed following the standard tutorial for construction of single-cell trajectories (https://cole-trapnell-lab.github.io/monocle3/docs/trajectories/); Bioconda, https://bioconda.github.io/; Bioconductor v.3.9, https://bioconductor.org/; Cell Ranger v.6.0.1, 10X Genomics, https:// support.10xgenomics.com/single-cell-gene-expression/software/ pipelines/latest/what-is-cell-ranger; data.table\_1.12.6, R Development Core Team, https://cran.r-project.org/package=data.table; dendsort\_0.3.3, https://cran.r-project.org/package=dendsort; dplyr\_0.8.5, R Development Core Team, https://cran.r-project.org/package=dplyr; ggplot2 3.3.2, R Development Core Team, https://CRAN.R-project. org/package=ggplot2; gridExtra\_2.3, R Development Core Team, https://cran.r-project.org/package=gridExtra; magrittr\_1.5, R Development Core Team, https://cran.r-project.org/package=magrittr; Matrix\_1.2-17, R Development Core Team, https://CRAN.R-project.org/ package=Matrix; pheatmap\_1.0.12, R Development Core Team, https:// cran.r-project.org/package=pheatmap; Python v3.7, Python Software Foundation, https://www.python.org/; R v.3.6, R Development Core Team, https://www.r-project.org/; RColorBrewer\_1.1-2, R Development Core Team, https://CRAN.R-project.org/package=RColorBrewer; reshape2\_1.4.3, R Development Core Team, https://cran.r-project.org/ package=reshape2; SAS v.9.4 for statistical analyses; Seurat v.3.1.2 and v.4.0.3, https://cran.r-project.org/web/packages/Seurat; Single-cell analysis code (https://github.com/reykajayasinghe/WashU\_Genesis\_Study); svav.3.40.0, https://bioconductor.org/packages/release/ bioc/html/sva.html; stringr 1.4.0, R Development Core Team, https:// cran.r-project.org/package=stringr; Subread v.2.0.1, https://sourceforge.net/projects/subread/; Tidyverse, https://www.tidyverse.org/; viridis\_0.5.1, R Development Core Team, https://github.com/sjmgarnier/viridis; viridisLite\_0.3.0, R Development Core Team, https:// github.com/sjmgarnier/viridis.

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# **Author contributions**

Z.D.C. conceived and designed the clinical trial, designed the correlative trial, performed data analysis and wrote the original paper. M.P.R. designed the correlative trial, performed data analysis and wrote the original paper. R.G.J. designed the correlative trial, performed data analysis and wrote original paper. K.S.-G. designed and conducted the clinical trial and reviewed and edited the original paper. S.M.L. conducted the clinical trial and reviewed and edited the original paper. I.A. conducted the clinical trial and reviewed and edited the original paper. G.A.M. conducted the clinical trial and reviewed and edited the original paper. M.M. conducted the clinical trial and reviewed and edited the original paper. P.S. conducted the clinical trial and reviewed and edited the original paper. D.S. conducted the clinical trial and reviewed and edited the original paper. D.P. conducted the clinical trial and reviewed and edited the original paper. I.M. conducted the clinical trial and reviewed and edited the original paper. G.M.-J. conducted the clinical trial and reviewed and edited the original paper. G.M. conducted the clinical trial and reviewed and edited the original paper. M.L.P.C. conducted the clinical trial and reviewed and edited the original paper. U.H. conducted the clinical trial and reviewed and edited the original paper. J.H. conducted the clinical trial and reviewed and edited the original paper. M.H.Q. conducted the clinical trial and reviewed and edited the original paper. N.H. conducted the clinical trial and reviewed and edited the original paper. T.L. conducted the clinical trial and reviewed and edited the original paper. I.G.-C. conducted the clinical trial and reviewed and edited the original paper. A.V.-H. performed data analysis and reviewed the original paper. E.S. performed data analysis and reviewed the original paper. I.G.-K. performed data analysis and reviewed the original paper. I.G. performed data analysis and reviewed the original paper. D.I. performed data analysis and reviewed the original paper. L.S.-D. performed data analysis and reviewed the original paper. S.K. provided biostatistical support, performed data analysis and contributed to the statistical section of the paper. F.G. provided biostatistical support, performed data analysis and contributed to the statistical section of paper. M.A.S. conducted the clinical trial and reviewed and edited the original paper. R.V. conducted the clinical trial and reviewed and

edited the original paper. J.F.D. conceived and designed the clinical trial, designed the correlative trial and reviewed and edited the original paper.

# **Competing interests**

Z.D.C.: research funding, BioLineRx; M.P.R.: no competing interests. R.G.J.: consulting fees, WUGEN. K.S.-G.: research funding, BioLineRx, Caelum Biosciences, GlaxoSmithKline, GlycoMimetics, Ionis Pharmaceuticals, Janssen Pharmaceuticals, Millennium Pharmaceuticals (Takeda), Sanofi; equity stock/ownership, Abbott, AbbVie; consulting Fees, GlaxoSmithKline; board or advisory committee membership, GlaxoSmithKline; speaking fees, Janssen Pharmaceuticals; expert witness, Celgene, G.A.M., M.M., P.S.: no competing interests. I.A.: consulting fees and board or advisory committee membership, Janssen, Celgene, Novartis, Pfizer, Takeda, Roche. D.S: consulting fees, Janssen, AbbVie, Sanofi, GlaxoSmithKline, BristolMyersSquibb, Pfizer; board or advisory committee membership, Janssen, Sanofi, GlaxoSmithKline. D.P.: no competing interests. I.M.: research funding, BioLineRx. G.M.J.: no competing interests. G.M.: research funding, AbbVie, Janssen-Cilag; consulting fees, AbbVie, Amgen, Bristol-Myers-Squibb/Celgene, Janssen-Cilag, Krka, Novartis, Roche, Takeda. M.L.P.C.: no competing interests. U.H.: consulting fees, Sanofi-Adventis. J.H.: no competing interests. M.H.Q.: research funding, Janssen Pharmaceuticals, BioLineRx, Angiocrrine, Neximmune. N.H.: board or advisory committee membership, Incyte, Kite/Gilead, American Gene Technologies. T.L., I.G.-C.: no competing interests. A.V.-H., E.S., I.G.-K., I.G., D.I., L.S.-D.: employment, BioLineRx. S.K., F.G.: no competing interests. M.A.S.: research funding, Incyte, Cellect, Inc., Equillium, Inc., Fortis, Seattle Genetics, Amgen, Celgene, PBD, Inc., Sanofi Genzyme, Janssen; consulting fees and board or

advisory committee membership, Advarra, Amgen, Astellas, CareDx, Dova pharmaceuticals, Equillium, Inc., FlatIron, Inc., GlaxoSmithKline, Gilead Sciences, Inc., Incyte, Inmagene Bio, Janssen, Novo Nordisk, Partners Therapeutics, Pfizer, Sanofi Genzyme, StemLine. R.V.: no competing interests. J.F.D.: equity stock/ownership, Magenta Therapeutics, WUGEN; consulting fees, Incyte, RiverVest Venture Partners; board or advisory committee membership, Cellworks Group, RiverVest Venture Partners, Magenta; research funding, Amphivena Therapeutics, NeoImmune Tech, Macrogenics, Incyte, BioLineRx, WUGEN; speaking fees, Incyte; patents (nos. PCT/US2019/035010 and PCT/US2019/035052), WUGEN, Magenta.

# **Additional information**

**Extended data** is available for this paper at https://doi.org/10.1038/s41591-023-02273-z.

**Supplementary information** The online version contains supplementary material available at https://doi.org/10.1038/s41591-023-02273-z.

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Extended Data Fig. 1 | Mobilization and Apheresis Schema. All patients underwent HSPC mobilization and collection on days 1–5 (and days 6–8, if needed). In the AM of days 1–5 (and days 6–8, if needed), all patients received G-CSF (10 mcg/kg) via subcutaneous injection indicated by the syringe in blue. (a) <u>GENESIS Trial</u>: In addition to G-CSF, in the PM of day 4 (and day 6, if needed), patients received either motixafortide (1.25 mg/kg) or placebo via subcutaneous injection indicated by the syringe in green. All patients began apheresis (4 blood volumes) on day 5 (and days 6–8, if needed) with the goal of collecting  $\geq$ 6 × 10° CD34+ cells/kg. Patients collecting to goal completed mobilization. (b) <u>Correlative Study</u>: A contemporaneous, cohort of demographically similar MM patients were prospectively enrolled on a parallel protocol and mobilized with G-CSF, as above, plus plerixafor (0.24 mg/kg) via subcutaneous injection on the PM of day 4 (and day 5–7, if needed) indicated by the syringe in yellow. All patients received plerixafor in addition to G-CSF, regardless of PB CD34+ cell count prior to apheresis. All patients began apheresis on day 5 (and days 6–8, if needed) with the goal of collecting sufficient CD34+ cells for ASCT, per institutional practice.



**Extended Data Fig. 2** | **Pre-specified Principal and Sensitivity Analyses of the Primary Endpoint.** Local and central lab CD34+ HSPC assessments of the apheresis product from each patient were analyzed for the primary endpoint using Cochran-Mantel-Haenszel (CMH) weights on an intent-to-treat (ITT) basis using N = 122 biologically independent samples examined over 1 experiment/ sample. In addition, a pre-specified sensitivity analysis was performed on a modified intent-to-treat (mITT) basis using N = 120 biologically independent samples examined over 1 experiment/sample and on a per protocol (PP) basis using N = 76 biologically independent samples examined over 1 experiment/ sample. The data are presented as CMH common proportions differences for each analysis with 2-sided 95% confidence intervals (CI) indicated by the error bars. Mixed\* analyses accounted for missing central lab values by replacing them with non-missing local lab values. Logistic regression (LR\*\*), multiple imputation missing not at random (MI MNAR) and missing at random (MI MAR) analyses were also performed.

# CMH Common proportions difference & 95% CI



**Extended Data Fig. 3** | **Flow Cytometry Gating Strategy.** CD34<sup>+</sup> HSPCs from day 1 apheresis products collected following treatment of patients with MM with G-CSF plus either placebo, plerixafor or motixafortide were purified by immunomagnetic selection and evaluated by multicolor FACS. Gating strategy used to define nine different CD34<sup>+</sup> HSPC subsets is shown.





and evaluated by flow cytometry. The expression of CXCR4 was determined by flow cytometry using anti-CXCR4 mAb clones 12G5 and 1D9. (**A**) Percent of HSPC subsets positive for 12G5+ and 1D9+ antibodies. (**B**) Mean Fluorescence Intensity (MFI) of antibody binding capacity (ABC) of 12G5 and 1D9 on HSPC subsets.

## Article



Extended Data Fig. 5 | Engraftment Kinetics by Total CD34+ HSPCs Infused and by Specific CD34+ HSPCs Subsets Infused. Pearson correlation with green line representing the regression curve and the shaded green area representing the 2-sided 95% CI presented with respective p-values. All statistical analyses were 2-sided. (a) time to platelet engraftment in days by total CD34+ HSPCs infused, (**b**) time to neutrophil engraftment in days by total CD34+ HSPCs infused, (**c**) time to platelet engraftment in days by total combined HSC, MPP, CMP and GMP CD34+ subset HSPCs infused and (**d**) time to platelet engraftment in days by total GMP CD34+ subset HSPCs infused. Each motixafortide+G-CSF patient indicated by circles and each placebo+G-CSF patient indicated by triangles.



**Extended Data Fig. 6** | **Gene Markers for CD34+ Cell Type Assignments.** Cell type markers from the literature used to annotate single-cell clusters. Dot plot shows the average expression of each gene for each cell-type population and size shows the percent of cells expressing that gene of interest.

		Hoalt	by C	ohor	+			Mvel	oma	Coh	ort	Pi	roport	tion c	of Cel	ls (20	) $(4$	o) (	60
		G-C	SF	Pleri	ι xafor	Motix	afortide	wyer	G-C	SF	511	PI	erixafo	r+G-C	SF	Moti	xaforti	de+G-	CSF
	PSC1 -	0.68	2 <mark>.0</mark> 4	0.88	1 <b>.5</b> 5	1 <mark>.8</mark> 2	4 <mark>.0</mark> 9	1.25	1.57	0.84	3.52	0.64	0.91	0.81	2.21	2.28	63.23	0.83	0.84
	PSC2 -							3.45		26.50		0.01	0.03	0.19		0.06		0.02	0.03
	LMPP_Ly1 -	0.61	0.05	1.¶0	0. <b>4</b> 5	0.43	0.82	2.47	1.16	0.03	0.22	0.51	0.22	0.18	0.88	0.88	2.40	0.86	0.21
	CLP_Ly2 -	4 <mark>.6</mark> 0	1 <mark>.7</mark> 8	10.84	10.35	10.73	6.93	3.58	2.51	0.23	3.06	6.15	7.51	2.46	3.21	2 <b>.9</b> 4	0.23	12.61	8.33
CLP	_Ly2_B_PC -	1 <b>.3</b> 9	0.75	0.81	0.11	2 <mark>.0</mark> 0	0.51	0.98	0.14	0.87	1.41	0.85	0.46	0.83	0.10	0.21	0.11	5.80	1.26
	MDP1 -	2 <mark>.6</mark> 2	0.30	20.75	5 <mark>.6</mark> 4	4 <mark>.3</mark> 5	8.39	1.00	0.84	0.21	4.00	2.07	1.67	1.15	2.01	2 <b>.0</b> 7	0.58	12.15	2.82
	MDP2 -	0.26	0.35	1 <b>.8</b> 2	0.48	2 <mark>.1</mark> 0	3 <mark>.7</mark> 3	1.68	0.78	1.52	3.27	1.26	0.77	5.56	1.68	3 <b>.3</b> 7	0.12	7.88	4.88
	GMP -	2 <mark>.6</mark> 0	2 <mark>.4</mark> 4	1 <b>.8</b> 2	1.48	3 <mark>.2</mark> 8	7.70	0.46	3.43	0.84	4.08	0.96	3.80	3.79	2.35	4.29	2.66	2.86	1.28
	Eo_B_Mast -	2 <mark>.1</mark> 0	1.41	1. <b>2</b> 5	1 <b>.6</b> 6	1.43	1 <b>.9</b> 4	1. <b>7</b> 9	1.37	1.24	1.27	0.60	1.21	1.08	1. <b>2</b> 5	3 <b>.1</b> 6	6.87	0.94	0.69
Type	HIST -							10.16	0.04	1.85	0.01		0.03	0.05			0.02	0.02	0.02
Cell	ERP -	0.31	0.34	0.07	0.29		0.04	2.01	1.50	2.05	0.20		0.71	1.74	0.19	1.26	0.20	0.51	0.06
	MEP -	26.55	<mark>29.33</mark>	6.30	6.43	6.27	7.23	6.31	16.73	4.35	10.64	5.67	17.07	14.50	15.63	20.18	7.55	9.38	13.95
	MKP -	2.05	6 <mark>.1</mark> 0	3 <mark>.8</mark> 5	1 <b>17</b> 9	3 <mark>.6</mark> 7	3 <mark>.0</mark> 8	0.87	1.62	2.39	2.66	1.95	5.51	6.58	3.02	4.76	1.78	2.99	6.38
	MLP -	52.03	45.37	47.50	9.31	20.93	26.19	0.68	7.92	0.69	5.33	59.84	46.79	11.41	16.73	10.01	3.51	24.54	36.32
	HSC6 -	1 <mark>.7</mark> 2	0. <b>2</b> 4	2 <mark>(3</mark> 5	0.88	1.18	0. <mark>9</mark> 3	0.17	1.00		0.94	0.99	0.59	0.81	0.91	0.64	0.04	2.81	2.95
	HSC5 -	1. <mark>3</mark> 2	0. <b>7</b> 8	0.74	0.12	5 <mark>.3</mark> 8	1.11	17.47	2.37	21.58	0.42	3.80	0.76	0.51	1.10	1.01	1. <b>3</b> 6	3.02	0.86
	HSC4 -	0.02	0.05	0.01	0.04	0.82	0.20	44.39	1.01	36.26	0.19	5.13	0.92	25.10	0.19	3 <b>.9</b> 5		0.03	0.24
	HSC3 -	0.07	1.10	0.04	54.83	2 <mark>.9</mark> 2	6 <mark>.8</mark> 9	0.15	1.05		0.88	0.10	1.28	0.18	2.68	0.66	0.59	0.18	0.11
	HSC2 -	0.04	5 <mark>.3</mark> 4	0.04	3.02	0.71	1 <b>.7</b> 6	0.44	20.59		43.07	0.41	2.46	8.04	28.48	9.47	6.63	0.16	0.49
	HSC1 -	1. <mark>0</mark> 3	2 <mark>.2</mark> 3	0.82	2 <mark>.3</mark> 7	32.48	18.97	0.68	34.77	0.05	14.84	9.56	7.29	16.55	17.88	29.19	2.12	12.91	18.78
		G-C	SF	Pleri	xafor	Motix	afortide		G-C	SF		PI	erixafo	r+G-CS	SF	Moti	xaforti	de+G-	CSF
		SF1	SF2	or1	or2	Je1	de2	10C	202	003	204	or1	or2	or3	or4	de 1	de2	de3	de4
		ö	ö	xaf	xaf	ortic	ortic	cek	cek	cek	cet	xaf	xaf	xaf	xaf	ortic	ortic	ortic	ortic
MLP=r	nulti-lineage progenito	or ()	G	leri	leri	(afc	cafc	Ыа	Pla	Pla	Pla	leri	leri	leri	leri	afc	afc	afc	(afc
MEP=r HSC=ł	megakaryocyte/erythro nematopoietic stem ce	oid proge	enitor	С_	С_	otix	otix		+	+	+	Ц Н	<u>с</u> +	<u>с</u> +	Ц Н	otix	otix	otix	otix
MDP=i	monocyte/dendritic ce	ll progen	nitor			Š	Š	CS	CS	CS	CS	Ŝ	SF	Ŝ	SF	Ň,	Ň+	+M(	Ň,
GMP=	granulocyte monocyte	progeni	tor					Ģ	Ģ	Ģ	Ģ	С С	С С	С С	С С	SF.	Ϋ́Γ	SF.	SF.
ERP=e LMPP= PSC=c	erythroid progenitor =lymphoid multipotent patient specific cluster	progeni	tor						S	ampl	е	U	J	0	U	С С	0- O	0- 0	С С

**Extended Data Fig. 7** | **Transcriptionally Defined HSPC Subsets with Distribution by Cohort and Mobilization Regimen.** Average cell-type proportions of each transcriptionally defined HSPC subset annotated by sample. Samples are grouped by mobilization regimen and cohort.





Extended Data Fig. 8 | See next page for caption.

**Extended Data Fig. 8** | **TNFα/NFκB, EGF and HBO1 (KAT7) Gene Scores.** For three different pathways a list of genes was curated, derived from GSEA and the literature, to evaluate changes in expression of related pathways (Methods). Each Gene Score dot plot shows the average gene expression score indicated by heat map color and overlayed with the numerical score. Dot size indicates the

percentage of cells expressing each transcriptional program. Each plot is also separated by mobilization regimen (left axis); by multiple myeloma (M) and healthy allo-donor cohorts (H) (right axis); and by the HSC1-6 or MLP population of interest (bottom axis).

#### Extended Data Table 1 | Correlative Study Demographics for GENESIS Trial and Contemporaneous MM Cohort

	GENESIS Tria	l Cohorts	Prospective, Parallel Correlative Study Cohort
	Motixafortide + G-CSF (N=24)	Placebo + G-CSF (N=13)	Plerixafor + G-CSF (N=14)
Mean age (+/- SD), y	63 (8.6)	58.6 (9.8)	68.5 (7.5)
Male sex, N (%)	18 (75)	10 (76.9)	11 (79)
Race/Ethnicity, N (%)			
African American	4 (16.7)	1 (7.7)	3 (21)
Asian	1 (4.2)	0 (0)	0 (0)
Caucasian	19 (79.2)	12 (92.3)	10 (71)
Median time from Dx to consent, mo	4.0	4.0	n/a
IMWG Response at screening	g, N (%)		
sCR	1 (4.2)	0 (0)	0 (0)
CR	4 (16.7)	1 (7.7)	7 (50)
VGPR	14 (58.3)	10 (76.9)	4 (29)
PR	5 (20.8)	2 (15.4)	3 (21)
Median # induction cycles (+/- SD)	4 (0.8)	4 (0.9)	4 (0.4)
Lenalidomide-containing induction, N (%)	24 (100)	13 (100)	14 (100)
Prior radiotherapy, N (%)	5 (20.8)	2 (15.4)	1 (7)

Among MM patients whose CD34+ HSPCs underwent additional correlative immunophenotypic and transcriptional profiling, similar demographics were observed between those mobilized on the GENESIS Trial with motivafortide+G-CSF (N=24) or placebo+G-CSF (N=13), and those mobilized with plerixafor+G-CSF (N=14) on the prospectively enrolled, parallel correlative study.

## Extended Data Table 2 | GENESIS Trial Primary and Secondary Endpoints

	Motixafortide + G-CSF	Placebo + G-CSF	OR (95% CI)	P-value	Plerixafor + G-CSF
% collecting $\geq 6 \times 10^6$ CD34+ cells/kg in $\leq 2$ apheresis	92.5%	26.2%	53.3 (14.1, 201.3)	<0.0001	78.6%
% collecting ≥6 x 10 <sup>6</sup> CD34+ cells/kg in 1 apheresis	88.8%	9.5%	118 (25.4, 549.4)	<0.0001	50.0%
% collecting ≥2 x 10 <sup>6</sup> CD34+ cells/kg in 1 apheresis	96.3%	64.3%	18.90 (4.5, 80.0)	<0.0001	100.0%
Median # of CD34+ cells/kg collected in 1 apheresis	10.8 x 10 <sup>6</sup> (range 0.5-39.4)	2.25 x 10 <sup>6</sup> (range 0.2-10.6)	-	-	5.47 x 10 <sup>6</sup>
Median # of CD34+ cells/kg infused for ASCT	5.2 x 10 <sup>6</sup> (range 2.2-26.7)	3.3 x 10 <sup>6</sup> (range 2.0-14.8)	-	-	3.14 x 10 <sup>6</sup>
Median time to neutrophil engraftment (days)	12 (95% Cl 11-12)	12 (95% Cl 11-12)	Not Estimable	0.9554	13
Median time to platelet engraftment (days)	18 (95% Cl 17-19)	17 (95% Cl 17-18)	0.95 (0.2, 5.7)	0.9554	18.5
Graft durability (D+100)	92.2%	91.9%	1.04 (0.2, 4.5)	0.96	-
	Motixafortide + G-CSF	Placebo + G-CSF	HR (95% CI)	P-value	
PFS (% at 1 year; median)	92.5%; NR	90.5%; NR	0.735 (0.2, 2.6)	0.6328	-
OS (% at 1 year; median)	97.5%, NR	100%; NR	Not Estimable	0.998	-

Primary and secondary endpoints for the GENESIS Trial stratified by mobilization regimen. The proportion of patients collecting to the stated CD34+ cell/kg targets are presented with odds ratio (OR), 2-sided 95% confidence intervals (CI) and p-values based on CMH analysis on an intent-to-treat basis. Median numbers of CD34+ cells/kg collected and infused are presented as median values with range (minimum, maximum); but without comparative statistics as this was not a pre-specified analysis. Time-to-engraftment was assessed by Kaplan-Meier analysis and is presented as the median time to event with 2-sided 95% CI and p-values. Graft durability was assessed by CMH, as previously described, and presented with OR, 2-sided 95% CI and p-value. Progression free survival (PFS) and overall survival (OS) was assessed using Cox's proportional hazard's model and is presented as proportion not meeting event at 1-year with hazard ratio (HR), two-sided CI and p-values. A hierarchical method for multiple endpoint testing was utilized to control for multiple comparisons. The final column represents the apheresis and clinical outcomes observed in the contemporaneous MM cohort mobilized with plerixafor+G-CSF on a separate, prospective protocol conducted in parallel to the GENESIS Trial. Statistical comparisons between primary/secondary endpoints for patients mobilized on the GENESIS Trial and those mobilized with plerixafor+G-CSF were not pre-planned; nor performed.

# nature portfolio

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Last updated by author(s): Feb 18, 2023

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	$\boxtimes$	For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
	$\boxtimes$	Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated
		Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.

# Software and code

Policy information about availability of computer code

Data collection	OnCore clinical research management software v1.0.0 was used at Washington University.
Data analysis	Commercially available R, SPSS, SAS and Xcel statistical and graphics software packages were used for data analysis and graphics. Monocle3 (https://cole-trapnell-lab.github.io/monocle3/) was used for pseudotime analysis. Analysis was completed following standard tutorial for constructing single-cell trajectories (https://cole-trapnell-lab.github.io/ Bioconda, (The Bioconda Team et al., 2018), https://bioconda.github.io/ Bioconductor v3.9, (Huber et al., 2015), https://bioconductor.org/ Cell Ranger v6.0.1, 10X Genomics, https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/what-is-cell-ranger data.table_1.12.6, R Development Core Team, https://cran.r-project.org/package=data.table dendsort_0.3.3, (Sakai et al., 2014), https://cran.r-project.org/package=dendsort dplyr_0.8.5, R Development Core Team, https://cran.r-project.org/package=dplyr ggplot2_3.3.2, R Development Core Team, https://cran.r-project.org/package=ggplot2 gridExtra_2.3, R Development Core Team, https://cran.r-project.org/package=gridExtra magrittr_1.5, R Development Core Team, https://cran.r-project.org/package=magrittr Matrix_1.2-17, R Development Core Team, https://cran.r-project.org/package=magrittr Matrix_1.2.17, R Development Core Team, https://cran.r-project.org/package=magrittr Matrix_1.2.17, R Development Core Team, https://CRAN.R-project.org/package=magrittr Matrix_1.2-17, R Development Core Team, https://CRAN.R-project.org/package=pheatmap Python v3.7, Python Software Foundation, https://www.rython.org/ R v3.6, R Development Core Team, https://CRAN.R-project.org/package=RColorBrewer reshape2_1.4.3, R Development Core Team, https://CRAN.R-project.org/package=RColorBrewer reshape2_1.4.3, R Development Core Team, https://CRAN.R-project.org/package=RColorBrewer reshape2_1.4.3, R Development Core Team, https://CRAN.R-project.org/package=RColorBrewer

Seurat v3.1.2 and v4.0.3, (Butler et al., 2018), https://cran.r-project.org/web/packages/Seurat single-cell analysis code, (https://github.com/reykajayasinghe/WashU\_Genesis\_Study). sva v3.40.0, (Huber et al., 2015), https://bioconductor.org/packages/release/bioc/html/sva.html stringr\_1.4.0, R Development Core Team, https://cran.r-project.org/package=stringr Subread v2.0.1, (Liao et al., 2013), https://sourceforge.net/projects/subread/ Tidyverse, (Wickham et al., 2019), https://www.tidyverse.org/ viridis\_0.5.1, R Development Core Team, https://github.com/sjmgarnier/viridis viridisLite\_0.3.0, R Development Core Team, https://github.com/sjmgarnier/viridis

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# Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

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# Human research participants

Policy information about studies involving human research participants and Sex and Gender in Research.

Reporting on sex and gender	The findings in this study should be applicable to all sexes and genders. Neither sex nor gender were criteria for participation in this clinical trial and there were no pre-specified or post-hoc analyses of these data based solely on sex or gender. Patient sex was recorded for demographic purposes and was self-reported by study participants. Multiple myeloma occurs more commonly in male sex as compared to female sex, thus the slightly higher proportion of males enrolled on study is expected and is representative of the overall demographics of the study population.
Population characteristics	Eligible patients to be enrolled on this study included those ages 18-78 years with a diagnosis of multiple myeloma in their 1st or 2nd complete or partial remission (per IMWG criteria) with an ECOG performance status of 0-1 and adequate organ function (defined in the protocol) who were otherwise considered to be eligible for autologous hematopoietic stem cell transplantation.
Recruitment	Research participant recruitment may vary based on institutional and individual provider practices. Therefore, there is the possibility for recruitment heterogeneity and selection-bias to occur on multi-institutional studies such as the GENESIS Trial. Patients on the GENESIS Trial were recruited at participating institutions based on the presence of objective criteria, including the requisite diagnosis of biopsy proven multiple myeloma and a set of pre-defined eligibility criteria in order to minimize selection bias. To the authors knowledge, all patients meeting these objective criteria were offered participation in the study by their treating providers. All patients who were willing to participate and who provided informed consented were formally screened and, if determined to be eligible, were enrolled on study. Therefore, while the authors cannot entirely rule out the possibility of selection bias and other forms of recruitment bias from impacting the results of this study, it is likely that the objective nature of the inclusion/exclusion criteria applied uniformly across participating sites along with the randomized, placebo-controlled, double-blinded nature of the study may have minimized the impact of such potential biases.
Ethics oversight	Local IRB/IEC as well as central IRB and Central EC approval of the study protocol and amendments, investigator brochure and amendments, informed consent and any other documents provided to the subject was required at all participating sites (See list below of site-specific IRB details). Site 1001 IRB: Washington University Institutional Review Board (IRB) 4590 Children's Place, Suite 2300 St. Louis, MO 63110 Site 1002 IRB: University of Miami IRB Sylvester Comprehensive Cancer Center Fox Building, Suite 300, 1550 NW 10th Avenue Miami, Florida 33136

Site 1003: Mayo Clinic IRB 200 First St SW Rochester, MN 55905

Site 1005: University of Oregon Oregon Health & Science University IRB Knight Clinical Research 3485 SW Bond Ave, Mailcode OC13CT Portland, OR 97239

Site 1006: Human Research Protection Program University of Kansas Medical Center IRB Fairway North Office Building at 4330 Shawnee Mission Parkway, Suite 3170, Fairway, KS 66205

Site 1007: University of Maryland IRB 620 W. Lexington St., Second Floor Baltimore, MD 21201

Site 1008: Western Cooperative Group IRB 1019 39th Ave SE, Suite 120 Puyallup, WA 98374

Site 1009: The Loyola University Chicago Health Sciences Campus IRB 1101 Wootton Parkway, Suite 200 Rockville, MD 20852

Site 1011: Mayo Clinic IRB 200 First Street SW Rochester, MN, USA

Site 1012: UCLA IRB 10889 Wilshire Blvd, Suite 830 Los Angeles, CA 90095-1406

Site 1013: University of Utah IRB Research Administration building 75 South 2000 East Salt Lake City, UT 84112

Site 1014: Western Cooperative Group IRB 1019 39th Ave SE, Suite 120 Puyallup, WA 98374-2115

Site 1015: University of Cincinnati IRB University Hall, Suite 300 P.O. Box 210567 Cincinnati, OH 45221-0567

Site 1016: The Weill Cornell Medical College IRB 1300 York Avenue, Box 89 New York, NY 10065

Site 1017: UT M.D. Anderson Cancer Center IRB 7007 Bertner Avenue, Unit 1637 Houston, TX 77030

Site 4041: Ethics Committee Catania 1 Azienda Ospedaliero-Universitaria "Policlinico-Vittorio Emanuele" Catania Via Santa Sofia 78 – 95123 Catania

Site 4042: Ethics Committee South Reggio Calabria Division Grande Ospedale Metropolitano "Bianchi-Melacrino- Morelli" Via Provinciale Spirito Santo, Palazzo Gangeri, 24 89128 - Reggio Calabria Site 5051: University of Cologne Ethics Committee 50931 Cologne

Site 8081: University of Debrecen, Clinical Center, Clinic of Internal Medicine, Hematology Scientific Council for Health - Ethical Committee for Clinical Pharmacology Széchenyi István tér 7-8. Roosevelt Office Building II. floor 214 office H-1051 Budapest, Hungary

Site 8082:

Central Hospital of Southern Pest, National Institute of Hematology and Infectious Diseases Scientific Council for Health - Ethical Committee for Clinical Pharmacology Széchenyi István tér 7-8. Roosevelt Office Building II. floor 214 office H-1051 Budapest, Hungary

All patients provided written informed consent prior to enrollment on the study. The study was conducted in accordance with the Guideline for Good Clinical Practice ICH E6(R2)- ICH Harmonized Guideline Integrated Addendum to ICH E6 (R1) (International Conference on Harmonization of Technical Requirements for the Registration of Pharmaceuticals for Human Use), Step 5, 14 June 2017; the Declaration of Helsinki: Seoul, 2008; the US Code of Federal Regulations (Title 21, CFR Part 11, 50, 54, 56 and 312) and/or EU Directives; and/or local country regulations and guidelines.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

# Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences Behavioural & social sciences Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see <u>nature.com/documents/nr-reporting-summary-flat.pdf</u>

# Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	Sample size determination for Part 2 of the GENESIS study was based on plerixafor public domain data (DiPersio et al 2009, DOI 10.1182/ blood-2008-08-174946). The primary study endpoint of the study was the proportion (%) of subjects mobilizing $\geq$ 6.0 x 10^6 CD34+ cells/kg with up to 2 apheresis sessions in preparation for auto-HCT after G-CSF + single administration of motixafortide/placebo. Subjects were randomized to treatment with motixafortide+G-CSF or to placebo+G-CSF using a 2:1 randomization ratio, respectively. According to DiPersio et al, it is assumed that the response rate of subjects randomized to treatment with G-CSF+Placebo will be 35%. It was also assumed that the minimal effect size required, in terms of proportions difference, is 35%. Accordingly, it was assumed that the response rate of motixafortide treated subjects will be 70% or more. In addition, a conservative assumption was made that 20% of the subjects randomized to treatment with BL-8040 will not adhere to treatment for whatever reason (e.g. early termination, lack of compliance) and therefore will have a placebo- like response rate of 35%. This assumption was made to ensure that the study will not be underpowered. Accordingly, the assumed combined response rate of 63% for subjects randomized to treatment with BL-8040 reflects the assumption that 80% of the BL-8040 treated subjects will have a response rate of 70%, while the remaining 20% of subjects (i.e., those who did not adhere to the planned treatment regimen) will have a response rate of 35%. According to these assumptions, a total of 147 subjects provides 89.8% power to at a two-sided alpha of 0.0466256. Incorporating an additional withdrawal rate of 20%, the sample size of Part 2 of the GENESIS study is inflated by 20% and therefore the study was designed to randomize a total of 177 subjects. The power at interim analysis (to be conducted at alpha=0.0108678), assuming an attrition rate of 20% and success rates of 70% and 35% for BL-8040 + G-CSF or placebo + G-CSF, respectively
Data exclusions	The data was analyzed on an intent-to-treat principle, therefore no data were excluded after patients were enrolled and randomized to treatment on protocol. The correlative studies were performed only on samples collected from patients treated at Washington University, therefore patients treated on the GENESIS Trial protocol at other institutions were excluded from these analyses, by definition.
Replication	Replication was not performed on clinical specimens from the GENESIS Trial nor correlative studies involving human subjects. This was due to the finite number of HSPCs per sample, the limited number of aliquots of apheresis product available from each patient and the need to reserve the maximum possible amount of apheresis product for clinical use for patients HSPC infusion.
Randomization	Randomization was performed in a 2:1 fashion on the GENESIS Trial. Patients mobilized on the correlative protocol were treated according to standard of care, with no randomization needed/performed. However, patients mobilized on the correlative protocol were demographically similar to patients in the GENESIS Trial in terms of co-variates relevant to HSPC mobilization, including age, diagnosis, number of prior lines of therapy and lenalidomide exposure. In this way, co-variates between the randomized, double-blind placebo-controlled cohorts on the GENESIS Trial and the parallel correlative study were controlled for to the degree possible given the experimental design.
Blinding	This was a double-blinded, placebo-controlled clinical trial, with both investigators and patients blinded to study treatment allocation until completion of the trial and database lock, as per the supplemental GENESIS Trial Protocol and SAP.

# Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems	Methods
n/a Involved in the study	n/a Involved in the study
Antibodies	ChIP-seq
Eukaryotic cell lines	Flow cytometry
Palaeontology and archaeology	MRI-based neuroimaging
Animals and other organisms	
Clinical data	
Dual use research of concern	

# Antibodies

Intibodies	
Antibodies used	For local laboratory CD34+ enumeration at each participating clinical site, CD34+ enumeration was performed in accordance with standard clinical protocols at the local institution, which commonly involve adherence to ISHAGE guidelines (PMID: 8817388) in a CLIA-certified or a commensurately credentialed lab based on the specific site location/country. For central laboratory CD34+ enumeration at each participating clinical site, CD34+ enumeration of apheresis and peripheral whole blood samples was performed using validated methods in accordance with standard clinical protocols using ISHAGE guidelines (PMID: 8817388) in two CLIA-certified central labs based on site location (EU or USA). In these assays, the following fluorochrome-labeled surface-marker antibodies were used: The BD Stem Cell Enumeration Kit from BD Biosciences (catalog #344563) uses antibodies to CD45 (clone 2D1) and CD34 (clone 8G12). For the correlative extended CD34+ immunophenotyping by flow, the following fluorochrome-labeled antibodies were used (clone, source designated and catalog number in parenthesis): CD45-BUV395 (HI30, BD Biosciences, catalog #563792), CD123-BUV737 (7G3, Biosciences, catalog #741769), CD49f-BV421 (GoH3, BioLegend, catalog #313624), CD14-BV650 (MSE2, BioLegend, catalog #301836), CD45RA-BV785 (HI100, BioLegend, catalog #304140), CD34-VioBright515 (REA1164, Miltenyi, 1:120), CD10-PECF594 (HI10a, Biosciences, catalog #562396), CD38-PE-Cy7 (HIT2, BioLegend, catalog #303516), CD90-APC (5E10, BioLegend, catalog #328114), CD303-APCVIO770 (REA693, Miltenyi Biotech, catalog #130-120-517), CD184-PE (1D9, BD, catalog #551510). and CD184-PE (12G5, Biosciences, catalog #555974). Also see Supplemental Table 1: Antibodies for correlative CD34+ HSPC immunophenotyping.
Validation	The BD Stem Cell Enumeration Kit is a direct immunofluorescence-based three-color flow cytometric in vitro diagnostic assay (https://www.bdbiosciences.com/content/dam/bdb/products/global/reagents/flow-cytometry-reagents/clinical-diagnostics/ multicolor-cocktails-and-kits-ivd-ce-ivds/344563_base/pdf/23-22014.pdf). All antibodies for correlative studies were purchased from commercial vendors (BD Biosciences, BioLegend, or Miltenyi Biotech) and used prior to their expiration date. All commercial antibodies underwent quality control testing before distribution as detailed online (https://www.bdbiosciences.com/en-us/products/ reagents/flow-cytometry-reagents/research-reagents/quality-and-reproducibility, https://www.biolegend.com/en-us/quality/quality-control, https://www.miltenyibiotec.com/US-en/products/macs-antibodies/Antibody-production-development-and-quality-

control.html#gref). Lot specific Certificates of Analyses are available online for each antibody listed in Supplemental Table 1 (https:// regdocs.bd.com/regdocs/qcinfo, https://www.biolegend.com/en-us/certificate-of-analysis, https://www.miltenyibiotec.com/US-en/

# Clinical data

Clinical trial registration	NCT03246529
Study protocol	The full GENESIS Trial Protocol has been submitted as a supplemental document with this manuscript.
Data collection	Patients were enrolled from January 22, 2018 through October 30, 2020 at 18 sites in 5 countries (USA, Italy, Hungary, Spain and Germany). These sites include: Washington University School of Medicine in St. Louis, Siteman Cancer Center; University of Miami, Sylvester Comprehensive Cancer Center; Mayo Clinic, including Rochester, MN and Jacksonville, FL sites; University of Oregon Oregon Health & Science University; University of Kansas Medical Center; University of Maryland, Marlene and Stewart Greenebaum Comprehensive Cancer Center; University of Florida, Loyola University Chicago Health Sciences Campus, University of California Los Angeles, School of Medicine; University of Utah, Huntsman Cancer Institute; University of Rochester Medical Center, University of Cincinnati, Weill Cornell Medical College, UT M.D. Anderson Cancer Center, Azienda Ospedaliero-Universitaria "Policlinico-San Marco"; Grande Ospedale Metropolitano "Bianchi-Melacrino- Morelli"; University Hospital of Cologne; Ramon y Cajal University Hospital; 12Hospital University 12 De Octubre; Hospital de la Santa Creu i Sant Pau; University of Debrecen, Clinical Center; and Central Hospital of Southern Pest.
Outcomes	The primary objective of the study was to demonstrate the superiority of one dose of motixafortide+G-CSF over placebo+G-CSF to mobilize $\geq 6.0 \times 10^{6}$ CD34+ cells/kg in up to 2 apheresis sessions in preparation for autologous hematopoietic cell transplantation (auto-HCT) in multiple myeloma subjects. This was assessed by the primary endpoint of the proportion of subjects mobilizing

resources/technical-documents/certificates.html#gref).

 $\geq$ 6.0x10^6 CD34+ cells/kg with up to 2 apheresis sessions in preparation for auto-HCT after G-CSF + single administration of BL-8040 or placebo + GCSF. The secondary objectives were to demonstrate the superiority of one dose of motixafortide+G-CSF over placebo +G-CSF to mobilize  $\geq$ 2.0x10^6 CD34+ cells/kg in 1 apheresis session; to demonstrate the superiority of one dose of motixafortide+G-CSF over placebo+G-CSF to mobilize  $\geq$ 6.0x10^6 CD34+ cells/kg in 1 apheresis session; to descriptively assess the comparability between the effects of motixafortide+G-CSF and placebo+G-CSF in time to neutrophil engraftment, platelet engraftment and the later of the two; and to descriptively assess the comparability between the effects of motixafortide+G-CSF and placebo+G-CSF on graft durability at 60 days, 100 days, as well as 6 and 12 months post-transplantation. These secondary endpoints were assessed by the proportion of subjects who collect  $\geq$ 2.0 x 106 CD34+ cells/kg in 1 apheresis session; the proportion of subjects who collect  $\geq$ 6.0 x 106 CD34+ cells/kg in 1 apheresis session; the proportion of subjects who collect  $\geq$ 2.0 x 106 CD34+ cells/kg in 1 apheresis session; the proportion of subjects who collect  $\geq$ 2.0 x 106 CD34+ cells/kg in 1 apheresis session; the proportion of subjects who collect  $\geq$ 6.0 x 106 CD34+ cells/kg in 1 apheresis session; the proportion of subjects who collect  $\geq$ 6.0 x 106 CD34+ cells/kg in 1 apheresis session; the proportion of subjects who collect  $\geq$ 6.0 x 106 CD34+ cells/kg in 1 apheresis session; the time from transplantation to platelet engraftment defined as the first of 3 consecutive measurements of platelet count  $\geq$ 20x10^9/L without platelet transfluxion support for 7 days following the conditioning regimen associated nadir; the time from transplantation to engraftment defined as the time to neutrophils and platelets engraftment, whichever comes later; and graft durability at 60 days, 100 days 6 months and 12 months post-transplantation, respectively.

# Flow Cytometry

#### Plots

Confirm that:

The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).

The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).

 $\square$  All plots are contour plots with outliers or pseudocolor plots.

A numerical value for number of cells or percentage (with statistics) is provided.

## Methodology

Sample preparation	For the local lab-based CD34+ enumeration, sample preparation was performed according to standard operating procedures. For the central lab-based CD34+ enumeration, sample preparation was performed according to CLIA-based standard operating procedures, including a lyse-no wash single platform flow cytometry method allowing for the direct measurement of absolute cell counts using BD TruCount <sup>™</sup> tubes. In the flow cytometry panel stem cells were identified based on CD34+ and CD45 expression. For the correlative flow cytometry studies, CD34+ HSPCs from N=51 patients (placebo+G-CSF N=13, plerixafor+G-CSF N=14, motixafortide+G-CSF N=24) were purified from apheresis product from apheresis day 1 via CD34+ immunomagnetic selection using an AutoMACS device (Miltenyi Biotech, Auburn, CA). CD34+ HSPCs were washed in phosphate-buffered saline (PBS) and stained 15 minutes at room temperature with a LIVE/DEAD Fixable Aqua Dead Cell Stain kit (Invitrogen, Carlsbad, CA). Cells were then washed in PBS supplemented with 0.5% bovine serum albumin and 2 mM EDTA and incubated for 10 min at room temperature with human Fc Block and Brilliant Stain Buffer (BD Biosciences; San Jose, CA). Samples were then incubated for 30 min at room temperature with pre-titrated saturating dilutions of the following fluorochrome-labeled antibodies (see Antibodies section for details). Fluorescence minus one controls were used to assess background fluorescence intensity and set gates for negative populations. After washing twice, samples were analyzed on a ZE5 (Bio-Rad, Hercules, CA) flow cytometer. Single stain compensation controls were obtained using UltraComp eBeads (Thermo Fisher Scientific) and data were analyzed using FCS Express (DeNovo Software, Pasadena, CA). The antibody-binding capacity (ABC) per cell of the different CD34+ HSPC subsets was determined for CD184 clones 12G5 and 1D9 using saturating concentrations of antibody and the Quantum Simply Cellular (QSC; Bangs Laboratories) system for fluorescence quantitation per the manufacturers
Instrument	Local Lab studies: local lab dependent. Central Lab studies: BD FACS Canto II Flow Cytometer (Becton Dickinson, Franklin Lakes, NJ, USA). Correlative studies: ZE5 flow cytometer (Bio-Rad, Hercules, CA)
Software	Local Lab studies: local lab dependent. Central lab studies: Sample Acquisition (BD FACSDiVa Software version 8 program of the instrument) analysis was performed by counting 150,000 CD45+ events or 300 seconds. Correlative studies: FCS Express (DeNovo Software, Pasadena, CA).
Cell population abundance	Cells were CD34 selected as detailed above via immunomagnetic selection. During the subsequent flow experiments on CD34 selected cells, CD34+ was gated on to exclude any potential remaining CD34 negative cells. The gating strategy for the correlative studies is further detailed in Extended Data Figure 3 within the submitted manuscript and as below. Cell population abundance for each CD34+ fraction is detailed in both absolute numbers and percentage of total in Figure 3, Extended Data Figure 4.
Gating strategy	Local lab studies: gating strategy for local lab-based CD34+ enumeration was performed per local standard operating procedures. Central lab studies: gating strategy for central lab-based CD34+ enumeration was performed per validated method protocol. After acquisition, samples were gated appropriately in adherence with ISHAGE and CLIA protocols/ guidelines. Correlative studies: ISHAGE guidlines were followed. In addition, fluorescence minus one controls were used to assess background fluorescence intensity and set gates for negative populations. After washing twice, samples were analyzed on a ZE5 (Bio-Rad, Hercules, CA) flow cytometer. Single stain compensation controls were obtained using UltraComp eBeads (Thermo Fisher Scientific) and data were analyzed using FCS Express (DeNovo Software, Pasadena, CA). The gating strategy for the correlative studies is further detailed in Extended Data Figure 3 within the submitted manuscript.

🔀 Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.