Highly potent human hematopoietic stem cells first emerge in the intraembryonic aorta-gonad-mesonephros region

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Hematopoietic stem cells (HSCs) are multipotent stem cells that emerge early during embryogenesis and maintain hematopoiesis throughout the entire lifespan of the organism (Dzierzak and Speck, 2008; Medvinsky et al., 2011). Although in various vertebrate species the first hematopoietic differentiation occurs in the yolk sac (Moore and Metcalf, 1970), a growing body of data suggests that the aorta-gonad-mesonephros (AGM) region plays a key role in the generation of definitive HSCs (Dieterlen-Lievre, 1975; Medvinsky et al., 1993; Müller et al., 1994; Medvinsky and Dzierzak, 1996; de Bruijn et al., 2002; Zovein et al., 2008; Boisset et al., 2010), possibly through hematopoietic transition of endothelial cells of the dorsal aorta, which has been most clearly shown in zebrafish (Bertrand et al., 2010; Kissa and Herbomel, 2010). The mouse AGM region is capable of autonomous initiation and expansion of HSCs (Medvinsky and Dzierzak, 1996; Cumano et al., 2001; Taoudi et al., 2008). The early umbilical cord and the placenta are also implicated in HSC development (de Bruijn et al., 2000; Gekas et al., 2005; Ottersbach and Dzierzak, 2005; Robin et al., 2009). In the mouse embryo, the first definitive HSCs are detected at approximately the same time in different tissues (Müller et al., 1994; de Bruijn et al., 2000; Kumaravelu et al., 2002; Gekas et al., 2005; Ottersbach and Dzierzak, 2005), thus their primary origin remains debatable (Medvinsky et al., 2011).

Hematopoietic stem cells (HSCs) emerge during embryogenesis and maintain hematopoiesis in the adult organism. Little is known about the embryonic development of human HSCs. We demonstrate that human HSCs emerge first in the aorta-gonad-mesonephros (AGM) region, specifically in the dorsal aorta, and only later appear in the yolk sac, liver, and placenta. AGM region cells transplanted into immunodeficient mice provide long-term high level multilineage hematopoietic repopulation. Human AGM region HSCs, although present in low numbers, exhibit a very high self-renewal potential. A single HSC derived from the AGM region generates at least 300 daughter HSCs in primary recipients, which disseminate throughout the entire recipient bone marrow and are retransplantable. These findings highlight the vast regenerative potential of the earliest human HSCs and set a new standard for in vitro generation of HSCs from pluripotent stem cells for the purpose of regenerative medicine.
It is thought that HSCs mature in intraortic cell clusters budding from the ventral wall of the dorsal aorta (Tavian et al., 1996, 1999; Jaffredo et al., 1998; Taoudi et al., 2008; Yokomizo and Dzierzak, 2010). The maturation of HSCs also occurs in deeper layers of the dorsal aorta (Rybtsov et al., 2011).

In this study, we describe spatiotemporal distribution of HSCs in the early human embryo and provide evidence that the AGM region is the first generator of highly potent HSCs in the human. Upon transplantation into immunodeficient mice, HSCs derived from the AGM region extensively propagate, migrate to different bones throughout the recipient, and provide progressively increasing long-term multilineage hematopoietic repopulation of the host animal with robust potential for retransplantation. Our study reveals that the first human HSCs possess an unprecedented self-renewal capacity. A better understanding of embryonic development of human HSCs may be instrumental for creating clinically relevant protocols for the production of HSCs from human embryonic and induced pluripotent stem cells (Kauffman, 2009).

RESULTS
Spatiotemporal distribution of HSCs in the early human embryo
AGM regions, yolk sacs, livers, umbilical cords, and placentas were obtained from human embryos between Carnegie stages (CSs) 12 and 17 (O’Rahilly and Müller, 1987). Cell suspensions prepared from these tissues were transplanted into irradiated NOD.Cg-Pkdcreid Il2rgtm1Wjl/Sz (NSG) mice (Shultz et al., 2005) within 3–4 h of termination of pregnancy. Cells from each individual tissue were split equally between two to four NSG recipient mice. In 27 independent experiments, HSCs were detected in 10 AGM regions and 3 yolk sacs by their capacity to provide human long-term (4–8 mo) multilineage hematopoietic repopulation and by their retransplantability into secondary recipients (Table I). In CS 14 and 15 human embryos, HSCs were detected exclusively in the AGM region (5 embryos). At CS 16 and 17, HSCs were found either solely in the AGM region (2 embryos) or both in the AGM region and the yolk sac (3 embryos). Thus, HSCs appear in the human AGM region at least 5 d earlier than in the yolk sac (Table I).

Out of 26 transplanted livers, none contained HSCs; however, two provided unilineage T cell engraftment (Table I and not depicted). In one case, exclusive T cell engraftment was also observed in a recipient of an AGM region (Table I and not depicted). In these three cases, human T cell contribution increased during the first 4–6 mo, and then decreased on a similar timescale (not depicted). Umbilical cords and placentas contained no HSCs at all stages tested (Table I).

Short tandem repeat (STR) analysis (Cotton et al., 2000) confirmed that the human cells detected in NSG recipient mice repopulated by AGM region, yolk sac, and liver cells originated from the donor embryo, with no evidence of contamination by maternal cells (Tables S1–S4).

The dorsal aorta is the source of HSC activity within the human AGM region
In the mouse embryo, definitive HSCs first appear in the dorsal aorta (de Bruijn et al., 2000; Taoudi and Medvinsky, 2007). We tested whether this is true for the human embryo. For these experiments, we used human embryos between CS 15 and 17. The human AGM region was subdissected into the dorsal aorta and the urogenital ridges, as previously described for the mouse AGM region (de Bruijn et al., 2000; Taoudi and Medvinsky, 2007). Cell suspensions prepared from these tissues were transplanted into irradiated NSG recipient mice. In 10 independent experiments performed, HSCs were detected in six dorsal aortas, but never in the urogenital ridges (see Materials and methods).

High-level hematopoietic repopulation by early embryonic HSCs
In NSG mice engrafted with human HSCs from the AGM region and the yolk sac, donor CD45+ hematopoietic cells first became detectable in the peripheral blood of recipients 1.5–3 mo after transplantation, with donor cell levels reaching

Table I. The first HSCs in the human embryo emerge in the AGM region

| Carnegie stage (days) | AGM region | Yolk sac | Liver | Umbilical cord | Placentas |
|----------------------|------------|----------|-------|----------------|----------|
| 12 (26)              | 0/2        | 0/2      | 0/2   | NA             | 1/2      |
| 13 (28)              | 0/2        | NA       | 0/1   | NA             | 1/2      |
| 14 (32)              | 1/7        | 0/7      | 0/7   | 0/4            | 2/7      |
| 15 (33)              | 4/5        | 0/3      | 0/5   | 0/4            | 1/2      |
| 16 (37)              | 2+ 1/6      | 1/4      | 0/6   | 0/6            | 1/3      |
| 17 (41)              | 3/5        | 2/4      | 2/5   | 0/3            | 1/3      |
| Total                | 10+ 1/27   | 3/20     | 2/26  | 0/17           | 7/19     |

Cell suspensions prepared from AGM regions, yolk sacs, livers, umbilical cords, and placentas obtained from CS 12–17 human embryos were individually transplanted into irradiated NSG mice. Data shown are the number of tissues that gave human hematopoietic repopulation compared to the total number of tissues transplanted. The numbers in parentheses indicate an approximate postovulatory gestational age in days accepted for each CS (O’Rahilly and Müller, 1987). NA, not assessed.

*Long-term embryonic multilineage repopulation.

*Long-term embryonic unilineage T cell repopulation.

*Transient maternal unilineage T cell repopulation.
repopulation in the peripheral blood by month 8 after transplantation (Notta et al., 2011).
expressed γδ TCRs (Fig. 2 C). Additionally, human AGM region– and yolk sac–derived HSCs produced CD94+CD3−NK cells and CD94+CD3+NKT cells at low frequencies (Fig. 2 D and Table III).

The percentage of human CD33+ myeloid cells in the recipient peripheral blood and spleen was ~20–25 times lower than that of human lymphoid cells; however, the recipient BM contained on average 30% (range, 13–70; n = 7) human granulocytes (CD33+CD66b+) and monocytes (CD33+CD14+; Fig. 2 E and Table III). Megakaryocytic lineage was also readily detectable by appearance of human FSC-H+CD41a+ platelets in the recipient peripheral blood (Fig. 2 F). The recipient BM contained ~8% human erythroid cells (CD235a+), of which one third coexpressed CD45, indicating their immature state (Fig. 2 G).

Additionally, the BM of engrafted recipients plated into methylcellulose medium supplemented with human cytokines gave rise to human erythroid colonies (burst-forming units–erythroid [BFU-E]; Fig. 2 H and Table IV), myeloid colonies (colony-forming units-myeloid [CFU-Myeloid]; Table IV), and mixed erythroid/myeloid colonies (CFU-Mix; Fig. 2 I and Table IV). The human origin of hematopoietic colonies was confirmed by flow cytometry analysis (not depicted).

**Early embryonic HSCs can be efficiently retransplanted**

We tested whether HSCs derived from the human AGM region and yolk sac can be serially transplanted. In all eight experiments with the AGM region and two experiments with the yolk sac, transplantation of BM from primary recipients into secondary recipients resulted in sustainable human long-term multilineage hematopoietic engraftment (Fig. 3). In contrast, secondary BM transplantations from all three recipients exhibiting long-term unilineage T cell repopulation were unsuccessful. This indicates that HSCs colonize the human embryonic liver after CS 17. T cell–restricted repopulation may originate either from previously undescribed long-term persisting embryonic T cell progenitors or from human counterparts of mouse lymphopoiesis-biased γ and δ subtypes of long-term repopulating hematopoietic cells, which show no capacity for retransplantation (Dykstra et al., 2007).

**Extensive amplification of early embryonic HSCs in primary recipients**

Because one HSC can efficiently restore hematopoiesis in vivo (Osawa et al., 1996; Notta et al., 2011), the frequency of HSCs can be estimated using a limiting dilution assay (Szilvassy et al., 1990). In 9 out of 10 experiments transplanting human AGM region cells into two to four NSG mice, one recipient per experiment was found engrafted with human hematopoietic cells. Both recipients were repopulated in only one experiment (not depicted). To estimate HSC numbers, we used software for small numbers of replicates and nonlinear situations (Hu and Smyth, 2009). Our data fit the single-hit Poisson model (χ2 = 0.108; P = 0.742), demonstrating consistency between all 10 experiments performed with the AGM regions containing HSCs. The model calculations showed that one human AGM region contains ~1.9 HSCs (95% CI, 1.0–3.3; Fig. 4 A). If all 23 CS 14–17 AGM regions are considered (including nonrepopulating ones), the frequency of HSCs is ~0.8 per one human AGM region, which is close to the number of definitive HSCs in the mouse AGM region (Kumaravelu et al., 2002).

Statistically, human hematopoietic repopulation in each engrafted NSG mouse must have arisen mainly from one human HSC. The progressive growth in levels of repopulation with human hematopoietic cells (Fig. 1) may either be the result of gradual enhancement of productivity of this one HSC or, alternatively, the result of amplification of HSC numbers in recipients. To address this issue, the BM of engrafted primary recipients was harvested 4–8 mo after transplantation in five independent experiments. BM from an individual recipient was transferred into 6–15 secondary recipients. In these experiments, 41 out of 44 secondary recipients transplanted were found engrafed with human HSCs. Because each primary recipient...

**Table III. Multilineage hematopoietic engraftment in primary recipient mice**

| Recipient no. | Blood | BM | Spleen | Thymus |
|---------------|-------|----|--------|--------|
|               | B     | T  | NK     | CD4+ | CD8− | CD4+CD8− | CD4+CD8+ | CD4+CD8− | CD4−CD8+ |
| 1             | 44    | 39 | 11     | 18   | 19   | 12       | 7        | 9        | NA      |
| 2             | 67    | 23 | 3      | 4    | NA   | NA       | NA       | NA       | NA      |
| 3             | 19    | 65 | 5      | 11   | 16   | 42       | 1        | 38       | 29      |
| 4             | 77    | 5  | 4      | 12   | 51   | 0.1      | 0.2      | 45       | 87      |
| 5             | 75    | 19 | 2      | 4    | 62   | 1        | 0.3      | 37       | 90      |
| 6             | 56    | 25 | 9      | 9    | 61   | 1        | 9        | 20       | 82      |
| 7             | 17    | 36 | 31     | 18   | 20   | 4        | 1.5      | 74       | 41      |
| 8             | 59    | 22 | NA     | 15   | 4    | NA       | NA       | 1        | 11      |
| 9             | 63    | 19 | NA     | 14   | 42   | 12       | 2        | 43       | 46      |
| 10            | 26    | 60 | 9      | 6    | 42   | 12       | 2        | 43       | 46      |

| % in gated human CD45+ cells |
|------------------------------|
| NA                          | NA | NA       | NA   |
| 18                          | NA | NA       | NA   |
| 19                          | NA | NA       | NA   |
| 12                          | NA | NA       | NA   |
| 14                          | NA | NA       | NA   |
| 20                          | NA | NA       | NA   |
| 42                          | NA | NA       | NA   |
| 43                          | NA | NA       | NA   |
| 46                          | NA | NA       | NA   |
| 46                          | NA | NA       | NA   |
| 46                          | NA | NA       | NA   |

Data shown are the percentage of B (CD19+), T (CD3+ or CD4+CD8+), CD4+CD8+, and CD8−CD4−, NK (CD94+CD3−), and myeloid cells (CD33+ or CD13+) in the human CD45+ cell population in the peripheral blood, BM, spleen, and thymus of NSG mice transplanted with AGM region and yolk sac cells. The recipient identification numbers are the same as in Table II. NA, not assessed.
recipient was engrafted with approximately one human HSC, these results suggest that human HSCs propagated in primary recipients.

We considered an alternative possibility that the human AGM region contained some HSCs that remained dormant in primary recipients but could be activated upon secondary transplantations. To test this, we selected nonrepopulated primary recipients (Fig. 5 A, mice a and b) from five independent experiments in which at least one recipient was engrafted with AGM region cells (Fig. 5 A, mouse c). The entire BM from two coxal bones, two femurs, and two tibiae was collected from these nonrepopulated primary recipients and transferred into secondary recipients. None of secondary recipients showed human hematopoietic engraftment, confirming that the AGM region contained only one or two HSCs. Secondary transplantations of BM from nonengrafted recipients transplanted with yolk sac, liver, umbilical cord, and placental cells were also unsuccessful (two to five independent experiments were performed for each tissue).

To estimate the number of daughter HSCs generated in primary recipients by a single HSC derived from the AGM region, we performed two independent secondary transplantation experiments using the limiting dilution assay. The BM from successfully engrafted primary recipients was harvested from the sternum, humeri, ulnae, radii, coxal bones, femurs, and tibiae, and then pooled and transplanted in serial dilutions into secondary recipients. In experiment 1, the lowest dose transplanted per secondary recipient was 1/75 of total BM from primary recipient. All secondary recipients showed human long-term multilineage hematopoietic engraftment. In experiment 2, the transplanted dose was decreased further to 1/300 of total BM per recipient. Again, all secondary recipients showed human long-term multilineage hematopoietic repopulation (Fig. 4 B). However, in NSG mice transplanted with the lowest BM dose the level of human contribution, as determined in the peripheral blood 2 mo after transplantation, was <3%. Although we did not reach the limiting dose, we were close to the extinction of HSC activity. Human HSCs were also detected in spleens of primary recipients (Fig. 4 C). Thus, one HSC from the human AGM region is capable of generating at least 300 daughter HSCs upon transplantation. Previous studies have demonstrated a

Table IV. Human CFU-C frequency in the BM of primary recipient mice

| Recipient no. | Human CFU-C no. per 10,000 input BM nucleated cells |
|---------------|-----------------------------------------------------|
|               | BFU-E                  | CFU-Myeloid         | CFU-Mix                  |
| 1             | 2.3 ± 0.6              | 13.0 ± 3.8          | 2.5 ± 0.4                |
| 2             | NA                     | NA                  | NA                       |
| 3             | 8.7 ± 1.5              | 34.7 ± 11.2         | 1.7 ± 0.6                |
| 4             | 7.6 ± 1.4              | 26.1 ± 4.0          | 1.7 ± 0.2                |
| 5             | 5.0 ± 1.4              | 31.0 ± 1.4          | 0.5 ± 0.7                |
| 6             | 6.3 ± 0.1              | 16.3 ± 0.4          | 1.6 ± 0.3                |
| 7             | 6.5 ± 0.7              | 33.0 ± 4.2          | 1.5 ± 0.7                |
| 8             | NA                     | NA                  | NA                       |
| 9             | NA                     | NA                  | NA                       |
| 10            | 5.6 ± 0.6              | 15.4 ± 2.0          | 1.8 ± 0.3                |

Data shown are the number ± SD of human CFU-Cs per 10,000 input BM nucleated cells. The BM was obtained from NSG mice transplanted with AGM region and yolk sac cells. The recipient identification numbers are the same as in Table II. NA, not assessed, as the recipients died.
very limited self-renewal capacity by HSCs from the human UCB and adult BM (Guenechea et al., 2001; McKenzie et al., 2006; Liu et al., 2010). Although the microenvironment in immunodeficient mice is not optimal for the homeostatic expansion of human HSCs (Manz, 2007), human AGM region–derived HSCs extensively self-renew and amplify in these suboptimal conditions, suggesting that their expansion potential in physiological conditions is likely to be higher.

**Extensive dissemination of daughter HSCs in primary recipients**

The engraftment of one or two HSCs from the early human embryo results in considerable amplification of the HSC pool. We then tested if daughter HSCs stay at one site where the lodging of the parental HSC had occurred or if they migrate and disseminate throughout the recipient BM. We used two primary recipients repopulated with different AGM regions. We harvested BM from each recipient from the following six bones: two coxal bones, two femurs, and two tibiae (Fig. 5 A, mouse c). BM cell suspensions from each bone were transplanted separately into secondary recipients (Fig. 5 A, mice 1–6). In experiment 1, all secondary recipients showed human long-term multilineage hematopoietic engraftment (Fig. 5 B). Consistent with this, in experiment 2, all secondary recipients except one were repopulated with human hematopoietic cells. Thus, daughter HSCs generated by one or two human AGM region–derived HSCs disseminate broadly throughout the entire recipient BM.

**Daughter HSCs are of the canonical phenotype**

Using conventional FACS, it is not technically feasible to determine the immunophenotype of one or two HSCs present in one human AGM region. However, robust amplification of early embryonic HSCs in primary recipients allowed us to determine the immunophenotype of daughter HSCs. The BM harvested from the sternum, humeri, ulnae,
T cell repopulation of maternal origin (Table I, Table S5, and not depicted). The placenta contains maternal blood, and T cells from human blood can readily expand in immuno-deficient mice (van Rijn et al., 2003). An older 15-wk-old placenta readily provided long-term multilineage hematopoietic repopulation of fetal origin in all three recipients (each received \( \approx 1/30 \) of total placental cells; Table S6 and not depicted). Thus, in contrast to the early CS 12–17 placenta, the midgestation human placenta does possess HSC activity. In the mouse embryo, definitive HSCs appear in the placenta concomitantly with the AGM region (Gekas et al., 2005; Ottersbach and Dzierzak, 2005).

**DISCUSSION**

Although the embryonic development of HSCs has been studied in various vertebrate species (Medvinsky et al., 2011), the generation of the first HSCs during human embryogenesis remains a largely unexplored topic. The present concept of the embryonic development of the human hematopoietic system is primarily based on morphological and in vitro studies (Tavian et al., 1996, 1999, 2001; Oberlin et al., 2002). By using functional in vivo long-term repopulation assays, we conducted systematic analysis of the spatiotemporal distribution of HSCs in the early human embryo. We have shown that the AGM region, more specifically the dorsal aorta, is the first generator of HSCs in the human. Interestingly, although in the mouse embryo definitive HSCs appear concomitantly in the AGM region, yolk sac, umbilical cord, and placenta (Müller et al., 1994; de Bruijn et al., 2000; Kumaravelu et al., 2002; Gekas et al., 2005; Ottersbach and Dzierzak, 2005), in the human embryo, the timing of HSC appearance in these sites is clearly radii, coxal bones, femurs, and tibiae of primary recipients engrafted with a single HSC from the human AGM region was sorted into four cell populations based on expression of human CD34 and CD38 antigens, and then transplanted into secondary recipients (Fig. 6 A). In two independent experiments, human HSCs were found exclusively within CD34+CD38−/lo cell fraction (Fig. 6 B). Thus, HSCs derived from the early human embryo generate numerous HSCs of the canonical phenotype (Bhatia et al., 1997; McKenzie et al., 2006).
resolved, possibly because of considerably longer development of the human embryo.

Our data indicate that early embryonic HSCs possess a considerably higher self-renewal capacity compared with UCB and adult BM HSCs (Guenechea et al., 2001; McKenzie et al., 2006; Liu et al., 2010). Upon transplantation into NSG mice, a single HSC from the human AGM region produces at least 300 daughter HSCs during a period of 8 mo; this unprecedented self-renewal capacity has not been previously described for human or for mouse AGM region–derived HSCs. It is likely because of this that the percentage of human leukocytes in the recipients’ peripheral blood progressively increased, reaching up to 90% of total blood leukocytes. To achieve the same effect with UCB or adult BM, considerably higher numbers of HSCs need to be transplanted (Liu et al., 2010). More recent data suggest that ~40% blood chimerism can be achieved 8 mo after transplantation of 10–20 UCB HSCs per one NSG mouse (Notta et al., 2011).

Using stochastic analyses, it has been shown that adult human HSCs replicate every 40 wk (Catlin et al., 2011). Our findings indicate that the extensive expansion of HSCs from the early human embryo occurs through replication every 3.5–4 wk or even more frequently, considering that HSCs also produce mature blood cells. This enormous regenerative capacity of early HSCs is consistent with them being founders of the adult hematopoietic system in the human.

We have shown that within the human AGM region HSCs emerge in the dorsal aorta. According to current views, it is possible that HSCs emerge in intraaortic cell clusters budding from the floor of the dorsal aorta (Jaffredo et al., 1998; Taoudi et al., 2008; Yokomizo and Dzierzak, 2010). In human, formation of such CD34+CD45+ cell clusters occurs between CS 12 and 16 (Tavian et al., 1996, 1999). Using limiting dilution analysis, we have shown that one human AGM region contains between one and three HSCs; however, the number of intraaortic cell clusters is much higher (Tavian et al., 1996, 1999), suggesting that not every cluster contains HSCs. Furthermore, although HSCs are still detected at CS 17, intraaortic cell clusters are reported to disappear by that stage (Tavian et al., 1996, 1999), indicating that the link between HSCs and hematopoietic cell clusters needs to be elucidated further. Notably, the maturation of definitive HSCs in the mouse embryo also occurs underneath the luminal layer of the dorsal aorta (Rybtsov et al., 2011).

As expected, HSC activity in the AGM region precedes that in the liver, which emerges there only after CS 17. Our data highlight the importance of in vivo long-term repopulation assays, as previous immunohistological and in vitro studies have suggested that HSCs appear in the human embryonic liver around CS 13 (Tavian et al., 1999). Recent transplantation experiments have shown that HSCs colonize the human embryonic liver around week 7–8 of development, which follows the stages we studied (Oberlin et al., 2010).

We demonstrate that in the human, in contrast to the mouse (Gekas et al., 2005; Ottersbach and Dzierzak, 2005), placenta acquires HSC activity long after the emergence of HSCs in the AGM region, indicating that in the human, placenta might be a secondary site for HSCs. Our data vary from a recent study’s observation that the human placenta harbors HSCs from week 6 of development (as postmenstrual gestational age was employed, it is comparable with CS 12 or 13; Robin et al., 2009). In that study, NOD.CB17-Pikdels (NOD/SCID) recipient mice, which are less receptive to human HSCs than NSG mice (McDermott et al., 2010), were used. The authors used a highly sensitive PCR analysis to detect repopulation with 6–8-wk-old placenta, identifying one human cell in 100,000 mouse cells. In the absence of flow cytometry analysis, a persistence of small numbers of some placenta-derived nonhematopoietic cells in recipients could not be excluded. However, the presence of multipotent HSCs in the placenta was convincingly shown by flow cytometry starting from week 9 of development (Robin et al., 2009).
In summary, we provide a systematic spatiotemporal picture of HSC emergence in the early human embryo and identify the AGM region as the primary source of HSCs with unprecedented self-renewal capacity. Future comparative studies of early human embryonic HSCs and their fetal, neonatal, and adult counterparts may shed light on the mechanisms of HSC self-renewal and aging.

MATERIALS AND METHODS

Human embryonic and fetal tissues. Human embryonic and fetal tissues were obtained immediately after elective termination of pregnancy. The procedure was performed using midtrimester and misoprostol. The study was approved by the Lothian Research Ethics Committee. Each patient gave informed consent in writing. The developmental stage of human fetuses was determined according to the Carnegie staging system (O’Rahilly and Müller, 1987). The postmenstrual gestational age of human fetuses was determined using medical ultrasonography. Tissue dissection was performed in buffer 1 (PBS with Ca++ and Mg++ [Sigma–Aldrich] supplemented with 7% heat-inactivated FBS [PAA Laboratories], 100 IU/ml of penicillin [InVitrogen], and 100 µg/ml of streptomycin [InVitrogen]). The dissection procedure was similar to that previously described for the mouse embryo (Medvinsky et al., 2008). For the enzymatic treatment of the human AGM region, yolk sac, liver, and umbilical cord, 1 mg/ml of collagenase/dispase (Roche), and 0.12 mg/ml of DNase I (Roche) were added to buffer 1 and incubated at 37°C for 40 min. For the enzymatic treatment of placental tissue, two previously published protocols were employed (Bárcena et al., 2009; Robin et al., 2009). In the majority of experiments with the human placenta, we omitted the Ficoll-Paque density gradient separation step to minimize cell losses. By the end of enzymatic treatment, tissue digests were washed with buffer 2 which, compared with buffer 1, was made with Ca++ and Mg++-free PBS. For preparation of placental cells, in contrast to other tissues, 2 mM of EDTA was added to this buffer. After the first wash with buffer 2, tissue digests were pipetted gently to prepare a single-cell suspension. Human AGM region, yolk sac, liver, and umbilical cord cells were washed an additional time with buffer 2. Placental cells were passed through a 40-µm nylon cell strainer (BD) and washed four more times. Finally, cells were resuspended in buffer 2 and kept on ice until transplantation. Throughout the manuscript “independent” refers to those experiments performed with different human embryos.

Long-term repopulation assays. NSG mice were used as recipients for human HSCs. Animals were housed and bred within the University of Edinburgh, Edinburgh, Scotland, UK. Experiments with animals were approved by the Animal Welfare Committee of the University of Edinburgh and were performed according to the provisions of the Animal (Scientific Procedures) Act 1986 under the project license granted by the Home Office. All researchers who performed experiments with the mice held personal licenses granted by the Home Office. Mice were kept under specific pathogen-free conditions. Acidified (pH 3.0) autoclaved water supplemented with 1.67 mg/ml of neomycin (Invitrogen) and γ-irradiated chow diet was provided ad libitum. Animals were exposed to 14-h light and 10-h dark cycle. Up to 6 h before transplantation with human cells, 6–8 wk-old female NSG mice received a sublethal total body irradiation dose of 3.5 Gy at a rate of 0.75 Gy/min from a 137Cs source (GSR D1 γ-iradiator; Gamma-Service Medical). Animals were transplanted with cells intravenously via the tail vein. The number of transplanted embryonic cells was expressed in embryo equivalents (e.e., defined as a unit of cells equivalent to the number present in one tissue. Starting from 6–8 wk after transplantation, NSG mice were bled by tail vein nicking every 1–2 mo, and human HSC contribution was assessed by flow cytometry analysis after labeling cells with anti-human CD45 antibody. At the end of each experiment (usually 4–8 mo after transplantation), all recipients (including those found nonengrafted after peripheral blood analysis) were killed by dislocation of the neck according to the Schedule 1 of the Animal (Scientific Procedures) Act 1986, and their BM was collected. BM cells were labeled with anti-human CD45 antibody, and flow cytometry analysis was performed to detect the presence of human hematopoietic cells. The peripheral blood, BM, spleen, and thymus of engrafted NSG mice were further analyzed by flow cytometry analysis. The BM and spleen from primary recipients were used in secondary transplantation experiments.

Recipient mouse tissues. To collect recipient mouse tissues, buffer 2 supplemented with 2 mM of EDTA was used. NSG mice transplanted with human AGM region, yolk sac, liver, umbilical cord, placental cells, or BM from primary recipients were bled by tail vein nicking, and erythrocytes were lysed using Pharm Lyse Buffer (BD) according to the manufacturer’s instructions. To analyze hematopoietic tissues, the spleen, thymus, long bones, coxal bones, and sternum were obtained. Spleen and thymus were gently mashed in a small volume of the buffer through a 40-µm nylon cell strainer. The BM was flushed out from the bones, and a single-cell suspension was prepared by gentle pipetting. A cell fraction associated with endosteum was obtained from remaining bones after treatment with collagenase/dispase and DNase I at 37°C for 40 min and added to the flushed-out BM fraction. The cells were spun down, resuspended in buffer 2, and kept on ice until transplantation. To calculate the proportion of total BM transplanted into secondary recipients, we used previously reported data on the distribution of BM cells in different mouse bones (Boggs, 1984).

Human CFU-C assay. Human CFU-Cs in the BM of NSG recipient mice were detected by plating in triplicates 10,000–25,000 BM nucleated cells in the MethoCult H4034 Optimum Medium (StemCell Technologies) according to the manufacturer’s recommendations. The BM from a non-transplanted NSG mouse was used as a negative control. Cells were incubated at 37°C in 5% CO2 and 95% humidity for 14 d. Hematopoietic colonies were enumerated microscopically. Some individual colonies were subject to flow cytometry analysis after labeling hematopoietic cells with anti–human CD45 and CD235a and anti-mouse CD45 and Ter119 antibodies.

Antibodies, flow cytometry analysis, and FACS. The following mouse anti-human monoclonal antibodies (all from BD) were used for flow cytometry analysis and/or FACS: CD3-APC, PE, and PerCP (clones SK7 and SP34-2), CD4-APC and APC-Cy7 (clone RPA-T4), CD8-PE and PE-Cy7 (clone RPA-T8), CD11b-PE-Cy7 (clone ICRF44), CD13-APC (clone WM15), CD14-APC and APC-Cy7 (clones M5E2 and M6P9), CD19-PE (clone HIB19), CD33-PE (clone W/MA), CD34-APC (clone 8G12), CD36-PE (clone HIT2), CD41a-FITC (HP8), CD45-Biotin, FITC, and V450 (clone HI30), CD66b-FITC (clone G10F5), CD94-APC (clone HP-3D9), CD235a-APC (clone GA-R2), IgM-Biotin (clone G20-127), αβ TCR-FITC (clone WT31), γδ TCR-PE (clone 11F2). Appropriate isotype controls, streptavidin-APC, rat anti-mouse CD45-APC (clone 30-F11), and Ter119-PE and FITC were also purchased from BD. The Human FcR Blocking Reagent (Milteny Biotech) and anti-mouse CD16/32 purified monoclonal antibody (clone 93; eBioscience) were used to prevent unwanted binding of antibodies to Fc receptors. All of the aforementioned antibodies and reagents were used at final concentrations recommended by manufacturers or determined by in-house titration. Cell labeling and washes were performed in Ca++ and Mg++-free PBS supplemented with 2% heat-inactivated FBS. For dead cell exclusion the buffer was supplemented with 1.5 µg/ml of 7-amino-acetoxyamine (eBioscience). A FACSCalibur (BD), LSXFortessa (BD), or CyAn ADP (Dako) instruments were used for flow cytometry analysis. FACS and postsort purity check were performed on a FACSAnnexII instrument (BD). Sorted cells were spun down, resuspended in buffer 2, and kept on ice until transplantation. Flow cytometry data were analyzed with FlowJo v7.6.1 software (Tree Star).

STR analysis. Genomic DNA was extracted from peripheral blood, spleen, and/or BM cells obtained from NSG mice repopulated with human hematopoietic cells and from embryonic or fetal donor tissues. For this, the QIAamp DNA Investor Kit (QIAGEN) and a QIAcube robotic instrument (QIAGEN) were used. DNA quantification was performed with the Plexor HY System (Promega) and an Mx3500P real-time PCR machine (Strategene).
DNA quantification values were determined using the Plexor Data Analysis Software (Promega). DNA samples and PCR positive and negative controls were amplified with the AmpFISTR SGM Plus PCR Amplification Kit (Applied Biosystems). PCR products were detected on a 3130 Genetic Analyzer (Applied Biosystems). Handling of raw data and genotyping were performed with GeneMapper ID v3.2.1 software (Applied Biosystems). All the reagents and instruments listed above were used according to manufacturers’ recommendations. DNA match probabilities were calculated based on allele frequencies in Scottish Caucasian population.

Limiting dilution analysis. Limiting dilution analysis was performed using ELDA software (Hu and Smyth, 2009) available at http://bioinf.wehi.edu.au/software/elda/.

Online supplemental material. Tables S1–S6 show STR analysis data confirming either embryonic or maternal origin of hematopoietic repopulation in recipients transplanted with human embryo or placenta-derived cells. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20111688/DC1.

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REFERENCES

Bárcena, A., M. Kapidzic, M.O. Muench, M. Gormley, M.A. Scott, J.F. Weter, C. Ferhat, and S.J. Fisher. 2009. The human placenta is a hematopoietic organ during the embryonic and fetal periods of development. Dev. Biol. 327:24–33. http://dx.doi.org/10.1016/j.ydbio.2008.11.017
Bertrand, J.Y., N.C. Chi, B. Santosu, S. Teng, D.Y. Stainier, and D. Traver. 2010. Haematopoietic stem cells derive directly from aortic endothelium during development. Nature. 464:108–111. http://dx.doi.org/10.1038/nature08738
Bhatta, M., J.C. Wang, U. Kapp, D. Bonnet, and J.E. Dick. 1997. Purification of primitive human hematopoietic cells capable of repopulating immune-deficient mice. Proc. Natl. Acad. Sci. USA. 94:5320–5325. http://dx.doi.org/10.1073/pnas.94.10.5320
Boggs, D.R. 1984. The total marrow mass of the mouse: a simplified method of measurement. Am. J. Hematol. 16:277–286. http://dx.doi.org/10.1002/ajh.840160309
Boisvert, J.C., W. van Cappellen, C. Andreu-Soler, N. Galjart, E. Dzierzak, and C. Robin. 2010. In vivo imaging of haematopoietic cells emerging from the mouse aortic endothelium. Nature. 464:116–120. http://dx.doi.org/10.1038/nature08764
Cashman, J.D., T. Lapidot, J.C. Wang, M. Doedens, L.D. Shultz, P. Landsorp, J.E. Dick, and C.J. Eaves. 1997. Kinetic evidence of the regeneration of multilineage hematopoiesis from primitive cells in normal human bone marrow transplanted into immunodeficient mice. Blood. 89:4307–4316.
Oberlin, E., M. Fleury, D. Clay, L. Petit-Cocault, J.J. Candelier, B. Ottersbach, K., and E. Dzierzak. 2005. The murine placenta contains hematopoietic stem and progenitor cells throughout development. Cell Stem Cell. 5:385–395. http://dx.doi.org/10.1016/j.stem.2009.08.020

Rybtsov, S., M. Sobiesiak, S. Taoudi, C. Souillhol, J. Senserrich, A. Liakhovitskia, A. Ivanovs, J. Freamton, S. Zhao, and A. Medvinsky. 2011. Hierarchical organization and early hematopoietic specification of the developing HSC lineage in the AGM region. J. Exp. Med. 208:1305–1315. http://dx.doi.org/10.1084/jem.20102419

Shultz, L.D., B.L. Lyons, L.M. Burzenski, B. Gott, X. Chen, S. Chaleff, M. Kohl, S.D. Gillies, M. King, J. Mangada, et al. 2005. Human lymphoid and myeloid cell development in NOD/LtSz-scid IL2Rγ^null mice engrafted with mobilized human hematopoietic stem cells. J. Immunol. 174:6477–6489.

Szilvassy, S.J., R.K. Humphries, P.M. Landsorp, A.C. Eaves, and C.J. Eaves. 1990. Quantitative assay for totipotent reconstituting hematopoietic stem cells by a competitive repopulation strategy. Proc. Natl. Acad. Sci. USA. 87:8736–8740. http://dx.doi.org/10.1073/pnas.87.22.8736

Taoudi, S., and A. Medvinsky. 2007. Functional identification of the hematopoietic stem cell niche in the ventral domain of the embryonic dorsal aorta. Proc. Natl. Acad. Sci. USA. 104:9399–9403. http://dx.doi.org/10.1073/pnas.070098410

Tavian, S., C. Gonneau, K. Moore, J.M. Sheridan, C.C. Blackburn, E. Taylor, and A. Medvinsky. 2008. Extensive hematopoietic stem cell generation in the AGM region via maturation of VE-cadherin’CD45+ pre-definitive HSCs. Cell Stem Cell. 3:99–108. http://dx.doi.org/10.1016/j.stem.2008.06.004

Tavian, M., L. Coulombel, D. Lutton, H.S. Clemente, F. Dieterlen-Liévre, and B. Péault. 1996. Aorta-associated CD34+ hematopoietic cells in the early human embryo. Blood. 87:67–72.

Tavian, M., M.F. Hallais, and B. Péault. 1999. Emergence of intraembryonic hematopoietic precursors in the pre-liver human embryo. Development. 126:793–803.

Tavian, M., C. Robin, L. Coulombel, and B. Péault. 2001. The human embryo, but not its yolk sac, generates lympho-myeloid stem cells: mapping multipotent hematopoietic cell fate in intraembryonic mesoderm. Immunity. 15:487–495. http://dx.doi.org/10.1016/S1074-7613(01)00193-5

van Rijn, R.S., E.R. Simonetti, A. Hagenbeek, M.C. Hogenes, R.A. de Weger, M.R. Canninga-van Dijk, K. Weijer, H. Spits, G. Storm, L. van Bloois, et al. 2003. A new xenograft model for graft-versus-host disease by intravenous transfer of human peripheral blood mononuclear cells in RAG2−/− gamma−/− double-mutant mice. Blood. 102:2522–2531. http://dx.doi.org/10.1182/blood-2002-10-3241

Wang, J.C., M. Doedens, and J.E. Dick. 1997. Primitive human hematopoietic cells are enriched in cord blood compared with adult bone marrow or mobilized peripheral blood as measured by the quantitative in vivo SCID-repopulating cell assay. Blood. 89:3919–3924.

Yokomizo, T., and E. Dzierzak. 2010. Three-dimensional cartography of hematopoietic clusters in the vasculature of whole mouse embryos. Development. 137:3651–3661. http://dx.doi.org/10.1242/dev.051094

Zovein, A.C., J.J. Hofmann, M. Lynch, W.J. French, K.A. Turlo, Y. Yang, M.S. Becker, L. Zanetta, E. Dejana, J.C. Gasson, et al. 2008. Fate tracing reveals the endothelial origin of hematopoietic stem cells. Cell Stem Cell. 3:625–636. http://dx.doi.org/10.1016/j.stem.2008.09.018