Reduction of endoglin receptor impairs mononuclear cell-migration

Zhenying Han1,2, Sonali Shaligram1,2, Marie E. Faughnan3,4, Dewi Clark3, Zhengda Sun5, Hua Su1,2*

1Department of Anesthesia and Perioperative Care, University of California, San Francisco, CA 94143, USA
2Center for Cerebrovascular Research, University of California, San Francisco, CA 94143, USA
3Toronto HHT Centre, Division of Respirology, Department of Medicine, St. Michael’s Hospital, University of Toronto, Ontario M5B 1W8, Canada
4Li Ka Shing Knowledge Institute, St. Michael’s Hospital, University of Toronto, Ontario M5B 1W8, Canada
5Department of Radiology, University of California, San Francisco, CA 94143, USA

*Correspondence: Hua Su, Department of Anesthesia and Perioperative Care, University of California, San Francisco, 1001 Potrero Avenue, Box 1363, San Francisco, CA 94143, USA. hua.su@ucsf.edu

Academic Editor: Akiko Mammoto, Medical College of Wisconsin, USA

Received: March 15, 2020 Accepted: May 11, 2020 Published: June 29, 2020

Cite this article: Han Z, Shaligram S, Faughnan ME, Clark D, Sun Z, Su H. Reduction of endoglin receptor impairs mononuclear cell-migration. Explor Med. 2020;1:136-48. https://doi.org/10.37349/emed.2020.00010

Abstract

Aim: To test if the impairment of mononuclear cell (MNC) migration in patients with hereditary hemorrhagic telangiectasia (HHT) is due to the reduction of the endoglin (ENG) receptor on the cell surface and oxidative stress.

Methods: MNCs of HHT patients and normal controls were subjected to migration assay. Fractions of MNCs were pre-incubated with antibodies specific to HHT causative genes ENG [hereditary hemorrhagic telangiectasia type 1 (HHT1)] or activin receptor-like kinase 1 [ALK1, hereditary hemorrhagic telangiectasia type 2 (HHT2)], AMD3100 or Diprotin-A to block ENG, ALK1 C-X-C chemokine receptor 4 (CXCR4) or CD26 (increased in HHT1 MNCs) before migration assay. The MNCs were allowed to migrate toward stromal cell-derived factor-1α (SDF-1α) for 18 h. The expression of CXCR4, CD26, superoxide dismutase 1 (SOD1) and glutathione peroxidase 1 (GPX1) in MNCs and nitric oxide levels in the plasma were analyzed.

Results: Compared to the controls, fewer HHT1 MNCs and similar number of HHT2 MNCs migrated toward SDF-1α. Diprotin-A pre-treatment improved HHT1 MNC-migration, but had no effect on normal and HHT2 MNCs. Pre-incubation with an anti-ENG antibody reduced the migration of normal MNCs. Diprotin-A did not improve the migration of ENG antibody pre-treated MNCs. Anti-ALK1 antibody had no effect on MNC-migration. AMD3100 treatment reduced normal and HHT MNC-migration. ENG mRNA level was reduced in HHT1 and HHT2 MNCs. ALK1 mRNA was reduced in HHT2 MNCs only. CD26 expression was higher in HHT1 MNCs. Pre-treatment of MNCs with anti-ENG or anti-ALK1 antibody had no effect on CD26 and CXCR4 expression. The expression of antioxidant enzymes, SOD1, was reduced in HHT1 MNCs, which was accompanied with an increase of ROS in HHT MNCs and nitric oxide in HHT1 plasma.

Conclusions: Reduction of ENG receptor on MNC surface reduced monocyte migration toward SDF-1α independent of CD26 expression. Increased oxidative stress could alter HHT MNC migration behavior.
Introduction

Hereditary hemorrhagic telangiectasia (HHT) or Rendu-Osler-Weber disease is an autosomal dominant vascular disease that affects approximately 1 in 5,000 people [1-3]. HHT is characterized by the telangiectases on the skin and mucosa, as well as organ arteriovenous malformation (AVMs) and can be complicated by hemorrhage, stroke and heart failure [4]. HHT is classified into hereditary hemorrhagic telangiectasia type 1 (HHT1), hereditary hemorrhagic telangiectasia type 2 (HHT2), and JP (juvenile polyposis)-HHT, depending on the causative gene mutations. Endoglin (ENG, HHT1) and activin receptor-like kinase 1 (ALK1, HHT2) are major causative genes for HHT, with at least 90% of HHT patients having a mutation in one of these genes [5]. The AVM phenotype in HHT patients and mouse models, e.g., in the skin [6] and in the brain [7-9] is associated with an increased burden of inflammatory cells. AVMs in HHT likely develop as an abnormal response to injury; the role of inflammation in this proposed mechanism remains undefined.

There is evidence that ENG-deficiency impairs monocyte homing to the injury site [10]. Recruitment of human monocytes to the infarcted murine heart and subsequent vessel formation are severely impaired when HHT1 monocytes are used [11]. Decreased homing is linked with impaired ability of the monocytes to respond to stromal cell derived factor-1α (SDF-1α). The mechanism has been attributed to increased CD26, also known as dipeptidyl peptidase-4, expression in HHT1 monocytes, a moiety that cleaves the amino-terminal dipeptide from SDF-1α interfering with C-X-C chemokine receptor 4 (CXCR4) binding [10]. Inhibition of CD26 restores the homing ability of HHT1 monocytes. Interestingly, Eng-deficiency in endothelial cells reduces leukocyte adhesion and transmigration [12] and impairs the endothelial cell-autonomous capacity to up-regulate SDF-1α expression in response to hindlimb ischemic injury [13]. Although underpowered to detect a significant difference, we also found that there is a 10% reduction in monocyte homing to the angiogenic focus in mice with Eng+/− bone marrow (BM) compared to mice with wild-type BM, two weeks after injection of an adeno-associated viral vector expression vascular endothelial growth factor [14]. Taken together, Eng deficiency appears to impair monocyte adhesion and migration. Whether loss of ALK1 affects these processes is not known.

It remains unclear why AVM lesions demonstrate more macrophages if ENG-deficiency impairs monocyte migration. We found Eng+/− mice had fewer cluster of cell differentiation 68+ (CD68+) microglia/macrophages 3 days after ischemic stroke and more CD68+ microglia/macrophages 60 days after stroke in the peri-infarct areas [15]. We have also showed that after angiogenic stimulation, the Eng deficient mice had fewer CD68+ microglia/macrophages at 2 weeks, similar numbers at 4 weeks, and more at 8 weeks in the brain angiogenic region compared with wild-type mice. These data suggest that delayed macrophage clearance or prolonged microphage infiltration could be a mechanism. The chemotactic activity of SDF-1α not only involves CD26, but also the upregulated expression and activity of integrin family members, which involved in cell adhesion [16-18]. It is not clear if integrin family play roles in the defect of HHT monocyte migration. However, above data indicate that blocking delayed macrophage homing to an AVM lesion could be a therapeutic strategy to reduce AVM severity.

In addition, we have also found that in an in vitro angiogenic niches mimicked by the endothelial cells and vascular smooth muscle co-culture system, HHT CD34+ mononuclear cells (MNCs) are more likely to be differentiated into macrophages than normal CD34+ MNCs, suggesting a pro-inflammatory phenotype of HHT MNCs [9].

In this study, we tested if blocking ENG or ALK1 receptor on the surface of MNC reduces MNC chemotactic migration and if HHT MNCs have a defect in removing oxidative species. Most MNCs used in this study were isolated using Ficoll (StemCell Technology, Vancouver, Canada) density gradient centrifugation, which contain lymphocytes and monocytes. Previous studies showed that only 10-20% MNCs were monocytes, upon activation, both monocytes and lymphocytes expressed ENG protein at the cell surface [19-21].

Keywords
Endoglin, ALK1, hereditary hemorrhagic telangiectasia, mononuclear cells, migration, oxidative stress
Materials and methods

Isolation of MNCs

Blood collected from HHT patients (Table 1), 15 with known ENG mutation (HHT1), 11 with known ALK1 mutation (HHT2) and 10 with unknown mutations [used for analyzing reactive oxygen species (ROS) in HHT MNCs] were used in this study. Healthy age- and gender-matched volunteers (N = 29) were used as controls. Patients and controls were recruited at the Toronto HHT Centre, an HHT Centre of Excellence at St. Michael’s Hospital and the University of Toronto. All patients provided written, informed consent and the study protocol was approved by the Research Ethics Board of St. Michael’s Hospital. We have also purchased concentrated normal monocytes (Trima residuals) from Blood Center of the Pacific (San Francisco, CA).

MNCs that contain lymphocytes and 10-20% monocyte were isolated from 50 mL of blood by Ficoll density gradient centrifugation. Blood were diluted two times with phosphate buffered saline (PBS) plus 2% fetal bovine serum, laid on top of Ficoll and centrifuged at room temperature (15-25°C) for 30 min at 400 x g with brake off. The layer of MNCs was transferred to a sterile 15 mL centrifuge tube. Three volumes of PBS was added to resuspend the cells. The cell suspension was then centrifuged at 400 g for 10 min at 20°C. After removing the supernatant, MNCs were resuspended in 6 mL to 8 mL PBS and centrifuged at 400 g for 10 min at 20°C. After removing the supernatant, MNCs were suspended in cell culture medium (RPMI 1640 Glutamax medium, supplemented with 10% FBS) for migration assay, ROS assay or frozen for quantitative reverse transcription polymerase chain reaction (RT-PCR) analysis. MNCs collected from different HHT patients were used for migration assay and ROS assay (Table 1).

Table 1. HHT patients

| Patients | Genotype | Mutations | Used in studies |
|----------|----------|-----------|----------------|
| 16       | ALK1     | c.586A>G  | 1. Migration    |
| 24       | ALK1     | no information | 2. qPCR for ENG, ALK1, CD26 and CXCR4 |
| 25       | ALK1     | c.772+2C>G | 3. NOx in plasma |
| 32       | ALK1     | c.867_868delinsTT (p.Gln290X) | |
| 33       | ALK1     | c.1, 403T>G (p.Met468Arg) | |
| 36       | ALK1     | c.329C>A (p.Ser110X) | |
| 29       | ENG      | c.690-2A>T | |
| 30       | ENG      | deletion of exon 2 | |
| 37       | ENG      | c.1, 205dupA | |
| 38       | ENG (ENG family) | no information | |
| 44       | ENG      | c.115delA | |
| 28       | ENG      | no information | |
| 45       | ENG      | no information | |
| CT002    | ENG      | c.657-658delCA | qPCR for SOD1 and GPX1 |
| CT003    | ENG      | c.1, 082_1, 085del (p.Thr361Serfs) | |
| CT004    | ALK1     | c.1, 107_1, 108delAG (p.Arg369fs) | |
| CT005    | ENG      | c.20dupC (p.Leu8fs) | |
| CT006    | ENG      | c.38T>G (p.Leu13Arg) | |
| CT007    | ALK1     | c.1, 231C>T (p.Arg411Trp) | |
| CT008    | ALK1     | c.1, 120C>T (R374W) | |
| CT009    | ENG      | Familial mutation | |
| CT011    | ALK1     | c.430C>T (p.Arg144X) | |
| CT012    | ENG      | c.588G>A (W196X) | |
| CT013    | ENG      | c.657-658delCA | qPCR for SOD1 |
| CT016    | ALK1     | c.1, 435C>T (p.Arg144X) | |
| CT001    | ENG      | c.655C>G (p.His219Asp) | |

qPCR: quantitative polymerase chain reaction; NOx: nitrogen oxide; GPX1: glutathione peroxidase 1
Migration assay
The membrane of the insert and the bottom of the transwells that were used for migration assay were pre-coated with human fibronectin (20 g/mL, ThermoFisher Scientific, Waltham, MA). Migration medium (RPMI 1640 from ThermoFisher Scientific, supplemented with 0.5% BSA from Sigma Aldrich, St Louse, MO) was added to the inserts of transwells, and incubated at 37°C for 10 min to equilibrate the temperature. The inserts were then transferred to the lower chambers that contained SDF-1α (200 ng/mL, Sigma Aldrich). MNCs were seeded to the insert at a density of 1 × 10^5/cm² in 100 µL migration medium and allowed to migrate for 18 h at 37°C. Transwell inserts were removed. The excess medium on the bottom and side of the inserts was scraped into the bottom chambers. Cells in the lower chambers were resuspended through pipetting 20 times and then transferred to tubes. The lower chambers were washed with 300 µL PBS once. The PBS was added to tubes with cells collected from the bottom chamber. The tubes containing migrated cells were spun at 1,000 rpm for 5 min. After removing the supernatant, the cells were resuspended with 200 uL PBS and counted using a hemocytometer.

To test the effect of blocking cell surface ENG or ALK1 receptor, prior to the migration assay, MNCs were incubated with an anti-ENG antibody (10 μg/mL, Abcam, Cambridge MA) or an anti-ALK1 antibody (10 μg/mL, R&D Systems, Minneapolis, MN) for 1 h in migration medium. After incubation, the antibody-containing media were removed and the cells were replenished with fresh migration media.

As controls, MNCs were incubated with AMD3100 (5 μg/m, Sigma Aldrich) for 30 min to block CXCR4 receptor or Diprotin-A (5mM, Sigma Aldrich) for 15 min to block CD26 prior migration assay.

The phenotype of migrated cells was analyzed by fluorescent-activated cell sorting (FACS) using macrophage markers: CD11b and CD14, endothelial marker: CD31, and pan lymphocyte marker: CD3. Antibodies used in the FACS were anti CD14-PE-Cy7 (1:100), anti-CD11b-V450 (1:100), anti CD31-V450 (1:100), and anti CD3-FITC (1:100). All antibodies were purchased from BD Biosciences (San Jose, CA). BD CompBead Anti-Mouse Ig Kappa (BD Biosciences) incubated with each antibody was used to optimize fluorescence compensation settings for multicolor flow cytometric analysis. The samples were analyzed on a FACS LSR II (BD Biosciences), and percentage of each antibody-stained cells was calculated using DIVA V 6.1.3 software.

Quantitative RT-PCR analysis
The mRNA levels of CD26, CXCR4, ENG, ALK1, and two antioxidant enzymes, superoxide dismutase 1 (SOD1) and GPX1 in MNCs were quantified by quantitative real-time-PCR. Total RNA was extracted from MNCs using RNeasy Mini Kit (Qiagen, Valencia, CA) and reverse-transcribed into cDNA using SuperScript® III First-Strand Synthesis System (Invitrogen, Carlsbad, CA). Real-time PCR was performed using TaqMan Fast Advanced Master Mix (Applied Biosystems, Foster City, CA). Gene-specific primers and probes purchased from Applied Biosystems were used: CD26 (Hs00897386_m1), CXCR4 (Hs00607978_s1), ALK1 (ACVRL1, Hs00953798_m1), ENG (Hs00923996_m1), SOD1 (Hs00533490_m1), GPX1 (Hs01028922_g1), HPRT (Hs02800695_m1), and GAPDH (Hs02758991_g1). All samples were run in duplicates or triplicate, and relative gene expression was calculated using the comparative threshold cycle (CT) and normalized to GAPDH or HPRT (ΔCT).

Measurement of mitochondrial ROS level in MNCs and nitric oxide level in plasma
The blood cells and plasma were separated by centrifugation (3,000 g for 10 min). CD34+ monocytes were isolated using CD34-antibody-coated Dynabeads (Life Technologies, South San Francisco, CA) according to manufacturer’s instructions. Total ROS in CD34+ MNCs of HHT patients were measured using the Total ROS Detection Kit (Thermo Scientific) following the manufacturer’s protocols. ROS positive cells and fluorescent intensity were quantified by microscopic analysis. NOx in the plasma of HHT patients using Nitric Oxide Assay kit (Thermo Scientific) following the manufacturer’s protocols.
Statistical analysis

Data are represented as mean ± SD. Prism 6 (GraphPad Software Inc., La Jolla, CA) was used for all the statistical analyses in this study. Sample sizes are shown in figure legends. Two-way analysis of variance (ANOVA) followed by Tukey’s post hoc test was used to compare the differences of MNC migration among groups. One-way ANOVA followed by Tukey’s post hoc test was used to analyze the differences of gene expression. Student t-test was used to compare the differences when there are two testing groups involved. A P-value equals or smaller than 0.05 was considered significant.

Results

The number of HHT1 MNCs migration towards SDF-1α is reduced

MNCs were isolated from the blood of HHT1 and HHT2 patients, and age and sex matched controls. MNCs migration towards SDF-1α was analyzed in a transwell system. Two-way ANOVA analysis showed that the difference among groups was significant (P < 0.001). Compared to controls (100.0 ± 7.0 cells), there were fewer HHT1 MNCs (60.3 ± 6.8 cells) migrated to the lower chambers (where the media contained SDF-1α, P < 0.001, Tukey’s multiple comparisons). Similar numbers of HHT2 (94.5 ± 21.2 cells) and normal MNCs migrated into the lower chambers (P = 0.23). Blocking CXCR4 receptor by pre-treating the cells with AMD3100 was equally effective in blocking HHT1, HHT2 and normal MNCs migration toward SDF-1α (P < 0.001 vs. untreated cells). Block of CD26 through Diprotin-A pre-treatment increased the number of HHT1 MNCs in lower chambers (82.0 ± 13.0, P < 0.001). Diprotin-A had no effect on normal and HHT2 MNCs (90.5 ± 19.1 cells, P = 0.94, Figure 1). Therefore, MNC-migration toward SDF-1α is impaired in HHT1, but this can be rescued by blocking CD26 receptor.

![Figure 1. Reduction of HHT1 MNC migration toward SDF-1α. N = 8 for control MNCs; N = 7 for HHT1 MNCs; and N = 6 for HHT2 MNCs. *: P < 0.001 compared to control; &: P = 0.005 compared to untreated HHT-1](image)

Blocking ENG receptor on MNC-surface reduce their migration toward SDF-1α

To test if blocking ENG or ALK1 receptor on MNC-surface could influence MNC migration toward SDF-1α, MNCs isolated from the concentrated normal monocytes (Trima residuals) purchased from Blood Center of the Pacific (San Francisco, CA) were incubated with an anti-ENG antibody or an anti-ALK1 antibody before being subjected them to migration assay. Two-way ANOVA analysis showed that the difference among groups are significant (P < 0.001). Compared to MNCs pretreated with control IgG (100.5 ± 0.9 cells), there were fewer ENG antibody pretreated MNCs (33.6 ± 18.1 cells) migrated to lower chambers (P < 0.001, Tukey’s multiple comparisons test, Figure 2A). Reduction of CD26 by Diprotin A pretreatment did not improve the migration ability of ENG antibody treated MNCs (P > 0.999). Pre-treatment of MNCs with ALK1 antibody did
not affect MNC migration toward SDF-1α ($P = 0.998$, Figure 2B). These data indicate that the availability of ENG receptors on MNC-surface is crucial for MNC migration toward SDF-1α.

The phenotypes of migrated cells were analyzed by FACS. Monocytes were detected by antibodies specific to CD11b and CD14. Lymphocytes were detected by an antibody specific to CD3. Same numbers of monocytes (CD11b+, or CD14+, or CD11b++CD14+) and lymphocytes (CD3+) were migrated toward to SDF-1α. Interestingly, ENG antibody or ALK1 antibody treatment reduced monocytes migration, but have no effect on lymphocyte-migration (Figure 3). Due to the large variation and limited sample sizes, we could not detect any difference among groups.

Pre-treatment of MNCs with anti-ENG or anti-ALK1 antibody had no effect on CD26 and CXCR4 expression

To test if antibody treatment influences the expression of CD26 and CXCR4, the mRNA levels of these genes were analyzed by quantitative RT-PCR. As control, the expression of these genes in HHT1 and HHT2 MNCs were analyzed first. Similar to published data [10, 22], ENG gene expression was reduced in HHT1 ($P < 0.001$) and HHT2 ($P < 0.001$) MNCs (Figure 4A). ALK1 gene expression was reduced in HHT2 MNCs ($P = 0.03$, Figure 4B).
CD26 gene expression was significantly increased in HHT1 MNCs only (HHT1 versus control: \( P = 0.002 \); HHT2 versus control: \( P = 0.0997 \), Figure 4C). CXCR4 expression was not significantly different among control, HHT1 and HHT2 MNCs (\( P = 0.97 \), Figure 4D).

![Figure 4](image)

We next analyzed CXCR4 and CD26 expression in ENG or ALK1 antibody treated MNCs. We found ENG or ALK1 antibody treatment did not alter the expression of CD26 (\( P = 0.34 \)) and CXCR4 (\( P = 0.24 \), Figure 5).

![Figure 5](image)

**Antioxidant enzyme is reduced and ROS is increased HHT MNCs, which are associated with an increase of NOx in the plasma of HHT patients**

We showed previously that HHT MNCs are more likely to differentiate into pro-inflammatory macrophages in angiogenic environment than normal MNCs in an *in vitro* angiogenic niche [9]. Macrophage mitochondrial oxidative stress has been shown to promote nuclear factor-κB-mediated inflammation in macrophage [23]. To test if HHT MNCs pro-inflammatory phenotype have any association with oxidative stress, we analyzed the mRNA levels of antioxidant enzymes, SOD1 and GPX1. ANOVA analysis showed that the SOD1 level was different among HHT1, HHT2 and control MNCs (\( P = 0.04 \)). Multiple comparison showed that HHT1 MNCs
expressed lower levels of SOD1 than control MNCs ($P = 0.05$). The difference between HHT2 MNCs and control MNCs ($P = 0.19$), HHT1 and HHT2 ($P = 0.92$) did not reach statistical significance. Of note, the lack of difference between HHT2 and control MNCs could be due to the small sample size ($N = 5$) of HHT2 MNCs. ANOVA analysis did not show significant difference of GPX level among HHT1, HHT2 and control MNCs ($P = 0.19$). Figure 6).

Interestingly, we found that the ROS was increased in HHT MNCs ($P = 0.01$ for ROS+ cells and $P < 0.001$ for fluorescent intensity, Figure 6A and B). Since these patients were not genotyped, it is not clear if HHT1 and HHT2 MNCs have different levels of ROS. The NOx (NO2- and NO3-) level is increased in the plasma of HHT1 patients (Figure 6C) compared to the plasma collected from controls ($P < 0.001$) and HHT2 ($P = 0.02$). NOx level in HHT2 and control plasma had no difference ($P = 0.3$, Figure 7). These data suggest that reduced levels of antioxidant enzyme in HHT MNCs reduce their ability to remove oxidative species in MNCs and plasma, which may contribute to a pro-inflammatory phenotype.

Discussion

It has been hypothesized that AVM formation involves an impaired response to injury that may include the dysfunction of peripheral blood mononuclear cells [22, 24]. The human AVM phenotype in HHT in the skin [6] and in the brain [7, 25, 26] is associated with an increased burden of inflammatory cells. Brain AVM lesions in HHT mouse models also have many macrophages [8, 9, 25]. However, there is evidence that ENG-deficiency reduces monocyte homing to the injury site [10, 11], which was linked with impaired ability of the monocytes to respond to SDF-1α.

Before MNCs are able to contribute to repair, they need to home to and retain in ischemic and damaged tissue. Directed migration (homing) of MNCs following tissue damage is regulated by the SDF-1α. MNCs that express the CXCR4 migrate toward the tightly regulated gradient of SDF-1α. This directed migration of monocytes can be inhibited by CD26, a moiety that cleaves the amino-terminal dipeptide from SDF-1α interfering with CXCR4 binding [10]. MNCs of HHT1 patients express elevated levels of CD26 and show...
impaired homing toward damaged tissue [24]. Impaired homing capacity of the MNCs might therefore contribute to the impaired angiogenesis and tissue repair observed in HHT1 patients due to impaired resolution of inflammation [27]. However, if ENG-deficiency reduces monocyte migration, it is unclear why AVM lesions contain more macrophages. We have previously demonstrated a pro-inflammatory phenotype of HHT1 MNCs with delayed and prolonged MNC homing and delayed clearance [9, 28], which could be the underlying mechanisms for macrophages accumulated in AVM lesions.

Although ENG expression is reduced in HHT2 MNCs, their migration toward SDF-1α was not significantly reduced. The reduced migration of HHT1 MNCs has been attributed to their increased expression of CD26. However, CD26 level is not increased in HHT2 MNCs. The low CD26 levels on HHT2 MNCs did not correlate with their migratory behavior capacity [10]. Another explanation could be the small sample size and large variation of HHT2 MNCs prevent us from detecting the defect. The migration ability of HHT2 MNCs should be investigated further in future study.

In this paper, we demonstrated that reduction of available ENG receptors on MNC-surface by specific antibody treatment reduced normal MNC-migration toward SDF-1α. The antibody treatment did not change CD26 and CXCR4 mRNA expressions in MNCs, which indicate that the antibody effect is independent of CD26 expression. The low CD26 levels on normal MNCs did not correlate with their migratory behavior capacity [10]. Therefore, inhibition of CD26 could not restore the migration of ENG Ab treated MNCs. One limitation here is that we did not analyze the CD26 and CXCR4 protein on cells surface. Pre-treatment of MNCs with anti-ALK1 antibody did not reduce MNC-migration toward SDF-1α, suggesting that ENG may have a direct effect on monocyte migration and the mechanisms underlying the accumulation of monocytes/macrophages in HHT1 and HHT2 AVMs may not be identical and need further study. The chemotactic activity of SDF-1α also upregulate expression and activity of integrin family members, which are involved in cell adhesion [16-18]. The role of integrin family in the defect of HHT monocyte migration should also be investigated in future study.

The chemotactic activity of SDF-1α also upregulate expression and activity of integrin family members, which are involved in cell adhesion [16-18]. The role of integrin family in the defect of HHT monocyte migration should also be investigated in future study.

One clinical study has shown that alveolar exhaled NO is elevated in HHT patients [29]. Uncoupled eNOS activity resulting in impaired myogenic response has been reported to represent an early event in HHT pathogenesis related to oxidative stress [30-33]. Increased levels of free radicals including superoxide, NO, and peroxynitrite have been reported to mediate vascular damage [34-39]. Increased ROS has been demonstrated to be one of the mechanisms for non-resolving inflammation [40], the one mechanism demonstrated to have clear relevance to HHT in both Eng [31] and Alk1 deficiency [32]. However, it was not clear previously that if ROS increases in HHT MNCs and if increased ROS drives monocyte pro-inflammatory phenotype. We demonstrated here that the level of an antioxidant enzyme, SOD1 in HHT MNCs was reduced, which was accompanied with an increase of ROS in HHT MNCs. In addition, the NOx level is also increased in the plasma of HHT patients. Since macrophage mitochondrial oxidative stress have been shown to promote nuclear factor-κB-mediated inflammation in macrophage [23], the reduction of antioxidant enzyme, increase of ROS in HHT MNCs and the NOx in the plasma of HHT patients may contribute to the pro-inflammatory phenotype of HHT MNCs.

The role of inflammation in HHT disease remains poorly understood [24]. The up-regulation of ENG in activated monocytes is reduced in HHT1 and HHT2 patients [41]. Therefore, it can be postulated that ENG gene expression might be necessary for proper macrophage performance. This theory is supported by clinical reports of brain abscess, osteomyelitis, septic arthritis, recurrent Staphylococcus aureus and other infections in HHT patients [41, 42]. Animal study showed that deletion of Eng in macrophages impaired the phagocytic activity of peritoneal macrophages [43], this could help to explain the higher rate if infectious disease seen in HHT patients. As we have discussed in previous sections in this paper that HHT1 monocytes are not able to homing to injury/angiogenic sites in response to SDF-1α immediately after the injury [9-11, 15], which may delay the clearance the damaged tissue. However, they enter continually and stay in the injury/angiogenic site for an extended period leading to an unresolved inflammation and delayed tissue repair [9, 15]. We have also showed previously that HHT MNCs have a pro-inflammatory phenotype. In this study, we demonstrated that HHT MNCs have increased ROS, suggesting that they are undergo oxidative stress. All of these indicate
that reduction of ENG in HHT MNCs impairs their response to tissue injury and infection. Large prospective studies are needed to characterize the association of altered HHT MNC-function and risk of infection.

The alternation of signaling pathways in HHT MNCs that are responsible to their altered functions are largely unknown. It has been shown that *ENG* heterozygous endothelial cells exhibit reduced ALK1-Smad1/5 signaling. Unexpectedly, these cells adapted their ALK5 expression with a decrease of 80% and therefore, they also have reduced ALK5-Smad2/3 signaling [44, 45]. We found in this study that ENG expression is reduced not only in HHT1 MNCs but also reduced in HHT2 MNCs. Whether, reduction of ENG on HHT MNCs is the only factor that contribute to the impaired function of HHT MNCs or there is other pathways, such as ALK5 signaling involved need to be study in future.

In summary, we demonstrated here that reduction of the availability of ENG receptor on the surface of MNCs is a mechanism for the reduction of HHT1 MNC-migration to SDF-1α. In addition, HHT MNCs and plasma have reduced anti-oxidant gene expression and increased ROS, which could be responsible for a pro-inflammatory phenotype in HHT patients.

**Abbreviations**

Ab: antibody  
ALK1: activin receptor-like kinase 1  
ALK1-Ab: ALK1 antibody treated MNCs  
ANOVA: analysis of variance  
AVM: arteriovenous malformation  
CD68+: cluster of differentiation 68+  
CT: threshold cycle  
CXCR4: C-X-C chemokine receptor 4  
ENG: endoglin  
ENG-Ab: ENG antibody treated MNCs  
FACS: fluorescent-activated cell sorting  
GPX1: glutathione peroxidase 1  
HHT: hereditary hemorrhagic telangiectasia  
HHT1: hereditary hemorrhagic telangiectasia type 1  
HHT2: hereditary hemorrhagic telangiectasia type 2  
MNCs: mononuclear cells  
NOx: nitric oxides  
PBS: phosphate buffered saline  
ROS: reactive oxygen species  
RT-PCR: reverse transcription polymerase chain reaction  
SDF-1α: stromal cell derived factor-1α  
SOD1: superoxide dismutase 1

**Declarations**

**Author contributions**

MEF and HS contributed conception and design of the study; ZH and SS performed the experiments, collected data; ZH, SS and HS analyzed and performed the statistical analysis; MF and DC recruited patients and collected blood samples from patients; ZH and HS wrote the first draft of the manuscript. All authors contributed to manuscript revision, read and approved the submitted version.
Conflicts of interest
The authors declare that they have no conflicts of interest.

Ethical approval
All protocols involving human samples were approved by the Research Ethics Board of St. Michael’s Hospital, University of Toronto, and the Committee on Human Research of the University of California, San Francisco (UCSF), in accordance with the Code of Ethics of the World Medical Association (Declaration of Helsinki).

Consent to participate
Informed written consent for study participation was obtained from all patients and healthy volunteers.

Consent to publication
Not applicable.

Availability of data and materials
Not applicable.

Funding
This study was supported by research grants from the National Institutes of Health (R01 NS027713, R01 HL122774, R01 NS112819 and R21NS083788) and the Michael Ryan Zodda Foundation to HS. MEF was also supported by the Nelson Arthur Hyland Foundation and the Li Ka Shing Knowledge Institute.

Copyright
© The Author(s) 2020.

References
1. Bayrak-Toydemir P, McDonald J, Markowitz B, Lewin S, Miller F, Chou LS, et al. Genotype-phenotype correlation in hereditary hemorrhagic telangiectasia: mutations and manifestations. Am J Med Genet A. 2006;140:463-70.
2. Letteboer TG, Mager JJ, Snijder RJ, Koeleman BP, Lindhout D, Ploos van Amstel JK, et al. Genotype-phenotype relationship in hereditary haemorrhagic telangiectasia. J Med Genet. 2006;43:371-7.
3. Sabbà C, Pasculli G, Lenato GM, Supressa P, Lastella P, Memeo M, et al. Hereditary hemorragic telangiectasia: clinical features in ENG and ALK1 mutation carriers. J Thromb Haemost. 2007;5:1149-57.
4. Guttmacher AE, Marchuk DA, White RI. Hereditary hemorrhagic telangiectasia. N Engl J Med. 1995;333:918-24.
5. McDonald J, Wooderchak-Donahue W, VanSant Webb C, Whitehead K, Stevenson DA, Bayrak-Toydemir P. Hereditary hemorrhagic telangiectasia: genetics and molecular diagnostics in a new era. Front Genet. 2015;6:1.
6. Braverman IM, Keh A, Jacobson BS. Ultrastructure and three-dimensional organization of the telangiectases of hereditary hemorrhagic telangiectasia. J Invest Dermatol. 1990;95:422-7.
7. Chen W, Guo Y, Walker EJ, Shen F, Jun K, Oh SP, et al. Reduced mural cell coverage and impaired vessel integrity after angiogenic stimulation in the Alk1-deficient brain. Arterioscler Thromb Vasc Biol. 2013;33:305-10.
8. Choi EJ, Chen W, Jun K, Arthur HM, Young WL, Su H. Novel brain arteriovenous malformation mouse models for type 1 hereditary hemorrhagic telangiectasia. PLoS One. 2014;9:e88511.
9. Zhang R, Han Z, Degos V, Shen F, Choi EJ, Sun Z, et al. Persistent infiltration and pro-inflammatory differentiation of monocytes cause unresolved inflammation in brain arteriovenous malformation. Angiogenesis. 2016;19:451-61.
10. Post S, Smits AM, van den Broek AJ, Sluijter JP, Hoefer IE, Janssen BJ, et al. Impaired recruitment of HHT-1 mononuclear cells to the ischaemic heart is due to an altered CXCR4/CD26 balance. Cardiovasc Res. 2010;85:494-502.

11. van Laake LW, van den Driessche S, Post S, Feijen A, Jansen MA, Driessen MH, et al. Endoglin has a crucial role in blood cell-mediated vascular repair. Circulation. 2006;114:2288-97.

12. Rossi E, Sanz-Rodriguez F, Eleno N, Düwell A, Blanco FJ, Langa C, et al. Endothelial endoglin is involved in inflammation: role in leukocyte adhesion and transmigration. Blood. 2013;121:403-15.

13. Young K, Conley B, Romero D, Tweedie E, O’Neill C, Pinz I, et al. BMP9 regulates endoglin-dependent chemokine responses in endothelial cells. Blood. 2012;120:4263-73.

14. Choi EJ, Walker EJ, Degos V, Jun K, Kuo R, Pile-Spellman J, et al. Endoglin deficiency in bone marrow is sufficient to cause cerebrovascular dysplasia in the adult mouse after vascular endothelial growth factor stimulation. Stroke. 2013;44:795-8.

15. Shen F, Degos V, Chu PL, Han Z, Westbroek EM, Choi EJ, et al. Endoglin deficiency impairs stroke recovery. Stroke. 2014;45:2101-6.

16. Cheng M, Qin G. Progenitor cell mobilization and recruitment: SDF-1, CXCR4, alpha4-integrin, and c-kit. Prog Mol Biol Transl Sci. 2012;111:243-64.

17. Brilha S, Wysoczanski R, Whittington AM, Friedland JS, Porter JC. Monocyte adhesion, migration, and extracellular matrix breakdown are regulated by integrin alphaVbeta3 in mycobacterium tuberculosis infection. J Immunol. 2017;199:982-91.

18. Zolota Z, Koliakos G, Paletas K, Kaloyianni M. NHE-1 and beta1 integrin dependent monocyte adhesion and migration after glucose, insulin or PPARgamma stimulation. Cell Adh Migr. 2011;5:258-65.

19. O’Connell PJ, McKenzie A, Fisicaro N, Rockman SP, Pearse MJ, d’Apice AJ. Endoglin: a 180-kD endothelial cell and macrophage restricted differentiation molecule. Clin Exp Immunol. 1992;90:154-9.

20. Lastres P, Bellon T, Cabañas C, Sanchez-Madrid F, Acevedo A, Gougos A, et al. Regulated expression on human macrophages of endoglin, an Arg-Gly-Asp-containing surface antigen. Eur J Immunol. 1992;22:393-7.

21. Schmidt-Weber CB, Letarte M, Kunzmann S, Rückert B, Bernabéu C, Blaser K. TGF-(beta) signaling of human T cells is modulated by the ancillary TGF-(beta) receptor endoglin. Int Immunol. 2005;17:921-30.

22. Cannavicci A, Zhang Q, Dai SC, Faughnan ME, Kutryk MJ. Decreased levels of miR-28-5p and miR-361-3p and increased levels of insulin-like growth factor 1 mRNA in mononuclear cells from patients with hereditary hemorrhagic telangiectasia. Can J Physiol Pharmacol. 2019;97:562-9.

23. Wang Y, Wang GZ, Rabinovitch PS, Tabas I. Macrophage mitochondrial oxidative stress promotes atherosclerosis and nuclear factor-kappaB-mediated inflammation in macrophages. Circ Res. 2014;114:421-33.

24. Dingenouts CK, Goumans MJ, Bakker W. Mononuclear cells and vascular repair in HHT. Front Genet. 2015;6:114.

25. Chen W, Guo Y, Bollen AW, Su H, Young WL. Reduced PDGFR-beta expression after regional Alk1 deletion and VEGF stimulation in the brain is associated with reduced mural cell coverage. Stroke. 2012;43:A3169.

26. Chen W, Guo Y, Jun K, Wankhede M, Su H, Young WL. Alk1 deficiency impairs mural cell recruitment during brain angiogenesis. Stroke. 2013;44:ATMP118.

27. Peter MR, Jerkic M, Sotov V, Douda DN, Ardelean DS, Ghamami N, et al. Impaired resolution of inflammation in the endoglin heterozygous mouse model of chronic colitis. Mediators Inflamm. 2014;2014:767185.

28. Shen F, Degos V, Chu PL, Han Z, Westbroek EM, Choi EJ. Endoglin deficiency impairs stroke recovery. Stroke. 2014;45:2101-6.

29. Gupta S, Zamel N, Faughnan ME. Alveolar exhaled nitric oxide is elevated in hereditary hemorrhagic telangiectasia. Lung. 2009;187:43-9.

30. Toporsian M, Gros R, Kabir MG, Vera S, Govindaraju K, Eidelman DH, et al. A role for endoglin in coupling eNOS activity and regulating vascular tone revealed in hereditary hemorrhagic telangiectasia. Circ Res. 2005;96:684-92.
31. Toporsian M, Jerkic M, Zhou YQ, Kabir MG, Yu LX, McIntyre BA, et al. Spontaneous adult-onset pulmonary arterial hypertension attributable to increased endothelial oxidative stress in a murine model of hereditary hemorrhagic telangiectasia. Arterioscler Thromb Vasc Biol. 2010;30:509-17.
32. Jerkic M, Kabir MG, Davies A, Yu LX, McIntyre BA, Husain NW, et al. Pulmonary hypertension in adult Alk1 heterozygous mice due to oxidative stress. Cardiovasc Res. 2011;92:375-84.
33. Jerkic M, Sotov V, Letarte M. Oxidative stress contributes to endothelial dysfunction in mouse models of hereditary hemorrhagic telangiectasia. Oxid Med Cell Longev. 2012;2012:686972.
34. Li JM, Shah AM. Endothelial cell superoxide generation: regulation and relevance for cardiovascular pathophysiology. Am J Physiol Regul Integr Comp Physiol. 2004;287:R1014-30.
35. Paravicini TM, Touyz RM. Redox signaling in hypertension. Cardiovasc Res. 2006;71:247-58.
36. Harrison DG, Widder J, Grumbach I, Chen W, Weber M, Searles C. Endothelial mechanotransduction, nitric oxide and vascular inflammation. J Intern Med. 2006;259:351-63.
37. Jo H, Song H, Mowbray A. Role of NADPH oxidases in disturbed flow- and BMP4- induced inflammation and atherosclerosis. Antioxid Redox Signal. 2006;8:1609-19.
38. McCormick ML, Gavrila D, Weintraub NL. Role of oxidative stress in the pathogenesis of abdominal aortic aneurysms. Arterioscler Thromb Vasc Biol. 2007;27:461-9.
39. Miller FJ Jr, Sharp WJ, Fang X, Oberley LW, Oberley TD, Weintraub NL. Oxidative stress in human abdominal aortic aneurysms: a potential mediator of aneurysmal remodeling. Arterioscler Thromb Vasc Biol. 2002;22:560-5.
40. Nathan C, Ding A. Nonresolving inflammation. Cell. 2010;140:871-82.
41. Sanz-Rodriguez F, Fernandez LA, Zarrabeitia R, Perez-Molino A, Ramirez JR, Coto E, et al. Mutation analysis in Spanish patients with hereditary hemorrhagic telangiectasia: deficient endoglin up-regulation in activated monocytes. Clin Chem. 2004;50:2003-11.
42. Aagaard KS, Kjeldsen AD, Tørring PM, Green A. Comorbidity among HHT patients and their controls in a 20 years follow-up period. Orphanet J Rare Dis. 2018;13:223.
43. Ojeda-Fernández L, Recio-Poveda L, Aristorena M, Lastres P, Blanco FJ, Sanz-Rodriguez F, et al. Mice lacking endoglin in macrophages show an impaired immune response. PLoS Genet. 2016;12:e1005935.
44. Lebrin F, Goumans MJ, Jonker L, Carvalho RL, Valdimarsdottir G, Thorikay M, et al. Endoglin promotes endothelial cell proliferation and TGF-beta/ALK1 signal transduction. EMBO J. 2004;23:4018-28.
45. Lebrin F, Mummery CL. Endoglin-mediated vascular remodeling: mechanisms underlying hereditary hemorrhagic telangiectasia. Trends Cardiovasc Med. 2008;18:25-32.