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Biomechanical forces promote blood development through prostaglandin E\textsubscript{2} and the cAMP–PKA signaling axis

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Blood flow promotes emergence of definitive hematopoietic stem cells (HSCs) in the developing embryo, yet the signals generated by hemodynamic forces that influence hematopoietic potential remain poorly defined. Here we show that fluid shear stress endows long-term multilineage engraftment potential upon early hematopoietic tissues at embryonic day 9.5, an embryonic stage not previously described to harbor HSCs. Effects on hematopoiesis are mediated in part by a cascade downstream of wall shear stress that involves calcium efflux and stimulation of the prostaglandin E\textsubscript{2}–cyclic adenosine monophosphate (cAMP)–protein kinase A (PKA) signaling axis. Blockade of the PGE\textsubscript{2}–cAMP–PKA pathway in the aorta-gonad-mesonephros (AGM) abolished enhancement in hematopoietic activity. Furthermore, Ncx1 heartbeat mutants, as well as static cultures of AGM, exhibit lower levels of expression of prostaglandin synthases and reduced phosphorylation of the cAMP response element–binding protein (CREB). Similar to flow-exposed cultures, transient treatment of AGM with the synthetic analogue 16,16-dimethyl-PGE\textsubscript{2} stimulates more robust engraftment of adult recipients and greater lymphoid reconstitution. These data provide one mechanism by which biomechanical forces induced by blood flow modulate hematopoietic potential.

The establishment of intra-aortic blood flow after initiation of the heartbeat coincides with a crucial period in development when a switch occurs from primitive to adult-type definitive hematopoiesis (Dzierzak and Speck, 2008). We and others have shown that the mechanical forces induced by blood flow play a fundamental role in the emergence and maintenance of hematopoietic stem cells (HSCs) and progenitors in the aorta-gonad-mesonephros (AGM) region (Adamo et al., 2009; North et al., 2009). Functional HSCs and precursors with potential for HSC formation (pre-HSCs) have been

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Abbreviations used: AGM, aorta-gonad-mesonephros; CREB, cAMP response element–binding protein; dmPGE\textsubscript{2}, 16,16-dimethyl-PGE\textsubscript{2}; embryo equivalent; HSC, hematopoietic stem cell; NFAT, nuclear factor of activated T cells; NO, nitric oxide; PGE\textsubscript{2}, prostaglandin E\textsubscript{2}; PKA, protein kinase A; PSp, paraaortic splanchnopleura; qRT–PCR, quantitative RT–PCR; WSS, wall shear stress.

*M.F. Diaz and N. Li contributed equally to this paper.
found to arise mainly at arterial sites of the embryonic vasculature (Gordon-Keylock et al., 2013). Mutant embryos of the mouse and fish that lack a heartbeat, and thereby have reduced blood flow, exhibit a dramatic reduction in intravascular hematopoietic clusters and definitive hematopoietic activity in the AGM, further implicating mechanical forces as critical regulators of HSC emergence and/or expansion (Adamo et al., 2009; North et al., 2009; Wang et al., 2011). Wall shear stress (WSS), or the frictional force parallel to the cells of the vessel wall, activates genes essential for arterial specification and definitive hematopoiesis in the developing embryo (Adamo et al., 2009). Nitric oxide (NO) signaling contributes to the induction of HSC formation by blood flow, and stimulation of this pathway either by mechanical forces or pharmacological treatment with NO donors can rescue hematopoiesis in embryos without a heartbeat (Adamo et al., 2009; North et al., 2009; Wang et al., 2011). In addition to NO, several other autacoids, including prostacyclins, are modulated by shear stress and influence fundamental properties of endothelial and smooth muscle function (Frangos et al., 1985; Alshihabi et al., 1996; Johnson et al., 1996; Topper et al., 1996; Smalt et al., 1997; Tsai et al., 2009). Their role in determination of hematopoietic fate remains poorly characterized.

Recently, several groups have shown that prostaglandin E2 (PGE2), a prostacynin-related prostanoid family member, regulates HSC and progenitor self-renewal, survival, trafficking, and engraftment potential and has led to the development of methods for expansion of hematopoietic cells for clinical use (North et al., 2007; Cutler et al., 2013; Hoggatt et al., 2013a,b; Porter et al., 2013). PGE2 is the gene that encodes the limiting enzyme in PGE2 production, COX2, and was recently identified in differential expression analysis as the second most highly up-regulated gene, second only to Fosb, in AGM-derived HSCs as compared with HSCs present in the fetal liver (McKinney-Freeman et al., 2012). Goessling et al. (2009) determined that the synthetic analogue 16,16-dimethyl-PGE2 (dmpPGE2) could enhance HSC formation in developing zebrfish embryos through cAMP–protein kinase A (PKA) modulation of Wnt signaling. Independent studies further implicate cAMP–PKA in initiation of vascular and hematopoietic differentiation of embryonic stem cells via recruitment of the transcriptional activator cAMP response element–binding protein (CREB) to the Etv2 promoter, resulting in up-regulation of vascular growth factor receptors and hematopoietic transcription factors including Flk1, Tie2, Scl/Tal1, and Gata2 (Yamamizu et al., 2012). Connections between these signaling pathways and fluid flow have been described in osteolinages of the bone but have not yet been investigated in blood development (Ogasawara et al., 2001; Ogawa et al., 2014).

Here, we demonstrate that WSS associated with embryonic blood flow potentiates development of definitive hematopoietic cells through the induction of developmental pathways known to be critical for hematopoiesis, including Wnt and Notch, as well as stimulating mechanosensors that trigger calcium flux. Signaling through calcium up-regulated expression of the COX2 gene, Pgo2. Subsequent increases in PGE2 biosynthesis and cAMP–PKA activity were necessary for expansion of nascent HSCs and progenitors. Phosphorylation of CREB, presumably by PKA, was associated with transcription of genes containing the cAMP response element. WSS and dmpPGE2 were each capable of promoting long-term multilineage hematopoietic reconstitution and lymphoid lineage potential from the paraaortic splanchnopleura (PSp) and AGM, respectively. This work directly links mechanical forces generated by blood flow to the regulation of biochemical and genetic pathways that define HSC potential during embryonic hematopoiesis.

RESULTS

Functional maturity of hematopoietic stem and progenitor cells is enhanced by WSS

Previously, we described specific characteristics of WSS that induce hematopoietic progenitor activity in two-dimensional adherent cultures of dissociated AGM (Adamo et al., 2009). In the present study, we hypothesized that WSS may play a role in specification of HSCs that support life-long adult hematopoiesis and so examined developmental time points that precede definitive HSC emergence in cells derived from the anlage of the AGM known as the PSp at embryonic day (E) 9.5 and from the AGM at E10.5. Runx1 and Myh, two master regulators of hematopoiesis, are transcriptionally up-regulated by shear stress typical of embryonic murine blood flow (5 dyn/cm²), as are other genes required for definitive hematopoiesis and lymphopoiesis, including Bcl11a, Etv6, Gata2, Gata3, Tjf, Rag1, and Rag2 (Fig. 1 A). Analysis of cell surface phenotype after WSS confirmed increases in two markers of hemogenic endothelium, CD144/VE-Cadherin and c-kit, in the live (DAPI−) population (Fig. 1 B). We observed a 5.2 ± 1.2-fold increase in the percentage of CD144+c-kit+ cells, a surface phenotype thought to distinguish a subset of endothelial cells with definitive HSC potential (Fig. 1 C; Eilken et al., 2009; Swiers et al., 2013).

E9.5 PSp has previously been found to produce primarily B1a lymphocyte progenitors when delivered into immunocompromised neonatal recipients (Yoshimoto et al., 2011) or intact under the adult kidney capsule (Godin et al., 1993) and requires whole organ culture for several days to acquire long-term multilineage repopulating activity in vivo (Cumano et al., 2001). To evaluate whether changes induced by WSS were capable of producing greater functional competence in stem and progenitor cells, we transplanted dissociated PSp from E9.5 embryos (21–29 somite pairs) cultured with or without WSS for 36 h into irradiated adult Rag2−/− Il2rg−/− recipients. Based upon the expectation that dissociated E9.5 PSp would provide absent to scant engraftment of adult recipients long term (Godin et al., 1993; Cumano et al., 2001), we transplanted each recipient with 16 embryo equivalents (e.e.). We detected low-level engraftment in the peripheral blood of 4 of 17 animals injected with E9.5 PSp cultured under static conditions, which dropped further by 10 wk in the surviving 10 animals, suggesting that the majority of engrafting cells were exhaustible short-term progenitors (Fig. 2 A and B). At 20 wk, static cultures contributed to <1.4% of total peripheral
blood leukocytes in all but one recipient (2.8% CD45.2+). In contrast, WSS produced higher initial engraftment levels in 6 of 15 recipients, which persisted long term in 5 animals (Fig. 2 B). PSp contributed to major adult blood lineages, including B lymphoid (B220+, CD19, and IgM), T lymphoid (CD3, CD4, and CD8), and myeloid (Mac1 and Gr1) populations (Fig. 2 C). WSS exposure promoted earlier emergence of B lymphocytes and sustained B lymphopoiesis in the periphery beyond 20 wk (Fig. 2, D and E). Greater B cell maturity was also apparent in the bone marrow (Fig. 2 F). Specifically, greater numbers of cells were in stages of pre-pro- and pro-B commitment (B220+ CD43-), with modest increases in late-stage pre-B and B development (B220+ CD43+) typical of adult-type bone marrow progenitors (Fig. 2 G; Hardy and Hayakawa, 1991). CD45.2+ bone marrow cells were subsequently transferred to secondary recipients and found to reconstitute four of seven WSS and two of five static recipients (Fig. 3, A and B). Importantly, peak chimerism was substantially higher for three of the WSS donors (4.2%, 12.2%, and 61.3%) as compared with static donors (0.9% and 2.1%). B and T cells were detectable as discrete populations and persisted long term in WSS recipients (Fig. 3, C and D). Collectively, these data suggest that WSS serves as a critical developmental signal that promotes commitment of precursors to the blood lineage and endows nascent HSCs and progenitors with the functional competence necessary to engrail the adult hematopoietic niche.

WSS is required for proper developmental signaling

Knockout of the Ncx1 gene that encodes the cardiac-specific sodium/calcium exchanger results in mutant embryos with no heartbeat and little to no hematopoietic progenitor activity (Lux et al., 2008). Development of mutant embryos proceeds up to 10 d of gestation, at which time cardiac defects result in embryonic lethality. Previously, we showed that hematopoietic activity could be rescued ex vivo after 36 h of WSS in cells derived from E9.5 Ncx1−/− PSp (Adamo et al., 2009). We therefore hypothesized that hemogenic endothelium was present in Ncx1−/− PSp and measured the frequency of cell surface markers typical of hematopoietic progenitors and progeny. We found that Ncx1−/− embryos harbor phenotypic hemogenic endothelium and hematopoietic cells (Fig. 4 A). Indeed, the frequency of CD144+ ckit+ CD45− hemogenic endothelium was elevated in mutant PSp by ~4.1 ± 1.1-fold (Fig. 4 B). As this contrasts with our observations of enhanced CD144+ ckit+ CD45− cell population in ex vivo WSS cultures, we speculate that accumulation of hemogenic endothelial cells in the mutant PSp could be a consequence of defective circulation, a phenomenon demonstrated previously to lead to pooling of primitive erythrocytes during development of zebrafish embryos (Iida et al., 2010). We thus analyzed gene expression in Ncx1−/− PSp and found decreases in Runx1, Gata2, Tal1, Etv2, and effectors of Notch (Hes1 and Hey1) and Wnt (Wnt3a and Left1), supporting a role for WSS in determination of signaling events in the PSp (Fig. 4 C).

We and others previously identified NO as a critical regulator of hematopoietic progenitor expansion in response to WSS and blood flow (Adamo et al., 2009; North et al., 2009; Wang et al., 2011). To more precisely define the signaling mechanisms triggered by WSS, we conducted global gene expression profiling of ex vivo cultures of AGM. Notch, Wnt,
Figure 2. Long-term multilineage repopulation of the blood system is enhanced by WSS. *Rag2<sup>−/−</sup> Il2rg<sup>−/−</sup> recipients received 16 e.e. of E9.5 PSp cultured in the presence of either static (<0.0001 dyn/cm<sup>2</sup>) or WSS (5 dyn/cm<sup>2</sup>) conditions for 36 h. (A) Donor contribution to recipient leukocytes in peripheral blood is distinguishable as a discrete CD45.2<sup>+</sup> CD45.1<sup>+</sup> population in WSS recipients (20-wk posttransplantation data shown). (B) Peripheral blood reconstitution is significantly greater from cells exposed to WSS (n = 17 static from six independent experiments, n = 15 WSS from seven independent experiments; two-way ANOVA: *, P = 0.03). (C) Representative flow cytometry plots from the WSS recipient in A show lineage<sup>+</sup> engraftment from CD45.2<sup>+</sup> donor-derived cells. (D) Lineages present in CD45.2<sup>+</sup> donor peripheral blood from individual recipients at 5, 10, and 20 wk after transplantation. B lymphopoiesis and long-term multilineage potential are bolstered by WSS. (E) CD45.2<sup>+</sup> donor PSp contribute to greater numbers of cells expressing B lineage markers during the posttransplant period (20 wk) when exposed to WSS (n = 17 static, n = 15 WSS; Mann–Whitney rank sum: *, P = 0.03). (F) Primary recipient bone marrow was analyzed at 20 wk after transplantation of cultured PSp with markers of B lineage maturation (two static and three WSS recipients are shown alongside adult donor strain). B cells within the bone marrow collected from WSS recipients resemble adult donor marrow. (G) Production of early and late B lineages in primary bone marrow was discriminated by B220 and CD43 cell surface expression in the CD45.2<sup>+</sup> donor population. Early B lineage phenotypes were marginally elevated by WSS (B220<sup>+</sup> CD43<sup>+</sup>; n = 3 static, n = 5 WSS; unpaired Student’s t test: *, P = 0.13). Data are represented as mean ± SEM.
microfluidic channels as described previously (Li et al., 2014), replicates were lysed, and RNA was processed for analysis by Illumina Mouse WG-6 v2.0 Expression BeadChips (45,200 transcripts). Unsupervised hierarchical clustering of unfiltered genes positioned 6- and 36-h cultures in two distinct groups, followed by segregation of static and WSS cultures (not depicted). Differential gene expression analysis ($P < 0.01$, two-fold threshold) revealed significant change in 1,435 unique and eicosanoid signaling have previously been found in proteomic analyses to respond rapidly to fluid shear stress in aortic endothelial cells (Wang et al., 2007). Informed by these studies and our own serial measurements of gene subsets, we selected an early time point (6 h) and one demonstrated previously to enhance hematopoietic activity (36 h) to perform global gene expression analysis (Adamo et al., 2009). In brief, WSS was applied to dissociated E10.5 AGM cultured within

Figure 3. WSS enhances long-term secondary engraftment. Secondary recipients received 20,000 CD45.2+ bone marrow cells sorted from primary recipients plus 200,000 CD45.1+ CD45.2+ Rag2−/− Il2rγ−/− competitor bone marrow cells. (A) Engraftment in the periphery of secondary recipients was monitored for up to 10 mo ($n = 5$ static, $n = 7$ WSS from independent primary recipients). Peak chimerism was found long term in two WSS and two static recipients, with substantially higher donor contribution in the WSS recipients. Long-term engraftment was defined as CD45.2 positivity in the periphery for >20 wk (denoted by the dashed line) at any level above background. (B) Plots of peripheral blood collected from recipients 8 mo after transplantation illustrate sustained contribution of donor-derived CD45.2+ marrow cells. (C) Analysis of lineages present in the CD45.2+ gate of the engrafted WSS recipient depicted in panel B alongside adult peripheral blood from the donor and recipient strains. (D) Lineages present in CD45.2+ donor peripheral blood from individual secondary recipients at 4, 24, 36, 48, and 68 wk after transplantation. Recipient strain is represented by an untransplanted adult CD45.1+ CD45.2+ Rag2−/− Il2rγ−/−; donor strain is adult Swiss Webster.
transcripts at 6 h, 347 at 36 h, and 109 common to both time points (Fig. 5 A and Dataset S1). By Ingenuity pathway analysis, these genes were found to encode enzymes and transcription regulators required for signaling through several overlapping pathways, including G protein–coupled receptors (GPCRs), calcium transport, nuclear factor of activated T cells (NFAT), phospholipase C, and PKA regulatory subunit (Chp1, Nfatc3, and Pdk3; Fig. 5 B). Components of the NFkB signaling pathway were also enriched, including the TRAF family member associated NFkB activator (Tank) and IKB regulatory subunit (Ikbkb). Several other kinases, enzymes, and transcription factors were found to contribute to signaling through CREB, such as the α regulatory subunit of cAMP-dependent protein kinase (Prkar1a), G protein–binding protein (Gobh), Cebb, and CREB-binding protein (Crebbp). Wnt/β-catenin regulators and ligands were also enriched (Akt3, Dkk1, Wnt1, and Wnt7b). Functional enrichment analysis of unfiltered genes by GSEA (Mootha et al., 2003; Subramanian et al., 2005) was used to construct a global network wherein overlapping gene sets clustered together (Merico et al., 2010) and connections between pathways could be visualized at 6 and 36 h in parallel (Fig. 5 C). Wnt signaling functionally clustered with the adaptive immune system and cell cycle, whereas Notch, TGF-β, and EGFR/CREB/MAPK clustered as distinct groups. Notably, several functional groups that emerged from network analysis are known regulators or effectors of prostaglandin production and signaling, including calcium, MAPK, CREB, Wnt, NFkB, and biological oxidation involving prostaglandin–endoperoxide synthase function of COX1 and COX2 (Ptges1 and Ptges2; Fig. 5 C and Dataset S2; Tsatsanis et al., 2006; Goessling et al., 2011).

**Calcium efflux propagates a signaling cascade downstream of WSS that amplifies prostaglandin synthesis**

PGE₂ has recently been documented to promote hematopoietic engraftment in mice and in human clinical trials (North et al., 2007; Hoggatt et al., 2009, 2013a,b; Cutler et al., 2013). Up-regulation of prostaglandin synthases Ptges1, Ptges2, Ptges, and Ptges3, as well as Wnt and Notch were verified by quantitative RT-PCR (qRT-PCR) in WSS-exposed populations of the E10.5 AGM, including the CD144⁺ CD45⁻ population identified as the hemogenic endothelium (Fig. 6, A and B). Importantly, we found that WSS promoted production and secretion of significantly higher levels of PGE₂ (Fig. 6 C). This enhanced PGE₂ production could be blocked by the COX1/2 antagonist indomethacin and the COX2-specific inhibitors NS-398 and CAY10404, suggesting that Ptgds2 up-regulation may be responsible for elevated PGE₂ production in WSS-exposed AGM (Fig. 6 D). In osteoblasts, calcium-dependent activity at focal adhesions is believed to regulate COX2 and PGE₂ synthesis (Ponik and Pavalko, 2004). We therefore hypothesized that WSS may also stimulate calcium flux upstream of PGE₂ production in the AGM. Live cell imaging with Fluo-4 AM confirmed sparks of intracellular calcium signaling in WSS-stimulated cells, as determined by increased intensity in
profiled global gene expression as described above. Analysis of differential gene expression revealed that PGE$_2$ contributed to transcription of CREB targets, as well as core components of Wnt/calcium signaling, including Creb1, Ctbp1, Shc1, Calm2, and Camk1 (Fig. 7A). In chondrocytes of the bone, WSS was recently shown to promote TOP-luciferase (LEF/TCF reporter) and CRE-luciferase (CREB reporter) activity (Ogawa et al., 2014). Whereas addition of exogenous PGE$_2$ was insufficient to drive activation of TOP-luciferase, PGE$_2$ was capable of promoting activity of the CRE-luciferase reporter. Consistent with these findings, we observed that WSS increased phosphorylation of CREB at serine 133, a post-translational modification required for CREB-mediated tran

scription (Fig. 7B). CREB phosphorylation can be mediated by PKA; thus, the increase in PKA transcript Prkaca by WSS.

WSS activates CREB and the cAMP–PKA signaling axis

To evaluate the contribution of PGE$_2$ to intracellular signaling, we subjected cells to WSS with or without indomethacin and cytoplasmic fluorescence (Fig. 6, E and F; and Videos 1 and 2). Although calcium flux was evident in single cells within static cultures, exposure to WSS induced more intense signaling across a greater number of cells. Sequestration of calcium by BAPTA-AM reduced WSS-dependent induction of Pge2 transcript (Fig. 6 G). In static cultures, pharmacological elevation of intracellular calcium concentration by pulse treatment of the calcium ionophore A23187 also stimulated accumulation of COX2 protein (Fig. 6 H). Observations in ex vivo cultures of AGM were further corroborated by reduction in COX2 protein levels in uncultured Ncx1$^-$/Psp (Fig. 6 I).

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Figure 6. PGE$_2$ production in the AGM is enhanced by calcium flux triggered by WSS. (A) E10.5 AGM-derived cells cultured for 36 h with WSS activate several pathways required for definitive hematopoiesis ($n = 5$ independent experiments; two-tailed Student’s $t$ test or Mann–Whitney rank sum: *, $P < 0.05$; **, $P < 0.01$). Klf2 and Klf4 are WSS-responsive genes that serve as positive controls for transcriptional activity. (B) Transcriptional up-regulation in WSS-exposed cells is apparent in sorted CD144$^+$ CD45$^-$ hemogenic endothelial cells ($n = 5$ independent experiments; unpaired
at 6 and 36 h (unpaired Student’s *t* test, *P* = 0.016 [6 h] and *P* = 0.004 [36 h]) suggested that modulation of PKA downstream of WSS could contribute to enhanced CREB activity. Furthermore, concomitant reduction in *Pkaa* transcript, the prohematopoietic CREB target *Etv2*, and P-CREB levels in uncultured *Ncx1*−/− Psp independently suggested a role for blood flow in coordinating PKA and CREB activity (Figs. 4 C and 7 C). Of six *Ncx1* mutants analyzed in the study, five expressed lower levels of P-CREB relative to wild-type littermates. Total CREB protein was also reduced in four of six mutants, likely indicative of complex signaling inputs known to regulate CREB expression and activity (Shaywitz and Greenberg, 1999). Binding of the EP2 and EP4 receptors by PGE2 stimulates the Gs-adenyl cyclase pathway, which leads to cAMP production and subsequent PKA activation (Zhang and Daaka, 2011; Yamamizu et al., 2012). As expected, AGM exposed to WSS for 36 h produced higher levels of intracellular and circulating cAMP compared with static cultures (Fig. 7 A). Importantly, antagonists of COX2 or PKA abolished enhancement in hematopoietic colony forming activity with ex vivo exposure to WSS, as did a blocking antibody to PGE2 (Fig. 7 E). The adenyl cyclase agonist forskolin amplified colony-forming potential in WSS-exposed cultures but not static cultures, suggesting that PGE2 cooperates with additional WSS-responsive signaling in initiation of the hematopoietic program. Consistent with this notion, inhibition of COX2 by indomethacin blocks WSS induction not only of CREB and MAPK signaling but also calcium, calmodulin, calcineurin, and NFAT, primary components of noncanonical Wnt/calcium signaling that function to control cell adhesion, migration, and ventral patterning (Fig. 7 F and Dataset S3; Kohn and Moon, 2005). Based on analyses of BrdU incorporation and Annexin V staining, WSS contributes significantly to survival of the CD144+ CD45− population (Fig. 8). However, PGE2 does not appear to influence cell proliferation or survival, but instead may serve to modulate fate selection through CREB. More detailed analyses of signal transduction will be required to understand the precise nature of how PGE2–PKA interfaces with other developmental signaling pathways to modulate hematopoietic-specific gene expression downstream of WSS in the AGM.

**PGE2 promotes engraftment of embryonic hematopoietic stem and progenitor cells into adult recipients**

The HSC enhancing ability of dmPGE2 was first identified in pharmacological screens of zebrafish embryos and has since been used for expansion of umbilical cord blood in phase I clinical trials (North et al., 2007; Cutler et al., 2013). To date, the effects of dmPGE2 in mouse and human have only been tested on committed hematopoietic stem and progenitor cells. We therefore evaluated the ability of transient dmPGE2 treatment to promote short- and long-term hematopoietic activity from E10.5 and E11.5 murine AGM. Colony formation assays showed an asymptotic dose-dependent increase in progenitor activity with up to 10 µM dmPGE2 (Fig. 9 A). 2-h dmPGE2 treatment also increased chimerism in the peripheral blood from donor AGM above mean levels in three of five recipients of E10.5 AGM and in six of eight recipients of E11.5 AGM (Fig. 9, B and C). Notably, dmPGE2 improved B and T lymphoid potential of hematopoietic progenitors/stem cells at both embryonic stages, although E11.5 produced the only HSC-like long-term engraftment of all lineages (Fig. 9, D and E). Collectively, our findings suggest that WSS and PGE2 function together to regulate developmental signaling that determines lineage potential of nascent hematopoietic cells in the earliest stages of definitive hematopoiesis.

**DISCUSSION**

Here we have shown that WSS activates developmental pathways that promote hematopoietic fate and potentiates long-term engraftment of embryonic hematopoietic stem and progenitor cells from the PSp and AGM. Prostaglandin, Wnt, and Notch signaling were all up-regulated in response to WSS. PGE2 production in particular was found to be required for WSS–enhanced progenitor activity, as its stimulatory effects could be inhibited by blocking antibodies or pharmacological inhibition of COX enzyme function. PGE2 was directly responsible for increased cAMP production and contributed to regulation of CREB through PKA. Mimicry of WSS by addition of dmPGE2 induced greater peripheral blood chimerism and enhanced lymphoid potential from E10.5 and E11.5 AGM, supporting a possible mechanism for transduction of mechanical cues into chemical signaling.

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**Student’s *t* test:** *, * *P* < 0.05; **, *P* < 0.003. (C) Prostaglandin production as measured by PGE2 release into media was increased over time by WSS (*n* = 3 independent experiments; two-way ANOVA: *, *P* = 0.001). (D) PGE2 biosynthesis was suppressed by addition of indomethacin or COX2-specific antagonists NS-398 and CAY10404. WSS-treated vehicle control produced significantly higher amounts of PGE2 than WSS samples treated with COX inhibitors and all static cultured samples at 60 h of culture (*n* = 3 independent experiments; two-way ANOVA: *, *P* < 0.02; **, *P* < 0.001). (E) WSS applied to E11.5 AGM-derived cells triggers elevated levels of intracellular calcium (*n* = 3 independent experiments, >5 replicates per experiment). WSS was initiated 3 s after image acquisition began. Abbreviated videos are available in Videos 1 and 2. Bar, 50 µm. (F) Quantification of Fluo-4 AM intensity by MetaMorph captures multiple spikes in calcium flux after application of WSS (red arrow on horizontal axis marks initiation of WSS at 3 s). Pastel traces represent calcium levels in individual cells (*n* = 150 cells), whereas bold traces (blue or red) represent the mean intensity of values collected from individual cells. (G) *P*tg2s transcript level was reduced by an inhibitor of calcium signaling (BAPTA-AM; *n* = 3 independent experiments at 3 h of WSS; one-tailed unpaired Student’s *t* test: *, *P* = 0.036; *P* > 0.26 for comparison between static vehicle- and BAPTA-AM–treated WSS group). (H) COX2 protein accumulates in response to transient treatment with A23187, a compound which increases intracellular calcium. Cells were pulse treated under static conditions with A23187 for 1 or 3 h, washed, and incubated in fresh medium for a total culture period of 18 h. A23187 was not removed from the 18-h sample (*n* = 3 independent experiments at 2 µM A23187). (I) COX2 protein levels are decreased in *Ncx1*−/− Psp (*n* = 6 mutant, *n* = 8 wild-type littermates analyzed). All data are represented as mean ± SEM.
and downstream gene activation within HSCs of the embryonic vasculature.

Prostaglandin signaling has emerged as an attractive target for enhancing hematopoietic function in cell therapy and as a medical countermeasure against radiological and nuclear threats (Goessling et al., 2011; Cutler et al., 2013; Hoggatt et al., 2013c). In animal models and humans, PGE₂ has been found to regulate survival, self-renewal, and trafficking of HSCs and progenitors.
(Hoggatt et al., 2009, 2013a). Ablation of Ptgs2 or the terminal prostanooid synthase Ptgs results in a variety of hematopoietic defects, impacting macrophages and platelets, and widespread impairment in inflammatory response (Uematsu et al., 2002; Cheng et al., 2006; Yu et al., 2007; Seta et al., 2009). Furthermore, PGE₂ has very recently been found to possess morphogen-like effects on bipotential endodermal precursors that determine fate selection between hepatic and pancreatic commitment (Nissim et al., 2014), raising the possibility that PGE₂ and downstream PKA activation could direct specification of hematopoietic lineages from precursors of the hemogenic endothelium, perhaps in part through modulation of Wnt signaling.

Our data support a model in which multiple pathways downstream of WSS must converge to balance developmental signaling essential for hematopoiesis. In addition to Wnt and Notch, other pathways altered by WSS corroborate findings from Goessling et al. (2011), in which pulse treatment of human and macaque CD34⁺ cells with dmPGE₂ induced pathways such as PKA, NFAT, and ERK/MAPK. Mechanosensation in hemogenic endothelium and other cells of the AGM involves release of second messengers into the cytosol, including intracellular calcium, that directly stimulate enzyme functions required for prostaglandin synthesis. In osteocytes, calcium flux triggers rapid PGE₂ synthesis and secretion through a biochemical cascade that includes PLC, DAG, PKC, and PLA₂ (Ajubi et al., 1999). Prostaglandin production has also been found in osteoblasts to be dependent on focal adhesions and transcriptional transactivation of Ptgs2 by transcription factors, including C/EBP β, AP-1, and CREB (Ogasawara et al., 2001; Ponik and Pavalko, 2004; Rangaswami et al., 2012). Prostaglandin synthesis is elevated in CD144⁺ CD45⁻ hemogenic endothelial cells, but PGE₂ signaling could also originate from other cell types in the AGM, such as committed CD144⁺/⁻ c-kit⁻ hematopoietic progenitors (unpublished data). As PGE₂ is released into the aortic lumen, activation of EP₁/EP₃ receptors expressed on hemogenic endothelium or committed hematopoietic stem and progenitor cells stimulates G protein–dependent adenyl cyclase activity and elevates intracellular cAMP levels. PKA, in the presence of cAMP, can then modulate CREB–dependent gene expression. Interestingly, genetic ablation of CREB phenocopies Wnt1 and Wnt3a knockout in differentiating myoblasts, suggesting that activation of CREB by PKA is a prerequisite for transduction of some noncanonical Wnt signals (Chen et al., 2005). We find that WSS stimulates CREB phosphorylation and up-regulates core components of noncanonical Wnt signaling, including calmodulin, calcineurin, NFAT, and calmodulin–dependent kinase, which are known to play an important role in self-renewal and repopulating activity of adult HSCs (Nemeth et al., 2007; Sugimura et al., 2012). Wnt receptor and ligand transcript levels were statistically unchanged by indomethacin or the PKA antagonist H89, suggesting that Wnt machinery at the cell membrane is regulated independently of PGE₂ but...
that target gene activation may be regulated by WSS-induced phosphorylation of CREB (unpublished data). The nature of the relationship between PGE$_2$ and NFAT will require further investigation, though it appears that PGE$_2$ may contribute to signaling activity through calcium, calmodulin, calmodulin-dependent kinase, and NFAT. CREB and NFAT are essential
for transduction of noncanonical Wnt signals; thus, regulation of the activity of these transcription factors may allow PGE2 to “fine tune” the dosage of canonical Wnt signaling required for expansion and maintenance of HSCs (Luis et al., 2011).

The identification of biomechanical cues that support hematopoiesis has begun to redefine our understanding of the types of signals that comprise the hematopoietic niche. Data from the current study demonstrate that PGE2 contributes essential signaling downstream of WSS to govern the expansion of hematopoietic populations with long-term reconstitution potential in the developing embryo. Moreover, we find that WSS-induced PGE2 acts through calcium and cAMP–PKA to regulate CREB for induction of prohematopoietic developmental programs. Further studies will be required to precisely address how biomechanical forces coordinate these and other signaling pathways in the PSp and AAM to define hematopoietic potential.

**MATERIALS AND METHODS**

**Mice.** Timed pregnancies were bred in house to E9.5–11.5 (C57BL/6) or Ncx1-/- or were purchased as pregnant females from Taconic at E9.5 for large-scale shearing experiments (Swiss Webster). Gestational age of embryos was determined by observation of a copulation plug on E0.5 and number of somite pairs. RAG2-/- E2Ry-/- mice were back-bred to C57BL/6 SJL (PepBoy) to produce CD45.1 RAG2-/- E2Ry-/- mice and were maintained by inbreeding to CD45.2 RAG2-/- E2Ry-/- Ncx1 knockout mice were maintained on a C57BL/6 background (minimum ninth generation). All animal experiments were performed according to the University of Texas Medical School at Houston and Children’s Hospital Boston guidelines for laboratory animals.

**Culture of PSp- and AGM-derived cells.** Embryos from C57BL/6 or Swiss Webster timed pregnancies were micro dissected for isolation of E9.5 PSp or E10.5 AGM regions. Tissues were dissociated either by treatment with Accutase (STEMCELL Technologies) at room temperature with gentle agitation for 20 min or, for WSS transplantation experiments, by 0.1% dispase (Gibco) at 37°C for 30 min. Cells were then resuspended in M5300 Myelocult medium (STEMCELL Technologies) enriched with Hepes (Invitrogen; 12.5 µl of 1 M Hepes per 500 µl Myelocult medium), nonessential amino acids (Gibco), sodium pyruvate (Gibco), 10 U/ml penicillin (Lonza), and 10 µg/ml streptomycin (Lonza) without addition of hydrocortisone. Cells were plated on tissue culture plastic coated with a 1:20 Matrigel–PBS solution (BD) at a concentration of 8 e.e. per 10 cm2 in prairie for culture on a dynamic flow system (Adamo et al., 2009) or in microfluidics (Li et al., 2014), as described previously. Microfluidics consisted of BD BioVIATM channel slided in line with a recirculating medium system driven by a single Harvard Apparatus PHD ULTRA 4400 with remote syringe pump, 0.27 PSI crack pressure check valves (Qosina), and female luer lock three-way stop cocks (Qosina). After 6.5 h of incubation, nonadherent cells were washed away and cells were cultured in enriched Myelocult medium with 10 ng/ml murine SCF (PeproTech) in static/low-flow conditions (<0.001 dyn/cm2) or in the presence of WSS (5 dyn/cm2). The shear stress pattern used to induce hematopoiesis is described in our previous work (Adamo et al., 2009). At the end of the culture period, adherent and nonadherent cells were collected by a 5-min incubation in Accutase, washed, resuspended in Myelocult, and transferred to functional or phenotypic assays. Cells analyzed immediately by flow cytometry were immunostained with antibodies available through BD, purified rat CD44 and APC c-kit (CD117); eBioscience, APC-Cy7 CD45.2; and Invitrogen. Alexa Fluor 488 anti-rat IgG. Colony formation assays were established in M3434 MethoCult medium (STEMCELL Technologies) at 100,000 cells per 1.5 ml media and counted after 10–14 d.

**Modulation of calcium and PGE2–PKA signaling.** Cells were cultured with pharmacological compounds or antibodies to block PGE2 production and/or signaling by incubation with 10 µM indomethacin, 10 µM NS-398, 1 µM CAY10404 (Cayman Chemical), or 1,1,1,9-tetrafluoro-2-(3-nitrophenyl)ethane (Abcam) for the duration of culture with or without WSS. H89 (Cayman Chemical) was applied at 10 µM to inhibit PKA activity. Forskolin (Sigma-Aldrich) was administered at 10 µM as an adenylyl cyclase activator to stimulate cAMP production. The stabilized synthetic analogue dmPGE2 (Cayman Chemical) was applied to freshly dissociated AGM cells for 2 h at 37°C from 1 to 100 µM for colony formation assays or at 10 µM for transplantation into adult recipient mice. Intracellular calcium was sequestered by 10 µM BAPTA-AM (Tocris Bioscience) and elevated by 2 µM of calcium ionophore A23187 (Tocris Bioscience).

**Gene expression profiling.** Total RNA was extracted with QIAGEN RNeasy kits for analysis of gene expression by Illumina Mouse WG-6 v2.0 Expression BeadChips or by TaqMan Assays. R/T of RNA for qRT–PCR was performed using Multiscribe DNA polymerase (Applied Biosystems), and PCR was performed in 10-µl reactions on a 7800 Real-Time PCR System (Applied Biosystems). All procedures were conducted according to the manufacturer’s instructions. Illumina data were checked for quality, background corrected, and quantile normalized with GenomeStudio (Illumina). Analysis of differential expression was conducted with GenomeStudio for probe sets filtered by spot detection (P < 0.01) and difference in mean signal intensity between treatment groups (dف > 20). Significantly changed genes were subjected to analysis of canonical pathways curated by Ingenuity (IPA, Ingenuity Systems). Gene set expression analysis (GSEA) was conducted with unfiltered genes using curated gene lists downloaded from MSigDB. Ranked gene list data were imported into Cytoscape for visualization of functional enrichments with Enrichment Map using edge-weighted force-directed layout. Heat maps were generated using genes selected from differential gene expression analysis in GenePattern. Expression data have been deposited for public access in the NCBI Gene Expression Omnibus (GEO) under accession no. GSE62463.

**Measurements of PGE2 and cAMP.** E10.5 AGM were dissociated and exposed to WSS or static culture conditions for up to 110 h. Cell lysates and/or medium was processed for measurement with the Prostaglandin E2 Express ELISA kit (Cayman Chemical) or the Direct cAMP EIA kit (Enzo Life Sciences). Standard curves were established in parallel for quantification of sample concentrations.

**Transplantation.** Microdissected PSp from 64 E9.5 embryos for WSS experiments were pooled and dissociated in 0.1% dispase–PBS with calcium and magnesium. PSp cells were cultured as described above and prepared for transplantation. After dissociation with 100 µg/ml collagenase (Cayman Chemical) was applied to freshly dissociated AGM cells for 2 h at 37°C from 1 to 100 µM for colony formation assays or at 10 µM for transplantation into adult recipient mice. Intracellular calcium was sequestered by 10 µM BAPTA-AM (Tocris Bioscience) and elevated by 2 µM of calcium ionophore A23187 (Tocris Bioscience).

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**Peripheral blood and bone marrow analysis.** Peripheral blood was collected at 4, 6, 10, and 20 wk after transplantation. Red blood cells were removed by 1% dextran sulfate–PBS–EDTA separation and treatment with RBC lysing buffer (Sigma-Aldrich). Leukocytes were immunostained for discrimination of CD45 allelic variants and detection of lineage markers, followed by analysis on a five-laser BD LSR II flow cytometer. Antibodies used from BD
BrdU was applied 3 s after initiation of image acquisition and thus represents static conditions. WSS or static culture was maintained for an additional 5 min at 37°C. Fresh MyeloCult medium was applied, and cells were placed in an environmental chamber maintained at 37°C, 5% CO2, 100% humidity, and 1% O2. Calcium imaging. Dissociated E11.5 AGM cells were plated at lower density (120,000 cells/cm2) for discrimination of calcium sparks in individual cells. In brief, adherent cells were washed with isotonic Tyrode's solution (139 mM NaCl, 3 mM KCl, 17 mM NaHCO3, 12 mM d-glucose, 3 mM CaCl2, and 1 mM MgCl2) and incubated with the fluorescent calcium indicator 5 μM Fluoro-4 AM (F14201; Invitrogen) in isotonic Tyrode's solution for 5 min at 37°C. Fresh MyeloCult medium was applied, and cells were placed in an environmental chamber maintained at 37°C, 5% CO2 for imaging. Successive images were collected at a 300-ms time interval for 60 s (201 images total) using MetaMorph v7.7.9.0 on an Olympus IX81 fluorescence microscope equipped with an Andor iXon X3 885 EMCCD camera. WSS was applied 3 s after initiation of image acquisition and thus represents static culture at start time. Images were subsequently analyzed for integrated intensity using manual selection of individual cell boundaries with MetaMorph software. Videos were compiled using ImageJ 1.46r software (National Institutes of Health) using 10 frames per second for visualization.

Assessment of cell death by Annexin V staining. E10.5 AGM were dissociated and exposed to WSS or control culture conditions for 36 h. After the culture period, adherent and nonadherent cells were collected with Accutase and washed in MyeloCult media. Cells were immuno stained on ice with each antibody in 2% FBS-PBS for 20 min, first with rat anti-mouse VE-Cadherin (BD) antibody and then with Alexa Fluor 488 anti-rat IgG (Invitrogen). Cells were washed and additional cell surface markers were detected with APC e−kit (CD117; BD) and APC-Cy7 CD45.2 (eBioscience). Annexin V binding buffer was used for subsequent washes and for incubation with PE-Cy7 Annexin V (eBioscience) at room temperature for 15 min. Cells were resuspended in binding buffer containing 1 μg/ml DAPI before analysis on a three-laser LSR II flow cytometer.

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Measurement of proliferation by BrdU incorporation. BrdU was added to a final concentration of 10 μM after culture for 34 h under WSS or static conditions. WSS or static culture was maintained for an additional 2 h, and then cells were washed in PBS and recovered by Accutase treatment. Cells were incubated sequentially in 2% FBS-PBS for 20 min with
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