Sorcin is involved during embryo implantation via activating VEGF/PI3K/Akt pathway in mice

Kanchan Gupta, Vijay Kumar Sirohi, Suparna Kumari, Vinay Shukla, Murli Manohar, Pooja Popli and Anila Dwivedi

Division of Endocrinology, CSIR-Central Drug Research Institute, Lucknow, Uttar Pradesh, India

Correspondence should be addressed to A Dwivedi: anila.dwivedi@rediffmail.com

Abstract

Our earlier studies have demonstrated the cyclic variation and also the altered expression of sorcin in endometrium during early-to-mid-secretory phase transition in women with unexplained infertility. The current study was undertaken to establish the functional role of sorcin in endometrial receptivity in mice. Results indicated that sorcin was highly expressed during the window of implantation in mice and functional blockage of sorcin caused significant reduction in number of implanted blastocyst. The receptivity markers (i.e. Integrin β3, HBEGF, IGFBP1, WNT4 and Cyclin E)) were found to be downregulated in sorcin knocked down uterine horn on day 5 as compared to untreated horn. The reduced attachment and expansion of BeWo spheroids on RL95-2 endometrial cells with sorcin knock down, in in vitro model of endometrium–trophoblast interaction further supported these findings. Uterine sorcin expression pattern during estrous cycle and in delayed implantation mice model suggested the upregulation of sorcin by estrogen. The functional blockade of sorcin induced the intracellular Ca²⁺ levels in endometrial epithelial cells (EECs), which indicated that altered Ca²⁺ homeostasis might be responsible for implantation failure. Sorcin silencing led to significant reduction in the expression of angiogenic factor VEGF and its downstream effector molecules i.e. PI3K, Akt and NOS. The migratory and invasive properties of HUVECs were abrogated by anti-VEGF or by adding culture media from sorcin blocked EECs, which indicated that sorcin might mediate angiogenesis during implantation. Taken together, sorcin is involved in the regulation of Ca²⁺-mediated angiogenesis via VEGF/PI3K/Akt pathway in endometrial cells and plays a crucial role in preparing the endometrium for implantation.

Key Words

- sorcin
- embryo implantation
- VEGF
- angiogenesis

Introduction

Embryo implantation involves a complex sequence of signaling events that are crucial for the establishment of pregnancy (Cha et al. 2012). A large number of identified molecular mediators have been postulated to be involved in early feto-maternal interaction (Kodaman & Taylor 2004, Yoshinaga 2008). These mediators include molecules associated with uterine receptivity e.g. cytokines, growth factors, adhesion molecules and extracellular matrix components, which play a crucial role in preparation and development of an appropriate endometrium for blastocyst adhesion and implantation (Lessey et al. 1992, Singh et al. 2010). In rodents, the implantation process initiates with an increased uterine vascular permeability at the sites of blastocyst apposition where VEGF plays significant role (Demir et al. 2010, Zhang et al. 2013). The cyclic changes in vascular endothelial growth factor
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Materials and methods

Animals and tissue preparation

Adult Balb/c mice aged 8–10 weeks were used in the study, and all experiments were performed according to the standards specified by the Institutional Animal Ethics Committee of Central Drug Research Institute, Lucknow. The estrous cycle was staged by examining the vaginal smears as previously described (Byers et al. 2012).

Delayed implantation model

Delayed implantation was induced in mice as previously described (Liang et al. 2010). Pregnant mice were treated with a subcutaneous injection of estradiol (100 ng/30 g body weight), progesterone (1 mg/30 g body weight) or a combination of the same doses of estradiol and progesterone, whereas mice of control group received the vehicle only. After 24 h of hormonal treatment, uterine samples were collected.
Bioneutralization of sorcin

Pregnant female mice underwent mini laparotomy under anesthesia on D4 of the window of implantation to deliver the 1.5μg of sorcin antibody (sc-100859; Santa Cruz Biotechnology) in one of the uterine horn, while the other horn (control) received normal mouse IgG, as described previously (Huang et al. 2012). On D5 and D10 of pregnancy, animals were killed and their uteri were photographed to record the number of implanted embryos.

In vivo sorcin knock down

Pregnant female mice underwent mini laparotomy under anesthesia on D3 of pregnancy to deliver 50nM of sorcin siRNA (sc-41017; Santa Cruz Biotechnology) in a final volume of 25μL with Lipofectamine RNAI MAX reagent (Thermo Fisher Scientific) per manufacturer’s instructions in one of the uterine horn. The other horn (control) received scrambled siRNA, as described (Zhang et al. 2012, Guo et al. 2014, Long et al. 2015). On D5 and D10 of pregnancy, animals were killed and their uteri were photographed to observe the number of implanted embryos. Uterine horns of D5 of pregnancy were dissected out and flushed with saline and stored in –80°C till further processing.

Immunohistochemistry

Immunohistochemical analysis was performed with minor modifications as described previously (Manohar et al. 2014) using ImmunoCruz ABC staining system (Santa Cruz Biotechnology). Briefly, formalin-fixed and paraffin-embedded uterine sections (5μm) were de-paraffinized and rehydrated. The sections were incubated with 0.5% H₂O₂ in PBS for 5 min to block endogenous peroxidase activity and blocked with 5% bovine serum albumin along with normal goat serum for 2h. The sections were then incubated with primary antibody anti-sorcin (sc-100859) in 1:300 dilutions for overnight at 4°C. For the negative control, IgG was added to the sections and kept for overnight at 4°C. Further, sections were incubated in biotinylated secondary antibody (1:500; ImmunoCruz ABC staining system) for 30 min followed by ABC enzyme reagent incubation. Staining was achieved with peroxidase substrate DAB for 5 min, counterstained by hematoxylin for 30s and mounted with DPX (Sigma-Aldrich). Images were captured under light microscope (ECLIPSE TE2000-S, Nikon) at 400× magnifications. The staining intensity of all these proteins in glandular epithelium, luminal epithelium and stromal compartment were quantified by image analysis software Image-Pro Plus 4.0 (Maryland, USA), and results were expressed as % image analysis score.

Primary endometrial epithelial cell culture

Uteri were taken out from adult female mice and tissue was collected in DMEM (Sigma-Aldrich), minced into 1 mm pieces and incubated with collagenase (1 mg/1 mL) and DNase (2 mg/mL) in DMEM for 2 h at 37°C with regular mixing. Digested tissue was mechanically dissociated through a 1 mL tip and resuspended in 2 mL of fresh DMEM. Supernatant was collected and centrifuged for 2 min at 600g and then the pellet was dissolved in DMEM. Cells were allowed to pass through a 40μm cell strainer placed on top of an open 50mL falcon tube. Epithelial cells retained in the strainer were collected in DMEM and then transferred to tissue culture flasks (75 cm², Corning). Epithelial cells were incubated at 37°C with saturating humidity and 5% CO₂.

Trophoblast spheroid adhesion assay

In order to study the effect of sorcin silencing on embryo implantation, we have quantified attachment of BeWo (human trophoblast cells, obtained from cell repository of National Centre for Cell Science, Pune, India) spheroids to the RL95-2 (human uterine EECs purchased from American Type Culture Collection (Manassas, VA, USA)) monolayer as previously described (Ho et al. 2012). RL95-2 cells were seeded into 96-well plates (5 × 10³ cells per well), and cultured for 24 h to form a monolayer. Cells were washed and transfected according to manufacturer’s protocol. Briefly, 20nM of sorcin siRNA was diluted in 50μL of culture media and added with 50μL of diluted Lipofectamine RNAI MAX Reagent (Invitrogen). The mixture was allowed to incubate at room temperature for 30 min then given to RL95-2 monolayer for 6 h followed by incubation of 24 h. Trophoblast spheroids were generated by agitation of BeWo cells on an orbital shaker at 110 rpm for 16 h. Spheroids (n = 30) were delivered to each well over a confluent monolayer of RL95-2 cells in triplicate and incubated for 1 h at 37°C in a humidified atmosphere with 5% CO₂. After 1 h, non-adherent spheroids were removed by centrifugation (with the cell spheroid surface facing down) at 12g for 5 min. Adhered spheroids were counted under a microscope and percent adhesion was calculated by the formula: Ratio of number of adhered spheroids/number of total spheroids delivered, multiplied by 100.
Trophoblast spheroid expansion assay

RL95-2 cells were seeded into 96-well plates (3 × 10^5 cells per well) and cultured for 24h to form monolayer. Then, trophoblast spheroids were transferred to the monolayer (3–5 spheroids/well). To delineate the margins of trophoblast spheroids after co-culture, the spheroids were stained with 10 µM of CellTracker Green CMFDA (Invitrogen) for 30min before adding to monolayer. Spheroids were photographed after 1 h and 24 h co-culture. The length (L) and width (W) of spheroids were measured. The size of spheroids was calculated by the formula: (L+W)/2. The expansion of spheroids was determined on the basis of the spheroid size at 1 h after co-culture. Images were captured using fluorescent microscope (Nikon Eclipse 80i, Nikon) to detect fluorescence.

Western blot analysis

Mouse uterine tissue samples/EECs were lysed in RIPA buffer (Sigma) supplemented with protease inhibitor cocktail (Sigma) and 1 mM PMSF and incubated overnight in −20°C. Supernatant was collected by centrifugation at 20,000g for 15 min at 4°C. The protein concentration was determined by Bradford method (Manohar et al. 2014). Equal amounts of protein (20 µg) were separated by gel electrophoresis and then transferred to Immuno-BlotTM PVDF membrane (Millipore). The membrane was blocked with 5% skimmed milk and then incubated with appropriate primary antibody (anti-sorcin: sc-100859; anti-VEGF: sc-7269; anti-PI3K: sc-1637; anti-p-PI3K: sc-293115; anti-Akt: #9272; anti-p-Akt: #9271; anti-VEGFR-2: sab4501645 and anti-NOS: sc-654) at 1:1000 dilution in 0.5% BSA in Tris buffered saline (TBS) and kept overnight at 4°C. The membrane was then washed twice with TBST for 5 min and incubated with a secondary peroxidase-conjugated antibody (1:3000) for 1 h. Antibody binding was detected using enhanced chemiluminescence detection system (GE Healthcare). After developing, the membrane was stripped and re-probed using another primary antibody of interest or β-actin (sc-1616; Santa Cruz Biotechnology) to confirm equal loading. Each experiment was repeated three times. Quantitation of band intensity was performed by densitometry using Quantity One software (v.4.5.1) and a Gel Doc imaging system (Bio-Rad).

Immunofluorescence imaging

Mouse primary EECs were grown on coverslips in 12-well plate and treated with E_2 (10 nM), P_4 (1 µM) or combination of E_2 (10 nM) + P_4 (1 µM) in primary EECs for 24 h to determine the expression level of sorcin.

In addition, the expression of sorcin and VEGF receptor-2 (VEGFR-2) was demonstrated in sorcin siRNA or scrambled siRNA-transfected primary EECs from mouse. Cells were fixed in paraformaldehyde (4%) for 20 min and permeabilized with 0.1% Triton X-100 followed by washing with PBS and blocked with 2% BSA. Next, cells were incubated with anti-sorcin (1:300) antibody for overnight followed by 1-h incubation with fluorescence-tagged secondary anti-mouse antibody (1:500), and then mounted on slides with SlowFade Gold Antifade reagent with DAPI (Molecular Probes, Life Technologies). Images were captured at 400× using Nikon microscope (Nikon Eclipse 80i, Nikon) to detect fluorescence and DAPI emissions.

Flow cytometric analysis

Primary EECs were seeded (2 × 10^5 cells/well) into 6-well plate and maintained overnight in phenol red-free media containing 10% stripped FBS. Next day, cells were transfected with sorcin siRNA for 6 h in serum-free media according to manufacturer’s protocol. Cells were washed with PBS and incubated in serum-free DMEM medium for 24 h. After 24 h, cells were collected by trypsinization and resuspended in PBS. Fluo-3-AM dye (2 µM) was added in each sample for 30 min at 37°C in the dark with continuous shaking. Cytosolic free Ca^{2+} measurement was performed using a FACScan flow cytometer (BD Biosciences, San Jose, CA, USA) with excitation and emission settings at 506 nm and 526 nm, respectively (Zhang et al. 2004). These experiments were performed three times.

Enzyme-linked immunosorbent assay

Primary EECs were seeded in 6-well plates (2 × 10^4 cells/well) and grown to confluence. Transfection of sorcin siRNA was given for 6 h, and then cells were washed with PBS. Cells were washed with PBS and incubated in serum-free DMEM medium for 24 h. The culture supernatant was collected to measure VEGF concentration using an ELISA kit as specified by the manufacturer (Abcam).

Cell proliferation assay in HUVECs

MTT assay was used to determine cell viability. Human umbilical vascular endothelial cells (HUVECs, purchased from Life Technologies) were seeded in 96-well plate at a density of 3 × 10^3 cells/well and allowed to adhere for
24 h. Conditioned media obtained from endometrial cell culture (control and sorcin siRNA transfected) were added (20% V/V) to these cells for 48 h. In additional group, a blocking anti-VEGF-A antibody was added in the conditioned media from scrambled siRNA-treated EECs. (Print et al. 2004). At the end of incubation, 100 µL of MTT (0.5 mg/mL) was added to the cells and incubated for 2 h at 37°C. Following incubation, the supernatant was removed, and 100 µL of dimethyl sulfoxide (DMSO) was added. The formazan crystals formed inside the viable cells were solubilized in DMSO, and the optical density was read with Microquant (Bio Tek) at 540 nm.

Wound healing assay

HUVECs were seeded in 6-well plate and allowed to attain 90–100% confluency. Conditioned media obtained from scrambled control or sorcin siRNA-transfected EECs (20% v/v) added to HUVECs, for 24 h. A vertical scratch was made in the middle of well using a 10 µL tip and washed with PBS. Images were captured at 24 h after supernatant addition. Migration rate was measured by calculating the percent wound closure using the following formula:

\[
\text{\% wound closure} = \frac{\text{wound at T} - \text{wound at Th}}{\text{wound at T}} \times 100; \text{ where } T \text{ denotes ‘time’}
\]

Cell migration/invasion assay

Cell migration assay was conducted in a 24-well format using cell culture inserts (BD Biosciences) with or without matrigel having a membrane pore size of 8.0 µm (Sirohi et al. 2017). Briefly, 2 × 10⁵ HUVECs were seeded into the top chamber. Conditioned media (20% v/v) collected from different groups of EECs were added into the top chamber for 48 h. At the end of the incubation, non-migrated cells present on the upper surface of the membrane were gently removed with a cotton swab. Next, cells on the lower surface of the membrane were fixed in chilled methanol and stained with 0.1% crystal violet. The cells migrated towards the lower surface of membrane were detected, photographed and counted under an inverted microscope (Nikon ECLIPSE TE2000-S).

Statistical analysis

All values were presented as the mean ± s.e.m., as determined from at least three independent experiments. Statistical significance was assessed by one-way ANOVA and Newmann–Keul’s test or Student’s t-test. P < 0.05 was considered statistically significant.

Results

Sorcin is upregulated during peri-implantation period and estrous phase

In order to check the differential expression of sorcin during early pregnancy, uterine samples were collected from non-pregnant (NP) mice and from pregnant mice on different days of pregnancy (D1, D4, D5, D6 and D7) and Western blotting was performed. Results indicated that sorcin expression was increased on D4 and D5 i.e. window of implantation and the level was maintained till D6 as compared to NP group (P < 0.001). On D7, the expression of sorcin was gradually decreased as compared to D5 (P < 0.001) (Fig. 1A). The expression of sorcin was higher at implantation sites but remained low at IIS as observed on D5, D6 and D7; the levels at IIS were comparable to NP uterine tissue (Fig. 1A). Significantly increased expression of sorcin on D4 and D5 revealed its possible role during embryo implantation and endometrial receptivity.

Study on estrous cycle showed that the expression of sorcin was significantly high (P < 0.001), during the estrus and metestruus phases as compared to that in proestrus phase (Fig. 1B).

Immunohistochemical localization of sorcin

To check the cell-specific expression and function of sorcin, immunohistochemistry was performed in uterine tissue. Sorcin expression was increased in the luminal epithelium (LE), glandular epithelium (GE) and also in the stromal cells (S) on D4 of pregnancy (Fig. 2) as compared to that in NP group. As analyzed on D5, in implantation sites, the expression of sorcin was significantly high (P < 0.001) in LE (~6-fold), followed by GE (~5.4-fold) and in stroma (~4-fold). The increased expression of sorcin in these cells was continued till D6, whereas on D7, expression of sorcin was detected to be high in stroma (P < 0.001) but was less in GE and LE (Fig. 2). Thus, sorcin may be involved in decidualization process as still high levels were detected in stroma during post-implantation.

Inhibition of sorcin caused suppression of uterine receptivity and failure of embryo implantation in mice

Further, to see whether sorcin is critical for embryo implantation, intrauterine blocking experiments were performed by sorcin-antibody or siRNA injection. Administration of anti-sorcin antibody at D4 p.c. in the uterine horn lumen reduced the number of implants.
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Effectively (~5-fold) as observed on D5 and D10, whereas uterine horn treated with control IgG showed normal number of implantations (Fig. 3A). Similarly, the number of implantation sites was significantly reduced (~6-fold) after the intrauterine injection of sorcin-specific siRNA given on D3 of pregnancy, as compared to that in scrambled siRNA-treated group (Fig. 3B).

The expression of protein level of receptivity markers like Integrin β3, IGFBP1, WNT4, Cyclin E and HBEGF was significantly downregulated ($P<0.001$) in siRNA-injected horn as compared to that in scrambled siRNA (control) group (Fig. 3C) as analyzed on D5. Significant reduction in receptivity markers suggested that sorcin inhibition suppressed the endometrial receptivity and thus reduced the implants number in pregnant mice.

Embryos recovered from the sorcin knocked down horn were morphologically dissimilar as compared over embryos collected from scrambled siRNA-transfected uterine horn (Supplementary Fig. 1, see section on supplementary data given at the end of this article).

We also checked whether control IgG has any adverse effect on implantations. No significant change was observed in control IgG-treated horn as compared to untreated horn (Supplementary Fig. 3).

**Sorcin silencing decreased the attachment and expansion of trophoblast spheroids on EECs monolayer**

To analyze whether sorcin is essential for attachment of blastocyst to the endometrial cells, the *in vitro* attachment assay was performed using endometrial epithelial RL95-2 cells monolayer and BeWo trophoblast spheroids. Transfection efficiency of sorcin siRNA in RL95-2 was confirmed by Western blot analysis. It showed ~2.5-fold reduction in sorcin expression after 24 h of knock down (Supplementary Fig. 2A). Transfection of sorcin siRNA to RL95-2 cells significantly decreased the percent attachment of BeWo spheroids to RL95-2 i.e. EEC monolayer. More than 70% of the spheroids were attached after 1 h of co-culture in control and scrambled groups but in sorcin siRNA-transfected group attachment was decreased up to 30% ($P<0.001$) (Fig. 4A). We next investigated whether the expansion of trophoblast spheroids on an EEC monolayer could be reduced by the treatment of EECs with sorcin siRNA. Fluorescent, phase-contrast and merged microscopic images have shown that spheroids attached to EEC monolayer in co-culture, began to flatten and spread on the EEC monolayer at 24 h (Fig. 4B). However, the decreased expansion (~3-fold) of trophoblast spheroids on the monolayer was observed on EECs treated with sorcin siRNA. Sorcin siRNA treatment caused reduction ($P<0.001$) in expansion as compared to that observed in scrambled siRNA control group. These *in vitro* results showed that sorcin is involved in endometrial receptivity and facilitates attachment and expansion of trophoblast on EECs.

**Estradiol upregulates the expression of endometrial sorcin**

Ovariectomized and delayed implantation mice models were used to examine whether sorcin expression is regulated by ovarian steroid hormones. In ovariectomized mice, sorcin expression was significantly induced by estrogen treatment, whereas progesterone alone had no effect. Combination of estrogen and progesterone also showed stimulatory effect on sorcin expression (Fig. 5A). Interestingly, similar results were found in delayed implantation model, and data indicated that progesterone
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alone had no effect on sorcin, but with estrogen surge, the sorcin expression was significantly upregulated in mice uterus (P<0.001) (Fig. 5B).

Immunohistochemical analysis suggested that expression of sorcin was high in LE, GE and stromal cells on the activation of implantation as compared to that in delayed implantation group (Fig. 5C).

Moreover, in in vitro experiment using primary culture of mouse EECs, the results of Western blot analysis and immunocytochemistry showed that sorcin expression was induced in E2-treated EECs and also the combination of estrogen and progesterone showed a stimulatory effect on sorcin expression (P<0.001) (Fig. 5D and E). This in vitro experimental data further support our in vivo data as shown in Fig. 5A. These results clearly indicate that sorcin is regulated by estrogen in mouse endometrium.

**Sorcin silencing increased the intracellular calcium level in mouse EECs**

Since sorcin is a calcium-binding protein, it may affect the calcium homeostasis in endometrium during embryo
implantation thus our next aim was to measure the effect of sorcin on intracellular Ca\(^{2+}\) in EECs. For this, mouse primary EECs transfected with sorcin siRNA or scrambled siRNA and knockdown efficiency of sorcin siRNA was confirmed by Western blot analysis. A significant reduction in sorcin expression (\(P<0.001\)) was observed 24 h after sorcin silencing (Supplementary Fig. 2B). The results of flow cytometric analysis revealed that silencing of sorcin in EECs significantly enhanced (~2.2 fold) the influx of intracellular calcium in these cells as compared to that in scrambled siRNA-transfected EECs (\(P<0.001\)) (Fig. 6A).

**Sorcin knock down significantly reduced the expression of VEGF, VEGFR-2 and its downstream signaling molecules**

Since angiogenesis is an important phenomenon during embryo implantation and is mediated mainly by VEGF (Zhang et al. 2001, Wang & Dey 2006, Demir et al. 2010, Bagheri et al. 2017), we measured the expression of VEGF and its associated downstream effector molecules like PI3K, AKT and NOS that are known to be involved in the process of angiogenesis. Quantification of VEGF level was analyzed in fresh conditioned media collected 24 h after
sorcin siRNA transfection in mouse EECs. Interestingly, VEGF level was found to be low (P < 0.01) in conditioned media collected from transfected cells as compared to control group (Fig. 6B). Further, we performed immunocytochemical analysis of VEGFR-2 in mouse EECs transfected with sorcin siRNA. Our results showed reduced expression of VEGFR-2 in the mouse EECs (Fig. 6C).

Figure 6
Effect of sorcin silencing on cytosolic free Ca$^{2+}$, VEGF level, VEGFR-2 expression in mouse primary EECs in vitro and on signaling molecules (PI3K, Akt and NOS) in D5 uterus in vivo. (A) Fluorescence was measured from cells loaded with 2 µM Fluo-3-AM dye. Exposure of cells to 10 µM ionomycin induced a Ca$^{2+}$ transient. (B) VEGF level in culture supernatants of mouse primary EECs measured by enzyme-linked immunosorbent assay. (C) Representative images of immunocytochemical analysis showing VEGFR-2 expression in EECs. (D) Representative images showing Western blot analysis of VEGF, VEGFR-2, PI3K, Akt and NOS in uterine horn collected on D5 pregnancy. Details have been given in ‘material and methods’ section. Results are expressed as mean ± s.e.m. P values are a P < 0.001 and b P < 0.01 vs scrambled control. A full color version of this figure is available at https://doi.org/10.1530/JME-17-0153.

Figure 7
The proliferation, migration and invasion assay of HUVECs. (A) The proliferation of HUVECs was detected in various groups; HUVECs were cultured with supernatants (20% V/V) from mouse primary EECs transfected with sorcin siRNA or scrambled siRNA. In additional group, a blocking anti-VEGF antibody was added in the supernatant from scrambled siRNA-treated EECs. Details have been given in ‘materials and methods’ section. (B) Analysis of migration of HUVECs was performed by scratch-wound healing assay. Representative images showing migration of HUVECs cultured for 24h in the supernatant of control EECs and in the supernatant of EECs with sorcin blockade (×200 magnification). Cell migration distance of the cells between the scratch edges was observed after 24h. (C) Transwell migration and matrigel invasion assays were done to analyse migration and invasion of HUVECs. The graph in the lower panel shows the relative percentage of migrated and invaded cells over control in three wells in each independent experiment (random fields were photographed and number of cells from three fields per well were counted). Results are expressed as mean ± s.e.m., n = 3. P values are a P < 0.001 vs scram control. A full color version of this figure is available at https://doi.org/10.1530/JME-17-0153.
Further, the effect of in vivo silencing (on D3) of sorcin on expression of VEGF, VEGFR and downstream effectors was analyzed by Western blotting on D5 in pregnant mice. Results revealed a significant reduction in the expression of VEGF, VEGFR-2, PI3K, AKT and NOS ($P<0.001$) in uterine horn transfected with sorcin siRNA (Fig. 6D). Above results indicated that sorcin mediates its angiogenic effect by regulating VEGF/PI3K/Akt signaling.

**Sorcin blockade suppressed endothelial cell proliferation, migration and invasion**

Next, to analyze whether sorcin affects angiogenesis in endometrium, we performed experiments on HUVECs. HUVECs cultured in medium from sorcin-silenced EECs contained low VEGF concentration as compared to control group ($P<0.01$) (Fig. 6B). To determine the angiogenic activity of the HUVECs, MTT assay, wound healing assay and matrigel migration/invasion assay were performed. The results revealed that 48 h after the addition of the culture medium from sorcin-silenced EECs, the proliferation, migration and invasion ability of HUVECs were significantly reduced ($P<0.001$) as compared to that of control group (Fig. 7A, B and C). When supernatant from control EECs was supplemented with anti-VEGF, similar results were observed with these parameters in HUVECs. Results indicate that the angiogenesis-inducing properties of sorcin (expressed in EEC) can be accounted for sorcin’s VEGF-inducing ability.

**Discussion**

To the best of our knowledge, present study is the first report to investigate the expression, regulation and physiological function of sorcin in mouse uterus during the period of uterine receptivity and embryo implantation. The dynamics of sorcin protein expression showed low expression on D1, which was found to be increased on subsequent days and reached to maximum on D5 in mice. In mice, window of implantation lasts for less than 24 h i.e. initiates from D4 evening of the pregnancy thereafter uterus becomes non-receptive (refractory) to implantation by the afternoon of D5 (Wang & Dey 2006). The findings indicated the possible role of sorcin during window of implantation in mice.

The spatio-temporal distribution study of sorcin in mouse uterus revealed that the expression of sorcin was significantly high in all compartments on D4 to D6. The expression was declined in LE and GE on D7, but was still detected high in stroma. The high expression of sorcin in the stromal cells on D7 indicated the possible involvement of sorcin in stromal cells-driven-event ‘decidualization’, which begins just after embryo implantation (Abrahamsohn & Zorn 1993, Wang & Dey 2006, He et al. 2015) in mice. Thus, sorcin appears to play a critical role during endometrial receptivity and subsequently may also be involved in decidualization process as it continued to be expressed in stromal cells after implantation.

To analyze whether sorcin is important for embryo implantation, we knocked down the expression of
sorcín. The intra-uterine injection of sorcín siRNA/sorcín antibody reduced the number of implantation sites in pregnant mice. Additionally, the downregulation of receptivity markers (Integrin β3, HBEFG, IGFBP1, WNT4 and Cyclin E) in sorcín knocked down uterine horn confirmed the functional role of sorcín in endometrial receptivity. The result of in vitro attachment assay further provided the direct evidence that sorcín renders endometrium receptive both for attachment and invasion process. Herein, EECs monolayer transfected with sorcín siRNA showed remarkable reduction in attachment of BeWo spheroids on EECs. Moreover, significant inhibition in spheroids outgrowth attached to EECs was seen after 24 h of co-culture. These in vivo and in vitro data clearly indicate the critical involvement of sorcín in embryo implantation.

Ovarian steroids play an important role in uterine tissue remodeling and making the endometrium receptive for embryo implantation (Aplin & Kimber 2004, Paulson 2011, Cha et al. 2012). Serum E₂ and P₄ levels vary throughout the estrous cycle (Wood et al. 2007). Therefore, it was interesting to analyse the E₂- and P₄-mediated regulation of sorcín expression in endometrium. In delayed implantation experiment, the sorcín expression was significantly increased when embryos were implanted on giving E₂ surge to terminate the delay in pregnancy on D7. Also, E₂ significantly promoted the expression of sorcín in the uterus of ovariectomized mouse. These results were consistent with the observation expression pattern of sorcín during estrous cycle, which was increased during estrogen-dominant estrus phase and declined during progesterone-dominating diestrus phase. The in vitro experiment on mouse EECs also showed that estrogen alone or in combination with progesterone enhanced the expression of sorcín in these cells. Collectively, evidences suggest that expression of sorcín is regulated by estrogen in uterus during estrous cycle and also during early pregnancy.

Calcium homeostasis is a necessary phenomenon during embryo implantation and the maintenance of calcium level is important at embryo–uterine interface (Yoshinaga 2008). Several calcium-binding proteins such as CaBP-9k, CaBP-28k and S100 protein family members have been reported to play a role in the maintenance of endometrial receptivity (Luu et al. 2004, Tong et al. 2010, Liu et al. 2012). The roles established for calcium-binding proteins emphasize their ability to enhance Ca²⁺ transport and increase the capacity of cells to store Ca²⁺ (Luu et al. 2004). Sorcín increases calcium accumulation in the endoplasmic reticulum and decreases cytoplasmic Ca²⁺ and maintains calcium level in fibroblasts cells (Lalioti et al. 2014). However, the role of sorcín-mediated calcium homeostasis during embryo implantation is not yet well known. In our study, mouse EECs after sorcín knock down, showed increase in cytoplasmic Ca²⁺ level. Sarco/endoplasmic reticulum Ca²⁺ATPase (SERCA) activation is reported to be involved in the induction of angiogenesis by promoting proliferation and migration of vascular smooth muscle cells (Shukla et al. 1997, Birkett et al. 1999). Sorcín activates SERCA, which further increases calcium accumulation in endoplasmic reticulum (ER) (Lalioti et al. 2014), whereas thapsigargin (SERCA inhibitor) inhibits angiogenesis via depletion of intracellular calcium pools in human umbilical vein endothelial cells (HUVECs) (Shukla et al. 2001). The depletion of sorcín is also known to decrease VEGF expression and cause inhibition of epithelial-to-mesenchymal transition as studied in breast cancer cells (Hu et al. 2014). Angiogenesis is an important phenomenon during embryo implantation and is mediated mainly by VEGF (Zhang et al. 2001, Wang & Dey 2006, Demir et al. 2010). Thus, we checked whether sorcín was involved in calcium-mediated angiogenesis in the uterus for successful establishment of pregnancy in mice. Interestingly, we found a significant decline in the expression of both VEGF and its receptor VEGFR-2 in the uterine horn receiving sorcín siRNA. The reduction in VEGF level was also found in conditioned media collected from sorcín siRNA-transfected EECs. Since the phenomenon of angiogenesis involves the proliferation and migration of endothelial cells, the effect of sorcín silencing on angiogenesis was checked on human umbilical vein endothelial cells (HUVECs) (Shukla et al. 2001, Print et al. 2004). The HUVECs cultured in media collected from sorcín-depleted EECs, showed a reduction in the proliferation, migration and invasion thereof. Similarly, administration of anti-VEGF antibody reversed the effect of sorcín on HUVECs. These results clearly indicate that sorcín regulates angiogenesis in the endometrium via regulating VEGF pathway, during embryo implantation.

VEGF mediates its action via VEGFR and activates PI-3-kinase/Akt-dependent activation of nitric oxide synthase (NOS), which plays an important role during embryo implantation (Morales-Ruiz et al. 2000, Ma et al. 2003, Koch et al. 2011). The higher NOS expression was detected in peri-implantation uterus of mice leading to higher nitric oxide (NO) production, which is required for successful embryo implantation (Novaro et al. 1997, Saxena et al. 2000). NO is known to act as a local vasodilator and an angiogenic mediator in murine
endometrium (Novaro et al. 1997, Saxena et al. 2000). In our study, the sorcin silencing inhibited the activation of PI3-K and Akt, which led to suppression of activation of downstream NOS required for NO production in uterus. The suppressed production of NO might adversely affect the angiogenesis at the implantation sites, which can lead to the implantation failure (Saxena et al. 2000). Thus, our studies revealed that sorcin is essential for maintaining the activation of VEGF-mediated signaling and production of NO in endometrium required for successful implantation in mice.

Conclusion

Taken together, the current study demonstrated that sorcin may have functional significance in the initiation and maintenance of the window of receptivity and subsequent embryo implantation. Attenuation of sorcin expression led to embryo implantation failure in both in vivo and in vitro settings. Sorcin is tightly regulated by E2 during window of implantation and exerts its effect via VEGF/PI3-K/Akt pathway and facilitates angiogenesis at the site of embryo implantation (Fig. 8). Further, the altered regulation of sorcin might cause imbalance in Ca2+ homeostasis and might result in inadequate functioning of endometrium leading to recurrent implantation failure. However, the role of sorcin in developing embryos needs to be explored further, which might delineate the absolute cause of pregnancy failure due to suppressed expression of sorcin. This study explored the molecular function of sorcin in establishment of pregnancy, thus provides a significant addition towards understanding the implantation biology and infertility. Future studies on clinical validation of sorcin in infertile patients are likely to open the opportunities for the development of novel strategies for infertility management.

Supplementary data

This is linked to the online version of the paper at https://doi.org/10.1530/JME-17-0153.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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