Epsin15 Homology Domains: Role in the Pathogenesis of Pulmonary Arterial Hypertension

Dan Predescu†, Shanshan Qin†, Monal Patel†, Cristina Bardita‡, Rabia Bhalli and Sanda Predescu*†

Division of Pulmonary Medicine, Critical Care and Sleep Medicine, Department of Internal Medicine, Rush Medical College, Rush University, Chicago, IL, United States

Intersectin-1s (ITSN) deficiency and expression of a biologically active ITSN fragment, result of granzyme B cleavage under inflammatory conditions associated with pulmonary arterial hypertension (PAH), are characteristics of lung tissue of human and animal models of PAH. Recently, we have shown that this ITSN fragment comprising two Epsin15 homology domains (EH\textsubscript{ITSN}) triggers endothelial cell (EC) proliferation and the plexiform arteriopathy in PAH. Limited evidence also indicates that the EH domains of endocytic proteins such as ITSN, upregulate compensatory endocytic pathways in cells with impaired vesicular trafficking. Thus, we sought to investigate whether the EH\textsubscript{ITSN} may be involved in this compensatory mechanism for improving the EC endocytic dysfunction induced by ITSN deficiency and possibly contribute to PAH pathogenesis. We used stably-transfected human pulmonary artery ECs expressing the Myc-EH\textsubscript{ITSN} (EC\textsubscript{EH−ITSN}) and ITSN knockout heterozygous mice (K0\textsubscript{ITSN}/−) transduced with the Myc-EH\textsubscript{ITSN}, in conjunction with functional assays: the biotin assay for caveolae internalization and 8 nm gold (Au)- and dinitrophenylated (DNP)-albumin perfusion of murine lung microvasculature. Pulmonary artery ECs of PAH patients (EC\textsubscript{PAH}), ITSN knockdown ECs (EC\textsubscript{KD−ITSN}), the monocrotaline (MCT)-induced mouse and rat models of PAH, as well as untreated animals, served as controls. ELISA via streptavidin-HRP or anti-DNP antibody (Ab), applied on ECs and lung lysates indicated greater than 30% increase in biotin internalization in EC\textsubscript{EH−ITSN} compared to EC\textsubscript{Ctrl}. Despite their endocytic deficiency, EC\textsubscript{PAH} internalized biotin similar to EC\textsubscript{Ctrl} which is twofold higher compared to EC\textsubscript{KD−ITSN}. Moreover, the lung microvascular bed of Myc-EH\textsubscript{ITSN}-transduced mice and MCT-treated animals showed greater than twofold increase in DNP-BSA transendothelial transport, all compared to untreated controls. Electron microscopy (EM) revealed the increased occurrence of non-conventional endocytic/transcytotic structures (i.e., caveolae clusters, tubulo-vesicular and enlarged endocytic structures, membranous rings), usually underrepresented. Most of these structures were labeled by Au-BSA, consistent with their involvement in the transendothelial transport. Furthermore, ITSN deficiency and EH\textsubscript{ITSN} expression alter the subcellular localization of the EH-binding protein 1 (EHBP1) and cortical actin.
organization, altogether supporting the increase occurrence/trafficking of the alternative endocytic structures. Thus, the EH\textsubscript{ITSN} by shifting the physiological vesicular (caveolae) transport toward the alternative endocytic pathways is a significant contributor to the dysfunctional molecular phenotype of E\textsubscript{GPAH}.

**Keywords:** intersectin, pulmonary arterial hypertension, endocytic deficiency, alternative transport pathways, non-conventional endocytic/transcytotic structures, EH-binding protein 1, cortical actin

**INTRODUCTION**

Patho-physiologically, pulmonary arterial hypertension (PAH) is defined by mean pulmonary arterial pressure exceeding the upper limits of typical values, i.e., ≥ 25 mmHg at rest, as result of extensive pulmonary vascular dysfunction (Stenmark et al., 2009; Sehgal and Lee, 2011). As a multiple etiology disease, PAH is triggered by diverse internal and/or external factors and it is perpetuated by a plethora of signaling events and altered cellular interactions at the level of lung vascular bed (Tuder et al., 2001, 2007, 2009). At the cellular level, PAH affects all segments of the pulmonary vascular bed (Chazova et al., 1995). Concentric and eccentric intimal thickening, medial hypertrophy, lumen obliteration and recanalization affect the pulmonary arteries. Modest venous hypertrophy due to increase in the intimal and adventitial thickness and capillary bed rarefaction are among the structural changes of the pulmonary vasculature in PAH, as well (Chazova et al., 1995). In most typical cases, an endothelial driven patchy proliferation of intima and formation of vascular lesions occur (Chazova et al., 1995). The plexiform lesions, hallmark of established PAH, are found at branching points in the small pulmonary arterioles; they are glomeruloid-like vascular structures, lumen-obliterating and composed predominantly of actively dividing and apoptosis-resistant endothelial cell (ECs) (Tuder et al., 1994; Sakao et al., 2005).

The plexiform arteriopathy, the rarefaction of lung capillaries and the venous hypertrophy, as leading drivers of PAH, depend on the status of pulmonary endothelium. The ECs that form the endothelial layer to separate the blood vessel lumen from the rest of the body were considered inert bystanders for an extended period (Tuder et al., 1994). However, the slow dividing and metabolically stable ECs, have the genomic fitness to switch to a migratory, proliferative, apoptotic-resistant and metabolically active phenotype in response to extra- and intracellular signals (Stevens and Gillespie, 2007). Thus, any dysfunction of ECs, permanently exposed to significant variations in physiological parameters (tension, stretch, shear) and to different chemical composition of the surroundings (blood on the luminal and the interstitial fluid on the abluminal side) is at the core of most vascular pathologies, PAH included. Current evidence indicates that paradoxical increase in EC apoptosis followed by exuberant proliferation, and formation of obliterative cellular lesions contribute to the severe vasculopathy in PAH (Long et al., 2015; Zhou et al., 2016). Additional factors such as defects in caveolae trafficking and cytoskeletal organization as well as increased vascular permeability have been linked to PAH pathology (Johnson et al., 2012; Prewitt et al., 2015; Zhou et al., 2016).

Historically, the EC was among the first cell type identified endowed with an abundant population of plasmalemmal vesicles (Palade and Bruns, 1968), re-named caveolae (Rothberg et al., 1992) – involved in endocytosis, one of the fundamental mechanisms by which the cells interact and respond to environmental clues. Moreover, caveolae function in endothelial transcytosis, a process that constitutively couples endo- and exocytosis on the opposite sides of the plasma membrane (Predescu et al., 1997). Endothelial transcytosis provides the morpho-functional basis for sustaining the magnitude, speed, and adaptability of the transendothelial transport as genetically established and functionally shaped to satisfy the general necessities and local requests (Predescu et al., 2007b).

Limited evidence indicates that the EH domains of endocytic proteins confine caveolae to the plasma membrane and upregulate compensatory endocytic pathways in cells with impaired vesicular trafficking such as pulmonary E\textsubscript{GPAH} (Yap et al., 2010; Stoeber et al., 2012). The EH domain which binds to peptides containing the sequence Asp-Pro-Phe (NPF) is implicated in the control of plasma membrane receptor endocytosis and the regulation of intracellular trafficking routes, with significant ramifications into transcriptional regulation and actin cytoskeletal remodeling (Polo et al., 2003).

Intersectin-1s (ITSN), a general scaffold and regulator of the endocytic machinery (Predescu et al., 2003; Knezevic et al., 2011; Patel et al., 2013) comprises two NH\textsubscript{2}-terminal EH domains involved in interactions with several endocytic proteins – Eps15, Stonin2, and SCAMP1 (Yamabhai et al., 1998; Fernandez-Chacon et al., 2000; Martina et al., 2001). Moreover, within the EH2 domain of I\textsubscript{ITSN} there is a well-conserved granzyme B cleavage site, IDQGD71GK (Loeb et al., 2006; Patel et al., 2013). Under inflammatory conditions associated with PAH, granzyme B cleaves I\textsubscript{ITSN} and generates the EH\textsubscript{ITSN}, a biologically active protein fragment, present in E\textsubscript{GPAH} and human lung tissue with plexiform lesions; its expression triggers EC proliferation and selection of a proliferative/plexiform EC phenotype leading to formation of complex pulmonary vasculopathy in a murine model (Patel et al., 2013, 2017). These observations suggest that the EH\textsubscript{ITSN} may facilitate the excessive growth of ECs in PAH lungs not only due to its endothelial proliferative potential but also due to upregulation of alternative transport pathways to compensate for caveolae dysfunction induced by I\textsubscript{ITSN} deficiency (Predescu et al., 2012). Considerations of impaired endocytosis in ECs and its consequences on the pathobiology of PAH are mostly absent; the effects of EH\textsubscript{ITSN} on vesicular trafficking and its role in the regulation of endothelial permeability are not known.
The concept that endocytosis is necessary for some forms of cell signaling (Vieira et al., 1996) has evolved into the hypothesis of endocytosis as a cellular process integrated with, and essential for the execution of various cellular programs like motility, cell fate determination, cellular reprogramming and biogenesis of miRNA, cell division, and transcription (Vieira et al., 1996; Polo and Di Fiore, 2006; Sigismund et al., 2012). Caveolae are multifunctional integrators of endocytosis and signaling (Sigismund et al., 2012). In pathological conditions, dysfunctional ECs also show altered intracellular trafficking and signaling of cell surface receptors such as the transforming growth factor beta receptor 1 (TGFβR1) implicated in the pathogenesis of PAH (Garcia-Rivas et al., 2017; Graf et al., 2018). Endocytic dysfunction and non-productive assembly of the endocytic machinery alter canonical signaling pathways with detrimental consequences on EC function (Mukherjee et al., 2006; Sorkin and von Zastrow, 2019; Bardita et al., 2015).

So far, no systematic studies have assessed the extent of endothelial endocytic dysfunction, and even fewer in vivo analyses measured the degree to which the transendothelial transport is affected. Thus, in this study we used stably-analyses measured the degree to which the transendothelial C

Materials and Methods

Materials

The reagents were obtained as follows: cholesterol, dimethyl dioctadecyl ammonium bromide, bovine serum albumin (BSA), H2O2, glycerol, benzamidine, phenylmethylsulphonyl fluoride (PMSF), sodium pyrophosphate, BSA, desipramine hydrochloride, protease inhibitors cocktail, poly-lysine solution (tissue culture grade), and Hank’s solution from Sigma-Aldrich (St. Louis, MO, United States); Micro BCA protein assay kit and reporter Abs were kindly provided by Dr. Melanie Pritchard (Monash University, Clayton, Australia). Breeding colonies were maintained in the Rush University animal facility. All mice, 6 - to 8-weeks-old, 20 g – 30 g, were kept under standardized housing and feeding conditions. All experiments were done under anesthesia using ketamine (60 mg/kg), acepromazine (2.5 mg/kg) + xylazine (2.5 mg/kg), in 0.1 – 0.2 mL sterile PBS. Mice were genotyped by tail snipping standard procedure (Patel et al., 2017). All experiments were repeated at least three times. No mouse mortality occurred during the study. MCT-mice were generated as previously described (Patel et al., 2013). Wild-type and MCT-treated rats were kindly provided by Dr. Jiwang Chen, University of Illinois at Chicago. All efforts were made to minimize suffering; in addition all studies using rodents adhered to APS's Guiding Principles in the Care and Use of Vertebrate Animals in Research and Training and were performed according with the Rush University IACUC approved protocol number 14-023.

Cationic Liposomes

Cationic liposomes were prepared using dimethyl dioctadecyl ammonium bromide and cholesterol by rotary evaporation of lipid solutions in chloroform, at 42°C, under vacuum (Miyawaki-Shimizu et al., 2006; Predescu et al., 2012). Briefly, the lipid film was rehydrated for 1 h at RT in 5% dextrose followed by sonication and sterile filtration through 0.45 µm membranes. Then, the suspension was extruded through the 50 nm polycarbonate filter mounted on the mini-extruder device, to assure unilamellar vesicles formation. DNA-liposomes complexes were prepared using 8 nmoles liposomes: 1 µg Myc-EHITSN DNA (amino acids 1 – 271) cloned into the pReceiver/Myc-M43 vector (GeneCopeia, Rockville, MD, United States), a ratio found in previous studies to induce efficient protein expression without pulmonary toxicity (Patel et al., 2017). Long-term Myc-EHITSN protein expression in mouse lungs was achieved by repeated Myc-EHITSN DNA-liposome delivery via retro-orbital injections, every 48 h, for 18 days. A mutant EHITSN-W263A fragment in which the tryptophan (W) 263 was substituted with alanine (A) was cloned into the same vector and used as control (Patel et al., 2017). At the end of
the treatment period, mice were anesthetized by i.p. delivery of anesthetic mixture; lung tissue was harvested and processed for histochemistry, morphological and biochemical analyses.

**Preparation of Lung Tissue Lysates**

For protein extraction, the mouse lung tissue was homogenized in a buffer containing 20 mM Tris–HCl, pH 7.4, 150 mM NaCl, 1 mM PMSF and protease inhibitors, using a Brinkman Polytron homogenizer (Brinkman, Oxford, CT, United States). Total lysates were prepared by adding SDS and Triton X-100 to a final concentration of 0.3% and 1%, respectively, with gently mixing for 2 h at 4°C. The resulting lysates were clarified by centrifugation (191,500 × g, 45 min, 4°C) in a Beckman Optima Max-XP ultracentrifuge with a TLA-55 rotor (Beckman, Indianapolis, IN, United States). Protein concentration was determined by the micro BCA method with BSA as standard.

**Tracer Preparation and Perfusion**

Colloidal gold particles with an average diameter of 8 nm, were prepared (Predescu et al., 1998). Gold-tagged BSA (Au-BSA) was obtained by stabilizing colloidal gold suspensions with BSA (800 µg/ml) followed by 200 µg/ml poly l-glutamic acid, as secondary stabilization. The stabilized solutions were kept as stock and diluted just before use in PBS to A520 = 1. BSA was also tagged with DNP (DNP-BSA) and perfused to mouse and rat lung microvasculature to assess the permeability to albumin (Predescu et al., 1994, 1997, 1998). Quantitative assessment of DNP-BSA transendothelial transport was done by ELISA via anti-DNP Ab using a standard curve generated with known amounts of the same tracer (Predescu et al., 1997).

**EC Culture and Transfection**

Pulmonary artery ECs (Lonza, Walkersville, Inc., MD, United States), and EC_PAH (kindly provided by Drs. Serpil Erzurum and Suzy Comhair, Lerner Research Institute, Cleveland Clinic, Cleveland OH, United States) passages 3 – 5, were grown in EBM-2 and medium 199-supplemented with 20% fetal bovine serum as previously described (Predescu et al., 2003; Patel et al., 2013). The EC_PAH were harvested and cultured as described in Comhair et al. (2012) from 3 PAH patients; CCF-005: Female, white, non-Hispanic, 47-year-old, idiopathic PAH; CC-027: Female, white, non-Hispanic 22-year-old, chronic thromboembolic PAH; CC-016: Male, white, non-Hispanic 27-year-old, idiopathic PAH). For stable transfection, Myc-ETH5TsN transiently transfected ECs were grown in 100 cm² dishes with 1 mg/ml G418. HAM’s F12 medium containing 1 mg/ml G418 was replenished every other day for about 2 weeks until stable transfected ECs (EC_Myc-ETH5TsN) began to grow. At that point the concentration of G418 in HAM’s F12 medium was switched to 800 µg/ml for a week and then maintained at 500 µg/ml. ECs transfected with the empty Myc-M43 vector (EC_Myc–M43) and with the mutant ETH5TsNA (EC_Myc–ETH5TsNA) were used as controls.

For ETH5TsN knockdown, the individual siGENOME duplex most efficient in knocking down ETH5TsN protein expression (siRNA_ETH5TsN), GGACAUUGUUGUACUGAAAU (sense sequence) and 5′ – UUUCAGUACAGACUUGCCUU (antisense), was selected from the SMART pool siRNA reagents and delivered in cultured ECs (Predescu et al., 2007a; Bardita et al., 2015), using DharmaFect1 transfection reagent according to the manufacturer’s instructions. The siCONTROL™ functional non-targeting siRNA sequence 5′- UAGGGAGUUAA ACACAAUU-3′ was used as control for potential secondary effects caused by siRNA transfection. The siGloCytochrome B siRNA was used as a control for efficient transfection and to evaluate any off-target effects caused by the introduction of siRNAs in ECs.

**Western Blot (WB), Immunoprecipitation and Densitometry**

Endothelial cells grown on Petri dishes were collected and solubilized in lysis buffer (20 mM Tris–HCl, pH 7.4, 150 mM NaCl, 0.3% SDS, 1% (w/v) Triton X-100, 1 mM Na3VO4, 1 mM PMSF, and protease inhibitors) for 1 h at RT, under gentle agitation. The resulting lysate was clarified by centrifugation in a Beckman Optima Max-XP ultracentrifuge with a TLA-55 rotor at 45,000 rpm for 45 min at 4°C. Protein concentration was determined using the micro BCA method with BSA as standard. Protein samples normalized for total protein were subjected to SDS–PAGE and transferred to nitrocellulose membranes. Strips of nitrocellulose membranes were incubated with the primary and reporter Ab and processed as described (Predescu et al., 2007a). The reaction was visualized using an enhanced chemiluminescent substrate and HyBlot CL films. Representative films were subjected to densitometry using NIH ImageJ1.37v software.

For immunoprecipitation studies, 250 µg total protein from ECs lysates were precleared using a rabbit IgG followed by protein A/G agarose beads, for 1 h each, at RT. The resulting supernatant was incubated overnight at 4°C with anti-EHBP1 or ETH5TsN Abs, followed by incubation 1 h, at RT with protein A/G. Beads were washed extensively, and after solubilization in electrophoresis sample buffer, the immunoprecipitates were resolved on 5 – 20% SDS–PAGE, electrotransferred to nitrocellulose membranes and analyzed by WB.

**Biotin Assay for Caveolae Internalization**

Control and transfected cells grown on plastic Petri dishes or glass coverslips were washed with ice-cold PBS and then incubated with 0.5 mg/ml cleavable biotin reagent (Predescu et al., 2003). Biotinylated cell surface proteins were internalized for 30 min, at 37°C. Biotinylated proteins still at the cell surface after 30 min were reduced with glutathione and the cells were further processed for morphological analysis or lysed for biochemical investigation. For imaging, ECs grown on glass coverslips were washed in PBS (5 × 2 min), fixed and permeabilized (methanol, 7 min, −20°C), quenched (PBS + 1% BSA; 1 h, RT), and incubated with NeutrAvidin Alexa Fluor 594 (1 h, at RT). ECs were rewashed (5 × 2 min), mounted on glass slides using ProLong Antifade reagent, examined, and photographed.
prepared as above. The amounts of biotin labeled proteins were assessed by ELISA via streptavidin-HRP. For quantitative assessment of biotin molecules in the EC supernatants, a standard curve was generated using known concentrations of BSA-biotin. The average number of biotin molecules present in each cell lysate was determined at a series of decreasing concentrations from the linear part of the curve obtained by successively diluting a standard volume (100 µl) from each lysate and normalized per mg total protein.

**Immunostaining**

Immunofluorescent staining of ECs grown on coverslips was performed as described previously (Bardita et al., 2015). ITSN1 Ab (Prestige, Sigma-Aldrich, United States), was used at 1:500 dilution and EHBPI Ab at 1:50 dilution in 0.1% BSA in PBS, whereas Alexa Fluor 350 Phalloidin was used at 6.6 nM, as per manufacturer’s instructions. Incubation of ECs with isotype-matched IgGs or omission of the primary Ab was used as controls for Ab specificity. Each set of experiments was performed in triplicate.

**Transmission Electron Microscopy (TEM)**

The EC monolayers were fixed in 2% PFA + 0.5% GA + 1% tannic acid in 0.1 PIPES, pH 7.2, for 30 min at RT; all specimens were post-fixed in 2% OsO$_4$ in acetate veronal buffer, pH 6.8, for 30 min on ice (Predescu et al., 2001, 2007b). Fixed ECs were stained for 30 min with 7.5% uranyl-magnesium acetate (UA), dehydrated through increasing concentrations of ethanol (50, 70, 90, and 100%), then infiltrated with a mixture of 100% alcohol: Epon 812 (1:1) for 1 h and embedded in Epon 812. Embedded specimens were cured for 72 h at 60°C. Blocks of ~0.5/0.5 cm cut off from polymerized resin were used to obtain sections 60 nm, which were stained with 7.5% UA for 5 min and saturated lead citrate for 3 min, examined and photographed in a Jeol-1220 EM.

The lung tissue was fixed in situ by perfusion of a fixative mixture (3% PFA, 2.5% GA in 0.1 PIPES, pH 7.2) for 15 min, and selected specimens were further fixed with the same mixture for 1 h at RT. All specimens were post-fixed in 2% OsO$_4$ in acetate veronal buffer, for 1 h on ice, stained for 1 h with 7.5% UA, dehydrated through increasing concentrations of ethanol and propylene oxide as above, and embedded in Epon 812. Embedded specimens were cured for 72 h at 60°C, sections of ~60 nm were stained with 7.5% UA for 5 min and saturated lead citrate for 3 min and then examined as above.

**RESULTS**

### Expression of Myc-EH$_{\text{ITSN}}$ Interferes With the ECs Endocytic Activity

The observations that the $E_{\text{CPAH}}$ and human PAH lung tissue with pleuriform lesions show reduced levels of ITSN and expression of the EH$_{\text{ITSN}}$ protein fragment (Patel et al., 2013), led us to evaluate the endocytic activity of $E_{\text{CPAH}}$ as well as of stable transfected EH$^-_{\text{ITSN}}$. Non-transfected ECs (EC$^-_{\text{Ctrl}}$), ITSN-deficient ECs (EC$^+_{\text{KD}}$–ITSN) and ECs transfected with the empty Myc-M43 vector (EC$^+_{\text{Myc–M43}}$) were used for comparison. The specificity of EH$_{\text{ITSN}}$ effects was addressed by transfection of ECs with the mutant EH$_{W263A}$. The substitution W263A reduces protein interactions of the EH domains with the NPF motif beyond detection (de Beer et al., 1998). Efficient expression of Myc-EH$_{W263A}$ and Myc-EH$_{\text{ITSN}}$, 48 h post-transfection is documented in Figure 1A. No immunoreactivity for Myc tag is detected in the EC$^+_{\text{Ctrl}}$ lysates, Figure 1A. As ITSN knockdown, 72 h post-siRNA$_{\text{ITSN}}$ transfection triggers EC apoptotic death, the EC$^+_{\text{KD}}$–ITSN were used 38 – 40h post-siRNA$_{\text{ITSN}}$ delivery, when ITSN protein expression is 50% lower compared to EC$^+_{\text{Ctrl}}$ (Figure 1B), and ECs are not yet apoptotic (Predescu et al., 2007a; Bardita et al., 2015). Controls and EC$_{\text{EH–ITSN}}$ were subjected to the biotin assay for caveolae internalization followed by detection of the internalized biotinylated proteins by fluorescent imaging via neutrAvidin Alexa Fluor 594. All experimental conditions used, EC$_{\text{Ctrl}}$ (Figure 1C), EC$_{\text{KD–ITSN}}$ (Figure 1D), EC$_{\text{EH–W263A}}$ (Figure 1E), EC$_{\text{EH–ITSN}}$ (Figure 1F), $E_{\text{CPAH}}$ (Figure 1G), and EC$_{\text{Myc–M43}}$ (not shown) revealed...
FIGURE 1 | Expression of Myc-EHITSN interferes with the ECs endocytic activity. (A) Lysates (70 μg/lane) of EC\textsubscript{Ctrl}, EC\textsubscript{EH}\textsubscript{−}W263A, and EC\textsubscript{EH}\textsubscript{−}ITSN were subjected to SDS–PAGE and WB analyses using anti-Myc Ab. (B) Expression of ITSN in EC\textsubscript{Ctrl} (lane a) and ECs transfected with the siRNA\textsubscript{Ctrl} (lane b) and EC\textsubscript{KD}−ITSN (lanes c, d, e). Three time points post-siRNA\textsubscript{ITSN} transfection (36, 48, and 72 h) are shown, n = 3; *p < 0.01. EC\textsubscript{Ctrl}, EC\textsubscript{KD}−ITSN, EC\textsubscript{EH}−W263A, EC\textsubscript{EH}−ITSN (F), and EC\textsubscript{PAH} (G), CCF-005 EC\textsubscript{PAH} shown) were subjected to biotin internalization assay followed by NeutrAvidin Alexa Fluor 594 staining. The arrows in panels C–G indicate the fine biotin-NeutrAvidin Alexa Fluor 594 punctate pattern, suggestive of biotin association with endothelial vesicular carriers. The arrowheads in D–G point toward large fluorescent puncta assumed to be the morphological intermediates of the alternative transport pathways. (H) The number of biotin molecules in the final supernates of ECs lysates, as indicated. Biotinylated cell surface proteins were detected by ELISA via streptavidin-HRP. Three different EC\textsubscript{PAH} preparations (CCF-005, CC-027, CC-016, as indicated) were used. Results are the averages ± SD of three different experiments performed in triplicates. *p < 0.001; **p < 0.01; ***p < 0.05 vs. EC\textsubscript{Ctrl}.

细小的斑点散布在细胞表面和胞浆 (Figures 1C–G, arrows)。两个显著的差异在生物素/neutrAvidin Alexa Fluor 594 标记中被发现：(i) 细小的斑点标记，表明生物素与内吞相关，与内皮细胞的微泡载体有关，由于减少了 Caveolae 数量 (由于 ITSN 缺陷)，(ii) 大的荧光斑点的数量在 EC\textsubscript{EH}−ITSN 和 EC\textsubscript{PAH} 中增加，与 EC\textsubscript{Ctrl} 相比。这种大的斑点与具有EH 域的补救端内吞途径（大内吞结构，膜环形结构，以及膜泡-管状结构）的 ability 相一致 (Figures 1F,G, 箭头)。一个轻微的增加在...
large fluorescent puncta is also detected in the EC\textsuperscript{KD–ITSN} (Figure 1D, arrowheads). No detectable differences were noted between the biotin/neutrAvidin staining pattern between EC\textsuperscript{Ctrl} and EC\textsuperscript{EH–W263A} (Figures 1C,E). Under control conditions, the large endocytic structures are underrepresented and thereby the large fluorescent puncta are only seldom detected (Figures 1C,E, arrowheads).

Quantification of internalized biotin by ELISA via streptavidin-HRP indicated a 26% increase in biotin uptake in the EC\textsuperscript{EH–ITSN} compared to EC\textsuperscript{Ctrl} (Figure 1H; \( n = 8; p < 0.001 \)); The lysates of three different EC\textsubscript{PAH} lines showed on average only 6% decrease in the number of biotin molecules compared to EC\textsuperscript{Ctrl} (Figure 1H, \( n = 9; p < 0.05 \)), while 50% ITSN knockdown (Figure 1B, inset) induced 75% inhibition of biotin uptake (Figure 1H, \( n = 5; p < 0.05 \)). The minimal inhibition in biotin internalization in EC\textsubscript{PAH}, a stable phenotype characterized by ITSN deficiency and EH\textsubscript{ITSN} expression, implies that despite impaired caveolae endocytosis caused by lack of ITSN, the functional upregulation of compensatory transport pathways accounts for a still highly efficient endocytic activity. This observation demonstrates a potential role of the EH\textsubscript{ITSN} in supporting EC\textsubscript{PAH} proliferation and overgrowth, and thus, the development of the disease.

To get more insights into the effects of the EH\textsubscript{ITSN} expression on the endothelial vesicular carriers and endocytic structures (morphological intermediates of compensatory endocytic pathways), we applied EM analysis of ultrathin sections prepared from randomly chosen Epon-embedded EC\textsubscript{PAH}. An obvious observation was the reduced number of caveolae in EC\textsubscript{PAH} (Figure 2B) by comparison to EC\textsubscript{Ctrl} (Figure 2A). The size of caveolae (50 – 80 nm external diameter), was not affected by the experimental manipulations performed; even the clustered caveolae preserved their dimensions. In addition, the increased presence of pleomorphic morphological structures/intermediates, deemed to function as alternative endocytic pathways was noticeable. Frequently, we detected enlarged endocytic structures (Figure 2C, asterisk), membranous tubules (Figure 2C, arrowhead), tubulo-vesicular elements (Figures 2C, arrow,G) open to the apical front of the EC or with no noticeable communication with the extracellular environment and membranous rings (Figures 2D–F), ranging between 100 and 300 nm diameter with a narrow lumen (20 ± 4 nm). Morphometric analyses showed numerical statistical significance, Table 1. The increased presence of these pleomorphic morphological structures / intermediates, usually underrepresented and thought to function in alternative endocytic pathways, was confirmed in EC-EH\textsubscript{ITSN} and EC\textsuperscript{KD–ITSN} as well, Table 1. Our data demonstrate ∼ 3.8-fold increase in the frequency of alternative endocytic structures in EC-EH\textsubscript{ITSN} and EC\textsuperscript{KD–ITSN} compared to EC\textsubscript{Ctrl}, while EC\textsubscript{PAH} show ∼ 4.5-fold increase. In addition, the number of caveolae decreases by more than 30% in all experimental conditions by comparison to EC\textsubscript{Ctrl}.

**Alternative Transport Pathways Are Upregulated and Involved in Transendothelial Transport in PAH Specimens**

To address the functional effects of ITSN deficiency and EH\textsubscript{ITSN} expression on ECs endocytic/transcytotic activity and on lung microvascular permeability in PAH we used our recently developed EH\textsubscript{ITSN}-transduced K\textsuperscript{0+ITSN+/−} mouse model of plexiform arteriopathy (Patel et al., 2017). Briefly, K\textsuperscript{0+ITSN+/−}
sections prepared from randomly chosen EH
ITSN interendothelial junctions (IEJs) and if the 8 nm Au-BSA tracer
tracer transport indicated an increase from 356 ng DNP-
when the DNP-BSA tracer was perfused through rat lung
MCT-induced mouse model of PAH (Figures 3A–C, D, E–G) or with no noticeable communication with
8 nm Au-BSA (Figures 3C, D, I, arrows, d1), tubules and tubulo-vesicular elements open to the lumen
(Figures 3C, E, G) or with no noticeable communication with
mice were selected by genotyping; conventional RT-PCR applied
on KO^ITSN+/− mouse lung samples, followed by densitometry,
showed a 50% decrease in ITSN mRNA level by reference to
wt-mice (Supplementary Figure S1A). Cyclophilin was used
as an internal control. Similarly, ITSN protein expression was
50% lower in the KO^ITSN+/− mice compared to wt-mice,
as documented by WB analyses of lung lysates, followed by
densitometry. Efficient expression of the EH^ITSN for 21 days was
achieved by repeated delivery of the Myc-EH^ITSN as described
under Methods and illustrated in Supplementary Figure S1B.
The PAH phenotype of these mice, right ventricular systolic
pressure (RVSP) and Fulton’s index were determined as in Patel
et al. (2017), and illustrated in Supplementary Figures S1C, D,
respectively. We have also used the MCT-treated mice and
rats – two widely accepted experimental animal models of PAH
(Stenmark et al., 2009). As previously reported (Patel et al., 2013),
the lung tissue of both MCT-mice and rats is deficient of ITSN
and shows expression of the EH^ITSN (Supplementary Figures
S1E, F).

First, the EH^ITSN-transduced KO^ITSN+/− mouse lung microvasculature was perfused with 10 mg/ml DNP-BSA
tracer for 10 min. Wt- and the KO^ITSN+/− mice were used as controls. Lung tissue was collected, homogenized in PBS
and subjected to biochemical quantification of the amounts of DNP-BSA tracer transported to the lung interstitia. ELISA
via anti-DNP Ab indicated a significant increase in DNP-BSA transport in the EH^ITSN-transduced KO^ITSN+/− mice relative
to controls (i.e., 1466 ng DNP-BSA/mg total protein/10 min vs. 325.3 ng DNP-BSA/mg total protein/10 min in wt- and controls
and 539 ng DNP-BSA/mg total protein/10 min in KO^ITSN+/− mice), consistent with endothelial barrier dysfunction, Figure 3A.
An increase in the DNP-BSA transport by comparison to wt-control mice (812 ng DNP-BSA/mg total protein/10 min vs. 325.3 ng
DNP-BSA/mg total protein/10 min) was also recorded in the MCT-induced mouse model of PAH (Figure 3A). Finally,
when the DNP-BSA tracer was perfused through rat lung microvasculature, the quantification of the transendothelial
tracer transport indicated an increase from 356 ng DNP-
BSA/mg total protein/10 min to 1528 ng DNP-BSA/mg total
protein/10 min (Figure 3A) for controls vs. MCT-treated rats.

We further analyzed by EM the morphology of
interendothelial junctions (IEJs) and if the 8 nm Au-BSA tracer
can escape through open IEJs. Detailed analysis of ultrathin
sections prepared from randomly chosen EH^ITSN-transduced
KO^ITSN+/− mouse lung specimens showed frequently edematous areas (Figures 3B, D), but no IEJs penetrated by the 8 nm
Au-BSA tracer (Figure 3B). Given the presence of the edematous areas, this observation suggested that the IEJs are leaky but their
opening is not wide enough to allow the paracellular transport
of the tracer. The electron micrograph in Figure 3B shows
indeed an IEJ whose luminal introit (arrow) is heavily labeled by
8 nm Au-BSA unable to penetrate the junction and nearby an
edematous area, suggestive of junctional leakiness. This finding
compellingly demonstrated that the increased permeability
for BSA-DNP in the lung vasculature of EH^ITSN-transduced
KO^ITSN+/− mouse is due to the leaky IEJs, restrictive to 8 nm
Au-BSA; the size of Au-BSA complexes ([the overall diameter of
Au-BSA complexes (gold + BSA molecules) is about 20 nm while
the BSA-DNP is only 6 nm], maximum molecular diameter).
A junctional opening of ~ 6 nm, allows water, ions and small
molecules to take the paracellular pathway to the interstitia,
data that are in agreement with previous reports (Simionescu
et al., 2002). We did not detect gold particles associated with the
abluminal exit of the junction.

Another interesting observation was the frequent detection
of pleomorphic morphological structures/intermediates thought
to function in alternative endocytic pathways, such as enlarged
endocytic structures, many of them fused with typical caveolae
and heavily labeled by gold particles (Figures 3C, D, I, arrows,
d1), tubules and tubulo-vesicular elements open to the lumen
(Figures 3C, E, G) or with no noticeable communication with
the extracellular environment (Figure 3G), membranous rings
(Figures 3I–1i–J), many of them loaded with 8 nm Au-BSA
particles or open to the lumen and loading the tracer (inset i1).
The rings ranged between 150 and 300 nm diameter and
displayed a narrow lumen (20± 4 nm). Morphometric analyses
revealed that the EH^ITSN administration causes 10-fold increase
in the number of alternative endocytic structures, while the
number of caveolae decreases by 26% compared to wt-mice
(Table 2).

To get insights into the contribution of the EH^ITSN
to this lung PAH phenotype, we also analyzed the MCT-
treated mice (Figure 4). Detailed analysis of ultrathin sections
prepared from randomly chosen MCT-mouse lung specimens
showed the same salient features identified in the EH^ITSN-
transduced KO^ITSN+/− mice: (i) caveolae heavily loaded with
8 nm Au-BSA (Figures 4A, C, F), (ii) IEJs restrictive to tracer
particles (Figures 4A, B), (iii) edematous regions specifying that

**TABLE 1 | Morphometric analyses of caveolae and alternative endocytic structures in cultured endothelial cell (ECs).**

| Experimental condition | Total number of caveolae | Caveolae open | Caveolae clusters | Total Rings | Rings per cell profile | Tubules/tubulo-vesicular features | Enlarged endocytic structures (secondary lysosomes, auto-phagosomes) |
|------------------------|--------------------------|--------------|------------------|-------------|-----------------------|----------------------------------|------------------------------------------------------------------|
| ECM                   | 624 ± 42                 | 368 ± 37     | 6 ± 0.6          | 3 ± 0.5     | < 0.1                 | 1 ± 0.2                          | 24 ± 3                                            |
| EC^PAH^               | 387 ± 22                 | 144 ± 31     | 12 ± 1.1         | 69 ± 9      | 9 ± 0.7               | 11 ± 0.6                         | 52 ± 8                                            |
| EC^EH^ITSN^           | 402 ± 34                 | 115 ± 29     | 11 ± 1.9         | 58 ± 11     | 7 ± 0.5               | 8 ± 0.3                          | 48 ± 7                                            |
| EC^K0^ITSN^           | 412 ± 18                 | 170 ± 26     | 14 ± 1.8         | 56 ± 12     | 7 ± 1.2               | 9 ± 0.8                          | 48 ± 9                                            |

Electron microscopy (EM) micrographs (178 for EC^Ctrl^, 237 for EC^PAH^, 198 for EC^EH^ITSN^ and 204 for EC^K0^ITSN^) with full EC profiles at a final magnification of × 30,000 were used for each experimental condition. Data were normalized per 100 μm² surface and expressed as mean ± SD. n = 3 independent experiments.
FIGURE 3 | Non-conventional endocytic intermediates function in the transendothelial tracer transport. (A) Quantification of BSA-DNP transendothelial transport by ELISA applied on lung lysates of wt-mice (a), K0/ITSN−/− mice (b), EH1TSN-transduced K0/ITSN+/− mice (c), and monocrotaline (MCT)-treated mice (d) as well as on lung lysates of wt-rats (e) and MCT-rats (f). Results are expressed in ng BSA-DNP/mg total protein/10 min. n = 3 experiments performed in triplicates Bars ± SD.

Bar a - wt-mice
Bar b - K0/ITSN−/− mice
Bar c - EH1TSN + K0/ITSN+/− mice
Bar d - MCT mice
Bar e - wt-rats
Bar f - MCT-rats

the EH1TSN expression interferes with endothelial junctional integrity, (iv) enlarged endocytic structures (Figures 4C,E), (v) membranous rings (Figures 4D,H), and (vi) tubulo-vesicular structures (Figures 4E–G). Note that the upregulated alternative endocytic structures are actively involved in uptake and transport of tagged BSA molecules. Morphometric analyses of caveolae and alternative endocytic structures in the MCT mouse model of PAH demonstrate that MCT treatment causes 12-fold increase in the number of alternative endocytic structures by comparison to the wt-mouse (Table 2). Moreover, the number of caveolae decreases by 22% compared to the wt-mouse (Table 2).

The functional effects of ITSN deficiency and EH1TSN expression on lung microvascular permeability were also assessed in the lungs of the MCT-rat model of PAH, by 10 min perfusion of 10 mg/ml DNP-BSA through lung microvasculature and quantifying by ELISA the amounts of DNP-BSA tracer transported to lung interstitial space. Biochemical quantitation of the tracer amounts transported to the lung interstitia indicated a 30% increase by comparison to untreated rats (Figure 3). EM analyses of ultrathin sections prepared from the lungs of the Monocrotaline (MCT)- rat model of PAH confirmed that ITSN deficiency and the presence of the EH1TSN induces a similar lung phenotype as in the EH1TSN-transduced K0/ITSN+/− mouse (i.e., leaky IEJs and increased occurrence of pleomorphic morphological structures functioning as morphological intermediates of alternative endocytic pathways),
A Functional Partnership ITSN-EHBP1 Contributes to Actin Organization and Modulation of Endothelial Transport

As ITSN may bind via its NH2-terminal EH domains the EHB1 to mediate actin reorganization, we examined the potential partnership ITSN/EHBP1 by biochemical and morphological approaches. First, anti-EHBP1 Ab incubated with lysates of EC<sub>Ctrl</sub> and EC<sub>EH−ITSN</sub> normalized to 250 µg total protein, brings down not only the EHBP1 (Figure 6A) but also ITSN (Figure 6B). When lysates of stable transfected EC<sub>EH−ITSN</sub> were incubated with the anti-EHBP1 Ab, the immunoprecipitates contains the EH<sub>ITSN</sub> fragment, as well (Figure 6B, arrowhead). Moreover, it appears that EHBP1 has a higher affinity for the EH<sub>ITSN</sub> by comparison to full-length ITSN, as the amount of immunoprecipitated full-length ITSN is slightly lower in EC<sub>EH−ITSN</sub> compared to EC<sub>Ctrl</sub>. To validate this observation, in parallel experiments we immunoprecipitated ITSN from EC<sub>Ctrl</sub> and EC<sub>EH−ITSN</sub> lysates, normalized to 250 µg total protein (Figure 6C) using anti-ITSN Ab. The immunoprecipitates were subjected to WB analyses using anti-ITSN Ab followed by densitometric analyses (NIH ImageJ). Accurate assessment of the amounts of ITSN immunoprecipitated by EHBP1 Ab in lysates of EC<sub>Ctrl</sub> by comparison to EC-EH<sub>ITSN</sub>, was done by expressing the data as ratio of ITSN immunoprecipitated by EHBP1 Ab/ITSN immunoprecipitated by ITSN Ab (Figure 6D). The amount of ITSN, immunoprecipitated by EHBP1 from the lysates of EC-EH<sub>ITSN</sub>, is about 20% lower compared to the amount of ITSN immunoprecipitated by EHBP1 from the lysates of EC<sub>Ctrl</sub>. The finding suggests that the EHBP1 binds preferentially the Myc-EH<sub>ITSN</sub> by comparison to full-length ITSN.

Double immunofluorescence anti-ITSN-Alexa Fluor 594 / anti-EHBP1-Alexa Fluor 488 reveals for ITSN a strong punctate pattern throughout the cytosol and at the plasma membrane (Figure 6E), suggestive of vesicle association and a punctate pattern that overlaps a fine diffuse staining for EHBP1 (Figure 6F). Consistent with the immunoprecipitation analysis, a pool of the two proteins co-localize in the cytosol (Figures 6G,H arrowheads) as well as in very close proximity of the plasma membrane or at the plasma membrane (Figures 6G,H arrows). Significant ITSN/EHBP1 colocalization is detected in the perinuclear area and nucleus (Figure 6G).

To determine the relationship linking the vesicles containing ITSN with the EHBP1 and cytoskeletal elements we applied three-color staining, anti-ITSN (Figure 7A-a1), anti-EHBP1 (Figure 7A-a2) and Phalloidin/Alexa Fluor 350 for filamentous (F)-actin detection (Figure 7A-a3). In EC<sub>Ctrl</sub> the endothelial dense peripheral actin band and several stress fibers are easily detected (Figure 7A-a3). Both ITSN and EHBP1 immunoreactivities are in close proximity of actin fibers (Figures 7A-a4–a8), and limited co-localization with actin is detected, suggesting a possible link of ITSN/EHBP1 complexes to actin fibers, (Figures 7A-a6,a9, arrows).

To mimic the disease conditions, we next investigated the distribution of ITSN, EHBP1 and F-actin in stably expressing EH<sub>ITSN</sub> ECs subjected to ITSN knockdown via the siRNA approach, for 38 h – 40 h as in Figure 1B. In these ECs the immunoreactivity for ITSN is considerably lower, Figure 7B-b1. The remaining ITSN and the stably expressed Myc-EH<sub>ITSN</sub> (recognized by the polyclonal ITSN Ab) are randomly distributed as discrete fluorescent puncta in the cytosol and at the plasma membrane, Figures 7B-b4,b7 and cytosol. ITSN knocked down altered, however, the distribution of both EHBP1 and actin. Instead of the discrete fluorescent puncta dispersed throughout the cytosol and at the plasma membrane, EHBP1 is found as

---

### Table 2: Morphometric analyses of caveolae and alternative endocytic structures in wt-, EH<sub>ITSN</sub>-transduced K0<sup>ITSN+/−</sup> and MCT-treated mice.

| Experimental condition | Total number of caveolae (per 100 µm<sup>2</sup>) | Caveolae Clusters | Membranous rings | Tubules/ tubulo-vesicular features | Enlarged endocytic structures (secondary lysosomes, auto-phagosomes) |
|------------------------|-----------------------------------------------|------------------|-----------------|-------------------------------|--------------------------------------------------|
| Wt-mice               | 3700 ± 55                                     | 3.2 ± 0.6        | 1.8 ± 0.5       | 0.8 ± 0.3                     | 9.3 ± 1.5                                         |
| EH<sub>ITSN</sub>-K0<sup>ITSN+/−</sup> mice | 2738 ± 32                                     | 31 ± 4.7         | 21.6 ± 2.4      | 24.2 ± 3.2                    | 76.3 ± 14                                         |
| MCT mice              | 2911 ± 51                                     | 22.1 ± 6.2       | 10.2 ± 2.4      | 10.4 ± 3.8                    | 139.5 ± 21                                        |

Aggregated areas of 625 µm<sup>2</sup> for wt-mice (164 arteriolar, 192 capillary and 107 venular profiles were examined), 817 µm<sup>2</sup> for the EH<sub>ITSN</sub>-transduced K0<sup>ITSN+/−</sup> mice (128 arteriolar, 255 capillary and 172 venular profiles were examined) and 417 µm<sup>2</sup> for the MCT-treated mice (125 arteriolar, 95 capillary and 102 venular profiles examined) were used for counting. The data are normalized per 100 µm<sup>2</sup> EC surface. All arterioles with diameters between 60 and 120 µm and venules up to 70 µm were included in the sampling. There are no detectable (less than 0.1%) secondary lysosomes in arterioles or capillaries under physiological conditions, while the venular endothelium displays less than 2%. Values are means ± SD; n = 3.
large fluorescent puncta accumulated in the perinuclear area, Figures 7B-b2,b5. However, on the endothelial background, the EHBPI immunoreactivity at the plasma membrane is very limited, Figures 7B-b5,b8. Phalloidin staining revealed the lack of the peripheral dense actin band, the presence of long stress fibers and membrane ruffles (Figure 7B-b3). It appears that ITSN expression is required for EHBPI presence at the plasma membrane. Only several large fluorescent puncta where ITSN colocalizes with the EHBPI were detected (Figures 7B-b6,b9 arrows), indicating a possible association of ITSN/EHBPI complexes to actin fibers. Interestingly, despite ITSN deficiency, ECs expressing the EH\textsubscript{ITSN} (Figures 7B-b1,b4) still display
nuclear immunoreactivity to ITSN Ab, most likely, the result of nuclear localization of the exogenously expressed EH_{ITSN}. The observation is very suggestive of potential functional implications of EH_{ITSN} localization in the nucleus.

These observations suggest that ITSN and EHBPI are components of a complex with essential functions in both the organization of cell membrane associated cytoskeleton and endocytosis. It appears that ITSN recruits the EHBPI to the plasma membrane where ITSN/EHBPI complexes can associate with actin and play a role in actin polymerization and formation of dense peripheral actin band. Moreover, lack of the peripheral actin band detected in the ITSN-deficient ECs stably expressing the EH_{ITSN} may favor the increased occurrence of alternative endocytic structures and their intracellular trafficking.

**DISCUSSION**

Hyperproliferation, apoptosis-resistance, dysfunctional endocytosis, altered intracellular protein trafficking and increased permeability are characteristics of pulmonary ECs in PAH (Sehgal and Lee, 2011; Long et al., 2015; Zhou et al., 2016). Several studies implicated caveolae and caveolin-1 in EC dysfunction and the pathogenesis of PAH. Mice deficient of caveolin-1 spontaneously develop PAH (Zhao et al., 2002, 2009). Heterozygous mutations in the caveolin-1 gene with distinct effects on caveolae assembly and caveolin-1 trafficking have been identified in patients with PAH (Copeland et al., 2017). It has also been reported that heterozygous null BMPR2 mutations promote caveolae trafficking defects (Prewitt et al., 2015).

Our studies also indicated that the lung tissue of human and animal models of PAH is deficient of ITSN and expresses a biologically active ITSN fragment which triggers the endothelial proliferative phenotype responsible for the severe plexiform arteriopathy (Patel et al., 2013). Deficiency of ITSN, a general endocytic scaffold and regulator of the membrane fission GTPase, dynamin, decreases significantly caveolae number (Knezevic et al., 2011; Predescu et al., 2012). Compelling evidence demonstrates that transendothelial transport/ transcytosis mediated by caveolae is a primary determinant of the basal lung endothelial permeability properties (Sun et al., 2011). Disruption of caveolae function alters the endothelial barrier integrity,

**TABLE 3** Morphometric analyses of caveolae and alternative endocytic structures in wt- and MCT-treated rats.

| Experimental condition | Total number of caveolae | Caveolae Clusters | Membranous rings | Tubules/ tubulo-vesicular features | Enlarged endocytic structures (secondary lysosomes, auto-phagosomes) |
|------------------------|--------------------------|------------------|------------------|-----------------------------------|---------------------------------------------------------------|
| wt rat                 | 4257 ± 82                | 5.4 ± 1.8        | 2.0 ± 0.8        | 1.1 ± 0.7                         | 10.1 ± 2.2                                                    |
| MCT rat                | 3180 ± 88                | 25.2 ± 3.3       | 19.1 ± 4.8       | 18.5 ± 4.1                        | 166.8 ± 16                                                   |

Aggregated areas of 924 µm² for wt-rats (181 arterioles, 212 full capillaries and 117 venular shapes examined) and 842 µm² for MCT-rats (155 arterioles, 195 capillaries and 202 venular profiles examined) were used for counting. The data are normalized per 100 µm² EC surface. All arterioles with a diameter between 60 and 125 µm and venules up to 75 µm were included in the sampling for both wt- and MCT-rats. Values are means ± SD; n = 3.
enhances vascular permeability and upregulates compensatory endocytic pathways and their morphological intermediates, usually underrepresented (Drab et al., 2001; Miyawaki-Shimizu et al., 2006; Predescu et al., 2012, 2013). Our present studies extend these observations as we found that prolonged EH1_{ITSN} expression cooperates with ITSN deficiency in EC_{PAH} and experimental animal models of PAH in decreasing caveolae number, impairing their transport function while boosting the upregulation of alternative transport pathways. Thus, due to these features, the experiments on EC_{PAH} cells, best reflect the human disease condition and also the 2-hit mouse model of plexiform arteriopathy we generated. We demonstrate using EC_{KD-ITSN} and EC_{PAH} that the non-conventional structures are involved in biotin internalization and can compensate for deficient endocytosis via caveolae. Pulmonary ECs of EH1_{ITSN}-transduced K0_{ITSN+/-} mice show decreased number of caveolae and increased occurrence of caveolae clusters, tubulo-vesicular and enlarged endocytic structures and membranous rings. Moreover, using gold tagged-BSA perfusion of the mouse and rat lung microvasculature, we demonstrate that these non-conventional structures are involved in the uptake, internalization and transendothelial transport. Our data prove that the upregulated alternative pathways may functionally rescue the transport and nutrient uptake, usually mediated by the endothelial caveolae. This observation demonstrate that EH1_{ITSN} expression, a 2nd component or 2nd “hit” mechanism generated in response to pathological risk factors (inflammation, increase CD8+ lymphocyte, granzyme B secretion), with no drug administration or hypoxia exposure, is required to drive not only formation of plexiform lesions but also to “bounce” the endocytic activity of dysfunctional ECs in order to support their hyper-proliferation and metabolic activity.

However, the lack of caveolae and as results the lack of the canonical caveolar signaling pathways responsible for controlling the understudied and less scrutinized molecular mechanisms governing the cross-talk between the caveolae and the non-conventional upregulated structures have detrimental consequences on EC function. This emerging phenotype characterized by loss of caveolae in favor of alternative pathways, cannot, however, substitute for endocytosis involvement in
FIGURE 7 | ITSN-EHBP1 interaction contributes to actin organization in ECs. ECCtrl (A) and EC-EH\textsubscript{ITSN} + siRNA\textsubscript{ITSN} (B) were subjected to tri-color fluorescent staining with anti-ITSN Ab - anti-rabbit IgG Alexa Fluor 594-conjugated (a1,b1), anti-EHBP1 Ab - anti mouse IgG Alexa Fluor 488-conjugated (a2,b2) and Phalloidin Alexa-Fluor 350 for detection of F-actin (a3,b3). Merged images anti-ITSN/Phalloidin (a4,b4), anti-EHBP1/Phalloidin (a5,b5) and anti-ITSN/anti-EHBP1/Phalloidin (a6,b6) were used to analyze the subcellular distribution of ITSN/EHBP1 relative to EC actin cytoskeleton. The magnifications a7–a9 (as indicated by boxed areas) highlight the significant co-localization of ITSN/EHBP1 immunoreactive puncta at the plasma membrane and its proximity. Arrows in a9 indicate ITSN/EHBP1 immunoreactive puncta very close to actin fibers. When EC-EH\textsubscript{ITSN} + siRNA\textsubscript{ITSN} were analyzed, the highly magnified images b7–b9 (boxed areas in b4–b6) revealed significantly less ITSN (b7) and EHBP1 (b8) immunoreactive puncta. The limited ITSN and EHBP1 immunoreactivity is still in very close proximity of actin fibers. Arrows in b9 indicate ITSN/EHBP1 immunoreactive puncta very close to actin fibers. n = 3.

cellular proliferation (Sigismund et al., 2012), and therefore the new feature – EC hyper proliferation – is first to be noticed. We have also shown that in pathological conditions, dysfunctional ECs show altered intracellular trafficking and signaling of cell surface receptors such as the TGF\textbeta\textsubscript{R1} (Bardita et al., 2015), implicated in the pathogenesis of PAH (Garcia-Rivas et al., 2017; Graf et al., 2018). Compelling evidence indicates the essential involvement of different endocytic routes (caveolae, clathrin-mediated, lipid rafts) in cellular signaling events with consequences on disease-linked phenotypes. Heterozygous
germline mutations in BMPRII, such as those observed in patients with familial PAH, contribute to the failure of the receptor to traffic routinely from the ER to Golgi and reach the cell surface; the intracellular withholding, lack of access to the extracellular ligand and endocytic internalization led to altered receptor signaling in response to BMPs (Sobolewski et al., 2008; Morrell, 2010). Multiple dysfunctions in the endocytic (anterograde and retrograde) vesicular trafficking, abnormal trafficking of relevant receptors (i.e., BMPRI, BMPRII, IL-6R, gp130) and vasoactive mediators via alternative pathways that bypass the Golgi apparatus, have been implicated in the pathogenesis of PAH, as well (Sehgal and Lee, 2011). Thus, the increasing occurrence of the alternative endocytic structures may alter the subcellular localization of cell surface receptors, their signaling, their fate and sorting with functional consequences for proteins involved (Di Fiore and De Camilli, 2001; Le Roy and Wranas, 2005; Bardita et al., 2015).

Our studies also demonstrate that ITSN-deficiency and the EHTSN presence promote actin reorganization leading to the widening and leakage of the intercellular junctions. The resulting loss of endothelial barrier integrity provides increased sensitivity for proliferative and inflammatory mediators to reach the medial and adventitial vascular layers, leading to cell proliferation and PAH progression. The studies reporting a link between a dysfunctional lung endothelial barrier and PAH pathogenesis are limited. Lung microvascular leak has been associated with lung cell hyperplasia and right ventricular hypertrophy in both MCT- and hypoxia-induced PAH (Sugita et al., 1983). More recently, Zhou et al. (2016) reported that animals with severe PAH display increased sensitivity to vascular permeability induced by activation of store-operated calcium entry. Additional studies demonstrated a role for endothelial BMPRII loss in causing increased vascular permeability and altered translocation across the vascular wall (Burton et al., 2011a,b; Kim et al., 2013). Our data are evidence that ECs, as the critical initiating cell type in PAH, can contribute to the PAH pathology not only by increased apoptosis followed by exuberant proliferation, but also by increased vascular permeability.

Actin reorganization, the formation of stress fibers and lack of cortical actin disturb not only the proper interaction of ECs with each other and their interaction with the micro-environment but also the normal endocytic membrane trafficking. We found that ITSN-deficiency and expression of the EHTSN alter the subcellular distribution of the EHBP1, a protein that functions near the plasma membrane and couples endocytosis to the actin cytoskeleton (Guilherme et al., 2004). EHBP1 is expressed in ECs and interacts with ITSN via the NH2-terminal EH domains. Within full-length ITSN, the EHTSN are under the inhibitory control of the SH3A–E domains that may account for the spatial and temporal coordination of cellular activities in specific location within the cell (Adams et al., 2000). The free EHTSN escapes the regulatory control of the SH3A–E, and as result, its interaction with the EHBP1 is augmented. ITSN deficiency and thus, the lack of its scaffolding properties, prevent the proper EHBP1 localization at its site of action. As a result, the organization of cortical actin cytoskeleton is impaired favoring the formation and motility of tubulo-vesicular and enlarged endocytic structures and membranous rings, and thus allow to compensate for deficient caveolae intracellular transport. We have previously shown that actin polymerization and remodeling sub-adjacent to the plasma membrane decrease caveolae endocytosis significantly in ECs (Klein et al., 2009). We have also shown that ITSN deficiency is a crucial player in Rac1 activation and vimentin filament organization (Jeganathan et al., 2016). This is significant as Rac1 and vimentin are essential regulators of the cytoskeleton structure with control over the epithelial-mesenchymal transition, cell migration and metastasis of lung cancer (Jeganathan et al., 2016). The observations raised the possibility that ITSN deficiency and EHTSN expression resulting in an EC proliferative phenotype and loss of junctional integrity may provoke ECs to acquire a mesenchymal phenotype. The mesenchymal EC phenotype is present in experimental PAH animal models and has significant contribution to tissue regeneration and vascular remodeling in PAH (Good et al., 2015; Ranchoux et al., 2015; Tang et al., 2018).

**CONCLUSION**

We show that the EHTSN besides contributing to the proliferation and overgrowth of ECs, triggers the upregulation of the morpho-functional alternative endocytic activity of dysfunctional ECsPAH, and hence, play a role in the development and maintenance ECsPAH phenotype.

**AUTHOR CONTRIBUTIONS**

DP and SP conceived and designed the research, analyzed the data, interpreted the results of experiments, and drafted the manuscript. DP, MP, SQ, CB, and RB performed the experiments. MP, SQ, and DP prepared the figures. SP edited and revised the manuscript.

**FUNDING**

This work was supported by National Heart, Lung, and Blood Institute, grants R01 HL127022 and HL60917.

**ACKNOWLEDGMENTS**

The authors thank Drs. Serpil Erzurum and Suzy Comhair (Lerner Research Institute, Cleveland Clinic) for providing the pulmonary artery endothelial cells isolated from PAH patients and Dr. Melanie Pritchard, School of Biomedical Sciences, Monash University, Clayton, Australia for providing the ITSN heterozygous knockout mice.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: [https://www.frontiersin.org/articles/10.3389/fphys.2018.01393/full#supplementary-material](https://www.frontiersin.org/articles/10.3389/fphys.2018.01393/full#supplementary-material)
Predescu, S. A., Predescu, D. N., Knezevic, I., Klein, I. K., and Malik, A. B. (2007a). Intersectin-1s regulates the mitochondrial apoptotic pathway in endothelial cells. J. Biol. Chem. 282, 17166–17178.

Predescu, S. A., Predescu, D. N., and Malik, A. B. (2007b). Molecular determinants of endothelial transcytosis and their role in endothelial permeability. Am. J. Physiol. Lung Cell Mol. Physiol. 293, L823–L842. doi: 10.1152/ajplung.00436.2006

Predescu, S. A., Predescu, D. N., and Palade, G. E. (1997). Plasmalemmal vesicles function as transcytotic carriers for small proteins in the continuous endothelium. Am. J. Physiol. 272(2 Pt 2), H937–H949.

Predescu, S. A., Predescu, D. N., and Palade, G. E. (2001). Endothelial transcytotic machinery involves supramolecular protein-lipid complexes. Mol. Biol. Cell 12, 1019–1033. doi: 10.1091/mbc.12.4.1019

Predescu, S. A., Predescu, D. N., Timblin, B. K., Stan, R. V., and Malik, A. B. (2003). Intersectin regulates fission and internalization of caveolae in endothelial cells. Mol. Biol. Cell 14, 4997–5010. doi: 10.1091/mbc.e03-01-0041

Prewitt, A. R., Ghose, S., Frump, A. L., Datta, A., Austin, E. D., Kenworthy, A. K., Sakao, S., Taraseviciene-Stewart, L., Lee, J. D., Wood, K., Cool, C. D., and Predescu, S. A., Predescu, D. N., and Palade, G. E. (1997). Plasmalemmal machinery involves supramolecular protein-lipid complexes. Mol. Biol. Cell 12, 1019–1033. doi: 10.1091/mbc.12.4.1019

Prewitt, A. R., Ghose, S., Frump, A. L., Datta, A., Austin, E. D., Kenworthy, A. K., Sakao, S., Taraseviciene-Stewart, L., Lee, J. D., Wood, K., Cool, C. D., and Voelkel, N. F. (2001). The pathobiology of pulmonary hypertension. Endothelium. Clin. Chest Med. 22, 405–418. doi: 10.1016/S0272-5231(05)70280-X

Tuder, R. M., Groves, B., Badesch, D. B., and Voelkel, N. F. (1994). Exuberant endothelial cell growth and elements of inflammation are present in plexiform lesions of pulmonary hypertension. Am. J. Pathol. 144, 275–285.

Tuder, R. M., Marecki, J. C., Richter, A., Fijalkowska, I., and Flores, S. (2007). Pathology of pulmonary hypertension. Clin. Chest Med. 28:23–vii. doi: 10.1016/j.ccm.2006.11.010

Vieira, A. V., Lameza, C., and Schmid, S. L. (1996). Control of EGF receptor signaling by clathrin-mediated endocytosis. Science 274, 2086–2089. doi: 10.1126/science.274.5295.2086

Yamabhai, M., Hoffman, N. G., Hardison, N. L., McPherson, P. S., Castagnoli, L., Cesareni, G., et al. (1998). Intersectin, a novel adaptor protein with two Eps15 homology and five Src homology 3 domains. J. Biol. Chem. 273, 31400–31401. doi: 10.1074/jbc.273.11.31400

Yap, C. P., Lasiecka, Z. M., Caplan, S., and Winckler, B. (2010). Alterations of EHD1/EHD4 protein levels interfere with L1/NgCAM endocytosis in neurons and disrupt axonal targeting. J. Neurosci. 30, 6646–6657. doi: 10.1523/JNEUROSCI.5428-09.2010

Zhao, Y. Y., Zhao, Y. D., Mirza, M. K., Huang, J. H., Potula, H. H., Vogel, S. M., et al. (2010). Alterations in caveolin-1 cause dilated cardiomyopathy and pulmonary hypertension in knockout mice. Proc. Natl. Acad. Sci. U.S.A. 99, 11375–11380. doi: 10.1073/pnas.12360799

Zhao, Y. Y., Zhao, Y. D., Mirza, M. K., Huang, J. H., Potula, H. H., Vogel, S. M., et al. (2009). Persistent ENOS activation secondary to caveolin-1 deficiency induces pulmonary hypertension in mice and humans through PKG nitration. J. Clin. Invest. 119, 2009–2018. doi: 10.1172/JCI33338

Zhou, C., Townley, M. L., Alexeyev, M., Voelkel, N. F., and Stevens, T. (2016). Endothelial hyperpermeability in severe pulmonary arterial hypertension: role of store-operated calcium entry. Am. J. Physiol. Lung Cell Mol. Physiol. 311, L560–L569. doi: 10.1152/ajplung.00057.2016

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2018 Predescu, Qin, Patel, Bardtis, Bhalli and Predescu. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.