Re-establishment of VWF-dependent Weibel-Palade bodies in VWD endothelial cells

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Type 3 von Willebrand disease (VWD) is a severe hemorrhagic defect in humans. We now identify the homozygous mutation in the Chapel Hill strain of canine type 3 VWD that results in premature termination of von Willebrand factor (VWF) protein synthesis. We cultured endothelium from VWD and normal dogs to study intracellular VWF trafficking and Weibel-Palade body formation. Weibel-Palade bodies could not be identified in the canine VWD aortic endothelial cells (VWD-AECs) by P-selectin, VWFpp, or VWF immunostaining and confocal microscopy. We demonstrate the reestablishment of Weibel-Palade bodies that recruit endogenous P-selectin by expressing wild-type VWF in VWD-AECs. Expression of mutant VWF proteins confirmed that VWF multimerization is not necessary for Weibel-Palade body creation. Although the VWF propeptide is required for the formation of Weibel-Palade bodies, it cannot independently induce the formation of the granule. These VWF-null endothelial cells provide a unique opportunity to examine the biogenesis of Weibel-Palade bodies in endothelium from a canine model of type 3 VWD. (Blood. 2005;105:145-152)

Introduction

von Willebrand factor (VWF) is a large, multimeric glycoprotein that mediates platelet adhesion to subendothelial tissue at the site of vascular injury. VWF is also the carrier protein for coagulation factor VIII (FVIII). Defects in VWF function or decreased levels of VWF cause von Willebrand disease (VWD). VWF is synthesized as pre–pro-VWF consisting of a 22–amino acid (aa) signal peptide, 741-aa propeptide, and 2050-aa mature VWF monomer. VWF undergoes extensive intracellular modification. In the Golgi, VWF dimers form amino-terminal multimers, and propeptide is cleaved, presumably by the enzyme furin. VWF is stored together with its propeptide in regulated secretory vesicles, Weibel-Palade bodies in endothelial cells and α-granules in megakaryocytes or platelets.

The VWF released from Weibel-Palade bodies consists of very high-molecular-weight VWF multimers that are the most active for binding to subendothelial tissue and in platelet-platelet interactions. Thus, the multimerization and regulated storage of VWF are important for initiation of clot formation following injury to a vessel. The large propeptide, VWFpp, is required for multimerization and regulated storage of VWF. Previous studies in our laboratory have demonstrated that VWF is sorted to regulated storage vesicles in a VWFpp-dependent manner. Our studies were carried out using the mouse pituitary model cell line AtT-20 and bovine aortic endothelial cells (BAECs) that have endogenous-regulated storage granules containing adrenocorticotrophic hormone (ACTH) or bovine VWF, respectively. We have demonstrated that the propeptide independently sorts to endogenous storage granules in these cells and co-traffics mature VWF multimers through a noncovalent association. However, because of the presence of endogenous-regulated secretory vesicles, these studies could not address the biogenesis of the regulated storage vesicle.

Although the biogenesis and exocytosis of Weibel-Palade bodies has been the subject of investigation, the mechanisms are not well defined. To pursue our studies on intracellular VWF trafficking and Weibel-Palade body biogenesis, we cultured endothelium from the Chapel Hill strain of canine (K9) type 3 VWD. Type 3 VWD is a severe bleeding disorder in humans, characterized by a complete absence of VWF. Natural animal models of type 3 VWD exist in pigs and dogs, as well as a VWF-null transgenic mouse model. The type 3 VWD dogs have been maintained in Chapel Hill for many years, but the molecular defect causing the disease had not been identified. Here we first identify the mutation in these dogs that results in premature termination of VWF protein synthesis. Endothelial cells were obtained from the aortas of VWD dogs, and the absence of Weibel-Palade bodies, VWFpp, and VWF were confirmed. We have examined the ability of expressed wild-type and mutant VWF, or VWFpp, to generate storage granules that recruit endogenous P-selectin to reestablish Weibel-Palade bodies in the VWD endothelial cells. The availability of a VWF-null, nontransformed endothelial cell has thus provided a beneficial model for investigating the mechanisms involved in Weibel-Palade body formation.
Materials and methods

Antibodies

Anti-VWF antibodies 280.1, AVW-1, AVW-5, and 105.4 and the polyclonal, Edwina, were produced by our laboratory. A rabbit polyclonal anti-human VWF was purchased from Dako (Carpinteria, CA). The monoclonals AVW-5 and 105.4 and the polyclonal antibodies recognize both human and canine VWF. The antibody 280.1 is specific for canine VWF and AVW-1 is human VWF specific. Anti-VWFp monoclonal antibodies 239.1 to 239.11, 242.4, 242.5, 275.1, and 275.2 were also produced by our laboratory; 239.1, 239.7, 239.8, 239.11, 242.4, and 242.5 recognize canine VWFp in addition to human, while the 275.1 and 275.2 only recognize canine propeptide. The anti-human platelet endothelial cell adhesion molecule (PECAM) antibody, PECAM 1.3, was also produced in our laboratory. Other antibodies used include rabbit polyclonal anti–P-selectin (BD Biosciences, San Jose, CA), and goat–anti-C3 (Sigma Chemical, St. Louis, MO). Nonimmune mouse and rabbit immunoglobulin G (IgG) were purchased from Jackson ImmunoResearch (West Grove, PA) and used as isotype controls. Secondary antibodies used for immunofluorescence detection included goat anti-rabbit and anti-mouse IgG (H+L) (F(ab’)2) fragments conjugated with either AlexaFluor-488 or AlexaFluor-568 (Molecular Probes, Eugene, OR), or donkey anti-rabbit, anti-goat, or anti-mouse IgG (H+L) (F(ab’)2) fragments (Jackson ImmunoResearch) conjugated with Texas Red (TXR), Red-X, or fluorescein isothiocyanate (FITC).

 Constructs

The human and canine VWF, VWF propeptide (VWFpp), and propeptide-deleted VWF (Δpro) expression vectors were constructed as previously described.21 Briefly, VWFpp expression vectors contain signal peptide and VWFpp sequence followed by a stop codon with the VWF sequence deleted. The Δpro expression vectors contain signal peptide followed directly by mature VWF sequence. The Y87S-VWF is a full-length VWF construct that encodes a point mutation in the D1 domain of the VWF propeptide as previously described.22 VWFppC3α vector expresses VWF signal peptide and VWFpp with a disrupted furin cleavage site followed directly by the α-chain of human complement C3 as previously described.19

 Sequencing

White blood cell DNA was purified from normal and VWD dogs from the colony maintained at the University of North Carolina, Chapel Hill. In particular, a segment of genomic sequence spanning from base 228 in exon 4 to base 532 in exon 5 of the VWF gene was amplified by polymerase chain reaction (PCR) using sense primer K9-VWF s228 to 250 (5’-CCA AAT GGC AAA AGA GTG AGC C-3’) and antisense primer K9-VWF a532 to 510 (5’-TTT TGG AGT CCT GAA TTC ATG C-3’). Directly sequencing the PCR product from 2 VWD dogs using the same antisense primer yielded a single sequence with a nucleotide deletion of the cytosine at base 255 (ΔC255) of normal canine VWF cDNA. The PCR products were cloned and sequenced in both directions to confirm the mutation.

 Cell culture and transfections

Endothelial cells were harvested by digestion with 1 mg/mL collagenase from the aortas obtained from normal and VWD dogs. The harvested canine aortic endothelial cells (AECs) were cultured in Medium 199 (Mediatech, Holy Hill, FL) supplemented with 10% fetal bovine serum (FBS) and grown on tissue culture dishes coated with 2% gelatin, at 37°C in an atmosphere of 5% CO2. Cells were transiently transfected by Nucleofection (Amaxa, Cologne, Germany) according to manufacturer’s instructions. Negative controls were either nontransfected or mock transfected with expression vector lacking the insert. When transfected cells reached confluency (3-7 days after nucleofection), conditioned media were harvested from the cells, centrifuged to remove debris, and frozen at -80°C for further analysis. Washed cells were either fixed for intracellular staining or used for secretion studies. To assess stimulated release of expressed VWF proteins, cells were washed with Hanks balanced salt solution (HBSS), followed by OptiMEM washes, and then incubated for 30 minutes with either 2 μM phorbol myristate acetate (PMA) or OptiMEM (control). Releasates were harvested, and secreted VWF protein was quantitated by either VWFpp or VWF enzyme-linked immunosorbent assay (ELISA).

 Confocal immunofluorescence microscopy

Cells were analyzed for the intracellular location of VWF, VWFpp, and other proteins by immunofluorescence using antibody staining and confocal laser scanning microscopy in the Imaging Core of the Medical College of Wisconsin using a Leica (Heidelberg, Germany) TCS SP2 confocal laser imaging system and software with 63 × or 100 × objective lenses as previously described.20 Cells were grown on 2-well chamber slides coated with 0.2% gelatin and cross-linked with 0.5% gluteraldehyde. Confluent cells were fixed using 3.7% (vol/vol) buffered formalin, permeabilized in 1% Triton X-100, and blocked with 2% normal goat serum or donkey serum in HBSS. Cells were incubated at 4°C overnight in primary antibodies diluted in HBSS containing 1% bovine serum albumin (BSA). The concentration of primary antibody varied from 2 to 5 μg/mL. Cells were then washed 3 times with HBSS. All immunostaining experiments used normal mouse and rabbit IgG for isotype controls. Cells were incubated in secondary antibodies diluted 1:1000 in HBSS containing 1% BSA previously described.20 Cells were mounted with cover glass using Vectashield (Vector Labs, Burlingame, CA).

 Multimer analysis

VWF in the conditioned medium of transfected endothelial cells was analyzed by electrophoresis through a 0.8% (wt/vol) HGT(P) agarose (FMC Bioproducts, Rockland, ME) stacking gel and 2% (wt/vol) HGT(P) agarose running gel containing 0.1% sodium dodecyl sulfate (SDS) for 16 hours at 40 V using the Laemmli buffer system with transfer and Western blotting as previously described.23 Submarine-agarose gel electrophoresis was performed as previously described.23

 Results

The VWD dog colony has been maintained in Chapel Hill for many years. Although the bleeding disorder in these dogs has been established as type 3 VWD, the molecular defect causing the disease had not been determined.26,28 Since our laboratory had previously identified the sequence of normal canine VWF, we carried out sequencing of the VWF gene using genomic DNA obtained from the VWD dog.21 In particular, a segment of genomic sequence spanning from exon 4 to exon 5 of the VWF gene was amplified by PCR. The 591–base pair (bp) PCR product included bases 228 to 323 in exon 4, the 286-bp intron 4, and bases 324 to 352 in exon 5. A comparison of the normal and VWD sequence is shown in Figure 1. Sequencing revealed a single (homozygous) nucleotide deletion (ΔC255) in exon 4 of the VWF propeptide (VWFpp), resulting in a frameshift mutation at amino acid 85 of the VWF protein. After 34 altered amino acids, the shifted reading frame results in a TAG stop codon at base 359, leading to premature termination of VWF protein synthesis. This mutation accounts for the complete absence of VWF in the canine VWD plasma.

Although no VWF is detected in the VWD dog plasma, we questioned whether any VWF or VWFpp protein could be detected in the endothelial cells of the VWD dogs. If these endothelial cells were indeed VWF-null they would provide a valuable tool for studying the intracellular trafficking of VWF. VWF trafficking and biogenesis of Weibel-Palade bodies have previously been studied by our laboratory and others using transformed model cell lines.17,20,21,29-31 A VWF-null, nontransformed endothelial cell has
nucleotide sequence results in a stop codon at base pair 359 in exon 5.

An altered VWD canine protein sequence at base pair 255 as highlighted in the box, resulting in a frameshift. The canine sequence. A single nucleotide deletion was found in the VWD canine acid sequence with normal canine sequence shown in bold font above the VWD intron 4, and exon 5. The protein sequence is shown above the corresponding nucleic sequenced, and the 2 sequences were compared, including sequence from exon 4, intron 4, and exon 5. The protein sequence is shown above the corresponding nucleic acid sequence with normal canine sequence shown in bold font above the VWD canine sequence. A single nucleotide deletion was found in the WVD canine sequence at base pair 255 as highlighted in the box, resulting in a frameshift. The altered VWD canine protein sequence is shown in italics. The altered VWD canine nucleotide sequence results in a stop codon at base pair 359 in exon 5.

not been available. We isolated and characterized the AECs from normal dogs to compare with those obtained from the VWD dogs. Harvested cells were grown on gelatin, immunostained as described in “Materials and methods,” and examined by confocal microscopy. Immunostaining for the endothelial cell marker PE-CAM-1 (CD31) as shown in Figure 2A demonstrated that a homogeneous population of normal AECs had been cultured. Immunostaining with a WVF polyclonal antibody confirmed that the normal canine endothelial cells also express WVF (Figure 2B). Immunostaining for WVFpp and WVF demonstrated colocalization of WVF with WVFpp in granules (Figure 2D-F), similar to the Weibel-Palade body distribution in human umbilical vein endothelial cells (HUVECs).43 Weibel-Palade bodies contain the membrane protein P-selectin.32,33 Dual-label immunostaining of canine AECs revealed that endogenous P-selectin was present in the granules containing WVF (Figure 2G-I). These data confirm that normal canine AECs contain characteristic Weibel-Palade bodies and are similar to human endothelial cells.

We next harvested and cultured endothelial cells from the VWD dogs. Immunostaining for PE-CAM-1 demonstrated a homogeneous population of VWD-AECs has been cultured (Figure 2J). We could not identify a significant difference in morphology between normal and VWD endothelial cells. We also stained for VWF, using the polyclonal antibody that detected the canine VWF in the normal endothelial cells, and no VWF was detected (Figure 2 K, N). No VWF was detected using monoclonal antibodies specific for canine VWF (data not shown). Although sequencing data predicted the potential synthesis of the first 85 amino acids of propeptide, we could not detect VWF mRNA by reverse transcriptase (RT)–PCR using canine specific probes (data not shown). The canine VWD-AECs were immunostained by using a mix of several monoclonal antibodies that recognize canine WVFpp, and no WVFpp expression was detected (Figure 2M). A monoclonal antibody specific for canine WVFpp also failed to detect WVFpp expression (Figure 2Q). These findings demonstrate that the AECs obtained from VWD dogs are WVF and WVFpp null at both a protein and an mRNA level.

Having established the absence of both WVFpp and VWF in the canine VWD-AECs, we examined the intracellular localization of P-selectin, a recognized marker of the Weibel-Palade body. The staining of P-selectin was faint and fairly diffuse. Although several very small granules were observed, most likely these were endosomes/lysosomes (Figure 2P). This would be consistent with previous findings that in the absence of VWF, in cells lacking a regulated secretory pathway, P-selectin is delivered to the lysosome and degraded.34 No apparent Weibel-Palade bodies are detected in the canine VWD-AECs.

Figure 1. A homozygous frameshift mutation results in premature termination of VWF protein synthesis. Genomic DNA obtained from normal and VWD dogs was sequenced, and the 2 sequences were compared, including sequence from exon 4, intron 4, and exon 5.
The VWF-null canine AECs provide a unique opportunity to study Weibel-Palade body biogenesis in a naturally occurring animal model of type 3 VWD. We tested the hypothesis that formation of Weibel-Palade bodies would be induced by expression of either wild-type human or canine VWF. Full-length VWF constructs were introduced by nucleofection of the canine VWD-AECs. Transfected cells were immunostained for VWFpp and VWF and examined by confocal microscopy. In cells expressing canine VWF (data not shown) or human VWF (Figure 3A-C), both VWFpp and VWF appeared to be colocalized in granules. In model cell lines, the recruitment of P-selectin to VWF-containing granules has been used to authenticate the formation of Weibel-Palade bodies. To confirm the formation of Weibel-Palade bodies we examined whether VWF-containing granules in nucleofected canine VWD-AECs would recruit endogenous P-selectin. In cells expressing full-length human VWF (data not shown) or canine VWF (Figure 3D-F), endogenous P-selectin colocalized with the VWF in granules, similar to that in the normal canine endothelial cell. These data indicate that the canine VWD-AECs will support the formation of Weibel-Palade bodies when VWF is expressed and the Weibel-Palade body distribution of P-selectin is reestablished.

Previous work in our laboratory demonstrated that the granular storage of VWF in the AtT-20 model cell line was not dependent on multimerization of VWF. To confirm this in an endothelial cell, we expressed 2 dimeric VWF proteins. The first protein is propeptide-deleted VWF (Δpro) that contains signal peptide followed directly by the mature VWF sequence. When canine Δpro (not shown) or human Δpro (Figure 3G-I) were expressed in the canine VWD-AECs, only diffuse VWF staining was observed, no apparent VWF-containing granules could be detected. The P-selectin distribution remained diffuse with the presence of some small granules (Figure 3H) and did not colocalize with VWF (Figure 3I). The second dimeric VWF protein we expressed is a full-length, propeptide-containing mutant with a point mutation in the D1 domain of the propeptide (Y87S) that our laboratory had previously characterized. Canine VWD-AECs expressing the Y87S-VWF demonstrated the presence of VWFpp and VWF colocalized in granules (Figure 3J-L), with some diffuse staining consistent with localization in the endoplasmic reticulum. Additionally, P-selectin was recruited to the granules in those cells that were expressing Y87S-VWF (not shown) verifying that multimerization is not required for granule formation.

To confirm that the propeptide is required for granule formation, we coexpressed Δpro together with VWFpp as 2 separate constructs (in trans). We observed that nucleofected cells received and expressed both Δpro and VWFpp rather than one or the other plasmid (not shown). Cells expressing both human VWFpp and Δpro proteins displayed the presence of VWF (and presumably VWFpp) in granules that recruited endogenous P-selectin (Figure 3M-O). These data indicate that only when Δpro is coexpressed with VWFpp, in either cis or trans, can granules be formed that recruit endogenous P-selectin. Together, these data confirm earlier findings that propeptide is necessary for granule formation, Δpro alone is not sufficient for formation of a Weibel-Palade body, and multimerization of the VWF is not required for granule biogenesis.

To assess the multimerization of expressed VWF proteins, we analyzed the culture supernatants of transfected cells by SDS–agarose electrophoresis and Western blotting (Figure 4A). No VWF multimers were detected in the supernatants of mock-transfected (lane 7) canine VWD-AECs. Both the wild-type canine and human full-length VWF were normally multimerized (lanes 2, 3). The lack of VWF multimerization of the Y87S-VWF and Δpro proteins was confirmed (lanes 4, 5). The conditioned media were also analyzed on submarine MetaPhor-agarose gels that better define lower molecular weight VWF proteins (Figure 4B). Both Y87S-VWF and Δpro proteins formed only a dimeric VWF species (lanes 4, 5). Examination of full-length human and canine VWF samples revealed that a higher molecular weight species (tetramer) migrated into the gel in addition to the VWF dimer (lanes 1, 3). Densitometry revealed a similar percentage of VWF dimer and tetramer in human and canine samples (lanes 1, 3; peaks 1, 2). Expression of wild-type VWF proteins in canine VWD-AECs results in normal multimerization and secretion of VWF.

Figure 3. Expression of human or canine VWF proteins in canine VWD aortic endothelial cells. (A–F) Wild-type VWF. Canine VWD aortic endothelial cells were transiently transfected with either human VWF (A–C) or canine VWF (D–F) by nucleofection. Transfected cells were fixed, immunolabeled, and examined by confocal microscopy as described in “Materials and methods.” Cells were immunolabeled with a mix of several mouse monoclonal anti-VWFpp antibodies (A), a rabbit anti-VWF (B), AW-1 mouse monoclonal anti-VWF (D), or rabbit anti–P-selectin (E). Immunostaining of cells expressing human VWF demonstrated a granular distribution of VWFpp and VWF (A–B). Merging the 2 images revealed colocalization of the proteins (C). Dual staining for VWF and P-selectin revealed that the canine VWF (D–F) colocalized with P-selectin in granules. Expression of canine VWF reestablishes the Weibel-Palade body distribution of endogenous P-selectin. (G–L) Dimeric VWF. Cells were transiently transfected with either a dimeric, propeptide-deleted VWF (Δpro) or dimeric, mutant Y87S-VWF. Cells were immunostained using the monoclonal anti-VWF, AW-1 (G), rabbit anti–P-selectin (H), a mix of several mouse monoclonal anti-VWFpp antibodies (J) or polyclonal anti-VWF (K). Cells expressing human Δpro (G–I) showed diffuse VWF staining with no apparent VWF-containing granules (G). P-selectin distribution was unchanged (H), and the 2 proteins did not appear to colocalize (I). A dimeric, Y87S-VWF mutant was also transiently expressed in the canine VWD aortic endothelial cells (J–L). The mutant Y87S VWFpp and VWF showed a granular staining pattern in addition to diffuse staining and were found to colocalize (J–L). VWF multimerization is not necessary for Weibel-Palade body formation. (M–O) Coexpression of VWFpp and Δpro. Cells were transiently transfected with Δpro and VWFpp in trans as 2 separate constructs. Cells were immunostained using AW-1 monoclonal anti-VWF (M) and rabbit anti–P-selectin (N). In cells expressing VWF, P-selectin was found to colocalize with VWF in granules (O). Coexpression of VWFpp and Δpro generates Weibel-Palade bodies in canine VWD aortic endothelial cells.
The propeptide is clearly necessary for formation of a VWF-containing Weibel-Palade body. Our previous studies in AtT-20 cells demonstrated that VWFpp independently traffics to ACTH-containing regulated secretory granules, but these studies could not address whether VWFpp could generate the granule. We exploited the absence of endogenous secretory granules in the canine VWD-AECs to examine whether the VWF propeptide is sufficient for Weibel-Palade body formation. Immunostaining and confocal microscopy of cells expressing either human VWFpp (Figure 5A-C) or canine VWFpp (Figure 5D-F) demonstrated mostly diffuse staining, although a number of small, VWFpp-containing granules were occasionally observed. We also examined the distribution of P-selectin in the VWFpp-expressing canine VWD-AECs. When either human VWFpp (Figure 5G-I) or canine VWFpp (data not shown) were expressed, the distribution of P-selectin was not altered. VWFpp and P-selectin did not appear to be colocalized, indicating that VWFpp alone is not sufficient for formation of a Weibel-Palade body that recruits endogenous P-selectin. These data suggest that in addition to VWFpp, the mature portion is necessary for the biogenesis of a P-selectin containing Weibel-Palade body.

Figure 4. Multimeric structure of expressed canine and human VWF proteins. (A) The multimeric structure of expressed VWF constructs was analyzed nonreduced on a 3% agarose–SDS gel. All samples were run on the same gel, and interfering (nonrelevant) lanes have been removed for clarity. No VWF was detected in the mock-transfected control (lane 7). Wild-type canine and human full-length VWF (lanes 2, 3) and normal human and canine plasmas (lanes 1 and 6) were normally multimerized. Neither Y87S-VWF (lane 4) nor Δpro (propeptide-deleted VWF, lane 5) formed high molecular weight multimers. (B) The structure of lower molecular weight VWF subunits was analyzed on a 3% MetaPhor-agarose submarine gel. No VWF was detected in the mock-transfected control (lane 2). Y87S-VWF and Δpro formed only a dimeric VWF species (lanes 4, 5). The density of the dimer and tetramer bands observed in human and canine VWF (lanes 1, 3) were calculated after scanning the immunoblot (peak 1 = dimer).

Figure 5. VWFpp alone does not induce Weibel-Palade body formation. The VWFpp and the fusion protein VWFpp*C3α were independently expressed in canine VWD aortic endothelial cells. Transfected cells were processed and examined as described in “Materials and methods.” Cells were immunostained with a rabbit anti-VWFpp (A, D), a mix of several monoclonal anti-VWFpp antibodies (G, J, M), mouse anti-PECAM (B, E), rabbit–anti-P-selectin (H, N), or polyclonal anti-C3 antibody (K). In cells expressing either human VWFpp (A-C) or canine VWFpp (D-F), the distribution of VWFpp was diffuse, although small VWFpp-containing granules were occasionally observed (D). Cells expressing human VWFpp were dual stained for VWFpp (G) and P-selectin (H): P-selectin distribution was unchanged compared with untransfected VWD endothelial cells and very little colocalization of VWFpp and P-selectin was observed (I). Cells expressing the VWFpp*C3α fusion protein showed both diffuse staining as well as granular storage of both VWFpp (J) and C3α (K). Dual-staining for VWFpp (M) and P-selectin (N) revealed very little colocalization of the VWFpp*C3α protein with P-selectin (O). While the VWFpp*C3α protein may induce granule formation in the canine VWD aortic endothelial cells, P-selectin is not recruited to the newly formed granule.
We hypothesized that the mature portion of the VWF molecule was providing the dense core of protein needed for condensation, aggregation, and granule formation. We sought to determine whether another protein could provide the necessary aggregation. We reported a VWFpp fusion protein, VWFpp*C3α, that contains signal peptide and VWFpp with a disrupted furin cleavage site followed by the α-chain of complement C3. This VWFpp*C3α fusion protein is stored in granules in the AtT-20 cell line and sorts to endogenous, VWF-containing granules in bovine aortic endothelial cells. Due to the presence of endogenous secretory vesicles in these cells we could not address the biogenesis of secretory vesicles. Dual-immunostaining for VWFpp and C3α of VWD-AEC expressing VWFpp*C3α (Figure 5J-L) occasionally showed the presence of a few VWFpp and C3α-containing granules, although the distribution of VWFpp and C3α in most cells was diffuse. VWFpp*C3α-transfected cells were also dual labeled for P-selectin and VWFpp. Some transfected cells demonstrated the presence of VWFpp-containing granules; however, the VWFpp did not appear to colocalize with endogenous P-selectin (Figure 5M-O). While the C3α fused to VWFpp may be sufficient for aggregation and granule formation, these granules do not recruit P-selectin, indicating that the mature portion of VWF may be necessary for the recruitment of P-selectin to the Weibel-Palade body.

To determine whether the granules formed by expression of VWF proteins are regulated secretory granules, we examined whether release of the expressed proteins could be induced by exposure to the secretagogue, PMA. Confluent transfected cells were exposed to either PMA or control for 45 minutes at 37°C. The incubation media was collected, and the amount of VWF (Figure 6A) or VWFpp (Figure 6B) was determined by ELISA. We observed a significant increase in VWF secretion in response to PMA (426% of control). The release of Y87S-VWF was also significantly increased (426% of control). Expressed Δpro did not appear to be released in response to PMA (123% of control). The VWF expression levels of cells exposed to control were equivalent (Figure 6A). Additionally, the VWF expression levels over the 24-hour period prior to agonist stimulation were found to be similar for wild-type VWF, Y87S-VWF, and Δpro, ranging from 3 to 6 mU/mL. The VWFpp concentration for wild-type VWF and Y87S-VWF samples were also in this range, indicating a VWFpp/VWF ratio of approximately 1:1. In contrast to full-length VWF, we did not observe a significant increase in secretion of VWFpp (99% of control) or VWFpp*C3α (100% of control) in response to PMA (Figure 6B). The prestimulation expression level of VWFpp*C3α was decreased when compared with VWFpp, 5 mU/mL and 13 mU/mL, respectively. Cells expressing both VWFpp and Δpro were responsive to PMA stimulation, and a significant increase in VWFpp secretion (and presumably VWF) was observed (Figure 6, 729% of control). In this case, the prestimulation expression levels of VWFpp and Δpro were not proportional. VWFpp concentration was almost 3 times greater than Δpro, 10 mU/mL and 3 mU/mL, respectively. This disproportionate expression could account for the seemingly increased agonist-induced release of VWFpp when it is coexpressed with Δpro. Together, these data indicate that full-length VWF, Y87S-VWF, and the trans expression of VWFpp and Δpro result in formation of Weibel-Palade bodies that undergo agonist-induced exocytosis, while Δpro, VWFpp, or VWFpp*C3α are not trafficked to the regulated storage pathway.

**Discussion**

In this study, we have identified the molecular defect causing the type 3 VWD in the dogs from the closed colony maintained in Chapel Hill. The ΔC255 deletion in exon 4 in the codon corresponding to serine 85 of the VWF propeptide results in 34 amino acids of altered protein sequence and premature termination of VWF synthesis. This mutation accounts for the absence of VWF protein in the plasma of the VWD dogs. Interestingly, this is the same mutation that was identified in Scottish Terriers with type 3 VWD. Although the sequence predicts the synthesis of the first 85 amino acids of VWFpp followed by 34 altered amino acids, we could not detect any truncated VWFpp protein, nor did we detect VWF mRNA in the VWD aortic endothelial cells by RT-PCR using canine VWF specific primers. Together these data indicate that the VWD dogs do not synthesize any VWF.

We have confirmed the absence of VWFpp and VWF in the AECS obtained from the VWD dogs. Examination of the distribution of P-selectin in the VWF-null aortic endothelial cells verified the absence of recognizable Weibel-Palade bodies in contrast to normal canine AECS. The canine P-selectin appeared to be degraded rapidly as indicated by the faint, diffuse staining and localization in small granules presumed to be lysosomes, similar to the P-selectin distribution observed in the VWF-deficient knockout mouse. A similar degradation of P-selectin was demonstrated in the T24 human epithelial cell line that does not synthesize VWF. We were unable to definitively identify the P-selectin-containing granules as lysosomes due to lack of availability of antibodies cross-reacting with canine lysosomal proteins. However, the degradation of P-selectin in the absence of a regulated secretory pathway is well-defined as previously described.

Many prior studies on VWF trafficking have been performed in transformed, non–endothelial cell lines. Several laboratories have demonstrated the colocalization of P-selectin with VWF in storage granules when VWF is coexpressed with P-selectin in non–endothelial cell lines. A VWF-null, nontransformed endothelial cell has not been available for exploring the intracellular processing and trafficking of VWF in its natural site of synthesis. We have developed a system to examine the formation of storage granules in a true endothelial cell that does not contain preexisting secretory vesicles, synthesizes P-selectin, and is responsive to secretagogues. Our results using the VWF-null endothelial cells corroborate many earlier findings. Our laboratory and other investigators have shown
that multimerization and storage of VWF are unrelated, independent processes.\textsuperscript{17,21,27,37} The formation of Weibel-Palade bodies that recruit endogenous P-selectin when dimeric Y87S-VWF is expressed in the canine VWD-AECs provides further evidence. The trans expression and granular storage of VWFpp and Δpro has been demonstrated in both AtT-20 and CV-1 cell lines.\textsuperscript{39} In the canine VWD cells, the coexpression of VWFpp and Δpro of either canine or human origin created Weibel-Palade bodies that recruited endogenous P-selectin and were released in response to PMA.

The question that has not been addressed in any prior studies is whether the VWFpp alone is sufficient for the formation of a Weibel-Palade body. Due to the presence of endogenous-regulated secretory vesicles in AtT-20 and bovine aortic endothelial cells, granule biogenesis could not be addressed. While the propeptide clearly sorts independently to these endogenous secretory granules, the ability to generate the storage granule is a different question. Here we demonstrate that VWFpp alone does not appear to generate a bona fide Weibel-Palade body. Canine VWD-AECs expressing VWFpp showed mostly diffuse staining consistent with its localization in the endoplasmic reticulum. Although many small granules were occasionally observed, the nature of these granules is not clear. The lack of P-selectin recruitment, diminished size, and lack of agonist-induced release of the granules formed by VWFpp in the VWF-null endothelial cells indicate that Weibel-Palade bodies are not formed. While VWFpp is clearly required in addition to mature VWF (Δpro), VWFpp independently is not sufficient for Weibel-Palade body biogenesis.

Two hypotheses have been proposed that would explain the formation of storage vesicles.\textsuperscript{40,41} One hypothesis focuses on the selective aggregation or condensation of proteins in the trans-Golgi to trigger granule formation. The second proposes that regulated secretory proteins contain targeting signals that interact with sorting “receptors” that escort the proteins into the storage pathway. We considered whether the “mature” portion (Δpro) of VWF is providing the aggregation needed for protein condensation and granule budding, or whether an alternative protein could substitute for Δpro. In a previous study we fused the α-chain of component C3 (C3α) to VWFpp to demonstrate the ability of VWFpp to traffic an unrelated, non–secretory-granule protein to the regulated storage pathway, but could not address the biogenesis of the granule.\textsuperscript{19} In the VWF-null canine endothelial cells, we observed a subset of cells that contained distinct granules, in addition to a significant number of cells with diffuse staining consistent with endoplasmic reticulum localization of VWFpp*C3α. In the cells that showed granule formation, canine P-selectin did not appear to colocalize with the VWFpp*C3α-containing granules. The lack of P-selectin recruitment can be rationalized by 1 of 2 explanations. The first explanation is that the VWFpp*C3α-containing granules are not regulated secretory vesicles. We did not observe any agonist-induced release of VWFpp*C3α. However, only a small percentage of cells expressing VWFpp*C3α contained granules, and the amount of released protein may not be within limits of detection. An alternative explanation for the lack of P-selectin recruitment is that P-selectin may be brought to the Weibel-Palade body in a VWF-dependent manner. Prior studies support this hypothesis as P-selectin can be diverted from ACTH-containing granules in AtT-20 cells (when expressed in the absence of VWF) to granules induced by VWF expression that do not contain ACTH.\textsuperscript{31} The mechanism for this VWF-dependent recruitment of P-selectin to the Weibel-Palade body remains to be clarified.

Although VWFpp alone does not appear to induce the formation of a Weibel-Palade body, it is clearly indispensable in the processing and intracellular trafficking of VWF. Evidently, both VWFpp and mature VWF protein are required for creation of the regulated secretory vesicle to which P-selectin ultimately traffics. Whether VWFpp provides the “sorting signal” and mature VWF provides the core of aggregation remains unresolved. With the exception of type 3 VWD, there have been no reports of patients that lack Weibel-Palade bodies. In the VWF-deficient mouse, the loss of Weibel-Palade body localized P-selectin resulted in defective regulated secretion and loss of leucocyte recruitment. Recently, P-selectin has been shown to interact with endothelial-released ultra-large VWF (ULVWF) multimers.\textsuperscript{42} P-selectin may serve to tether the ULVWF to endothelial cells. In addition to the impact on P-selectin, the loss of Weibel-Palade bodies due to defective VWF would have an impact on several other proteins, including tissue-type plasminogen activator (t-PA), interleukin 8 (IL-8), and potentially factor VIII that can be stored with VWF.\textsuperscript{43,47} Thus, defining the complex mechanism of Weibel-Palade body biogenesis has significant biologic implications.

References


