Introduction

Tumor endothelial marker 8 (TEM8) was originally discovered by serial analysis of gene expression (SAGE) in endothelial cells of colon carcinomas [1]. The analysis revealed markedly elevated levels of TEM8 mRNA in these cells, and normal endothelial cells appeared to have little of it. Further, these studies identified three splice variants of the gene, V1, V2, and V3. The variants encode proteins of 564, 368, and 333 residues, respectively. Topologically, V1 and V2 are type-1 integral membrane proteins with a single transmembrane helix. V2 is identical to V1 up to residue 364, but the last four residues of V2 are unique, a consequence of differential splicing. V3 sequence diverges from V1 and V2 right before the beginning of the transmembrane helix. Thus, V3 has a portion that is identical to the extracellular portions of V1 and V2, but its 13-residue carboxyl terminal segment is unique.

Using a human cDNA library for expression cloning, Bradley et al. [2] independently identified TEM8 V2 as an anthrax toxin receptor. Later studies showed that V1 also functions as an anthrax toxin receptor, and that V3 does not [3]. Thus, these findings were consistent with the original reports that V1 and V2 are integral membrane proteins and V3 is a secreted form [1, 2]. However, TEM8 is not the only cell surface protein that anthrax toxin uses to enter cells. Capillary morphogenesis gene 2 protein (CMG2), a protein similar to TEM8, was found to be an avid receptor as well [4]. Indeed CMG2 proved a stronger anthrax toxin receptor than TEM8 [5]. CMG2, like TEM8, has multiple splice variants. At least two membrane-bound forms of it, the 488- and 489-residue proteins, both strongly support anthrax toxin entry [4, 5]. CMG2 was originally identified by way of its enhanced expression in human umbilical vein endothelial cells, and evidence suggests it is a player in angiogenesis [6].

The only definitively known function of TEM8 is its role as an anthrax toxin receptor, and commensurate with that function it is designated ANTXR1. Likewise, the confirmed role of CMG2 is that of anthrax toxin receptor, and therefore it is designated ANTXR2. Each protein has a von Willebrand Factor A (vWA) domain in its extracellular portion, and within this domain each protein also has a metal ion dependent adhesion site (MIDAS). Both features are important for anthrax toxin protective antigen (PA) binding [4, 5, 7].

Tumor endothelial marker 8 (TEM8; ANTXR1) is one of two anthrax toxin receptors; the other is capillary morphogenesis gene 2 protein (CMG2; ANTXR2). TEM8 shows enhanced expression in certain tumor endothelia, and is thought to be a player in tumor vasculature formation. However, a comprehensive expression profile of individual TEM8 variants in normal or cancerous tissues is lacking. In this work we carried out an extensive analysis of all splice variants of human TEM8 in 12 digestive tissues, and 8 each fetal and adult tissues, 6 of them cognate pairs. Using variant-specific primers, we first ascertained the status of full-length transcripts by nested PCR. We then carried out quantitative analysis of each transcript by real-time PCR. Three splice variants of TEM8 were reported before, two single-pass integral membrane forms (V1 and V2) and one secreted (V3). Our analysis has revealed two new variants, one encoding a membrane-bound form of the receptor and the other secreted, which we have designated V4 and V5, respectively. All tissues had V1, V2, V3, and V4, but only prostate had V5. Real-time PCR revealed that all variants are present at different levels in various tissues. V3 appeared the most abundant of all. To ascertain its functionality for anthrax toxin, we expressed the newly identified form V4 in a receptor-negative host cell, and included V1 and V2 for comparison. Cytotoxicity, toxin binding, and internalization assays showed V4 to be as efficient a receptor as V1 and V2.
Anthrax toxin is secreted by *Bacillus anthracis*, a Gram-positive, encapsulated, spore-forming rod [8,9]. The toxin comprises three proteins: protective antigen (PA), edema factor (EF), and lethal factor (LF). None of the proteins alone is toxic to cells; the toxic assemblies are PA+EF and PA+LF [8,9]. EF is a calmodulin-dependent adenyl cyclase that harms cells by producing excessive amounts of cAMP [10]. LF is a Zn-dependent metalloprotease that targets MAP kinase kinases in cytosol; LF can cleave six out of seven of these kinases [11,12,13,14,15]. This cleavage is detrimental to cells, and its consequences include apoptosis in at least human endothelial cells and the mouse macrophage line RAW264.7 [16,17,18], as well as in human melanoma cells [19]. PA itself has no toxic activity, but is crucial because it delivers EF and LF inside cells by receptor-mediated endocytosis, a process that requires functioning ANTXR1 and ANTXR2 [2,4,8,9]. Upon binding the receptors PA is cleaved by furin, releasing a 23-kDa amino-terminal fragment [20,21]. The receptor-toxin complex is then internalized. Upon exposure to acidic pH in an endosome the heptamer changes its conformation, subsequentially translocating EF and LF into cytosol [8,9]. Acidification of the endocytic pathway is indispensable for the intoxication process [25].

The natural ligands for TEM8 and CMG2 have not been precisely identified. It is known, however, that TEM8 interacts with cleaved C5 domain of collagen z3 [26], while CMG2 can interact with collagen IV and laminin [6]. But the functional significance of these interactions remains obscure. The suggested native role for the two proteins is in neo-angiogenesis, a complex process crucial not only for normal development, but also for many pathological states, such as tumor development and wound healing [9,6,27,28]. Notably, TEM8 shows enhanced expression in endothelia of certain colon carcinomas, suggesting its role in tumor vasculature formation [1,29]. TEM8 expression has also been studied in some phagocytes [30].

We have carried out extensive variant-specific expression analysis of human TEM8 by nested as well as real-time PCR. We report here that TEM8 transcripts show ubiquitous expression, albeit at different levels. We also report identification of two new transcripts, one of which encodes a fully functional anthrax toxin receptor.

### Materials and Methods

#### Reagents and Cells

Human prostate and fetal brain Smart RACE Marathon Ready cDNAs, human multiple tissue cDNA panels, and plasmid pRESHyg3 were purchased from Clontech (Palo Alto, CA). The template in Marathon Ready cDNAs is double-stranded, but that in the tissue panels is first strand synthesized (reverse transcribed) RNA-DNA hybrid. The reagents for PCR (polymerases, dNTPs, and buffers) were from Clontech, Invitrogen (Carlsbad, CA), and Takara (Japan). Primers were made by the FDA Core Facility (Bethesda, Maryland), Sigma-Genosys (Houston, Texas), or Invitrogen (Carlsbad, CA). Restriction enzymes and the ligation kit were from New England Biolabs (Ipswich, MA). Supercompetent XL1-Blue cells were from Stratagene (La Jolla, CA). Functionally receptor-negative cell line PR290 and its parental line, WTB111, have been reported before [3].

#### Primers

For TEM8 V1, the primers were based on the reported longest original transcript (NCBI accession number, NM_032208; 5540 bp; coding sequence, 144–1836; 564 aa). For nested PCR, the primary and secondary (nested) PCR primers were designed to partially overlap (Table 1). The primers s21 and as 1842 amplify an 1848-bp V1 fragment containing the coding sequence. For analysis of V2 and V3, the sense primers were the same as for V1, but the antisense primers were specific, each representing the unique 3’ region of V2 (NCBI accession number, NM_053034; 1454 bp; coding sequence, 144–1247; 368 aa), or V3 (NCBI accession number, NM_018153; 2143 bp; coding sequence, 144–1142; 333 aa). All primers were assessed for any secondary structures and self-complementarity using the Oligonucleotide Calculator (http://www.basic.northwestern.edu/biotools/oligocalc.html).

The primers were resuspended in 1 mL of Mill-Q water (Millipore) and incubated at 70°C for 15–20 min to ensure thorough dissolution and inactivation of any accidental contamination with DNase. Appropriate dilutions were made, typically 1:25, to measure absorbance at 260 nm. All primers were then diluted to a working concentration of 10 μM.

#### Toxins

PA+FP59 was the toxin for cytotoxicity assays. FP59 is a recombinant toxin that carries the PA-binding amino terminal domain of LF and the catalytic domain of *Pseudomonas aeruginosa* exotoxin A. Unlike PA+LF, PA+FP59 can kill all cells. PA was also purchased from List Biological Laboratories (Campbell, CA).

### Table 1. A directory of primers used to analyze different splice variants of ANTXR1 (TEM8).

| Primer       | Var | Sequence                  | PCR   |
|--------------|-----|----------------------------|-------|
| TBV1-S3      | 1–5 | 5’TGTGCTTCCGGGGAGTTGCGAGGGAGCG 1 |       |
| TBV1-S21     | 1–5 | 5’GGAGGGAGCGGAGGGGAAAAGGACCC 2 |       |
| TBV1-A51744  | 1   | 5’GTTGGAGAGGTGGGAGGCGG 2     | 1°C   |
| TBV1-A51738  | 1   | 5’GGAAGGTGGGAGGGGAGAGGAGGAGG 2 |       |
| TBV1-A51728  | 1   | 5’TGGGAGAGGTGGGAGGAGGAGGAGG 2 | 1°C   |
| TBV1-A51716  | 1   | 5’GAGGAGGGAGGAGGAGGAGGAGGAGG 2 |       |
| TBV1-S1721   | 1   | 5’CCCAGCGCCACGGCGCCCTAC 1    |       |
| TBV1-S1726   | 1   | 5’CTACCCGCACACCCCTATCCG 2    |       |
| TBV1-A53695  | 1   | 5’GGCGCGATCCGGTCGTCGTAAGGCA 1 |       |
| TBV1-A53670  | 1   | 5’GACTCCCTGAGCTTACCTGTATCTGG 2 |       |
| TBV1-A51859  | 1,4,5| 5’ACATCCCTCCTGAGGCTTATGAGGACCC 1 |       |
| TBV1-A51842  | 1,4,5| 5’TGGAGAGGCGGAGAGGAGGAGGAAGTTGG 2 |       |
| TBV1-A51919  | 1,4,5| 5’GTTGGTAAGGCTTCACCTGATATCCAC 1 |       |
| TBV1-A51903  | 1,4,5| 5’GCTCACCATGACATCTCGTTCATAC 2 |       |
| TBV2-A51273  | 2   | 5’GGCGCGATCCGGAGAGTATCTGGGAGGAGGAGGAGG 1 |       |
| TBV2-A51251  | 2   | 5’TCTGGGAGTCAGTCATGATGAGGAGGAGGAGGAGG 1 |       |
| TBV3-A51374  | 3   | 5’CCACCGGTCGATGAGGAGGAGGAGGAGGAGGAGG 1 |       |
| TBV3-A51352  | 3   | 5’TATCAGGGAATATGAGGAGGAGGAGGAGGAGGAGG 2 |       |

*S, sense; AS, antisense. 1, primary PCR. 2, secondary (nested) PCR. The 5’ sense primers 53 and 521 are common to all TEM8 variants. The pairs of sense and antisense primers for primary and nested PCR had partial overlaps. V2-a5121 and V2-a51273 are 30 nucleotides long. All other primers are 27 nucleotides long. The expected amplicons after 2° PCR are: V1, 1848 bp (with as1842); V2, 1260 bp; and V3, 1358 bp. doi:10.1371/journal.pone.0043174.t001

Additional information:

- **New ANTXR1/TEM8 Transcripts and Expression**
- **Table 1. A directory of primers used to analyze different splice variants of ANTXR1 (TEM8).**
Nested PCR

The following steps were taken to control for systematic and random experimental variations in PCR results: 1) For comparison of TEM8 transcript expression levels, all reagents and supplies for PCR were from the same source. 2) The PCR reaction mixture was always 50 μL, prepared by mixing 5 μL of cDNA template with 43 μL of master mix. The master mix contained (per sample): 1 μL each of 10 pM primers (10 pmol), 5 μL of 10X polymerase buffer, 1 or 4 μL of dNTP stock (2.5 mM or 10 mM each), 1 unit of DNA polymerase (1 μL for Clontech’s Advantage-2 polymerase; 0.25 μL for TaKaRa Taq polymerase), and 33 or 36 μL of PCR grade water. 3) The same type of 200-μL tubes were used for all PCR reactions. 4) All PCR cycles were done in the same machine under identical conditions (MyCycler, Bio-Rad, Hercules, CA). 5) The primary PCR protocol was: 1) 94°C for 25 sec. 2) 72°C for 3 min. 3) 6 times to step 1. 4) 94°C for 25 sec. 5) 72°C for 3 min. 6) 29 times to step 4. 7) 70°C for 5 min. 8) 4°C until removed. 6) The secondary PCR differed only in the number of cycles: step 3 was 4 times to step 1, and step 6 was 24 times to step 4. For step-wise amplification, the secondary PCR was done in increments of 5 cycles after the first 5. To do so, step 6 repeats were 4, 9, 14, 19, and 24. At the end of each repeat, 4 μL aliquots were removed and the tubes put back for the next 5 cycles. The samples were immediately subjected to agarose gel analysis, or were frozen at –20°C for later analysis.

Real-time PCR

The Quantifast SYBR Green PCR Kit was used for qPCR (Qiagen, Gaithersburg, MD). The qPCR machine used was AB7300 (Applied Biosystems/Life Technologies, Carlsbad, CA). The reactions and protocols were performed as directed by the manufacturers. The Quantifast SYBR green PCR master mix has ROX passive reference dye, which is necessary for Applied Biosystems real-time PCR cyclers. Each reaction mixture contained 12.5 μL of 2X Quantifast SYBR green PCR master mix, 8.5 μL of RNase-free water, 1 μL each of the sense and antisense primers, and 2 μL of template cDNA. The PCR protocol comprised initial 5 min at 95°C to activate HotStarTaq Plus DNA polymerase, and then 40 cycles of 30-sec denaturation at 95°C and 60-sec annealing/extension at 70°C. Data acquisition was performed during combined annealing/extension step. Melting curve analysis was done immediately after the PCR. The SDS v1.2 software (Applied Biosystems) was used for data analysis. Real-time analysis of β-actin was included as the normalization standard. Ct values were determined using the Applied Biosystems software as directed. TEM8 splice variant expressions were compared in terms of ΔCt values, calculated by subtracting the β-actin Ct values from the TEM8 variant Ct values. Two independent real-time PCRs were done to get the average values.

Electrophoresis

All analytical and preparative gels were made with Ultrapure agarose (Invitrogen, Carlsbad, CA). The gels were 1% in 1X TAE buffer, and the electrophoresis was also in 1X TAE buffer. For analytical gels, 3 μL of primary or secondary PCR samples were used. For quantitative comparison, analytical gels were made fresh, and all PCR reaction mixtures for fetal and adult tissue samples were loaded on the same gel. Such gels were run for the same length of time. For visualization the gels were stained in water containing 0.2 μg/mL of ethidium bromide solution, made by mixing 3 μL of ethidium bromide (10 mg/mL stock) in 150 mL of deionized water. The gels were stained for 30 min and destained for 45–60 min with two changes of deionized water. The gels were then photographed for permanent records using Chemi-Doc Imager (Bio-Rad, Hercules, CA). To control for exposure and other visualization variations, all photography was performed at constant camera and other settings.

Purification of PCR Products

Based on results from analytical gels, those mixtures that had unique bands were purified directly using a PCR product purification kit (Qiagen, Germantown, MD). Those mixtures that had two or more PCR bands were subjected to preparative gel electrophoresis. Long-tooth combs were used to make preparative gels. Sample over-loadings were avoided to retain electrophoretic resolution of DNA fragments that had physical proximity. The bands of interest were then cut out of gels and the amplified DNA fragments purified using a gel extraction kit (Qiagen, Germantown, MD). The final elution volume depended on the amount of products, as judged from the analytical gels, and varied from 30 to 100 μL. The purified DNA fragments were analyzed on gels to assess the degree of purity and recovery. When purity was insufficient or recovery low, the PCR reactions were carried out again, sometimes in duplicate or triplicate, to obtain sufficient amounts for purification.

Sequencing and Analysis

Purified TEM8 amplicons were sent to a commercial facility for sequencing (UT Austin, DNA Core Facility, Austin, TX). Multiple sense and antisense primers were used to get overlapping sequences, and from these the consensus sequences were assembled with Lasergene software (DNA Star, Madison, WI). The sequences were analyzed by BLASTn, BLASTp, and BLAST-2 (NCBI, Bethesda, MD). BLAT (University of California, Santa Cruz) was also used to determine differential splicing of TEM8 variants, as well as precise determination of splicing boundaries. Multiple DNA and protein sequences were aligned using ClustalW2 (http://www.ebi.ac.uk/Tools/msa/clustalw2/).

Cloning

The fetal brain SmaI-XbaI fragment of TEM8V4 amplicon obtained with primers s21+r1842 was cloned into the SnuI-NheI sites of pRESHyg3 by routine recombinant DNA techniques. The final construct was confirmed by sequencing. Ligation of XbaI ends with NheI ends reconstituted the TGA stop codon of TEM8V4. Plasmid-encoded TEM8 V1 and V2 constructs have been reported before [3].

Transfection

On day 1, PR230 cells were seeded in 10-cm plates in 10 mL of DMEM. On day 2, when cells were about 30% confluent, cells were transfected with plasmid DNA encoding TEM8 V1, V2, or V4 using Cellfectin as directed by the supplier (Invitrogen, Carlsbad, CA). Briefly, 3 μg of plasmid DNA was mixed with 20 μL of Cellfectin. The medium was replaced with 8 mL of fresh DMEM, the transfection mixture added drop by drop, and thoroughly mixed. Following incubation for 6 h, the medium was replaced with 10 mL of fresh DMEM. Two days later the cells were passed 1:5 in DMEM containing 250 μg/mL of hygromycin B. PR230 alone and transfected with the vector were also subjected to selection. Full selection took nearly two weeks. Cells were kept at 37°C in humidified chambers with 5% CO2. The selected cells were stored in liquid nitrogen.

Cytotoxicity Assays

On day 1, cells were seeded in 96-well plates at a density of 4000 (WTB111) or 5000 (PR230) cells/well in 150 μL of growth
medium (DMEM containing 5% FBS, 2 mM glutamine, and 50 μg/mL gentamicin). On day three, various dilutions of PA+FP59 were added in triplicate. FP59 concentration was constant at 50 ng/mL, but PA concentration varied by 5-fold serial dilutions (2000, 400, 80, 16, 3.2, 0.64, and 0.128 ng/mL). On day 5 (40 h), MTT was added (final concentration, 0.5 mg/mL) to assess cell viability. The incubation with MTT was for 45–60 min. The medium was removed and cells lysed in a lysis solution (isopropanol containing 0.1 N HCl and 0.5% SDS; 100 μL/well). Colorimetric readings were taken at 570 nm, with 650 nm as the reference, in Synergy 2 plate reader (Biotek, Winooski, VT).

PA Binding and Internalization Assays

WTB111, PR230 alone, and PR230 expressing V1, V2, or V4 were seeded in 24-well plates at the density of 150,000/well in 1 mL of DMEM. The next day regular growth medium was replaced with 1 mL of binding medium containing 1 μg/mL of PA. The binding medium was DMEM without bicarbonate but supplemented with L-glutamine (2 mM), penicillin (50 u/mL), streptomycin (50 μg/mL), and 0.1% BSA (0.5 mg/mL). Incubation with the toxin was at 4°C for 2 h. The toxin medium was then aspirated, cells washed five times with binding medium, and lysed in 150 μL of RIPA buffer (Thermo Scientific, Rockford, IL). The buffer composition was 25 mM Tris-Cl, 150 mM NaCl, 1% NP40, 1% sodium deoxycholate, 0.1% SDS, pH 7.6, and it was supplemented with a commercially available cocktail of protease inhibitors (Roche Applied Science, Indianapolis, IN). The lysates were shaken for 5 min, centrifuged for 20 min (16,000 × g, 4°C), and the supernatants recovered for further analysis.

Binding at 37°C was performed identically, except that the binding medium was prewarmed to 37°C before adding PA (1 μg/mL). The incubation with the toxin was for 1 h. The cells were then placed on ice, toxin medium removed, washed, and lysed as described above.

Western Blotting

Total protein in cell lysates was determined using a commercially available Bradford assay kit (Bio-Rad, Hercules, CA). Lysates from the 4°C PA binding assays were subjected to 7.5% SDS-PAGE to resolve PA63 and PA63. The lysates from the 37°C binding assays were subjected to SDS-PAGE in a 10–15% gradient gel to resolve PA63, PA63, and the PA63 heptamer. Equal amounts of total protein (40 μg/mL) were used for SDS-PAGE. The proteins were then transferred to nitrocellulose filters. The filters were incubated at room temperature for 1 h in blocking solution (5% nonfat dry milk in PBS-Tween20) at room temperature, the solution removed, and the filters washed 3 times with PBS-Tween20. Next the filters were incubated for 3 h at room temperature with anti-PA antibody (rabbit polyclonal, 1:5,000 dilution in PBS-Tween20). The antibody was removed, filters washed 3 times, and incubated with HRP-conjugated goat anti-rabbit antibody (1:10,000 dilution in PBS-Tween20). Following removal of secondary antibody, the filters were washed again, processed using an ECL Western blotting kit, and exposed to X-ray films (CL-Xposure) for image development (Thermo Scientific, Indianapolis, IN).

Results

Identification of a New Membrane-bound Variant of ANTXR1 (TEM8)

The reported original TEM8 (ANXR1) splice variants encode proteins of 564, 368, and 333 residues, respectively designated V1, V2, and V3 [2,3]. V1 and V2 are single-pass integral membrane proteins and V3 secreted. Both V1 and V2 function as anthrax toxin receptors, while V3 does not [2,3]. TEM8 was originally discovered as one of nine tumor endothelial markers that showed enhanced expression in colon tumor endothelia [1]. Later findings showed that TEM8 mRNA also shows elevated expression in certain other tumor endothelia [29]. However, a detailed expression analysis of each of the three transcripts has been lacking. For this work we analyzed the expression of each variant in a large number of human tissues using commercially available cDNA panels. Using variant-specific primers, we analyzed the presence of full-length TEM8 transcripts, which required nested PCR.

For V1 transcript analysis, the primary PCR primers were s3+ras1859 (Table 1, Figure 1A). We have based all primer and other numbering on the original 5540-nucleotide transcript for V1 (accession number, NM_002208; ORF, nucleotides 144-1835), originally reported when the gene was first discovered [1]. Initially we ran a pilot analysis using human fetal brain and prostate Marathon Ready cDNA. But PCR with these primers resulted in no amplification. We think this was partly because of low concentration of template cDNA, but could not rule out the possibility that the V1 transcript may be a rare one. We therefore opted to do nested PCR with primers s21+ras1842 (Table 1, Figure 1A). This resulted in definite amplification of a unique fragment from the fetal brain cDNA (Figure 1B). The unique amplicon from fetal brain appeared smaller than the expected 1848-bp fragment (nucleotides 21–1868). Indeed sequencing revealed this fragment to be 1740 bp, 108 bp shorter than the expected V1 amplicon. We designated it TEM8 variant 4 (V4).

We analyzed the V4 sequence by BLAST, BLAST-2, and BLAT to precisely determine its splicing boundaries. This analysis revealed that differential splicing of V4 relative to the reported V1 transcript is within exon 18, the last one. The intron 17/exon 18 boundary is the same for V1 and V4. Apparently V4 differential splicing comes about by recruiting new splice donor and acceptor sites within exon 18, resulting in exclusion of the 108-bp segment from the coding sequence of V1. This alternative splicing retains the V1 reading frame, and therefore V4 differs from V1 only in that it has an internal 36-residue in-frame deletion spanning residues 522–557 (Figure 2A, B & C). Thus, V4 is a new 528-residue membrane bound form of TEM8/ANTXR1 (GenBank, JX424835).

PCR with the same primers amplified multiple fragments from the prostate template (Figure 1B), including one identical to the unique fetal brain amplicon, V4. Sequencing confirmed that the prostate fragment corresponding to V4 is indeed the same; that is, it lacks the same 108-bp segment from the coding region. We also isolated the amplicon immediately above the V4 band (labeled V1 in Figure 1B), sequenced it, and found that it indeed is V1; it had the 108-bp segment missing in V4.

Identification of TEM8 V5, a New Secreted Variant

As already stated, nested PCR analysis of pooled prostate cDNA with the primer sets s3+ras1859 and s21+ras1842 resulted in multiple amplicons, one of them identical to the 1740-bp fetal brain fragment. However, prostate had a unique smaller amplicon of about 1300 bp. Sequencing revealed this amplicon to be 1368 bp long. Analysis of this sequence by BLAST, BLAST-2, and BLAT revealed that the amplicon represents a new splice form of TEM8, with the same start codon as the other variants. However, differential splicing of this variant comes about by recruiting an alternative splice donor site within exon 12, ten nucleotides upstream of the one for V1. This site then splices with a new acceptor site within exon 18, as shown in Figure 2A. The overall
consequence of this is that exons 13–17 are altogether skipped and the reading frame is shifted. The ORF in this sequence is 358 residues, which diverges from the other forms after residue 314, right before the transmembrane helix (Figure 2B). We have designated this TEM8 splice variant V5 (GenBank, JX424839). Like V3, V5 does not have the transmembrane helix that V1, V2, and V4 do. It also does not have any other predicted transmembrane helix. We therefore conclude that V5 is a secreted form of TEM8. The divergent carboxyl-terminal segments of V3 and V5 are unique, but otherwise they are identical to the membrane-bound TEM8 proteins. Figure 2C schematically shows the relative topological positioning of the three previously reported variants of TEM8, as well as the two new ones reported in this work.

Prostate also had an amplicon slightly below the 1740-bp one (Figure 1B). But we have not established its identity; repeated attempts at isolating and sequencing it failed. Nonetheless, we cannot rule out that it may be yet another variant of TEM8.

**Analysis of Full-length TEM8 V1, V4, and V5 by Nested PCR**

Nest    ed PCR results with human fetal brain and prostate revealed a highly differential profile of TEM8 expression in these two developmentally and histologically distinct tissues. To ascertain a more precise expression profile of each TEM8 transcript, we analyzed a large number of normal human tissues. To do that we used one normalized cDNA set representing 12 digestive system tissues. A second set comprised cDNAs from 8 fetal and 8 adult tissues, 6 of them cognate pairs. For this analysis we used the same V1-specific primers that revealed V4 and V5.

The most prominent band following nested PCR was confirmed to be V4, not V1. As shown in Figure 3, V4 showed ubiquitous expression in digestive, as well as the fetal and adult tissues tested. Upon primary PCR, there appeared noticeable differences in the variant’s expression levels. However, this was not a quantitative analysis, and therefore we could not rely on the differences in expression by this approach. To more precisely determine the expression levels we did real-time PCR.

**Specific Analysis of Full-length TEM8 V1 Transcript by Nested PCR**

As described above, nested PCR primers specifically meant to amplify the 1848-bp V1 fragment, which carries the complete coding sequence for this variant, failed to do so, except in prostate. Instead they revealed a new ubiquitous transcript that encodes a 528-residue membrane-bound form of the protein, as well as a putative secreted form unique to prostate (Figures 1 and 3). This initially led us to conclude that V1 transcript may be rare, and that colon carcinoma endothelial cells from which it was first identified may be among the rare tissues that have it [1].

Nonetheless, we allowed the possibility that unexpected absence of V1 from so many tissues may be an artifactual anomaly, and therefore decided to further assess its apparent absence. To do so, we made sense and antisense primers specific for the 108-bp region unique to V1, but absent from V4 and V5, as well as V2 and V3. We used these primers in conjunction with s3 and s21, as well as other sense and antisense primers to amplify V1 (Table 1). Indeed these attempts resulted in definite bands of the expected sizes in all tissue cDNA templates (Figure 4). Sequencing confirmed that these fragments carried the 108-bp segment unique to V1. We therefore conclude that the absence of V1 upon first analysis was anomalous. However, we are unable to explain why the antisense primers as1859 and as1842, both specific for the reported V1 transcript, in conjunction with s3 and s21 failed to amplify the expected V1 amplicon in any of the 28 tissues represented in the three cDNA panels.

**Nested PCR Analysis of Full-length TEM8 V2 Expression**

ANTXR1/TEM8 V2 encodes a 368-residue membrane-bound form of the protein, and this is also the form first identified as an anthrax toxin receptor [2]. We analyzed this transcript using specific primers that amplify a 1258-bp segment carrying the
entire coding sequence. As shown in Figure 5A and B, this variant appeared to show highly selective expression. However, quantitative analysis by real-time PCR showed that the variant is actually present in all tissues, but its levels differ from tissue to tissue.

**Nested PCR Analysis of Full-length ANTXR1 V3 Expression**

Splice variant 3 of ANTXR1/TEM8 encodes a 333-residue secreted form of the protein [1]. Consistent with that, the variant fails to support anthrax toxin entry when expressed in a receptor negative host [3]. We analyzed expression of this variant using specific primers designed to amplify a 1358-bp cDNA fragment (Table 1). V3, like V4, showed wide expression (Figure 6). All fetal and adult tissues tested had it, as did all digestive tissues.

**Quantitative Analysis of TEM8 Transcripts by Real-time qPCR**

The major goal of nested PCR was to ascertain the status of full-length splice variants of TEM8. Although this analysis revealed some remarkable differences in variant expression from tissue to tissue, notably for V2, the approach is imprecise for quantitative comparisons. To more precisely ascertain the expression levels, therefore, we carried out real-time quantitative PCR as described in Material and Methods. Consistent with real-time qPCR requirements, the target amplicons for this analysis were much smaller (~231 bp for V1, 281 bp for V2, 234 bp for V3), and 218 bp for V4 – and the primers used for this work are listed in Table 2. Initially we did several pilot studies to set the proper conditions and to verify the authenticity of target amplicons. Results from one such experiment are shown in Figure 7. As is clear, the target fragments amplified. We then used the same conditions to perform qPCR on the tissue cDNA panels.

The results in Figure 8 show that V1 is ubiquitous, consistent with the nested PCR results. However, its levels vary from tissue to tissue, at times markedly. Of the digestive tissues, jejunum had the greatest and colon the least amounts of V1, as suggested by the difference of 5 in their ΔCt values. Among the fetal tissues, kidney showed the greatest and thymus and liver the least of V1. Of the adult tissues, skeletal muscle showed much greater expression than brain. There were differences in V1 expression among the cognate pairs of fetal and adult tissues. Fetal brain, for example, had much greater expression of V1 than did adult brain. But there appeared no definitive pattern; some fetal tissues had more of V1 than their cognate adult tissues, and vice versa. Of all 28 tissues analyzed, adult brain had the lowest expression of V1.

Quantitative analysis results for V2 differed from the nested PCR results in that the qPCR revealed presence of V2 in all tissues, whereas nested PCR did not. V2 showed differences in...
expression from tissue to tissues, but generally these variations were not as remarkable as for V1 (Figure 8).

Judged by the ΔCt values, V3 generally showed the strongest expression of all variants (Figure 9). Placenta and fetal spleen showed the most robust expression, while adult brain, fetal liver, duodenum, and transverse colon expressed the least. Of the paired fetal and adult tissues, V3 appeared at greater levels in fetal brain than adult brain.

The newly identified splice variant that encodes a membrane-bound form of TEM8, V4, was present in all tissues, consistent with the nested PCR results. Of the digestive tissues, ileum showed significantly stronger expression than did esophagus (ΔCt difference ~4). Other differences with ileum varied from 1 to 3. Among the fetal and adult tissues, fetal kidney appeared to have markedly greater level of V4 than did adult kidney. The variant also showed stronger expression in fetal brain than did adult brain.

Figure 3. Analysis of TEM8 V1, V4, and V5 expression in human tissue panels. A & B, Digestive tissue cDNA panel. C & D, Fetal and Adult tissue cDNA panels. The primer pairs used were as for the prostate and fetal brain analysis (Figure 1). There was little or no amplification following 1° PCR, but 2° PCR resulted in definite bands. The number of 2° PCR cycles is shown on the left of each set. The results show that only V4 amplified; V1 and V5 transcripts were absent. Digestive tissue panel: C, cecum; AC, ascending colon; DC, descending colon; TC, transverse colon; D, duodenum; E, esophagus; Ilc, ileocecum; Ile, ileum; J, jejunum; L, liver; R, rectum; St, stomach. Fetal and adult tissue panel: B, brain; H, heart; K, kidney; L, liver; Lu, lung; P, pancreas; Pl, placenta; S, skeletal muscle; Sp, spleen; T, thymus.

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Figure 4. Amplification of V1 with unique V1-specific antisense primers. The sense primer was the same as for primary PCR of other variants, but the antisense primers were specific for the 108-bp segment absent in V4, as well as other variants. Because the amplification was quite robust upon primary PCR, nested PCR was not required. A, Digestive tissue panel: C, cecum; AC, ascending colon; DC, descending colon; TC, transverse colon; D, duodenum; E, esophagus; Ilc, ileocecum; Ile, ileum; J, jejunum; L, liver; R, rectum; St, stomach. B, Fetal and adult tissue panels: B, brain; H, heart; K, kidney; L, liver; Lu, lung; P, pancreas; Pl, placenta; S, skeletal muscle; Sp, spleen; T, thymus.

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We were unable to amplify the specific V5 fragment by nested or real-time PCR in any of the 12 digestive tissues, 8 fetal tissues, and 8 adult tissues.

**Analysis of V4 Function as an Anthrax Toxin Receptor**

The extracellular portion of V4 is identical to those of V1 and V2, both functional anthrax toxin receptors. It has also been shown that TEM8 functions as an anthrax toxin receptor even when its intracellular portion is removed [3]. We therefore reasoned that V4 would also prove a functional toxin receptor, despite the 36-residue in-frame internal deletion in its cytoplasmic domain. To verify, we tested V4 as an anthrax toxin receptor in parallel with the previously reported forms V1 and V2. V4 cDNA was cloned into the mammalian expression vector pIREShyg3 as described in Materials and Methods. V1 and V2 cloning and expression has been described before [3]. The plasmid-encoded receptors were expressed in PR230, a functionally receptor-negative Chinese hamster ovary cell line [3]. As shown in Figure 10, V4 supported anthrax toxin entry as efficiently as V1 and V2, and the toxin IC50 for PR230 expressing any of the three receptors was nearly the same. The host cell alone was resistant. WTB111, the parent of PR230, was included as a control for toxicity. A modified form of anthrax toxin, PA+FP59, was used for these assays. FP59 is a recombinant toxin that carries the PA-binding amino-terminal domain of LF and the ADP-ribosyltransferase domain of *Pseudomonas aeruginosa* exotoxin A [31]. The target for exotoxin A is the eukaryotic elongation factor 2, which the toxin inactivates by ADP-ribosylation, thus arresting protein synthesis and killing the cell.

**Assessment of PA Binding, Processing, and Internalization**

The overall efficiency of intoxication depends on a series of sequential events that underlie the toxin endocytosis. The notable ones are PA binding to receptors, cleavage by furin to release a 20-kDa amino-terminal fragment, PA63 heptamer formation on the cell surface, heptamer internalization, and routing to an acidic compartment, where the heptamer undergoes conformational changes [9,22,25,32,33]. It is not inconceivable that various receptor forms could differ in their capacity to support these events, and yet the toxin IC50 values may be nearly the same. We therefore assessed PA binding, cleavage by furin, and oligomerization in PR230 cells expressing V1, V2, or V4.

The binding assays were done at 4°C and 37°C as described in Materials and Methods. At 4°C membrane trafficking is essentially arrested, and consequently no PA enters cells. But receptor-bound PA63 is still cleaved by furin to release a 20-kDa amino-terminal fragment. The PA63 heptamer that forms on the cell surface, heptamer internalization, and routing to an acidic compartment, where the heptamer undergoes conformational changes [9,22,25,32,33]. It is not inconceivable that various receptor forms could differ in their capacity to support these events, and yet the toxin IC50 values may be nearly the same. We therefore assessed PA binding, cleavage by furin, and oligomerization in PR230 cells expressing V1, V2, or V4.

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4°C, but a strong doublet of oligomers appears when the experiment is done at 37°C. With these cells we have consistently observed that two bands representing PA oligomers appear. Such doublets have been reported before [3,34]. It is also known that aside from the heptamer, PA63 also forms octamers, and such assemblies involve intermediate stages of dimers and tetramers [35]. However, the lower order assemblies do not appear to be SDS-resistant [3,9,22,23,24,33].

The 4°C binding assays revealed that V1, V2, and V4 equally support PA binding and its cleavage by furin, as is clear from the PA83 and PA63 bands in Figure 11B. We next assessed the three receptors’ capacity to support entry of PA. To do that we incubated cells with PA at 37°C, followed by processing as for the 4°C assays. As shown in Figure 11C, all three receptors bound PA83 equally well at this temperature as well. Further, the subsequent cleavage of PA83 to yield PA63 was also equally efficient for all three receptors. And the SDS-resistant oligomer formation was also efficient for all three receptors. The oligomers reflect successful entry of the toxin. Overall these results are fully consistent with the finding that all three receptors make PR230 cells sensitive to PA+FP59 to the same extent. Thus, when expressed in PR230, V4 as an anthrax toxin receptor is as efficient as V1 and V2.

Discussion

Discovered as a gene that showed elevated expression in endothelial cells of colon carcinomas, TEM8 was suggested to be a player in tumor vasculature development [1]. Histological studies that employed in situ mRNA hybridization also showed the gene’s enhanced expression in tumor endothelia [29]. However, the relative expression levels of the three reported TEM8...
transcripts remained undetermined. Therefore it was unclear whether all three were elevated in tumor endothelia or only one of them. Indeed at that time it was unclear whether TEM8 is expressed in normal tissues. Moreover, in situ mRNA hybridizations are not nearly as sensitive as PCR, and thus often fail to detect transcripts expressed at low levels. Variant-specific expression profile of the gene remained undetermined altogether. Once determined as an anthrax toxin receptor [2], it was clear that the

![Figure 7. Quantitative analysis of TEM8 splice variants. A, Real-time PCR protocol. B, Agarose gel analysis of amplicons. β-actin was used as the normalization standard. The expected amplicons were: V1, 231 bp; V2, 281 bp; V3, 234 bp; V4, 218 bp. Several experiments were done to set the appropriate conditions, and to ensure that the real-time PCR worked satisfactorily and that the expected fragments were amplified. Different tissue cDNAs were used for this purpose. St, stomach; R, rectum; L, liver; J, jejunum. We got no reliable results with V5, and therefore excluded it from real-time PCR analysis. doi:10.1371/journal.pone.0043174.g007](image)

![Figure 8. Quantitative analysis of TEM8 splice variants V1 and V2. The analysis was carried out according to the real-time PCR protocol in Figure 7, and as described in Materials and Methods. ΔCt values are the averages of two independent real-time PCRs, calculated by subtracting the Ct values for each variant from the Ct values for β-actin. Digestive tissue panel: C, cecum; AC, ascending colon; DC, descending colon; TC, transverse colon; D, duodenum; E, esophagus; Ilc, ileocecum; Il, ileum; J, jejunum; L, liver; R, rectum; St, stomach. Fetal and adult tissue panels: B, brain; H, heart; K, kidney; L, liver; Lu, lung; P, pancreas; Pl, placenta; S, skeletal muscle; Sp, spleen; T, thymus. doi:10.1371/journal.pone.0043174.g008](image)
gene shows wide expression, given that the toxin can bind and enter a wide variety of cell types [8,36]. It has also been reported that TEM8 shows differential expression in mouse tissues [37].

The major focus of this work was to ascertain an expression profile of all splice variant mRNAs of TEM8 in human tissues. The analysis employed variant-specific primers and nested PCR. To further ensure this analysis is indeed of those TEM8 transcripts that encode the three reported variants, we used primers designed to amplify fragments that carry the complete coding regions. Our findings show that all tissues analyzed have the previously reported

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**Figure 9.** Real-time PCR analysis of TEM8 splice variants V3 and V4. Digestive tissue panel: C, cecum; AC, ascending colon; DC, descending colon; TC, transverse colon; D, duodenum; E, esophagus; Ilc, ileocecum; Ile, ileum; J, jejunum; L, liver; R, rectum; St, stomach. Fetal and adult tissue panels: B, brain; H, heart; K, kidney; L, liver; Lu, lung; P, pancreas; Pl, placenta; S, skeletal muscle; Sp, spleen; T, thymus.

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**Figure 10.** Assessment of TEM8 functionality as an anthrax toxin receptor. A, Sensitivity of cells expressing V1, V2, or V4. The plasmid-encoded receptors were expressed in the receptor-negative host PR230 as described in Materials and Methods, followed by cytotoxicity assays with PA+FP59. B, Toxin concentrations required for 50% cell death (IC₅₀, ng/mL) for PR230 expressing V1, V2, or V4. The means of 2–3 independent experiments are shown in parentheses. The greater sensitivity of the parental cells, WTB111, is probably due to expression of both receptors, especially CMG2/ANTXR2, which has about 10 times greater affinity for anthrax toxin protective antigen than does TEM8 [40,4,33].

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**Table:**

| Cell or Receptor | IC₅₀ (ng/mL) |
|------------------|-------------|
| PR230            | >2000       |
| WTB111           | 2.5, 5, 1.5 (3)* |
| T8V1             | 13, 17 (15) |
| T8V2             | 12, 14 (13) |
| T8V4             | 12, 10, 22 (14) |

*a, mean in parentheses
variants, V1, V2, and V3. Although nested PCR showed selective expression of V2, the real-time PCR showed that the variant is indeed expressed in all tissues analyzed. We therefore conclude that V2 is not selectively expressed, and its absence in some cases was due to failure of nested PCR.

Analysis of V1 transcript initially proved problematic; two separate sets of primers specific for the reported V1 sequence failed to amplify it. We are unable to explain this anomaly, but the same primers amplified two novel splice variants of TEM8. One of these, V4, encodes a 528-residue membrane-bound form of TEM8, and the other, V5, a 358-residue protein (Figures 1, 2 & 3). The V5 protein sequence diverges after residue 314, and it does not have the TEM8 hydrophobic transmembrane helix, nor any other predicted one. We therefore conclude that the putative V5 protein is a secreted form of TEM8.

The newly discovered variant V4 shows broad expression, as evidenced by both nested and real-time PCR (Figures 3 & 9). The V4 sequence differs from V1 in that it lacks a 108-bp cDNA segment, resulting in an in-frame deletion of 36-residues (residues 522–557) in the cytoplasmic domain. We therefore exploited this difference to analyze V1 transcript using primers specific for the 108-bp V1 segment. This approach worked; separate sets of sense and antisense primers specific for this region gave the expected amplicons (Figures 4 & 8).

V1 and V2 are proven anthrax toxin receptors [3,7]. We have shown that V4 too is a functional anthrax toxin receptor. The receptor confers the same degree of sensitivity on PR230 as V1 and V2 (Figure 10). PR230 is a functionally receptor-negative cell line that has been described before [3]. Consistent with these results, anthrax toxin protective antigen efficiently binds V4, is cleaved by furin, is internalized, and forms the SDS-resistant oligomers (Figure 11B, C).

Our results demonstrate that TEM8 expression is not restricted to any particular type of tissue. However, whether the normal and tumor endothelial cells show different expression patterns for all five variants remains to be seen. Another point to note is that we used pooled tissue cDNA panels representing different individuals. Thus, it is impossible to tell whether the gene exhibits individual-specific variations. As an example, we found V5 in one pooled prostate cDNA panel, but not in another (not shown). This suggests the possibility that the gene’s expression profile may even vary from individual to individual. A statistically significant

Figure 11. V1, V2, and V4 capacities for anthrax toxin protective antigen (PA, 83 kDa) binding, processing, and internalization. A, PA binding to normal cells (WTB111). At 4 °C, PA83 binds the receptors and is cleaved by furin to yield PA63. At 37 °C, PA83 not only binds and is cleaved by furin, but enters cells and forms SDS-resistant oligomers. B, PA binding and processing at 4 °C to PR230 expressing V1, V2, or V4. C, PA binding, processing, internalization, and SDS-resistant oligomer formation at 37 °C. Two blots of a typical experiment are shown. doi:10.1371/journal.pone.0043174.g011
number of individual samples would be required to discern such differences, if any.

An intriguing expression pattern of TEM8 in neuroblastoma cell lines has been reported. In these cells the gene is found as fusion transcripts with various segments of novel neuroblastoma gene (NNG1) [38,39]. Oberthuer et al. identified 4 such transcripts, each representing the first 14 exons of TEM8 and various segments of NNG1. Revealed by sequencing, the transcripts encode fusion proteins of 398, 411, 412, and 420 residues, all comprising the first 363 residues of TEM8. However, the functional significance, if any, of these fusion transcripts is unclear.

The putative role of ANTXR1/TEM8 in neo-angiogenesis remains dubious. But in mice the gene is dispensable, as evidenced by normal development of TEM8−/− knockout mice [40]. Indeed, the other anthrax toxin receptor, ANTXR2/CMG2, is also dispensable for mouse development, as shown by studies with CMG2−/− mice [40]. Note that CMG2 has also been reported a player in angiogenesis [6,41]. However, these findings do not necessarily rule out angiogenic role for TEM8 and CMG2 during mouse development. If the two genes are indeed players during mouse developmental angiogenesis, then there exist other players that compensate for the loss of TEM8 and CMG2. In as much as ANTXR1/TEM8 shows enhanced expression in at least some tumor endothelial cells [1,29], the gene does appear to be a player in tumor vasculature development. Evidence also suggests that TEM8 promotes endothelial cell adhesion and migration, which it seems to achieve by interacting with cell matrix proteins [42].

Many recent findings have significantly unraveled the endocytic pathway of TEM8 and the protein’s interactions with other factors. Wei et al. discovered that LRP6, a single-pass integral membrane protein related to the LDL receptor, interacts with TEM8, as well as the other PA receptor, CMG2 [43]. Anthrax toxin endocytosis is largely via the clathrin-mediated pathway, evidently for both TEM8 and CMG2 [3,32,34]. Both receptors cluster in lipid rafts, and TEM8 is palmitoylated and ubiquitinated, modifications that evidently trigger its endocytosis [44]. TEM8 V1 interacts with actin, and this interaction affords more PA binding to this form of ANTXR1/TEM8, although it does not necessarily afford greater sensitivity of cells to anthrax toxin [34,43]. Some other classical players in endocytosis, such as Cbl and β-arrestins 1 & 2, are also important for PA heptamer formation. In contrast, Grb2 and Dab2 appear unimportant [34]. Surprisingly, AP2, the predominant adaptor for clathrin-mediated vesicle budding at the plasma membrane, seems less relevant to PA entry than AP1, which largely functions in vesicle budding from endosomes and trans-Golgi network [34].

At the molecular and cellular level, the native functions of ANTXR1/TEM8 remain obscure. The relative roles of the receptor’s various isoforms are even more elusive. Is it possible, for example, that relative enhancement or attenuation of TEM8 variants defines the gene’s role in tumor angiogenesis. Is the gene important in vascular formation during other disease states, such as wound healing and other forms of tissue repair? Lui et al. have shown that TEM8 and CMG2 are not needed for mouse developmental angiogenesis [40]. However, whether these findings can be extended to human development is unknown. They further reported that CMG2 is the major anthrax toxin receptor and that toxin binding to it is 10 times stronger than to TEM8. Evidence also suggests that TEM8 exists in different forms on the cell surface, and that these forms are influenced by actin and the actin binding protein transgelin [46]. Clearly much more research is needed to precisely elucidate the role of TEM8 and CMG2 at the molecular level in disease state and normal vasculature formation.

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Author Contributions

Conceived and designed the experiments: GJC. Performed the experiments: MV RK. Analyzed the data: SHL GJC. Contributed reagents/materials/analysis tools: SHL. Wrote the paper: MV GJC.

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