Evaluation of Global Differential Gene and Protein Expression in Primary Pterygium: S100A8 and S100A9 as Possible Drivers of a Signaling Network

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Abstract

**Purpose:** Pterygium is a wing shaped fibrovascular growth on the ocular surface, characterized by fibrosis, angiogenesis, extracellular matrix remodeling, and inflammatory infiltrates. Epidemiologic studies have linked pterygium formation to various chronic inflammatory conditions, such as ultraviolet radiation, sawdust exposure, and dry eye disease. The purpose of this study is to identify proteins that are differentially expressed in primary pterygium by using a combination of gene microarray and proteomic platforms.

**Methods:** Paired pterygium and uninvolved conjunctiva tissues of four patients were evaluated for differences in global gene transcript levels using a genechip microarray. Proteins extracted from another four pairs of tissues were quantified by iTRAQ approach. Western blot and immunofluorescent staining on additional patients were used to validate dysregulated protein expression obtained from microarray and proteomics data. In addition, primary conjunctival fibroblasts were treated with recombinant S100A8, S100A9 or both. Transcript level changes of a panel of potential target genes were evaluated by real time-PCR.

**Results:** The following were up-regulated at both protein and transcript levels S100 A8 and A9, aldehyde dehydrogenase 3 family, member1 (ALDH3A1) and vimentin (VIM). Conversely, serpin peptidase inhibitor clade A member 1 (SERPINA1) and transferrin (TF) were down-regulated. Upon adding S100A8, S100A9 or both, the inflammatory chemokine CXCL1, matrix proteins vimentin, biglycan, and gelsolin, as well as annexin-A2, thymosin-β4, chymase (CMA1), member of Ras oncogene family RAB10 and SERPINA1 were found to be up-regulated.

**Conclusions:** We identified 3 up-regulated and 2 down-regulated proteins by using a stringent approach comparing microarray and proteomic data. On stimulating cells with S100AB/9, a repertoire of key genes found to be up-regulated in pterygium tissue, were induced in these cells. S100A8/9 may be an upstream trigger for inflammation and other disease pathways in pterygium.

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Introduction

Pterygium is an ocular surface disease which characterized by abnormal epithelial and fibrovascular proliferation, invasion, and matrix remodeling. The lesion leads to astigmatism, ocular irritation, dryness, and reduced vision once it involves the visual axis. Despite its prevalence, the exact pathogenesis of pterygium remains unknown. Evidence from epidemiology and histopathology studies support the concept that pterygium arises as a result of ultraviolet (UV) radiation-damage [1–4]. UV radiation induces damage in DNA and other cellular substrates [5–7]. In addition, several studies have pointed to an immune response-inflammatory component in pterygium pathology [8–11].

Previously, microarray techniques have been used to compare the genome wide gene expression profiles between primary pterygium and normal conjunctiva tissues [12–14]. Extracellular-matrix-related, proinflammatory, angiogenic, fibrogenic, and oncogenic genes were found to be dysregulated in primary pterygium [13]. About 1724 genes were significantly different between primary and recurrent pterygium by genome-wide microarray scanning [15]. Our group reported 51 genes with at
least 2 fold changes between primary and recurrent pterygium [14]. More recently, proteomic techniques have been applied in pterygium research. Using two-dimensional electrophoresis and mass spectrometry (2D-MS), Kanamoto et al. identified 11 differentially expressed proteins between primary and recurrent pterygium [16]. Using a similar assay technique, peroxiredoxin 2 was found to be detected in primary pterygium tissue [17].

The S100A8 and S100A9 are calcium binding secreted proteins that have been previously identified to be up-regulated in various ocular surface conditions including pterygium, dry eye, meibomian gland dysfunction and corneal neovascularisation [18–21]. In living tissues, these two proteins may exist as homodimers or heterodimers and play a big role in regulation of oxidative damage [22–28] as well as initiation of immune responses [24,29–32]. Because of their possible role as a master regulator of immune response and inflammation, S100A8/S100A9 deserve further study. The relationship of S100A8 and S100A9 and downstream inflammatory molecules has not been evaluated in the context of the ocular surface.

Since there are advantages to an unbiased molecular profiling method in a disease with uncertain etiology, we aimed to identify dysregulated molecules in primary pterygium tissue compared to normal uninvolved conjunctiva by proteomics evaluation and comparing findings with gene microarray analysis. We further investigated the ability of S100A8, S100A9 and S100A8/A9 to regulate the expression of a panel of genes involved in pterygium pathology using primary cell cultures.

Methods and Materials

1. Patients and Ethics Statement

Primary nasal pterygium and uninvolved conjunctival tissues from the superior temporal quadrant of the same eye were collected and stored immediately in RNAAlater RNA Stabilization Reagent (Qiagen) for RNA extraction, or immediately frozen in liquid nitrogen for protein extraction. All procedures adhered to the tenets of the Declaration of Helsinki and were approved by the Institutional Review Board of the Singapore Eye Research Institute. Written informed consent was obtained from all participating patients.

2. Microarray Gene Expression Evaluation

RNA was extracted from 4 pairs of tissues using the following protocol: Tissues were minced into small pieces, and ground into powder using pestle and mortar with liquid nitrogen. The tissue powder was transferred into TRIzol (Invitrogen) and passed through 22G needles several times (Becton Dickinson). Chloroform was added to each homogenized sample, mixed well and centrifuged at 13,000 rpm for 15 min at 4°C. The aqueous layer was transferred to a new tube and an equal volume of 70% ethanol added. Subsequently the samples were purified using RNeasy RNA extraction kit (Qiagen) according to manufacturer’s protocol. RNA concentration and quality were assessed using the ND-1000 spectrophotometer (Thermo Scientific).

The microarray experiment was performed on the Agilent Sureprint G3 human GE 8 × 60 K (ID: G6851-60510) single-colour chip by a commercial service (Molecular Genomics Pte Ltd, Singapore) according to the Agilent protocol. In summary, one-hundred nanograms of total RNA was labeled with Low Input Quick Amp Labeling Kit (One-Colour Microarray), then converted to double-stranded cDNA using oligo-dT primers and transcribed by T7 RNA polymerase to produce cyamine 3-CTP labeled cRNA. Sixty nanograms of labeled cRNA was hybridized onto Agilent SurePrint G3 human GE 8 × 60 K microarray for 17 hrs at 65°C, 10 rpm in Agilent hybridization oven. After hybridization, the microarray slide was washed and scanned on Agilent High Resolution Microarray scanner. Raw signal data was extracted with Agilent Feature Extraction Software (V10.7.1.1), and analyzed by GeneSpring 12.5 (Agilent, CA).

The threshold of raw signals was set to 1.0 and data was transformed to log2 base 2 values. Per chip normalization, performed by baseline transformation to the median of all samples, was performed by percentile shift to 75th percentile. A box plot of normalized data for the 8 samples showed a similar distribution. Subsequent quality control (QC) on the probe set to remove faulty or suspicious probes was done by filtering on flags (probes flagged as ‘not detected’ and ‘compromised’) and filtering on expression (removal of probes with very low raw signal intensity (<20)). This QC processing identified 27714 positive signals. The microarray data reported in this study had been deposited in NCBI Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo) with GEO series record number GSE51995.

3. ITRAQ-LC-MS/MS Proteomics

Protein extraction and quantification. Four pairs of tissues were weighed and washed with phosphate buffered saline (PBS), and then put into 100 μl of lysis buffer (75 mM NaCl, 1 mM EDTA, 0.2% CHAPS, 50 mM Sodium Fluoride, 1 mM sodium orthovanadate and 1× protease inhibitors cocktail prepared in PBS). Ten freeze-thaw cycles involving snap-freezing with liquid nitrogen and thawing on ice water were subsequently performed. Samples were then ground using a mini grinder, and 400 μl of the lysis buffer was added. Ground samples were further homogenized via sonication. Triton-X (0.2%) and IGEPA (0.2%) were added after sonication. Homogenized samples were spun at 16,000 g for 20 minutes at 4°C, the supernatant was concentrated to approximately 50 μL by a 0.5 ml 3 KDa ultra-centrifugal filter (Amicon, USA) and the protein concentration was measured by Bradford assay.

Mass spectrometry and data analysis. Samples were reduced, alkylated, digested and labeled with iTRAQ following the manufacturer’s instruction. The combined samples were cleaned and fractionated with strong cation exchange (SCX) chromatography on a PolySULFOETHYL A column (200 × 4.6-mm, 5 μm, 200 Å; PolyLC Inc., Columbia, MD, USA). Forty-one fractions were first reduced volume and reconstituted in 1% formic acid/5% acetonitrile for desalting by solid phase extraction. C18 purification was performed on Pierce C18 Spin Columns (Thermo Scientific). The desalted samples were lyophilized and reconstituted in 20 μl of 0.1% formic acid/2% acetonitrile. LC-MS/MS analysis was performed on a QStar Elite quadruple time of flight mass spectrometer system (AB Sciex) coupled to a Tempo nanoLC (AB Sciex) with autosampler.

Database search and relative analysis was performed on ProteinPilot™ software (AB Sciex) version 4.0.8085, rev 148085 with the Paragon Algorithm™ version 4.0.0.0, rev 148083 using UniProtKB/Swiss-Prot database release 2013_12 (containing 49,243,530 entries) against the human genome (20274 sequences) [33–34]. Database search was performed by setting cysteine modification by MMTS as a fixed modification [35] and tryspin as the digesting enzyme. None of the special factors was selected. The unused Protocore [conf] in the software was set to 1.3 (95% confidence). Protein level changes were compared between paired samples of pterygium and conjunctiva, and iTRAQ fold change of >1.3 was considered to be up-regulated in pterygium, while a ratio of <0.77 was considered down-regulated. This is based on
fact that the systematic variation of our analytical system has been determined consistently to be within 30% across several studies [36–41].

4. Primary Pterygium and Conjunctival Fibroblast Cell Culture

Primary pterygium and conjunctiva fibroblasts were cultured as previously reported [42–43]. Briefly, pterygium and conjunctiva tissues were immediately submerged in 1 ml of DMEM/F12 (Gibco) medium containing 1× antibiotic-antimycotic (1× A/A) (Invitrogen, USA) and transferred to lab on ice. Culture plates (60 mm) were coated with 1 ml of fetal bovine serum (FBS) (Gibco) for 10 min at 37°C. Each specimen was cut into pieces of approximately 1–2 mm² and placed onto the FBS-coated plates. Sixty microliters of FBS with 1× A/A was dropped over each piece. Explants were incubated overnight at 37°C with 5% CO². The next morning, 120 μl of DMEM/F12 with 10% FBS and 1× A/A were dropped over each explant, and 400 μl more of the same medium was added later in the evening. About 3–5 days later, cells started to migrate from explants and the volume of medium was increased to 2 ml and changed every other day. Fibroblasts were then sub-cultured at day 18–21 using TrypL Express (Life Technologies). Cells from passage 1 to passage 3 were used for experiments. Cell population homogeneity was observed as previously described [44].

5. Western Blot

Western blots were performed as described [44]. In summary, total protein (30 μg) from primary cultured pterygium and conjunctiva fibroblast cells were electrophoresed in 10–12% SDS-polyacrylamide gels, and then transferred to polyvinylidene difluoride (PVDF) membranes at 4°C overnight. The membranes were subsequently incubated with primary antibodies to ALDH3A1 (Abcam, UK, Cat. No. AB76976, 1:1000) and GAPDH (Abcam, UK, AB37168, 1:1000) for 2 hrs at room temperature. Membranes were then incubated with secondary antibodies conjugated to horseradish peroxidase for 1 hr at room temperature. Immunoreactivity was visualized with chemiluminescence substrate (SuperSignal West Pico or West Dura; Pierce).

6. Immunofluorescent Staining

Pterygium and conjunctival tissues were sectioned with a Microm HM550 cryostat (Microm, Walldorf, Germany) at 8 μm thickness. Sections were fixed with 4% paraformaldehyde for 20 min, washed with PBS, permeated in PBS containing 0.15% Triton X-100 for 15 min, blocked with 4% BSA in PBS containing 0.1% Triton X-100 (Sigma) for 1 hr, and then incubated with ALDH3A1 antibody (Abcam, UK, Cat. No. AB76976, 1:200 dilution in blocking solution) at 4°C overnight. After washing with PBS containing 0.1% Tween-20, the sections were incubated with Alexa Fluor 488-conjugated secondary antibody at room temperature for 45 min. Slides were then mounted with UltraCruz Mounting Medium (Santa Cruz). For negative controls, non-immune reactive serum was used in place of the specific primary antibody. Sections were observed and imaged using a Zeiss Axioplan 2 fluorescence microscope (Zeiss, Oberkochen, Germany).

7. S100A8, S100A9 and S100A8/A9 Treatment of Conjunctival Fibroblast Cells

The maximum non-toxic dose of recombinant S100A8, S100A9 and combined S100A8/S100A9 for primary conjunctiva fibroblasts treatment were pre-determined to be 40, 25 and 20 μg/ml respectively. Primary conjunctiva fibroblast cells were seeded (200,000 per well) in a 12-well culture plate (Greiner bio-one) and cultured in serum-free DMEM/F12 medium (Gibco) for 24 hrs. Cells were washed with PBS and treated with S100A8, S100A9 and S100A8/A9 proteins in fresh media for 6 or 24 hrs. Control cells were given fresh medium without recombinant protein.

8. Real Time PCR

RNA was extracted with RNeasy mini kit (Qiagen) according to the manufacturer’s instructions. Reverse transcription was performed using SuperScript III first strand synthesis supermix (Invitrogen) according to manufacturer’s protocol. Real time relative quantitative polymerase chain reaction (qPCR) was performed with SYBR Green qPCR Mastermix (Roche). Primers used for real time PCR are listed in Table 1. GAPDH was used as the endogenous control gene. PCR cycles were performed on the Roche LightCycler 480 with the following cycle conditions: 10 min at 95°C, 45 cycles of 10 s at 95°C, 15 s at 60°C and 20 s at 72°C, followed by melting curve analysis. The delta-delta Ct method was used for data analysis and experimental reproducibility was confirmed using 3 biological replicates. Statistical significance was evaluated at p<0.05. Pooled standard deviations of the fold changes were calculated for 3 independent experiments which used cells cultured from 3 different patients.

Results

1. A Panel of Genes Found to be Dysregulated at both Transcript and Protein Level

Fold change analysis was performed using the microarray data from the 4 paired pterygium and conjunctiva tissues, a list of differentially expressed genes (fold change ≥2, p<0.05) was delineated (GEO series record number GSE51995). Some of the significantly dysregulated transcripts in pterygium are shown in Table S1. Other up-regulated transcripts (Table S1) include chemokine ligand CXCL1, biglycan (BGN), and gelsolin (GSN), annexin-A2 (ANXA2), thymosin-β4 (TMSB4X) and member of Ras oncogene family (RAB10), whereas the transcripts down-regulated in pterygium were those of Serpin peptidase inhibitor, clade A member 1 (SERPINA1) and chymase (CMA1). At least 3 out of 4 pairs of the transcripts were regulated in the direction shown although p<0.05 was only for GSN and SERPINA1. We confirmed that in the dataset (http://www.ncbi.nlm.nih.gov/geo/, series accession GSE2513) from our previous study [14] these transcripts were also dysregulated in the direction shown (Table S1). In the previous dataset, the p-values were <0.05 for 3 of these genes: BGN, GSN and TMSB4X, and these were 0.003, 0.033 and 0.014 respectively. In the current dataset, however, none of the evaluated genes, passed the Benjamini-Hochberg False discovery rate correction (p<0.05) when the paired student’s T-test was used, suggesting heterogeneity amongst different biological samples. The iTRAQ LC-MS/MS experimental design is shown in Figure S1. The complete raw data of the iTRAQ analyses are shown in Table S2 and S3 respectively. Only proteins identified with 95% confidence were included. After relative quantification analysis, all the proteins found to be significantly different between pterygium and conjunctiva at least in 3 pairs of tissues are shown in Table S4. Inter-patient heterogeneity of protein expression was also observed across many proteins. When the list of dysregulated proteins in proteomics was compared against that from the microarray data, a panel of overlapping genes was generated and summarized in Table 2. Although not all functionally important
molecules are dysregulated at both the transcript and protein levels, we decided to report and emphasize these for simplicity.

Aldehyde dehydrogenase 3 family, member 1 (ALDH3A1), vimentin (VIM), S100A8 and S100A9 were found to be up-regulated in primary pterygium compared to healthy conjunctiva control. SERPINA1 and transferrin (TF) were down-regulated at both protein and transcript levels in pterygium (Table 2).

2. Validation of Over-expression of ALDH3A1 by Western Blot and Immunofluorescent Staining

On immunofluorescent staining in pterygium and conjunctiva tissues (Figure 1A, left and middle columns), ALDH3A1 was found to be present in both conjunctival and pterygium epithelium, with a stronger expression in pterygium than in conjunctiva. The localization of the ALDH3A1 was in the cytoplasm and cell periphery of the epithelium, and present in all layers of the epithelium. Some fibroblast staining was visualized in the stroma of both pterygium and conjunctiva but comparison of staining intensity was difficult (data not shown). No fluorescent signals were detected in the negative controls (Figure 1A, right column). Levels of ALDH3A1 in primary cultured pterygium and conjunctiva fibroblast cells and epithelial cells were analyzed by western blot. Results showed that ALDH3A1 was more highly expressed in pterygium fibroblast cells than conjunctiva fibroblast cells (Figure 1B). However, we were not able to show similar up-regulation in cultured epithelial cells (data not shown).

3. S100A8, S100A9 and S100A8/S100A9 Up-regulate Disease-inducing Genes

Previously, our group has reported that S100A8 and S100A9 were localized in the superficial layer of both pterygium and normal conjunctiva epithelium, with higher levels in pterygium than uninvolved conjunctiva [21]. To test our hypothesis that S100A8/S100A9 are master regulator of many other dysregulated

Table 1. Primers used in Real Time PCR.

| Gene Symbol | Gene Name                          | NCBI accession number | Primer Sequence       |
|-------------|------------------------------------|-----------------------|-----------------------|
| TMSB4X      | Thymosin beta 4, X-linked          | NM_021109.3           | F: AGACAGACTGCTCTCGTA R: CTGCTGTCCTCCTCCGT |
| CKL1        | Chemokine(C-X-C motif) ligand 1    | NM_001511.3           | F: CCGACCCGGCGTCTCTC R: CCTCTCCTCCTCCTCCGT |
| ANXA2       | Annexin A2                         | NM_001002857.1       | F: CCTACACCAAGGTGCAT R: TCAACGAGTGAAGCCCTC |
| SERPINA1    | Serpin peptidase inhibitor, clade A member 1 | NM_000295.4 | F: GTCAGGACACCCAGGAGA R: TATTCATACGACGACCA |
| CMA1        | Chymase 1                           | NM_001836.3           | F: CTGGAGAGAACAGGTTGT R: TTTTGTCTCCTGGATG |
| GSN         | Gelsolin                            | NM_000177.4           | F: TGTTGGCAGACACTTCTCC R: CTTCACAGGATTTGCT |
| BGN         | Biglycan                            | NM_001711.4           | F: CCGACCCGGCGTCTCCTTC R: GGGAGCGGGAGAGGTTTC |
| VIM         | Vimentin                            | NM_003380.3           | F: CCAACCTTCCTCCTCCTAC R: GTGATGCTGAGGTTGATG |
| RAB10       | Ras-related protein Rab-10         | NM_016131.4           | F: CACTCCTTCTGCTGCTCC R: GAGGACATCTGGGGACATTT |
| GADPH       | Glyceraldehyde 3-phosphate dehydrogenase | NC_000012.11 | F: CCCACACACATGACTTCACC R: CTTCTCCTGCCGCTTGG |

Table 2. Genes dysregulated in both transcriptomics and proteomics.

| Gene Symbol | Description                              | Microarray | Proteomics |
|-------------|------------------------------------------|------------|------------|
|             | p F p < 0.05                             | p < 0.05   | p < 0.05   |
| ALDH3A1     | aldehyde dehydrogenase 3 family, member 1 | 0.01 2.9   | 8.24       |
| VIM         | vimentin                                 | 0.45 1.1   | 4.64       |
| S100A8      | S100 calcium binding protein A8          | 0.47 1.2   | 3.98       |
| S100A9      | S100 calcium binding protein A9          | 0.64 1.2   | 5.93       |
| SERPINA1    | serpin peptidase inhibitor, clade A member 1 | 0.01 3.0   | 0.29       |
| TF          | transferrin                              | 0.24 1.5   | 0.48       |

*averaging obtained in duplicated analyses of 4 pairs of tissues.

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genes, recombinant human S100A8, S100A9 or both were used to treat conjunctiva fibroblast cells. A panel of dysregulated genes in pterygium (mentioned above) was selected as possible downstream effectors of the S100 proteins. The transcript level changes of these genes in cultured conjunctival fibroblasts, induced by addition of recombinant S100A8, S100A9 or both were evaluated by qPCR.

Inflammation-related genes TMSB4X, CXCL1 and ANXA2, were significantly increased upon addition of S100A8, S100A9 singly or in combination (Figure 2). TMSB4X transcript was significantly up-regulated by any of these three interventions at 6 hrs and continued to increase 24 hrs after treatment, while a significant up-regulation of TMSB4X by S100A9 was observed only at 24 hrs after treatment (Figure 2A–C). CXCL1 showed a marked fold change (of more than 700 folds) upon S100A8 and S100A9 treatment at 6 hrs, but this was less marked at 24 hrs. CXCL1 expression was not responsive to S100A8/A9 combination at 6 hrs but showed an increase at 24 hrs (Figure 2D–F). ANXA2 transcript increased significantly at 6 hrs after any of these interventions, and the increased expression persisted through 24 hrs, but this transcript was relatively more sensitive to the addition of the S100A8/A9 combination than single proteins.

Extracellular matrix genes GSN, BGN and VIM increased upon S100A8, S100A9 or combined treatment in conjunctiva fibroblast cells (Figure 3), but the magnitude of change (1.5 to 20 fold) was generally not as large as the transcripts for inflammation-related genes.

The expression levels of enzyme-coding genes SERPINA1 and CMA1 were up-regulated by S100A9 (Figure 4B and 4E respectively) and combination of S100A8 and S100A9 (Figure 4C and 4F), but not by S100A8 (Figure 4A and 4D). Intracellular trafficking gene RAB10 was up-regulated by S100A8 and S100A9 at 6 hrs (Figure 4G and 4H), while the increase persisted 6 hrs through 24 hrs for the combination treatment (Figure 4I).

Discussion

In this study we reported a list of differentially expressed genes by genome-wide microarray analysis using pairs of primary pterygium and autologous conjunctiva tissues, and identified 10 dysregulated proteins by iTRAQ-LC MS/MS proteomics. Five genes were found to be dysregulated in pterygium tissue in the same directions using both approaches. In addition, we found that S100A8, S100A9 and combination could up-regulate genes related to inflammation, extracellular matrix and matrix enzymes in cultured primary human conjunctiva fibroblast cells.

A microarray study conducted by John-Aryankalayi et al compared two primary and one recurrent pterygium with autologous conjunctiva tissues [13]. Previously, we performed a
microarray study with 4 pooled samples of conjunctiva tissues and 10 un-pooled samples of primary pterygium tissues [14]. Different with all previous studies, our current microarray utilized 4 pairs of primary pterygium and conjunctiva tissues from the same eyes. The differentially regulated genes that were further evaluated in the current study were also found to be dysregulated in the same direction in the previous dataset uploaded to GEO (http://www.ncbi.nlm.nih.gov/geo/, series accession GSE2513) [14].

To date, there have been only two published studies which identified dysregulated proteins in pterygium using proteomics [16–17]. Both published method utilized 2-D PAGE for protein separation and detection. While 2-D PAGE has powerful resolution, it is generally incapable of resolving proteins of extreme PI and molecular weight. Furthermore, sensitivity of the analytical system is largely dependent on the ability to visual protein spots on 2-D gels. In contrast, this study used a more modern technology involving isobaric tagging for relative and absolute quantification (iTRAQ). Coupled to LC/MS/MS, this method is capable of analyzing up to 8 samples in a single analysis thus providing a more reliable comparative analysis by eliminating the reproducibility issue encountered in 2-D PAGE. One of the 2-D PAGE based studies identified 11 proteins differentially expressed between recurrent and primary pterygium [16], but did not examine differences between primary pterygium and normal control tissue. The other study reported the over-expression of only 1 protein, peroxiredoxin 2, in primary pterygium [17].

The dysregulated genes identified were known to have biological functions relevant to inflammation or cell stress.
ALDH3A1 catalyzes the oxidation of a wide variety of endogenous and exogenous aldehydes to the corresponding carboxylic acid, and is associated with cell proliferation and drug metabolism [45]. This enzyme is a corneal crystallin and plays a critical role in protecting the cornea from UV-induced oxidative stress by directly absorbing UV-radiation [46–48], scavenging free radicals or producing NADPH [45]. A previous study had found ALDH3A1 to be among the 25 most abundant transcripts in an un-normalized un-amplified pterygium cDNA library [13]. Over-expression of ALDH3A1 has also previously been found in tumor cells and increased ALDH3A1 expression can lead to cell proliferation [45]. The dysregulation of ALDH3A1 in pterygium tissue may be attributed to abnormal control of oxidative stress or proliferation. ALDH3A1 in cultured pterygium and conjunctiva epithelial cells showed no significant difference, most likely is related to differences in ALDH3A1 in cultured and native epithelium, the latter having matrix and fibroblast interactions.

Vimentin is a type III intermediate filament, a major cytoskeletal component of mesenchymal cells, and present in cells at the focal areas of a surgical pterygium tissue [49]. It has been reported to be involved in corneal wound healing and fibrosis [50], which are processes that may be implicated in pterygium pathology. In fact, abnormal vimentin expressing cells were reported in pterygium tissue [51].
The S100A proteins are a family of small calcium-binding dimeric proteins with pleiotropic functions, and the S100A8/A9 heterodimer play an important role in inflammatory regulation [24,52]. S100A proteins are abundantly expressed on the ocular surface [18] and have been suggested to be involved in the pathogenesis of various ocular surface diseases, such as dry eye [20], corneal angiogenesis [19] and pterygium [20–21].

SERPINA1, also known as α1-proteinase inhibitor or α1-antitrypsin, is a protease inhibitor. Decreased levels of SERPINA1 at both transcript and protein levels were noticed in circulating hematopoietic cells, induced by granulocyte colony-stimulating factor or chemotherapy [53]. TF is an iron-binding blood plasma glycoprotein that regulates the level of free iron in biological fluids [54].

Pterygium pathogenesis is associated with increased UV exposure [1–4] and S100A protein is known to be induced by UV radiation [55–56]. Functions of the genes induced by addition of S100A proteins may suggest that they propagate the pterygium disease. It is possible that the S100A proteins may serve as a master regulator to activate this network of genes (Figure 5). In our proposed hypothesis, SERPINA1 and ALDH3A1 may also be dysregulated under the influence of UV radiation, but may not be regulated through S100A8/A9. Nevertheless, Figure 5 shows how these pathways may be linked. For example, the ANXA2 is involved in resistance to UV-induced cell death [57]. Although we

Figure 4. Real-time PCR quantification of SERPINA1, CMA1 and RAB10 transcripts fold change in human conjunctiva fibroblast after S100A proteins treatment. Please refer to Figure 2 for details.
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used fibroblasts in the study, other cell types such as dendritic cells, may have important responses to S100A proteins in pterygium and should be studied further.

A strength of the current study is that comparisons were done using un-pooled samples from individuals, avoiding noise related to inter-individual differences. As the amount of RNA extracted from small surgical samples of pterygium and conjunctiva was limited, earlier genome-wide gene expression studies required pooling of samples from different patients [12,14–15]. Another shortcoming of sample pooling is that some of the differentially expressed genes may be obscured.

A major limitation of the protein studies is that we did not do a more definitive evaluation of small molecules like cytokines using a technique such as bead-based indirect immunofluorescent assay, as such we may not be able to verify the expression of some chemokines and cytokines. Another limitation of this study is that pterygium and conjunctiva tissues have different epithelium/stroma ratio which definitely introduce variations. This should be considered when interpreting the list of dysregulated molecules.

In conclusion, several genes were dysregulated at the transcript and protein levels in pterygium. The dysregulated molecules were consistent with previous experiments reported. The S100A8 and S100A9 have been found to be up-regulated in pterygium tissue. These may control the progression of pterygium via a network of other molecules which have also been found to be dysregulated in pterygium.

**Figure S1** Four pairs of pterygium and conjunctiva samples were used for two 8-plex iTRAQ LC-MS/MS. E1 and E2 represented two 8-plex experiments. P: pterygium; C: conjunctiva. i and ii: each pair of samples was duplicated.

**Table S1** Dysregulated gene identified by microarray.

**Table S2** iTRAQ-LC MS/MS Results from Plate E1.

**Table S3** iTRAQ-LC MS/MS Results from Plate E2.

**Table S4** Dysregulated proteins identified by iTRAQ-LC MS/MS.

**Author Contributions** Conceived and designed the experiments: AH LT YPL. Performed the experiments: AH WWL KPL SCJK MQT. Analyzed the data: AH WWL KPL YPL LT. Contributed reagents/materials/analysis tools: AH LT YPL. Wrote the paper: AH WWL YPL LT.
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