Prostaglandin D synthase isoforms from cerebrospinal fluid vary with brain pathology

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Abstract. Glutathione independent prostaglandin D synthase (Swissprot P41222, PTGDS) has been identified in human cerebrospinal fluid and some changes in PTGDS in relation to disease have been reported. However, little is known of the extent that PTGDS isoforms fluctuate across a large range of congenital and acquired diseases. The purpose of this study was to examine changes in PTGDS isoforms in such a population. Spinal fluid from 22 healthy study participants (normal controls) with no classifiable neurological or psychiatric diagnosis was obtained and PTGDS isoforms were identified by specific immunostaining and mass spectrometry after denaturing 2D gel electrophoresis. The PTGDS isoforms in controls consisted of five charge isoforms that were always present and a small number of occasional, low abundance isoforms. A qualitative survey of 98 different people with a wide range of congenital and acquired diseases revealed striking changes. Loss of the control isoforms occurred in congenital malformations of the nervous system. Gain of additional isoforms occurred in some degenerative, most demyelinating and vasculitic diseases, as well as in Creutzfeldt-Jakob disease. A retrospective analysis of published data that quantified relative amounts of PTGDS in multiple sclerosis, schizophrenia and Parkinson’s disease compared to controls revealed significant dysregulation. It is concluded that qualitative and quantitative fluctuations of cerebrospinal fluid PTGDS isoforms reflect both major and subtle brain pathophysiology.

1. Introduction

The polyunsaturated fatty acid arachidonic acid (20:4, n-6) is oxygenated and reduced in the two-step, cyclooxygenase reaction to generate prostaglandin G\( _2 \) (PGG\(_2 \)) or prostaglandin H\( _2 \) (PGH\(_2 \)), which in turn is converted to generate signaling molecules by an array of PGH\(_2 \) metabolizing enzymes. One such enzyme catalyzes the conversion of PGH\(_2 \) to prostaglandin D\( _2 \) (PGD\(_2 \)), a prostaglandin involved in pain, sleep, and smooth muscle contraction/relaxation, and a potent inhibitor of platelet aggregation [16,35]. This enzyme is glutathione independent prostaglandin D synthase (Swissprot P41222, PTGDS, also known as PGDS). PTGDS has been identified in cerebrospinal fluid (CSF) [9, 11,20,25,34,39] and its isoforms have been reported as altered in neurological diseases [11,17,18,24,26–28].

The cause of these changes is not known, but altered PTGDS may indicate the source of cell pathophysiology: PTGDS is normally produced by meningeal and glial cells [6,32,36]. Consequences of altered PTGDS are also not known, but can be predicted to include variations in prostaglandins that may impact brain function because of their many cellular activities.

Our aim was to examine PTGDS and its many isoforms [11] in human cerebrospinal fluid (CSF) and determine, qualitatively, the range of variation in different diseases. To accomplish this, we surveyed PTGDS in CSF from patients with various congenital and acquired neurological diseases as well as from well-defined controls. Our prediction was that disorders
that affected the cellular source of PTGDS or its downstream prostaglandin products would correlate with discernable deviations of this protein. We defined a profile of PTGDS isoforms from controls that was invariant and, when compared to this baseline, we found gains or losses of PTGDS isoforms in many diseases of the nervous system.

2. Methods

2.1. CSF

Samples from a total of 120 different persons were obtained from four sources, based on nationally accepted clinical diagnostic criteria: A) samples from 42 persons with different congenital or acquired diseases were obtained from the CSF bank at Duke University department of neurology [1,2], under the direction of R.E.A.: alobar holoprosencephaly, schizencephaly with ventriculoperitoneal shunt, Arnold-Chiari malformation, myelomeningocele, hydranencephaly, neurometabolic disorder (negative biochemistry), immunodeficiency and metabolic disorder, glutaric aciduria Type 1, propionic academia, methyl malonic aciduria, HMG CoA lyase deficiency, Dandy Walker syndrome, prematurity with intraventricular hemorrhage, Lowe’s syndrome, Pelizaeus-Merzbacher disease, Wilson’s disease, Leigh’s disease, Moyamoya with Down’s syndrome, congenital myopathy, tuberous sclerosis, Rett’s syndrome, Kawasaki syndrome, faccio-scapulo-humeral dystrophy, Down’s syndrome with argininosuccinic aciduria, X-linked agammaglobulinemia, systemic lupus cerebritis, Niemann-Pick disease with dementia, schizophrenia (X2), Churg-Strauss syndrome, Wegener’s granulomatosis, neurosyphilis, Alzheimer’s disease, subdural hematoma, subarachnoid hemorrhage, normal pressure hydrocephalus, amyotrophic lateral sclerosis with dementia, multiple sclerosis, cerebral sarcoidosis, subacute cerebellar degeneration, B12 deficiency, and pneumococcal menigitis; B) three patients with MPTP-induced parkinsonism, from R.S. Burns, Springfield, Illinois; C) samples from 10 patients with suspected Creutzfeldt-Jakob disease referred to MGH, confirmed with a positive 14-3-3-γ test [22], and 7 of these were pathologically verified at later autopsy; D) samples of 24 study participants with Alzheimer’s disease, 9 with Parkinson’s disease, 8 with multiple sclerosis, 2 with schizophrenia, and 22 with no classifiable psychiatric or neurological disorder after extensive evaluation (“normal” controls) were collected by RPC and MGH as part of an ongoing ethically approved study at HMRI. All samples were collected by routine lumbar puncture, and stored frozen in aliquots at −70°C before assay.

2.2. Protein assay

Concentrations of protein in CSF were determined using a micro titer plate-based Coomassie protein assay using human serum albumin (Sigma), 0–100 µg/ml as a standard [40]. Briefly, 5 µl of diluted CSF (10 X and 100 X) was added to a 96 well microtiter plate in triplicate. Coomassie dye (BioRad, Hercules, CA) was diluted (5 X) and 200 µl added to each well. After 5 min, the OD at 595 nm was obtained using a microplate reader (Molecular Devices, Sunnyvale, CA) and protein concentrations in each sample were computed using Softmax software from Molecular Devices.

2.3. Denaturing 2DE

CSF proteins were precipitated by ethanol (9 mL: 1 mL ethanol:CSF) and re-solubilized in a solution of β2-mercaptoethanol, Triton X100, 8 M urea, CHAPS and carrier ampholyte solution of pH 3–10, and heated to 95°C for five minutes. Based on the total protein assay, 30 µg of volume-reduced, denatured CSF protein per sample was applied to 16 cm, carrier ampholyte gels for isoelectric focusing [12], followed by SDS PAGE on 14%T gels (16 cm by 20 cm), with detection by silver diamine staining [19]. Charge and mass standards are determined for the x and y axes respectively by pl calibration from published maps and co-migration of molecular weight markers [38]. Approximately 1250 protein species were detected per sample. Immunodetection of PTGDS was achieved on identically run 2DE gels that were blotted to PVDF membrane for immunostaining. A rabbit polyclonal primary antibody specific for PTGDS [11] was employed, followed by secondary horseradish peroxidase conjugated anti-rabbit anti-IgG reagent (Sigma), and these were visualized on the blot either with chromogen (chloronaphthol or diaminobenzidine) or by ECL chemiluminescence (Amersham Bioscience) and Kodak XAR5 film exposure. The constitutive isoforms of PTGDS from the control samples were defined and qualitative changes in samples from different congenital and acquired diseases were evaluated. Changes were noted if either a loss or gain of immunoreactive proteins were observed for the samples from disease-affected partici-
Fig. 1. (A) is a western blot of CSF after 2D gel separation as described in the Methods, from a study participant control with no classifiable psychiatric or neurologic disorder. The immunostain was specific for PTGDS, based on a synthetic peptide-derived antibody against the PTGDS sequence EAQVSVQPNFQQD. Molecular weight markers and pI distribution are noted on the y and x axis respectively. The invariant isoforms #2-6 are indicated in continuous lined rectangles and the variant, faint isoform #1 and variant “at-the-edge” basic #7 are indicated at the sides, in broken lined rectangles. (B) is the image of the silver stained total CSF protein from the same sample. PTGDS isoforms are identified in the same manner as in 1A.

pants when compared with spot patterns and intensities from control gels.

Primary structural identification of proteins was completed using a scaled-up version of the 2D gel protocol described in the previous section. Briefly, 2–4 fold more total protein was separated on a 2D gel, directly stained with Sypro Ruby, and visualized using the Dark Reader (Claire Biochemicals) blue light box. Spots for proteins of interest were excised with a scalpel and subsequently prepared for mass spectrometry analysis. An aqueous solvent-based methodology for production and subsequent extraction of trypsin-derived peptides from gel plugs, was used [7]. Tryptic peptides were analyzed with an ion trap mass spectrometer (LCQ Deca, Thermo Electron) equipped with a nanoflow solvent delivery system and polymeric reversed-phase peptide trap (PepFinder Kit, Thermo Electron) to identify PTGDS isoforms. Peptides were identified from the mass spectra using Bioworks 3.1 (Thermo Electron), which correlates the experimental tandem mass spectra against theoretical tandem mass spectra from human protein via the Sequest algorithm [37]. Protein information was derived from the SWISS Prot database (Release 39) obtained from the Swiss Institute of Bioinformatics and contained 7459 human protein entries. Confidence in correct identifications by Bioworks was based on several factors: (repeated MS/MS analysis of the sample, repeated MS/MS of peptides within one run of the sample, MS/MS spectra quality, and a Bioworks Xcor and δCn greater than fairly stringent criteria [33]).

3. Results

3.1. PTGDS identification

Figure 1 illustrates the identification of the typical isoforms of PTGDS from a control CSF sample after immuno (1A) or silver (1B) staining. There were five isoforms that were invariably present in all controls, labeled #’s 2–6. There was an additional fainter blob, #1, that was not always present, presumably related to its low abundance, and a more basic isoform, #7, at the basic edge of the pH range for IEF that was sometimes at the end. Another observation was the appearance in 3 of the 21 controls of very faint immunoreactive isoforms at a somewhat higher molecular weight (28–34 kD) in the basic region (pI 5.8–7.5), as indicated in Fig. 2. The same results were obtained with immobilized pH gradient IEF (data not shown). There was no observable qualitative change of these PTGDS immunoreactive spots with loading up to 200 µg total protein.

Isoform #’s 2–6 were consistently identified as PTGDS after excision, trypsin digestion, and LC/MS, performed as described in the Methods. Two peptides (APEAQSVSQPNFQQDK and AQGFTEDTIVFLPQTDK, both sequenced as doubly charged ions) were consistently derived from all five invariant normal isoforms. An example of the MS$^2$ spectrum from isoform 2 is illustrated in Fig. 3. Notably, the only other proteins identified from these excised blobs were occasional immunoglobulin fragments, proteins known to have similar mass and charge, that were not immunoreactive with the antibody.
PTGDS in controls and patients with diseases of the nervous system

Fig. 2. This figure depicts the position of all PTGDS isoforms in CSF from control and different disease samples. The five black filled regions are invariant, while the gray filled regions are variable in controls, as described in the text. The higher molecular weight, white-filled regions are additional EAQSVQPNFQQD immunoreactive isoforms that appear in select disease states, as described in the text. The numbers refer to pI/MW(kD).

3.2. Qualitative assessment of PTGDS

In 22 healthy controls there were no qualitative differences for the 5 PTGDS isoforms across an age range of 19–97 yrs or between the genders, 10 F:12 M, as reported in Table 1. Two isoforms that were occasionally present were the acidic #1 that was often too faint to be confidently identified and the #7 that was at the variable basic edge of the isoelectric focusing. No sexual dimorphism was seen (data not shown).

In the disease groups, 28/98 had no gain or loss of the invariant 5 isoforms, as itemized in Table 1. However, there were dramatic changes in CSF from the other 70 persons (Table 1) and these are summarized in Fig. 2. All isoforms # 2–6 (invariably present in controls) were absent from the CSF in hydranencephaly and one or more invariant forms were lost in schizencephaly, Arnold-Chiari malformation, myelomeningocele, Dandy-Walker syndrome, Wilson’s disease, Leigh’s disease and moyamoya disease with Down’s syndrome. There were gains in additional PTGDS-immunoreactive spots in 64/98 samples studied, itemized in Table 1. An example of additional immunoreactive PTGDS species in the CSF from a participant with multiple sclerosis is illustrated in Fig. 4, compared to the same region 2D gel from a typical healthy control. In this analysis, many additional immunoreactive spots are visible in a “satellite” distribution around the normal isoforms.

4. Discussion

Identification of PTGDS in these experiments has relied on two independent techniques. The synthetic peptide-derived polyclonal antibody used throughout the study was highly specific, with no cross-reactivity to any other protein on the gels of the control samples, as seen in the western blot in Fig. 1(A) compared with the silver stained total proteins in Fig. 1(B). The characterization of isoform #’s 2–6 by mass spectrometry from excised 2D gel blobs was of high confidence, as described in the methods, and as illustrated in Fig. 3.

This study is by no means comprehensive, but provides clear evidence of altered PTGDS isoforms in select disease types. Congenital malformations, two out of eleven people with mutations and one with severe hemorrhage led to loss of normal PTGDS isoforms. Additional PTGDS-immunoreactive isoforms appeared in most cases of multiple sclerosis, Creutzfeldt-Jakob disease and many neurodegenerative disorders such as Alzheimer’s and Parkinson’s diseases. It will be important to define the biochemical nature of the altered PTGDS-immunoreactive additional isoforms.

We know from our work and others that the variants of PTGDS are heavily glycosylated and all shift to a number of lower molecular weight basic forms after N- and O-linked deglycosylation [10,11,21]. It is also known that altered glycosyl transferases can lead
Fig. 3. The MS² spectrum, with annotation, is presented for the doubly charged peptide APEAQVSVQPNFQQDK that was eluted from a gel plug of PTGDS spot #3, and processed as described in the methods. The y⁵−¹⁴ ions are indicated, as are six of the b ions.

Fig. 4. The region of a western blot from a 2D gel of CSF from a control (no classifiable neurological or psychiatric disease) and a participant with multiple sclerosis, probed with anti-PTGDS as described in Methods. The additional immunoreactive isoforms around and above the normal PTGDS in the CSF from the MS participant are clearly visible. This result was typical for CSF from all CJD and MS participants, most cases of vasculitis and more than half of the neurodegenerative cases (see Table 1).

to variable glycoforms of PTGDS [8,23,31]. While the isoforms we observe may be differentially glycosylated, several other possibilities exist to explain the changes in IEF and gel mobility seen for normal and disease-derived isoforms. Genetic variants of PTGDS have been reported from brain [3,29,30] and these may contribute to altered isoforms. Another possibility is that PTGDS is a small molecule carrier [5] and, although our 2D gels are denaturing, it is still possible that varied carrier states of retinoic acid, retinaldehyde, bilirubin or biliverdin could exist and lead to isoform variability. Casein kinase II phosphorylation of PT-GDS has been reported [3,4] and thus isoforms may be altered by phosphorylation. In fact, it is not known whether any of the many possible post-translational modifications may be responsible for altered PTGDS isoforms in addition to the glycosylation. Further structural identification of the additional immunoreactive spots found in the 64 samples from people with different diseases will be required to more substantially test whether they are PTGDS variants and, if so, to characterize the modifications.

Qualitative assessment of the 22 controls revealed the PTGDS isoforms to be remarkably consistent. This
Table 1
Clinical data of study participants and their PTGDS correlates

| Etiology*                        | # of cases and gender ratio (F:M) | Age ranges | Normal PTGDS | Loss of Normal PTGDS | Gain of additional PTGDS immuno-reactive isoforms |
|----------------------------------|----------------------------------|------------|--------------|----------------------|--------------------------------------------------|
| Congenital malformation          | 7 (4:3)                          | 1 d–27 y   | 0            | 6 lose 1             | 6                                                |
| Inborn error of metabolism and other mutations | 15 (10:5) | 1 m–38 y   | 12           | 1 loses all          | 2 lose 1                                          |
| Infectious                       | 12 (6:6)                         | 30–76 y    | 2            | 0                    | 10                                               |
| Inflammatory                     | 14 (10:4)                        | 4–65 y     | 1            | 0                    | 13                                               |
| Degenerative                     | 36 (18:18)                       | 37–91 y    | 6            | 0                    | 30                                               |
| Hemorrhagic                      | 3 (1:2)                          | 16 d–55 y  | 1            | 1                    | 1                                                |
| Metabolic                        | 2 (1:1)                          | 7–8 y      | 2            | 0                    | 0                                                |
| Toxic                            | 3 (1:2)                          | 28–33 y    | 3            | 0                    | 0                                                |
| Nutritional                      | 1 (1:0)                          | 67 y       | 1            | 0                    | 0                                                |
| Unclassified                     | 5 (1:4)                          | 24–79 y    | 0            | 0                    | 5                                                |
| Controls (no classifiable disease)| 22 (10:12)                      | 19–97 y    | 22           | 0                    | 0                                                |

*Congenital = alobar holoprosencephaly, schizencephaly with ventriculoperitoneal shunt, Arnold-Chiari malformation, myelomeningocele, hydranencephaly, Dandy Walker syndrome, Lowe’s syndrome; Inborn errors of metabolism/mutations = glutaric aciduria Type 1, propionic academia, methyl malonic aciduria, HMG CoA lyase deficiency, Pelizaeus-Merzbacher disease, Wilson’s disease, Leigh’s disease, congenital myopathy, tuberous sclerosis, Rett’s syndrome, fascio-scapulo-humeral dystrophy, Down’s syndrome with argininosuccinic aciduria, X-linked agammaglobulinemia, Moyamoya with Down’s syndrome, Niemann-Pick disease with dementia; Infectious = neurosyphilis, pneumococcal meningitis, Creutzfeldt-Jakob disease (10 cases); Inflammatory = Kawasaki syndrome, systemic lupus cerebritis, Churg-Strauss syndrome, Wegener’s granulomatosis, multiple sclerosis; Degenerative = Alzheimer’s (25 cases), Parkinson’s disease (9 cases), amyotrophic lateral sclerosis with dementia, subacute cerebellar degeneration; Hemorrhagic = prematurity with intraventricular hemorrhage, subdural hemATOMAT, subarachnoid hemorrhage; Metabolic = neurometabolic disorder (negative biochemistry), immunodeficiency and metabolic disorder; Toxic = MPTP-induced parkinsonism (3 cases); Nutritional = B12 deficiency; Unclassified = schizophrenia (4 cases), normal pressure hydrocephalus.

Table 2
Relative changes in PTGDS protein quantity in CSF from patients with multiple sclerosis, Parkinson’s disease, and schizophrenia compared to 100 control subjects [13–15]

| PTGDS          | Multiple sclerosis | Parkinson’s disease | Schizophrenia |
|----------------|--------------------|---------------------|---------------|
| pI 5.2, invariant #3 | Not significant     | Decreased p < 0.0005 | Not significant |
| pI 6.7, invariant #5 | Increased p < 0.01  | No change           | No change     |
| Higher MW (30–45) | Yes, in all        | Yes, in one third   |               |

What may be the explanation for the changes in PTGDS forms? Loss of the five control isoforms likely represents a serious cellular defect in PTGDS regulation. It is interesting that the only conditions we have observed total or partial loss of the invariant five forms were in congenital malformations of the brain/meninges or other severe brain pathologies (Table 1). Since the sites of normal PTGDS synthesis are the glial cells and meninges, perhaps this is not surprising in conditions that damage these cells. The PTGDS gene may still be expressed, but processed differently. Such loss of the normal 25 kD isoforms has already been reported in extracts from many other tissues with different size and charge products compared to the normal CSF pattern [11]. To explain the major charge/mass shifts of immunoreactive PTGDS, one must invoke alternative splicing, translational processing or other post-translational modifications.

The cause of the gain of additional isoforms, such as illustrated in Fig. 4, will only be elucidated with definition of their structure. Changes in the regulation of glycosylation likely lead to the altered expressed forms, but there are many alternative possibilities, including phosphorylation, acylation, glycation or oxidation.

What consequences may result from the gain or loss of PTGDS isoforms? It is likely that many downstream
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Alterations in PTGDS represent pathology at the cellular level and will produce fluctuations in prostaglandins that correlate with symptoms in these diseases. This study was qualitative. However, we have published quantitative data from diseases that we studied prior to characterizing PTGDS. We quantified isoforms that we now know as PTGDS, from 54 patients with schizophrenia [13], 20 with idiopathic Parkinson’s disease [14], and 22 with multiple sclerosis [15], compared to 100 controls. In those studies we reported differential up/down regulation of isoforms. We have extracted this data in the context of PTGDS from those references and summarized it in Table 2, where the isoforms previously described by pl and MW as (5.2, 25 kD), (5.4, 25 kD), (6.7, 24 kD) correspond to invariant PTGDS isoforms (#3), (#4), (#5), respectively.

This quantitative measure of PTGDS dysregulation in disease further supports the qualitative data from this study. Combining the qualitative and quantitative data, there are multiple lines of evidence supporting the prediction that altered PTGDS reflects underlying meningeal/glial pathophysiology. The only losses of PTGDS occur in congenital malformations of the leptomeninges and massive structural brain damage. The appearances of additional PTGDS isoforms occur in diseases with the common feature of glial pathology and at locations known to be the main sources of PTGDS mRNA [6,32,36], exemplified by multiple sclerosis. The significant quantitative changes in schizophrenia, multiple sclerosis and Parkinson’s disease may be indicators of more subtle changes in glial pathology in these conditions. Further study of PTGDS should illuminate understanding of these brain disorders.

5. Conclusions

PTGDS, a heavily glycosylated protein, was found to have five isoforms in human CSF from all controls who had no classifiable psychiatric or neurological disease \((n = 22)\). The same proteins were present in a wide variety of nervous system disorders, including many different inborn errors of metabolism, infections, neoplasms and degenerative disease. However, losses of these invariant PTGDS isoforms were observed in congenital malformations of the nervous system, and additional isoforms were detected in malformations, Creutzfeldt-Jakob disease, vasculitic, demyelinating and hemorrhagic disorders. Review of the quantitative data of three invariant PTGDS isoforms from prior published work reveals that statistically significant differential up/down regulation of PTGDS occurs in multiple sclerosis, Parkinson’s disease and schizophrenia compared to controls [13–15].

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