Glycosylation of Erythrocyte Spectrin and Its Modification in Visceral Leishmaniasis

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Abstract

Using a lectin, Achatinin-H, having preferential specificity for glycoproteins with terminal 9-O-acetyl sialic acid derivatives linked in a2-6 linkages to subterminal N-acetylgalactosamine, eight distinct disease-associated 9-O-acetylated sialoglycoproteins was purified from erythrocytes of visceral leishmaniasis (VL) patients (RBCVL). Analyses of tryptic fragments by mass spectrometry led to the identification of two high-molecular weight 9-O-acetylated sialoglycoproteins as human erythrocytic α- and β-spectrin. Total spectrin purified from erythrocytes of VL patients (spectrinVL) was reactive with Achatinin-H. Interestingly, along with two high molecular weight bands corresponding to α- and β-spectrin another low molecular weight 60 kDa band was observed. Total spectrin was also purified from normal human erythrocytes (spectrinN) and insignificant binding with Achatinin-H was demonstrated. Additionally, this 60 kDa fragment was totally absent in spectrinN. Although the presence of both N- and O-glycosylations was found both in spectrinN and spectrinVL, enhanced sialylation was predominantly induced in spectrinVL. Sialic acids accounted for approximately 1.25 kDa mass of the 60 kDa polypeptide. The demonstration of a few identified sialylated tryptic fragments of α- and β-spectrinVL confirmed the presence of terminal sialic acids. Molecular modelling studies of spectrin suggest that a sugar moiety can fit into the potential glycosylation sites. Interestingly, highly sialylated spectrinVL showed decreased binding with spectrin-depleted inside-out membrane vesicles of normal erythrocytes compared to spectrinN suggesting functional abnormality. Taken together this is the first report of glycosylated erythrocytic spectrin in normal erythrocytes and its enhanced sialylation in RBCVL. The enhanced sialylation of this cytoskeletal protein is possibly related to the fragmentation of spectrinVL as evidenced by the presence of an additional 60 kDa fragment, absent in spectrinN which possibly affects the biology of RBCVL linked to both severe distortion of erythrocyte membrane and impairment of erythrocyte membrane integrity and may provide an explanation for their sensitivity to hemolysis and anemia in VL patients.

Introduction

The erythrocyte membrane is supported by a well-structured cytoskeleton. This cytoskeleton comprises of a network of different proteins maintaining the structural integrity and rigidity of the red blood cell (RBC) and of the RBC membrane [1]. Spectrin is a major cytoskeletal protein present as tetramers of α- and β-subunits associated with other cytoskeletal proteins forming a lattice that governs erythrocyte membrane properties. Alterations of spectrin have been associated with several congenital anomalies like hereditary hemolytic anemia and hereditary elliptocytosis leading to cellular distortion [2]. Biochemical modifications of spectrin, mainly glycation and oxidation, have been observed in diabetes mellitus indicating erythrocyte membrane changes [3–4]. Therefore, the status of the cytoskeletal proteins in disease may be affected by genetic abnormalities or metabolic or other stress inducing changes in cytoskeletal protein structure. Different levels and/or pattern of terminal sialic acid (SA) and its O-acetylation of cell surface expressed sialoglycoconjugates have occupied a pivotal position in inducing changes in different diseases [5–8]. The presence of different derivatives of SA on *Leishmania donovani* [9–13], different immune cells [14,15] and RBC [16–18] of patients with visceral leishmaniasis (VL) [RBCVL] and their role [19–26] have been demonstrated. VL caused by the intracellular kinetoplastid protozoa *L. donovani* accounts for an estimated 12 million infected humans with an incidence of 0.5 million cases per year [27–28]. Approximately 50% of the world’s VL cases occur in the Indian subcontinent.

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Along with other signature manifestations, VL is almost always associated with anemia [17–18]. However alteration of the RBC membrane architecture as one of the causes leading to anemia remains poorly understood.

We have detected the exclusive presence of eight distinct disease-associated 9-O-acetylated sialoglycoproteins (9-O-AcSGPs) on RBCVL [17], using the preferential specificity of a snail lectin, Achatinin-H for glycoproteins with terminal 9-O-acetyl sialic acid (9-O-AcSA) derivatives linked in α2-6 linkages to subterminal N-acetylgalactosamine (GalNAc) [29]. Interestingly, normal erythrocytes (RBCN) are devoid of such 9-O-AcSGPs. Antibodies directed against 9-O-acetylated sialic acids have also been demonstrated in VL [30,15]. Moreover enhanced pattern of altered sialylation demonstrated a direct correlation with the degree of complement-mediated hemolysis of RBCVL, providing a plausible basis for anemia associated with VL [18].

Taking into consideration the involvement of 9-O-AcSGPs in VL erythrocyte pathology, we report the presence, purification and identification of sialylation, N- and O-glycosylation of two high molecular weight 9-O-acetylated sialoglycoproteins as human erythrocytic spectrinVL and totally absent in spectrinN. In summary, this is the molecular identification of VL-associated proteins is a key to understanding of the status of glycosylation of spectrin in RBCVL. The identification of erythrocytic spectrin by mass spectrometry as 9-O-acetylated sialoglycoprotein, prompted us to explore the status of total spectrin in RBCVL (spectrinVL). Accordingly, spectrins were separately purified by the method as described by Ungewickell et al. [31]. The yield of spectrinVL, purified from ghost membrane (1.489±0.064 mg) of RBCVL was 0.231±0.017 mg. Purified spectrin from RBCN (spectrinN) demonstrated only two bands corresponding to α- and β-spectrin on SDS-PAGE (Fig. 2A, lane 1). In contrast, purified spectrinVL exhibited an additional 60 kDa fragment along with α- and β-spectrin bands (Fig. 2A, lane 2). Purified spectrinVL was further allowed to bind with Achatinin-H-Sepharose 4B. Achatinin-H bound spectrinVL demonstrated three similar bands (Fig. 2A, lane 3). These observations confirmed the presence of 9-O-AcSA on α-, β-spectrin and 60 kDa band of RBCVL, which was indicative of alteration of spectrin in VL.

Western blot analysis of purified spectrinVL separately by the method as described by Ungewickell et al [31] also showed reactivity with Achatinin-H reconfirming the identity of all three bands as spectrin containing 9-O-AcSA (Fig. 2B). In contrast, similar analysis of spectrinN demonstrated the absence of all these three bands suggesting lack or undetectable 9-O-AcSA on RBCN.

SDS-PAGE analysis (5 and 7.5%) of spectrinN and spectrinVL demonstrated slight variation in their electrophoretic mobility (Fig. 2C) suggesting some changes in VL. Two dimensional gel electrophoresis of purified spectrinVL reveals that individual spots corresponding to α-spectrin and β-spectrin and 60 kDa band have multiple isoelectric points (pI) (Fig. 2D) suggesting microheterogeneity of each spot possibly due to the differential sialylation of the same protein resulting into different pI.

Identification of 60 kDa band of erythrocytic α-spectrin in RBCVL

The 60 kDa band was identified as a fragment of α-spectrin (g: 119573202) by mass-spectrometric PMF analysis (Fig. 3A) as well as sequence determination of fragments from PSD spectra. Representative MS/MS spectra of tryptic fragments of m/z = 1257.6 (Fig. 3B) and m/z = 1709.8 (Fig. 3C) are shown. The 24 detected and annotated tryptic fragments matched the α-glycosidase F which evidenced a shift of the N-terminal section of α-spectrin with sequence coverage of 18.4% for the entire spectrin sequence and of 36.1% for the N-terminal exons (Table 1).

Erythrocytic spectrinN and spectrinVL are glycosylated

The identification of sialic acids in spectrinVL, prompted us to explore the status of glycosylation of spectrin in RBCN. The presence of comparable N- and O-glycosylation was demonstrated by a shift in the respective protein bands corresponding to α- and β-spectrin following enzymatic deglycosylation of neuraminidase treated purified spectrinN and spectrinVL due to their reduced molecular mass (Fig. 4A).

In parallel, neuraminidase treated 60 kDa fragment was also exposed to N- and O-glycosidase F which evidenced a shift of the band corresponding to a reduction of molecular weight by ~13.12 kDa and ~3.12 kDa indicating presence of both N- and O-glycosidic bonds (Fig. 4B). The 60 kDa fragment also demonstrated a shift of ~1.25 kDa after desialylation. Hence, deglycosylation accounted for about ~16.24 kDa of the total mass of 60 kDa.

The existence of both N- and O-glycosylation was further confirmed by binding with Sepharose/agarose bound specific lectins using iodinated spectrinVL and spectrinN (Fig. 4C–D). The

**Results**

Eight distinct disease-associated 9-O-AcSGPs exclusively induced on RBCVL

Disease-associated 9-O-AcSGPs were purified from RBCVL of clinically confirmed untreated VL patients (n = 30) using Achatinin-H-Sepharose 4B affinity matrix (Fig. 1A). Total membrane protein (1.20±0.187 mg) obtained from 2×10¹⁰ RBCVL yielded 0.432±0.025 mg of purified 9-O-AcSGPs separated as eight distinct bands on SDS-PAGE. Purification of the same from RBCN (2×10¹⁰) of normal healthy individuals (n = 30) by the same procedure yielded undetectable amount of protein.

Identification of spectrin by mass spectrometry

Molecular identification of VL-associated proteins is a key to their significance in the disease pathology. With this aim, peptide mass fingerprint (PMF) analysis and sequencing of tryptic fragments of the two high molecular weight bands of 9-O-AcSGPs by MALDI-TOF/PSD-MS led to the identification of erythrocytic α- and β-spectrin with sequences coverage of 33.6% and 22.7% respectively (Fig. 1B–C). These tryptic fragments were mapped on to the NCBI database sequences of human erythrocytic α- and β-spectrin sequences gi: 119573202 and gi: 67782321 respectively (shown in red color in Fig. S1).

SpectrinVL is specifically a 9-O-AcSGP present only in RBCVL

The identification of erythrocytic spectrin by mass spectrometry as 9-O-acetylated sialoglycoprotein, prompted us to explore the status of total spectrin in RBCVL (spectrinVL). Accordingly, spectrins were separately purified by the method as described by Ungewickell et al. [31]. The yield of spectrinVL, purified from ghost membrane (1.489±0.064 mg) of RBCVL was 0.231±0.017 mg. Purified spectrin from RBCN (spectrinN) demonstrated only two bands corresponding to α- and β-spectrin on SDS-PAGE (Fig. 2A, lane 1). In contrast, purified spectrinVL exhibited an additional 60 kDa fragment along with α- and β-spectrin bands (Fig. 2A, lane 2). Purified spectrinVL was further allowed to bind with Achatinin-H-Sepharose 4B. Achatinin-H bound spectrinVL demonstrated three similar bands (Fig. 2A, lane 3). These observations confirmed the presence of 9-O-AcSA on α-, β-spectrin and 60 kDa band of RBCVL, which was indicative of alteration of spectrin in VL.

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binding of $^{125}$I-spectrin$_N$ with immobilized Concanavalin A (ConA), Ricinus communis agglutinin (RCA), Helix pomatia agglutinin (HPA) and Ulex europaeus agglutinin (UEA) clearly suggested the existence of N-glycosylation. Similarly the binding of Dolichos biflorus agglutinin (DBA) and Jacalin reflected the presence of O-glycosylation in $^{125}$I-spectrin$_N$. These lectins also showed affinity towards the N- and O-glycosylated sugars present in $^{125}$I-spectrin$_L$.

Immobilized ConA and UEA showed comparable binding with $^{125}$I-spectrin$_N$ and $^{125}$I-spectrin$_L$ suggesting equivalent glycosylation levels having $\alpha$-Man (mannose), $\alpha$-Glc (glucose) and $\alpha$-L-Fuc (fucose) (Fig. 4C). However immobilized RCA, HPA, DBA and Jacalin showed higher binding towards $^{125}$I-spectrin$_N$ as compared to $^{125}$I-spectrin$_L$ suggesting the presence of more terminal $\beta$-D-Gal (galactose) (GalNAc, $\beta$-Gal), $\alpha$/$\beta$-D-GalNAc, $\alpha$-GalNAc and $\beta$1-3GalNAc sugars in spectrin$_N$ than spectrin$_L$.

The occurrence of Man $\alpha$ (1–3), (1–6) and (1–2) Man, Gal $\beta$ (1–3) GalNAc and Gal $\beta$ (1–4) GlcNAc (N-acetylglucosamine) in spectrin$_N$ and spectrin$_L$ was further demonstrated based on the comparable binding with Galanthus nivalis agglutinin (GNA), peanut agglutinin (PNA) and Datura stramonium agglutinin (DSA) respectively using DIG-glycan differentiation kit (Fig. 4E–F).

**Erythrocytic spectrin$_L$ is highly sialylated**

Isoelectric focusing (IEF) of spectrin$_L$ demonstrated four distinct bands within a pH range of 4.6–5.21 (Fig. 5A), which showed a considerable shift of their pH to a range of 6.25–7.95 after neuraminidase treatment indicating the presence of sialic acids. Furthermore the homogeneous shifts of the individual bands demonstrated the homogeneity of the proteins. In contrast shift in pH of spectrin$_N$ before and after neuraminidase treatment was less.
marked suggesting lower degree of sialylation. IEF of 60 kDa fragment showed two distinct bands, which demonstrated a shift in their pI after desialylation. This indicated that 60 kDa fragment comprised of two fragments of similar mass having sialic acids. Lane M shows the pI markers.

Enhanced presence of SA in spectrinVL as detected by biochemical and glycoanalytical methods

The presence of total sialic acid was demonstrated in spectrinVL/N using DIG-glycan detection kit (Fig. 5B–C). SpectrinVL showed enhanced (3-fold) sialylation as compared to spectrinN. The variable expression of linkage-specific sialic acids was demonstrated (Fig. 5D–E). SpectrinVL showed ~2.7–2.8 fold enhanced binding with Sambucus nigra agglutinin (SNA) and Manicka amurenis agglutinin (MAA) compared to spectrinN.

Sialylation were further demonstrated by binding with iodinated spectrinVL and spectrinN with Sepharose/agarose bound Wheat germ agglutinin (WGA), SNA and MAA. SpectrinVL showed significantly higher binding (~2.4 fold) with all three lectins than spectrinN (Fig. 5F). Achatinin-H also showed higher binding with spectrinVL whereas negligible with spectrinN (Fig. 5F).

With an attempt to search for the presence of sialylated tryptic fragments, the α- and β-subunits of spectrinVL were partially digested with trypsin separately. This controlled digestion yielded many fragments though we might have missed many smaller fractions (Fig. 5G). Theses fragments were allowed to bind with SNA and MAA agarose separately. SNA and MAA bound fragments were analysed on SDS-PAGE. Approximately 25 fragments of α-spectrinVL showed α2,6 linked and 18 of them had α2,3 linked terminal sialic acid. β-spectrinVL showed comparatively less number of α2,6 and α2,3 linked terminal sialic acids containing fragments (Fig. 5H).

Glycosidically bound sialic acids (SA) liberated from spectrinVL when separated on a TLC (thin layer chromatography) plate

Figure 2. Purification and characterization of spectrin. A. Purification of spectrinVL and spectrinN. A representative SDS-PAGE (7.5%) profile of SpectrinN (2.0 μg, lane 1) and spectrinVL (2.0 μg, lane 2), purified from RBCN and RBCVL as described by Ungewickell et al [31]. Purified spectrinVL was further passed through an Achatinin-H-Sepharose 4B affinity column and 9-O-acetylated sialic acid containing spectrinVL (2.0 μg, lane 3) was purified as described in Materials and Methods. Lane M shows molecular weight standards. B. Presence of 9-O-AcSA as detected by Western blot analysis. Equal amounts (2 μg) of purified spectrinVL and spectrinN were transferred onto nitrocellulose membrane after SDS-PAGE (8.5%). The blots were incubated overnight at 4 °C with Achatinin-H and processed as described in Materials and Methods. C. Equal amount (2 μg) of purified spectrinVL and spectrinN were separated both on 5 and 7.5% SDS-PAGE under similar conditions. D. Two dimensional (2D) gel electrophoresis of spectrinVL. A representative 2D (pI range 4–7, 4–15% gradient) profile of purified spectrin (100 μg) from RBCVL after staining with Coomassie is shown.

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demonstrated the presence of Neu5Ac (N-acetyl neuraminic acid) and Neu5,9Ac2 (5,9-diacetyl neuraminic acid) as compared with standard Neu5Ac and free SA purified from bovine submandibular mucin (BSM) (Fig. 6A). The presence of these derivatives was also demonstrated in the chromatogram of the liberated SA from spectrinVL by fluorimetric high-performance liquid chromatography (HPLC) (Fig. 6B). The Neu5,9Ac2 peak of spectrin VL completely disappeared on saponification. In contrast, spectrinN showed undetectable Neu5,9Ac2 suggesting disease-associated modification. BSM-derived SA showing 40% Neu5,9Ac2 was used as standard. Each fraction corresponding to Neu5Ac and Neu5,9Ac2 of spectrinVL was collected after fluorimetric-HPLC and was subsequently confirmed by MALDI-TOF-MS which yielded their expected molecular ion signals having m/z at 448.1 (Fig. 6C) and 490.1 (Fig. 6D) respectively.

Molecular modelling of glycosylated residues

In α-spectrin, four potential N-glycosylation sites were identified all of which contains the consensus sequence, Asn-Xaa-Ser/Thr (Table 2). However, only one potential O-glycosylation site was found at Thr-817 position with a score above the threshold value (0.35). However, in β-spectrin, two potential N-glycosylation sites were found and no potential O-glycosylation site (Table 2).

Structural verification of the predicted models of the modules containing N- and O- linked glycosylation sites revealed that the backbone conformations were satisfactory as the allowed phi-psi combinations were above 90% in the allowed region of Ramachandran’s plot. Verify 3-D results showed that the models have an average 85% of the residues with 3D-1D score greater than 0.2 which indicates good quality 3D structural parameters. From ERRAT analysis it was observed that most of the models.
have an overall quality factor greater than 90% indicating good structural quality.

Solvent accessible surface areas was calculated by ACCESS and the probable glycosylation sites shows that the residues are exposed enough needed for glycosylation at the sites (Table 2). Further, modelling of a representative sugar (β-GlcNAc) into one each of N- and O- glycosylation sites showed that the sugar moiety can go into the available space around the amino acid residues (Fig. 7).

SpectrinN and spectrinVL showed slight variations in their secondary structures

The CD (Circular Dichroism) spectra of spectrinN in far-UV (ultra violet) region showed that protein contains 51.71% of α-helix, 9.17% of β-sheet and 39.12% of random coil (Fig. 8A). The similar trend was observed in the secondary structure prediction using GOR 4 [32]. The sequences of α- and β-spectrinN was taken as weighted averages; the values of α-helix, β-sheet and random coil are 71.61%, 4.98% and 23.28% respectively. In parallel, the values for α-helix, β-sheet and random coil are 77.9%, 3.03% and 18.71% respectively as predicted from the modelled structures using MODELYN [33]. However, spectrinVL demonstrated a slight increase of α-helicity (63.05%) and a minimal decrease of β-sheet (5.68%) structure suggesting higher degree of sialylation possibly playing a role for such minute changes in its secondary structure.

Binding of 125I-spectrinN and 125I-spectrinVL to spectrin-depleted inside-out membrane vesicles (IOV)

In order to further demonstrate the modified structure of spectrinVL in comparison to spectrinN we have compared the binding status of iodinated spectrinVL and spectrinN with spectrin-depleted IOV from normal RBC-ghost (IOVN). The binding of 125I-spectrinVL to spectrin-depleted IOVN increases with increasing amount of 125I-spectrinVL (Fig. 8B). In contrast, under identical condition, 125I-spectrinN showed much higher binding towards spectrin-depleted IOV N. Such differences in binding signifies that minute structural modifications due to enhanced sialylation in spectrinVL possibly make it less available for interacting with other associated proteins in the spectrin-depleted IOVN of RBCN.

Discussion

VL is often complicated by anemia. The exclusive presence of 9-O-AcSGPs on erythrocytes of active VL has been correlated to RBC hemolysis [18,20]. The functional attributes of erythrocytes

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**Table 1.** The 24 tryptic fragments of 60 kDa band determined by MALDI-TOF-MS analysis.

| Mass [M+H]+ | Sequence range | Deviation from theoretical mass | Missed cleavage | Sequence |
|------------|----------------|---------------------------------|-----------------|----------|
| 1316.8079  | 17–27          | 0.14                            | 0               | VLETAEEIQER |
| 1378.8979  | 49–59          | 0.19                            | 0               | LEDSYHLOVK |
| 1534.9679  | 49–60          | 0.17                            | 1               | LEDSYHLQVKR |
| 1374.8279  | 119–130        | 0.21                            | 0               | FTMGHASSHEEK |
| 1452.9379  | 199–210        | 0.17                            | 1               | KFEDQSELVAK |
| 1324.8279  | 200–210        | 0.15                            | 0               | FEDQSELVAK |
| 1216.6979  | 239–248        | 0.13                            | 0               | QNEVNAAWER |
| 1237.7579  | 366–374        | 0.14                            | 0               | LQATYYWHR |
| 2240.4279  | 390–411        | 0.28                            | 0               | TAAINADELTPDVAGGEVLLDR |
| 2532.3879  | 426–448        | 0.27                            | 0               | FQSDADETGOLQVLNANHEASDEV |
| 1339.7179  | 484–494        | 0.16                            | 0               | DSEQVDSWMSR |
| 1355.7279  | 484–494        | 0.17                            | 0               | DSEQVDSWMoxSR |
| 2476.5179  | 495–517        | 0.29                            | 0               | QEAFLENEGLGNSLGSAEALOK |
| 1709.9579  | 518–531        | 0.23                            | 0               | HEDFEAFAQEEK |
| 2839.5879  | 678–701        | 0.21                            | 1               | QKGTLHEANQQLOFENNAEDLQR |
| 2583.4879  | 680–701        | 0.27                            | 0               | GTQLHEANQQLOFENNAEDLQR |
| 2604.5279  | 740–762        | 0.28                            | 0               | QQDQDILTDLAAAYEFGPDSDK |
| 1377.8679  | 837–848        | 0.12                            | 0               | VILENASHEPR |
| 1647.9479  | 859–873        | 0.20                            | 0               | MVEEGHFAAEDVSR |
| 1663.9579  | 859–873        | 0.22                            | 0               | Mox.VEEGHFAAEDVSR |
| 1191.7279  | 876–885        | 0.14                            | 0               | SLNQNMESLR |
| 2447.4879  | 917–939        | 0.29                            | 1               | KHEAFLLDLNSFGDSMoxK |
| 1968.0279  | 940–956        | 0.26                            | 1               | KHEAFLLDLNSFGDSMoxK |
| 1840.0479  | 941–956        | 0.18                            | 0               | HEAFLLDLNSFGDSMoxK |

The tryptic fragments matched the N-terminal portion of human erythrocytic α spectrin as compared to the protein sequences of the NCBI sequence database. The identification was confirmed by complete de novo sequencing of two fragments (shown in Figure 3 B–C). Mass [M+H]+ denotes the mono-isotopic masses of the fragment ions; sequence range refers to the alignment of the sequence of the denoted fragments with the α-spectrin reference sequence [gi: 119573202]; deviation from theoretical mass is the mass difference between the measured mass and the mass calculated from the corresponding database sequence; missed cleavage refers to the missed tryptic cleavage sites in the identified fragment; sequence is the fragment sequence in one-letter code, Mox is oxidized methionine.

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are determined by the structural integrity of the membrane, which is often described in terms of alterations in the membrane characteristics like osmotic fragility, fluidity and hydrophobicity. Any kind of perturbation in the milieu of the erythrocytes like oxidative changes or ligand specific interaction culminates in changes in the membrane characters generally associated with pathological conditions [25]. Therefore, we considered it worthwhile to unravel the molecular determinants and implications of 9-O-AcSGPs on RBCVL.

The major observation of this study is the demonstration of glycosylation in normal spectrin purified from RBCN, presence of higher degree of sialylation in spectrin purified from RBCVL, and fragmentation of spectrinVL as a 60 kDa 9-O-AcSGP. Therefore, it may be envisaged that enhanced sialylation of spectrinVL is possibly responsible for the generation of this fragmented O-acetylated sialic acid-containing spectrinVL. Altered binding of highly sialylated spectrinVL with spectrin-depleted inside-out membrane vesicles of RBCN possibly suggested functional abnormality. Membrane characteristics of RBCVL were observed by enhanced hydrophobicity, fragility, fluidity as compared to RBCN hinting towards membrane damage [25].

We have purified eight distinct 9-O-AcSGPs from RBCVL using Achatinin-H as an affinity matrix indicating linkage specific terminal 9-O-AcSA in these sialoglycoproteins. Distinct multiple spots of individual 9-O-AcSGP suggested microheterogeneity possibly due to differential sialylation. As 9-O-AcSGPs are
exclusively present on RBCVL, their identification through mass spectrometry was necessary to assess their possible implication in the disease pathology.

The analysis of two high molecular weight 9-O-AcSGPs by MALDI-TOF-MS evidenced a match with the NCBI entry of human erythrocytic α and β-spectrin with sequence coverage of 33.6% and 22.7% respectively. The amino acid sequences of tryptic fragments deduced from MS analysis confirmed the identification. Sequencing of two tryptic fragments and database-dependent Mascot as well as database-independent Sequit analyses made the identification unambiguous.

Figure 5. Presence of Neu5Ac and Neu5,9Ac2 in spectrinVL by biochemical methods. A. Enhanced sialylation demonstrated by IEF. Equal amounts (3.0 µg) of purified spectrinVL, 60 kDa band and spectrinN before and after removal of sialic acids were analyzed by IEF within a pH gradient of 3–10 and the respective bands visualized by silver staining. Lane M shows the pI markers. B–C. Enhanced sialylation in spectrinVL. Equal amount (1.0 µg) of purified spectrinVL and spectrinN was analyzed by using DIG-glycan detection kits and total sialylation was compared based on the densitometric scores of spots (B). Representative bar graph of densitometric scores of corresponding spots (C). D–E. Detection of linkage-specific terminal sialic acids in spectrinVL. Equal amount (2.0 µg) of spectrinVL and spectrinN was dot blotted on NC-paper and analyzed by DIG glycan and differentiation kit using SNA and MAA lectins following manufacturer’s protocol (D). Densitometric scores of corresponding spots are shown as bar graph (E). F. Binding of 125I-spectrinVL,N with several sialic acid binding lectins. To demonstrate the presence or absence of terminal sialic acids, a fixed concentrations of 125I-spectrinVL,N were analyzed by binding with Sepharose/agarose bound WGA, SNA, MAA, Achatinin-H (25 µl bead volume) having specificity towards linkage specific sialic acids as described in materials and methods. Bound radioactivity of 125I-spectrinVL,N was measured by Gamma-counter and represented as bar graphs. G–H. Detection of sialylated tryptic fragments in spectrinVL. The α and β subunits of purified spectrinVL were digested separately by restricted amount of trypsin. Such controlled digested and extracted tryptic fragments were dried and redissolved and an aliquot was separated in SDS-PAGE (7.5%–15% gradient) (G). Subsequently the presence of sialic acids on resulting tryptic fragments was analyzed by binding with SNA-agarose and MAA-agarose separately and followed by electrophoresis on SDS-PAGE (7.5%–15% gradient) (H) as described in Materials and Methods. Lane M shows the molecular weight standers.
parallel arrangement. The presence of both N- and O-glycosylation was indicated by shifts in molecular mass after the respective glycosidase treatments of neuraminidase-treated spectrinVL and spectrinN. Binding with several lectins specific for N- and O-glycosylation also supported the presence of such glycosylation in spectrinVL and spectrinN.

Molecular modelling studies also supported both N- & O-glycosylations of α-spectrin. However, only N-glycosylation was found in β-spectrin. Modelling the sugar moiety to the predicted glycosylation sites suggested that glycans could fit into these sites without any steric clashes, thus signifying the probability of glycosylation of spectrin.

Cell surface sialic acids have been widely associated with different pathological conditions. Enhanced presence of sialic acids in spectrinVL has been convincingly exhibited by lectin binding, which was further confirmed by TLC, fluorimetric-HPLC and MALDI-TOF-MS. Demonstration of pI of spectrinVL in acidic region and a huge shift of pI after neuraminidase treatment
established enhanced sialylation compared to spectrinN. More importantly, exclusive presence of Neu5,9Ac2 in spectrinVL suggested disease-associated enhanced sialylation in VL. Enhanced sialylation in spectrinVL compared to spectrinN possibly causes structural modification of spectrinVL. Such structural changes were perhaps the basis for the reduced capacity of spectrinVL to complex with other associated cytoskeletal proteins in normal environment as demonstrated by its less binding with spectrin-depleted IOVN.

Interestingly, purified 60 kDa fragment demonstrated the presence of two distinct bands in IEF and each of the bands depicted a distinct shift in their pI after removal of SA showing the presence of two sialylated proteins of similar molecular mass. In contrast spectrinN having comparable glycosylation, showed complete absence of such fragmentation, which suggested that alteration of spectrin mainly enhanced sialylation may be associated with VL pathology.

The cleaved 60 kDa fragment which belongs to α-spectrin contains two potential N-linked glycosylation sites, at Asn-633 & Asn-657 and one O-linked glycosylation site at position Thr-817 with sufficient surface accessibility. The remaining portion of α-spectrin although contains two potential N-linked glycosylation sites, but no O-linked glycosylation site was found. On the other hand, we could not identify any potential O-linked glycosylation sites in the β-spectrin. Therefore, it may be envisaged that the exclusive presence of O-linked glycosylation site in the N-terminal region of α-spectrin with high surface accessibility tends to have higher sialylation for interaction with each other. All these factors combined may play important role in the cleavage of α-spectrin to 60 kDa fragment in VL.

Production of erythrocytes requires synthesis of red cell proteins specially cytoskeleton proteins. During terminal differentiation of erythroid progenitor cells in culture it retains the key components of the endoplasmic reticulum protein translocation, glycosylation,

Table 2. Potential glycosylation sites and solvent accessibility values of spectrinN.

| SpectrinN | Glycosylation sites | Accessible surface area [Å²] |
|----------|---------------------|-------------------------------|
| α-spectrin | Asn-Lys-Thr [633–635] | 16.94 |
|           | Asn-Val-Thr [657–659] | 75.37 |
|           | Asn-Thr-Ser [1625–1627] | 93.70 |
|           | Asn-Leu-Ser [2077–2079] | 52.24 |
|           | Thr-817 | 142.89 |
| β-spectrin | Asn-Val-Thr [194–196] | 45.05 |
|           | Asn-Phe-Thr [197–199] | 21.97 |

*N-glycosylation sites (Asn) are shown as the consensus sequence of three amino acids and O-glycosylation site (Thr) is shown as the single amino acid. Sequence numbering is done according to the human alpha spectrin, erythrocytic 1, isoform CRA_b (gi: 119573202, taken from NCBI protein sequence database).

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Figure 7. Space filling structural representation of GlcNAc in spectrinN. Sugar moiety are colored by atoms (C = green, O = red, N = blue and H = white). The protein model is represented as conolly surface. A. N- glycosylation of α-spectrin is shown in yellow color at position Asn-1625. B. N- glycosylation of β-spectrin is shown in yellow color at position Asn-194. C. O- glycosylation of α-spectrin is shown in blue color at position Thr-817. doi:10.1371/journal.pone.0028169.g007
and protein folding machinery, chaperones, calreticulin and Hsp90 for red cell glycoprotein biosynthesis [34].

Non-enzymatic glycation and oxidation of spectrin were reported under several physiological conditions [3]. Such non-enzymatic changes are a result of glyco-oxidation, where the oxidative stress within, surpasses the antioxidant defense system of the cell [35]. The associated biochemical alterations affect the structure, aggregation and integrity of the membrane and membrane-associated proteins. Such changes have been witnessed in erythrocytic spectrin of subjects suffering from long-term diabetes mellitus. Here the elevated glucose concentration increases oxidation and advanced glycation end product formation of structural and membrane proteins of erythrocyte [36]. The demonstration of spectrinVL with enhanced sialylation in VL patients raised questions regarding the basis of these modifications. Therefore, it may be envisaged that oxidative modification of spectrin affects membrane morphology of the erythrocytes. Enhanced fragility, membrane fluidity and hydrophobicity of RBCVL as compared to RBCN were demonstrated earlier [25]. Hence, the evidence of altered spectrin reported here may provide an explanation for the known-impaired stability of erythrocytes in VL.

The presence of elevated levels of serum sialic acids in cardiovascular diseases and their relation to evaluated myocardial cell damage have been documented and it has been suggested that either the shedding or secretion of cell membrane sialic acids determines their accumulation in serum [37,38]. Furthermore, the importance of elevated serum sialic acids and soluble sialyltransferases in the diagnosis of Down-syndrome affected pregnancy and oral cavity cancer has been documented [39,40]. The presence of sialyltransferases in human serum may provide a possible way of changes in serum proteins with terminal ω2-6 sialic acid [40]. Bulai et al. have characterized a transport system and demonstrated the uptake of free sialic acids into human erythrocytes [41]. Therefore, free sialic acids could be transported across the membrane into RBC through a sialic acids transport system. Interestingly, the presence of enhanced sialic acids in the serum of VL patients probably hinted towards a possible mechanism of transport of these free sialic acids under the influence of sialyltransferases. Hence presence of sialic acids in VL serum could essentially serve as a source for the erythrocyte sialic acids, which could use a transport system for their entry. The uptake of sialic acids was monitored by measuring free sialic acids and ManNAc produced by cytosolic sialate pyruvate-lyase in human erythrocytes that indicated the presence of a sialic acid transport system [42]. Furthermore, in VL, peripheral hematopoietic cells have increased sialic acids [26] that could be shed in the serum and be transported across the erythrocyte membrane. The presence of serum sialyltransferases in VL patients and testing this hypothesis as well as the elucidation of the mechanisms of enhanced sialylation of spectrin in RBCVL demands extensive studies and will be a subject of future investigations.

Taken together the current study provides evidence for the first time not only for glycosylation of spectrinN but also enhanced sialylation in diseased condition i.e. spectrin in RBCVL. Additionally, we have demonstrated fragmented spectrinVL which could be triggered by such enhanced sialylation. Therefore, we hypothesize that the higher sialylation along with exclusive presence of 9-O-AcSGPs trigger membrane damage and may serve as an important factor leading to anemia-associated with VL. Hence the study successfully dissects one of the causal mechanisms leading to anemia, a common manifestation in VL.

Materials and Methods

Clinical samples

Blood sample of clinically confirmed active VL patients (n = 30; 21 males, 9 females, median age: 30 years) based on microscopic demonstration of Leishmania sp. amastigotes in splenic aspirates

Figure 8. Physicochemical study of structural modification of spectrinVL. A. CD-spectra. Far-UV CD spectra of spectrinN and spectrinVL in phosphate buffer (20 mM, pH 7.0) indicating the molar residue ellipticity as a function of wavelength along with the buffer only. B. Binding of 125I-spectrin to spectrin-depleted IOVN. Various concentrations of 125I-spectrinVL/N were incubated with a constant amount of spectrin-depleted-IOVN followed by determination of specific binding as described in Materials and Methods.

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were collected from School of Tropical Medicine, Kolkata and immediately processed for the separation of RBC\textsubscript{VL} at Indian Institute of Chemical Biology. The diagnosis was validated by two in-house techniques, in which the increased presence of linkage-specific 9-O-AcSGPs was quantified by erythrocyte binding assay [17] and anti-9-O-AcSGPs antibodies were detected by enzyme-linked immunosorbent assay (ELISA) using BSM known to contain a high percentage of 9-O-AcSAAs, as coating antigen [30,23]. The hematological parameters evidenced anemia in these patients and ruled out any other blood cell disorder (Table 3). Existence of high level of sialic acid in VL serum was observed.

Peripheral blood from normal human donors from endemic (n = 15) and non-endemic areas (n = 13) was processed similarly to obtain RBC\textsubscript{N} for the study. The Institutional Human Ethical Committee had approved the study and samples were taken with the consent of the donors, patients, or in case of minors from their parents/guardians.

### Purification of 9-O-AcSGPs from RBC\textsubscript{VL}

RBC\textsubscript{VL} (2 \times 10\textsuperscript{10}) after Ficoll-Hypaque (Amersham Pharmacia, Uppsala, Sweden) gradient separation comprising of 85–90% 9-O-AcSGP-positive cells were washed consecutively thrice with sodium chloride (NaCl, 0.15 M). Erythrocyte ghosts were prepared by sequential lyses of RBC\textsubscript{VL} using 5.0 mM, 2.5 mM and 1.25 mM ice-cold phosphate buffer, pH 7.0. The ghost membranes were solubilized in solubilizing buffer containing Tris-HCl (0.05 M), 1% (w/v) detergent concentration (CHAPS:BOG (9:1), MgCl\textsubscript{2} (1.0 mM), CaCl\textsubscript{2} (1.0 mM), dithiothreitol (DTT, 0.2 mM), phenylmethyl sulfonyl fluoride (PMSF, 20 \textmu M), protease inhibitor cocktail, pH 7.2, sonicated (three pulses, 10 sec each) in ice-mixture and incubated at 4 \textdegree C for 1 hr [43]. After centrifugation at 8200 \times g, 4 \textdegree C the supernatant was collected and dialyzed against Tris-HCl (0.05 M, pH 7.2) saline (TBS) containing 0.03 M Ca\textsuperscript{2+} (TBS-Ca\textsuperscript{2+}), 0.01% (w/v) detergent (CHAPS: BOG 1:1), sodium azide (0.02%). The dialyzed protein was processed for affinity chromatography and the protein content was quantified by Lowry method [44].

RBC\textsubscript{VL} ghost membrane protein fraction (1.85 mg) was passed through Achatinin-H-Sepharose-4B affinity column (2.0 mg/ml) equilibrated with TBS-Ca\textsuperscript{2+} containing sodium azide (0.02%) at 4 \textdegree C as described elsewhere [18]. After extensive washing, Achatinin-H-bound 9-O-AcSGPs were eluted at 25 \textdegree C with TBS containing sodium citrate (0.04 M, pH 7.2), dialyzed against TBS at 4 \textdegree C and stored at −70 \textdegree C for future use. As the binding of Achatinin-H towards 7-O- and/or 3-O-AcSA cannot be ruled out, therefore, presence of such linkages in O-acetylated sialoglycoproteins are also possible.

### Protein/peptide mass spectrometry

The identification of the glycoprotein was done by mass spectrometry using Bruker-Daltonics MALDI-TOF mass spectrometer Reflex IV (Bruker Daltonics, Bremen, Germany). The samples were prepared by dried-droplet procedure using 2,5-dihydroxybenzoic acid (DHBA) as matrix. Calibration was done externally with a mixture of Angiotensin I, Angiotensin II, Substance P, Bombesine, ACTH clip 1–17 and ACTH clip 18–39. Subsequently, peptide samples were used for analysis. Tryptic fragments were generated by overnight in-gel digestion of two high molecular hands in ammonium hydrocarbonate (5 mM) using 10 ng trypsin (Promega, Mannheim, Germany) per sample. Sequence analysis of selected tryptic fragments was done with an Ultraflex III MALDI-TOF-TOF mass spectrometer (Bruker Daltonics). PSD spectra were acquired using default LIFT method for MS/MS spectra acquisition with manually adjusted laser energy accumulating data from 1500–2000 laser shots. Spectra annotation was done using the FlexAnalysis 3.0 (Bruker Daltonics) software. PMF analyses and MS/MS ion searches were done with Mascot (Matrix Science Ltd., London, UK). Database searches through Mascot with PMF and MS/MS data were done with the BioTools 3.1 software (Bruker Daltonics). For database searches the following parameters were used. Taxonomy: Homo sapiens; database: NCBI; enzyme: trypsin; variable modifications: oxidation on methionine and one missed cleavages. Database searches for PMF spectra were done at the fragment mass tolerance ±0.3 Da. For the MS/MS searches mass tolerances for precursor was ±0.2 Da and 0.4 Da for fragment masses were used. The identification of the fragments and thereby of the protein was confirmed by database-independent de novo sequencing using the FlexAnalysis software [45].

### Purification of spectrin

The spectrin\textsubscript{N} from RBC\textsubscript{N} (1.25 mg total ghost membrane protein) was purified following the method of Ungewickell et al. with slight modifications [31]. Briefly, the ghosts were washed twice and resuspended in 3 volume of sodium phosphate (0.3 mM, pH 7.2) containing ethylene diamine tetraacetic acid (EDTA; 0.2 mM), (extraction buffer) and incubated for 20 min at 37 \textdegree C. The fragmented ghosts were pelleted by centrifugation at 80000 \times g for 1 h at 2 \textdegree C. Water-soluble proteins in the supernatant were immediately applied to a Sepharose 4B column (90 \times 2 cm), eluted with Tris (25 mM), EDTA (5 mM), NaCl (0.1 M), pH 7.6 (Tris/EDTA/saline buffer) at 4 \textdegree C. The column was eluted at 10 ml/h and 4-ml fractions were collected. Protein in the effluent was monitored by absorbance at 280 nm. Fractions containing purified spectrin dimer were pooled, concentrated and dialyzed overnight at 4 \textdegree C against TBS-Ca\textsuperscript{2+} buffer containing sodium azide. In parallel, spectrin\textsubscript{VL} was similarly purified from

### Table 3. Diagnostic features of patients with active visceral leishmaniasis (VL).

| Parameters | Patient\textsubscript{VL} (n = 30) | Normal (n = 30) |
|-----------|-----------------|----------------|
| Age (yr)  | 20–40           | 20–40          |
| Weight (Kg)| 32–40           | 50–60          |
| Duration of illness (mo) | 4–6 | Not applicable |
| RBC count | 1.0–2.5 \times 10\textsuperscript{12}/l | 4.0–6 \times 10\textsuperscript{12}/l |
| Leukocyte count (x/mm\textsuperscript{3}) | 3–4 \times 10\textsuperscript{3} | 5–10 \times 10\textsuperscript{3} |
| Hemoglobin concn. (g/dl) | 4–5          | 10–12         |
| Reticulocyte count (%) | 4–5          | 1–3           |
| Spleen size (cm) | 7–10            | Not palpable   |
| Splenic aspirate score\textsuperscript{a} | 3.5–4.2       | Negative       |
| RBC-ELISA | 0.95–1.38       | 0.19–0.24      |
| 9-O-AcSGP RBC\textsuperscript{b} | 85–90%         | 0.08–0.12%     |
| BSM-ELISA\textsuperscript{c} | 0.85–1.2        | 0.19–0.24      |
| Parasite ELISA\textsuperscript{d} | 1.1–1.8        | 0.21–0.28      |
| Serum sialic acid content (mg/dl)\textsuperscript{e} | 77.05±3.6     | 57.42±3.49    |

\textsuperscript{a} Determined by flow cytometry using FITC-Achatinin-H [18].
\textsuperscript{b} Anti-9-O-AcSGP antibody was detected by using BSM as coating antigen as described elsewhere [30].
\textsuperscript{c} Specific antibody was detected by using parasite lysate as coating antigen as described elsewhere [16].
\textsuperscript{d} Sialic acid content in serum was estimated by thiobarbituric acid method [54].

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RBCVL. Additionally, the purified total spectrinVL was further passed through an Achatinin-H-Sepharose 4B affinity column and 9-OAc-containing spectrinVL was purified as described above.

**Electrophoresis**

**2D Gel electrophoresis.** Purified spectrinVL was processed with 2D-clean up kit according to manufacturer’s protocol (Bio-Rad, USA). The precipitated protein was dissolved in rehydration buffer containing urea (6 M), thiourea (2 M), CHAPS (2%), DTT (50 mM), carrier ampholyte cocktail (2%), EDTA (0.1 mM), bromophenol blue and quantitated by Quick Strat Bradford. Protein (100 μg/100 μl) was rehydrated on IEF strips (pI 4–7, 7 cm) for 8 hrs and ran in PROTEAN IEF cell. Strips were equilibrated consecutively in two steps of 30 min each in equilibration buffer containing urea (6 M), SDS (2%), Tris-HCl (0.375 M, pH 8.8), glycerol (20%), DTT (2%) followed by same buffer with iodoacetamide (2.5%). The second dimension was carried out on gradient (4–15%) SDS-PAGE and stained by Biosafe Coomassie brilliant blue [46].

Purified spectrinN, spectrinVL or affinity-purified spectrinVL and gel eluted 60 kDa fragment and were analysed by SDS-PAGE (5 and 7.5%) in a minigel apparatus (Bio-Rad, USA) [25] and the gels were stained. N- or O-linked glycosylation of spectrinVL, spectrinN and 60 kDa fragment was demonstrated after deglycosylation with specific glycosidases using deglycosylation kit (Roche Applied Science, Mannheim, Germany) according to the manufacturer’s protocol [26].

Western blot analysis [26] of spectrinVL and spectrinN was performed by semidry method at 15 V for 20 min. After blocking, the membranes were incubated with Achatinin-H (100 μg/ml) in the presence of Ca²⁺ (0.03 M). Subsequently, the Achatinin-H probed membrane was incubated with polyclonal rabbit anti-Achatinin-H antibodies (1:400) at 4°C. Both the blots were developed using HRP-conjugated goat anti-rabbit IgG (1:5000, Cell signaling) and detected using diaminobenzidine (Sigma, St. Louis, MO) as substrate.

To obtain pure 60 kDa protein, Coomassie-stained bands corresponding to 60 kDa were excised from the electrophoresis gels and the proteins eluted using an Electro-Eluter Model 422 (Bio Rad, USA). IEF of purified spectrinVL, gel-eluted 60 kDa fragment and spectrinN was performed in capillary tubes within a pH range 3.0–10.0 using Mini-PROTEAN II tube cell apparatus (Bio-Rad, USA) and silver stained. The samples were desialylated overnight with *Arthrobacter ureafaciens* neuraminidase (0.2 μM/μg) at 37°C and processed similarly. The isoelectric points were determined from the pl of known proteins used as standards [26].

**Analysis of carbohydrates**

**DIG-glycan detection.** Equal amounts (1.0 μg) of spectrinVL and spectrinN were dot blotted on nitrocellulose paper (NC-paper) and total sialic acid content was analyzed by using DIG-glycan detection kit (Roche Applied Science, Mannheim, Germany) following manufacturer’s protocol [8]. Densitometric measurement of spots was done by using ImageQuantTL software (GE Healthcare).

**DIG-glycan differentiation.** The detection of terminal sugars on equal amount (2.0 μg, dot blotted on NC-paper) of spectrinVL and spectrinN was analyzed [26] by DIG-Glycan differentiation kit (Roche Applied Science, Mannheim, Germany) using several plant lectins SNA (specific for α2–6 linked Neu5Ac), MAA (specific for α2–3 linked Neu5Ac), GNA (specific for terminal Man (α1–3), (α1–6) and (α1–2) Man), PNA (specific for Gal (β1–3) GalNAc) and DSA (specific for Gal (β1–4) GlcNAc) according to the manufacturer’s protocol. Densitometric measurement of spots was done as above.

**Immunobilized lectin binding assays to 125I-spectrinVL/N.** To analyze the terminal sugar linkages, spectrinVL and spectrinN were separately iodinated with 125I (Bhabha Atomic Research Centre, Mumbai, India) yielding specific activity of spectrinN 1.96×10⁶ cpm/μg and of spectrinVL 1.3×10⁶ cpm/μg respectively. Fixed concentrations of 125I-spectrinVL/N were incubated separately with Sepharose/agarose bound lectins (25 μl bead volume) of different sugar-linkage specificity. Con A (specific for α-Mannose and α-Glucose), RCA (specific for β-D-Gal (GalNAc, β-Gal)), HPA (specific for α- or β-D-GalNAc), UEA (specific for α-L-Fuc) were used to demonstrate the presence of α-glycosylation. Similarly Jacalin (specific for β1,3GalNAc) and DBA (specific for α-GalNAc) were used to illustrate the presence of O-glycosylation. The lectins WGA (specific for GlcNAc and Neu5Ac), SNA, MAA, Achatinin-H (specific for 9-O-acetyl sialic acid residue) were used to show the presence of sialic acids. Unbound radioactivity was removed using TBS-bovine serum albumin (1 mg/ml) and bound 125I-spectrinVL/N was measured by Gamma-counter (Electronic Corporation, India) [47].

**Identification of sialoglycopeptides from spectrinVL containing α2,6 and α2,3 linked sialic acids.** Purified spectrinVL (150 μg) was run in 5% SDS-PAGE and spectrin & β-spectrin bands were digested partially by trypsin (200 ng) using in-gel tryspin digestion kit (Pierce, Rochford, USA) following the manufacturer’s protocol. Digested and extracted tryptic fragments were dried under speed-vac and redissolved in 0.1% TFA (100 μl). An aliquot of the redissolved tryptic fragments were separated on 7.5%–15% gradient SDS-gel. Remaining portion was neutralized by Tris-HCl (pH 8.0) and an aliquot was incubated separately with SNA-agarose (10 μl) and MAA-agarose (10 μl) for overnight at 4°C under mild shaking. The mixture was centrifuged at 5000 rpm for 10 min, supernatant collected as unbound fraction. Pellet was suspended in cold phosphate buffered saline (0.02 M, pH 7.0) and centrifuged at 5000 rpm for 10 min. Washed pellet was boiled with SDS-PAGE sample buffer, centrifuged at 5000 rpm for 5 min and the supernatant was ran in 7.5%–15% gradient gel.

**TLC.** Glycosidically bound sialic acids were released from purified spectrinVL (60 μg) by hydrolysis with propionic acid (4 M) for 4 h at 80°C. Liberated SA was subsequently passed onto Dowex 50WX8 (100–200 mesh) cation and Dowex 2×8 (200–400 mesh) anion exchange columns. Free sialic acids eluted from these columns were separated on TLC plates (Merck KGaA, Germany) in 1-propanol/H₂O (7:3 v/v) and developed by spraying with orcinol/HCl/FelCl₃ with heating at 180°C for 20 min [8,26]. Commercially available sialic acids (Neu5Ac, Sigma) along with those released from BSM served as standards. In parallel, sialic acids released from spectrinN were similarly processed.

**Fluorimetric HPLC.** An aliquot of liberated sialic acids was derivatized with 1,2-diamino-4,5-methyleneedioxybenzene (DMB) and DMB-SA was separated on an RP-18 column (LichroCART 125-4 HPLC-cartridge, 5 μm; Merck, Germany) and detected at excitation and emission wavelengths of 373 nm and 448 nm respectively as described elsewhere [8,26,48]. In parallel, sialic acids purified from BSM were run for comparison.

**MALDI-TOF MS.** Each fraction corresponding to different forms of sialic acids was collected from fluorimetric HPLC and subsequently analyzed by MALDI-TOF-MS (Applied Biosystem, USA) using DHBA as matrix as described previously [8,26,48]. Positive ion mode was used for analysis. The acquired spectra were accumulations of 1000 laser shots.
Molecular modelling of spectrin

Prediction of glycosylation sites. The protein sequences of human erythrocyte α- and β-spectrin were collected from NCBI database (gi: 119573202 and gi: 67782321 respectively). Prediction of N-glycosylation sites were performed with NetNGlyc 1.0 server (http://www.cbs.dtu.dk/services/NetNGlyc/). Predictions of O-glycosylation sites were done with NetOGlyc 3.1 server [49]. Probable glycosylation sites above a threshold value of 0.35 were selected for solvent accessibility calculation.

3-D structural modelling of spectrin modules. 3D structure of the modules containing the potential glycosylation sites were modelled using Swiss Model software [50]. The quality of the models was validated using Structural Analysis and Verification Server (http://saves.mbi.ucla.edu). The quality of the models was validated using Structural Analysis and Verification Server (http://saves.mbi.ucla.edu).

Solvent accessibility of probable glycosylation sites. Solvent accessibility surface area of all amino acid residues of the modules were calculated using ACCESS software [51]. The probable N- and O-linked glycosylation sites were identified by their percentage of surface exposure. Sites falling within the identified segments of α- and β-spectrin sequences by MASCOT program with significantly high intensity were eliminated from the list of probable sites as their non-glycosylated status was confirmed.

Attachment of carbohydrates to probable glycosylation sites. The structures of the modules attached to the carbohydrate at assigned Asn and Thr of the identified N- and O-glycosylation sites were optimized using molecular modelling software suite InsightII (2005) of Accelrys (San Diego, CA) by repeated energy minimization and molecular dynamics simulations with DISCOVER module. Energy minimization was performed alternatively with steepest descent and conjugate gradient methods (200 steps each using eff91 force field). Molecular dynamics simulation run was done with 10,000 steps of 1 fs after 1000 steps of equilibration with a conformation sampling of one in 100 steps at 300 K. At the end of the molecular dynamics simulation, the lowest potential energy conformation was picked using ANALYSIS module of Insight II for further energy minimization. The molecular dynamics simulation followed by energy minimization was performed on the glycosylation site residues attached with the sugar moiety while keeping the rest of the protein molecule fixed by applying positional constraints. This process was continued until satisfactory conformational parameters were achieved [26].

Physicochemical studies

CD spectra of spectrin. Far-UV CD spectra (between 190 nm and 250 nm) measurement of equal amount (0.05 µg/µl) of spectrinα and spectrinβ were performed at 25°C on a JASCO J-715 spectropolarimeter using a quartz cuvette of path length 1 mm under continuous flushing of nitrogen gas. The spectra shown are the average of ten data collected in continuous scan mode. The individual secondary structural contents in terms of α-helix, β-sheet, and random coil were analyzed from the far-UV CD spectra using the K2D2 software [52].

Preparation of spectrin-depleted inside-out vesicles (IOVs) and binding with 125I-spectrinαβ/N. Ghosts from RBCN were incubated for 30 min at 37°C in 30 vol of EDTA (0.25 mM), PMSF (25 µg/ml), pH 8.0, centrifuged at 50,000 g for 25 min and the inside-out vesicles (IOVs) were suspended in the buffer containing sodium phosphate (10 mM), KCl (130 mM), NaCl (20 mM), EDTA [1 mM], Na2S2O3 (0.5 mM), DTT [1 mM], pH 7.5 (Buffer A). Spectrin-depleted-IOVαβ was stored at a concentration of 1 mg/ml for overnight at 4°C [53].

To demonstrate the binding of spectrinαβ with spectrin-depleted-IOVαβ, 125I-spectrinαβ (0–10 µg/ml) were incubated for 90 min at 0°C in a buffer A [100 µl] containing 20 µg/ml spectrin-depleted-IOVαβ protein [53] and centrifuged at 50,000 g for 25 min at 4°C. Membrane-bound 125I-spectrin was washed with Buffer A and the radioactivity was counted by a Gamma-counter (Electronic Corporation, India). Nonspecific binding at each 125I-spectrin concentration was determined by the use of heat-denatured (70°C, 15 min) spectrin, and this value (10–20% of total counts) was routinely subtracted [53].

Estimation of sialic acid (SA) in serum. Estimation of total SA in serum was carried out colorimetrically by the thiobarbituric acid method after hydrolysis with 0.1 N sulfuric acid at 80°C for 1 hr [54]. The absolute value of sialic acid in serum was obtained from standard curve of authentic Neu5Ac.

Results are expressed as means ± S.D for individual sets of data. Each experiment was performed at least 3 times.

Supporting Information

Figure S1 Sequence of α-spectrin and β-spectrin with the identified and annotated fragments in red and the sequenced fragments underlined. (TIF)

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Author Contributions

Conceived and designed the experiments: S. Samanta Chhabinath Mandal PW Chitra Mandal. Performed the experiments: S. Samanta DD AG SM. Analyzed the data: S. Samanta DD AG SJ MF Chitra Mandal. Contributed reagents/materials/analysis tools: BS S. Sundar PW. Wrote the paper: S. Samanta Chhabinath Mandal PW Chitra Mandal.

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