

The α TSR Domain of *Plasmodium* Circumsporozoite Protein Bound Heparan Sulfates and Elicited High Titers of Sporozoite Binding Antibody After Displayed by Nanoparticles

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Introduction: Malaria is a devastating infectious illness caused by protozoan *Plasmodium* parasites. The circumsporozoite protein (CSP) on *Plasmodium* sporozoites binds heparan sulfate proteoglycan (HSPG) receptors for liver invasion, a critical step for prophylactic and therapeutic interventions.

Methods: In this study, we characterized the α TSR domain that covers region III and the thrombospondin type-I repeat (TSR) of the CSP using various biochemical, glycobiological, bioengineering, and immunological approaches.

Results: We found for the first time that the α TSR bound heparan sulfate (HS) glycans through support by a fused protein, indicating that the α TSR is a key functional domain and thus a vaccine target. When the α TSR was fused to the S domain of norovirus VP1, the fusion protein self-assembled into uniform S₆₀- α TSR nanoparticles. Three-dimensional structure reconstruction revealed that each nanoparticle consists of an S₆₀ nanoparticle core and 60 surface displayed α TSR antigens. The nanoparticle displayed α TSRs retained the binding function to HS glycans, indicating that they maintained authentic conformations. Both tagged and tag-free S₆₀- α TSR nanoparticles were produced via the *Escherichia coli* system at high yield by scalable approaches. They are highly immunogenic in mice, eliciting high titers of α TSR-specific antibody that bound specifically to the CSPs of *Plasmodium falciparum* sporozoites at high titer.

Discussion and Conclusion: Our data demonstrated that the α TSR is an important functional domain of the CSP. The S₆₀- α TSR nanoparticle displaying multiple α TSR antigens is a promising vaccine candidate potentially against attachment and infection of *Plasmodium* parasites.

Keywords: malaria, Plasmodium, α TSR domain, receptor binding domain, S₆₀ nanoparticle, malaria vaccine, norovirus

Introduction

Malaria is a deadly, transmissible human disease caused by unicellular protozoan parasites that are members of the *Plasmodium* genus in the family *Plasmodiidae*. The World Health Organization (WHO) estimated that about 40% of the global population is at risk of *Plasmodium* infection and that *Plasmodium* infection resulted in approximately 241 million malaria cases in 2020, claiming around 627,000 lives worldwide.¹ Thus, malaria remains a major threat to global public health. Various countermeasures have been attempted to prevent malaria, including control of mosquito vectors carrying *Plasmodium* parasites, generations of antimalarial drugs, and development of vaccines.² However, due to the resistance against insecticide by mosquitos and the diminished effects of antimalarial medicines owing to the resistance developed

by the *Plasmodiidae* parasites, current efforts focus on vaccine development,^{3,4} mostly targeting *Plasmodium* sporozoites that occur in the early stage of human infection.^{5,6}

Plasmodium protozoans that infect humans exhibit a common life cycle with an early development stage in the liver, followed by further proliferation in host bloods. During a mosquito bite by an infected female anopheline mosquito, *Plasmodium* sporozoites are transmitted from the mosquito into the dermis of a human host. The sporozoites then make their way into the liver via the bloodstream and invade hepatocytes, where they undergo proliferation known as schizogony, resulting in merozoites.^{3,7} The merozoites are released back into the bloodstream to invade erythrocytes where the merozoites proliferate asexually.^{8,9} Some merozoites differentiate into gametocytes, which is the end stage of the parasites' life cycle in humans.^{10,11} This asexual blood stage involves fast and iterative invasion of erythrocytes, leading to symptoms of malaria disease. Development beyond the gametocyte stage occurs in mosquito, where the male and the female gametes fuse for sexual reproduction.¹²

Sporozoites appear in the first stage of human infection, representing a bottleneck in the *Plasmodium* life cycle, as only 10 to 100 sporozoites are transmitted during a mosquito bite¹³ and they are exposed to host antibodies before they invade the liver.¹⁴ In addition, since the sporozoites occur before the symptomatic stage, they are ideal targets for antimalarial drugs and vaccines. The sporozoites are coated densely and homogeneously with circumsporozoite proteins (CSPs)¹⁵ with multiple functions. The CSP is 414 amino acids in length, consisting of an N-terminal domain, region I with a protease cleavage site, a repeat region, region III with conserved sequences, a region II plus that is homologous with the thrombospondin type-1 repeat (TSR) superfamily, and a glycosylphosphatidylinositol (GPI) anchor site at the C terminus.¹⁶ CSPs are associated with the plasma membrane via the GPI anchors.¹⁷

The CSP is essential for sporozoite development, including host cell targeting and invasion. It interacts with heparan sulfate proteoglycans (HSPGs) as receptors for sporozoites to be arrested in the liver, facilitating liver infection,^{15,18–20} likely using the TSR (region II plus) as the HSPG-binding domain,^{20–23} although contradictory data were reported.^{16,24} The CSP is an immunodominant, protective antigen.²⁵ Immunization with irradiated sporozoites resulted in strong protection against malaria infection in rodents,²⁶ monkeys,²⁷ and humans²⁸ by eliciting CSP-specific antibodies that inhibited sporozoite infectivity. Irradiated sporozoites also elicited T cell responses that destroy the exo-erythrocytic forms of the parasites, including sporozoites.²⁹ In addition, polymorphic T-cell epitopes were mapped to the TSR region, such as CS.T3 that induces a CD4⁺ T-cell response correlated with protection.³⁰ Thus, the CSP and TSR are excellent vaccine targets.

Among the malaria vaccine candidates tested to date, the most clinically advanced one is RTS, S/AS01 (Mosquirix) that was developed by GlaxoSmithKline (GSK).^{5,6} This vaccine contains the tandem repeats, region III, and TSR of *P. falciparum* CSP that are fused with the hepatitis B surface antigen that can self-assemble into virus-like particles (VLPs),^{5,6} although the morphology or structure of the resulted vaccine candidates have not been reported. In a Phase III clinical study, RTS S/AS01 vaccine revealed an average efficacy of about 39% against clinical malaria in children aged 5–17 months over a 4-year period, which, however, diminished over time.^{5,6,31,32} The continued absence of an effective malaria vaccine indicates an urgent need of an immunoprophylactic approach with improved efficacy and long-lasting immunity.

In this study, we first provided evidence that the α TSR domain covering CSP region III and TSR¹⁶ binds heparan sulfate (HS) after fusion with GST protein, showing it as an important functional domain and a vaccine target. We then took advantage of our S₆₀ protein nanoparticle platform^{33–35} to display the α TSR antigens of *P. falciparum* CSP for improved immunogenicity as a vaccine candidate. The *P. falciparum* α TSR is selected, because its crystal structure is known¹⁶ and *P. falciparum* is the most prevalent and most deadly species of *Plasmodium* parasites.^{2,36} The resulted S₆₀- α TSR nanoparticle was produced, bound HS glycan receptors, structurally elucidated, and is highly immunogenic, eliciting high α TSR-specific antibody titers in mice. The α TSR-specific antibody bound the CSPs of *P. falciparum* sporozoites at high titer, which could potentially block the receptor-binding function of CSPs. Thus, the S₆₀- α TSR nanoparticle may be a promising malaria vaccine candidate.

Materials and Methods

Plasmids for α TSR-Hisx6 and GST- α TSR Protein Expressions

A DNA fragment encoding the α TSR domain¹⁶ of *P. falciparum* 3D7 strain (GenBank AC#: CAB38998.2, from E309 to C375, 67 residues) was codon-optimized to *Escherichia coli* (*E. coli*) and synthesized via GenScript (Piscataway). This DNA fragment was cloned into the pET-24b vector (Novagen) for expression of α TSR-Hisx6 protein (Figure 1A, top

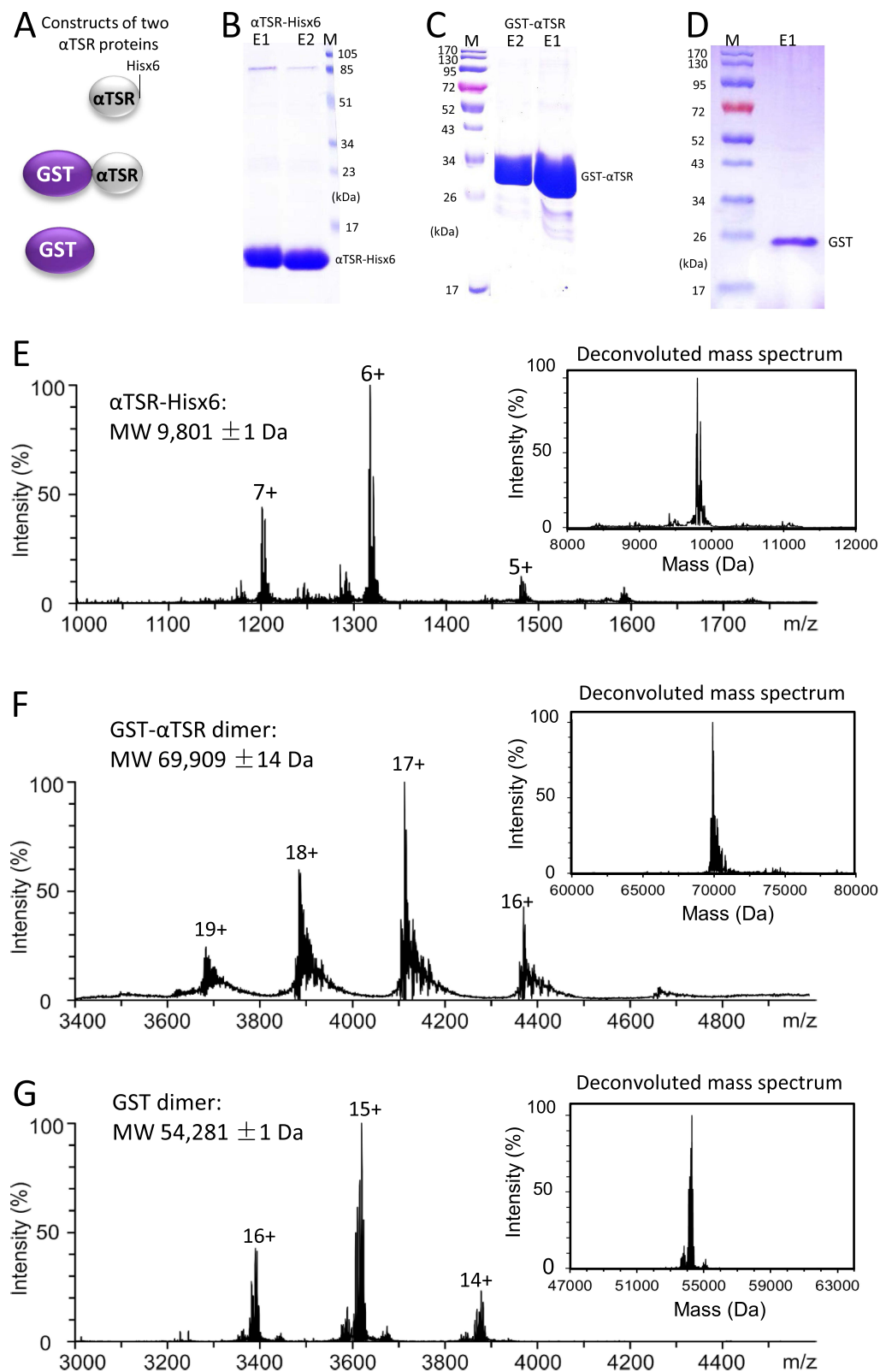


Figure 1 Generation and characterization of two α TSR proteins. **(A)** Schematic constructs of the α TSR-Hisx6 (top panel), the GST- α TSR (middle panel), and the GST (bottom panel) proteins. **(B–D)** SDS-PAGE showing the purified α TSR-Hisx6 protein at ~ 10 kDa **(B)**, the GST- α TSR protein at ~ 35 kDa **(C)**, and the GST at ~ 26 kDa **(D)**, respectively. Lanes E1 and/or E2 are elution fractions from the purification resin. M is protein markers with indicated molecular weights. 16% gel was used in **(B)**, whereas 10% gels were used for **(C)** and **(D)**. **(E–G)** Molecular weight (MW) determinations of the α TSR-Hisx6 **(E)**, GST- α TSR **(F)**, and GST proteins by electrospray ionization (ESI) and corresponding deconvoluted (insets) mass spectra, revealing 9801 Da, and 69,909 Da, and 54,281 Da, respectively.

panel). The DNA fragment was also cloned into the pGEX-4T-1 vector (GE Healthcare Life Sciences) for production of GST (glutathione s-transferase)-tagged α TSR (GST- α TSR) protein (Figure 1A, middle panel). In addition, GST was also expressed using the pGEX-4T-1 plasmid alone (Figure 1A, bottom panel).

DNA Construct for S- α TSR Protein Production

The DNA fragment encoding the α TSR antigen was cloned into the previously generated, pET-24b-based plasmid for production of the S_{R69A}-VP8* fusion protein that self-assembles into the S₆₀-VP8* nanoparticle,³⁴ in which the VP8*-encoding sequences were replaced with the α TSR encoding DNA fragment. The resulting new plasmid expresses a Hisx6-tagged S- α TSR fusion protein. This new pET-24b-based plasmid was also modified to express a tag-free S- α TSR fusion protein by inserting a stop codon in front of the Hisx6-encoding sequences.

Recombinant Protein Production and Purification

Recombinant proteins were expressed using the *E. coli* (BL21, Arctic strain) system after induction by 0.25 mM isopropyl- β -D-thiogalactopyranoside (IPTG) at 13°C overnight as described elsewhere.^{34,37,38} Soluble Hisx6-tagged proteins were purified using the Cobalt resins (Thermo Fisher Scientific), while GST and GST-tagged protein were isolated using the Glutathione Sepharose 4 Fast Flow medium (GE Healthcare Life Sciences) according to the instructions of the manufacturers. The tag-free S- α TSR protein was purified by a two-step procedure, including a precipitation of the target protein from the bacteria lysate using ammonium sulfate [(NH₄)₂SO₄], and an ion exchange chromatography (see below).³⁵ (NH₄)₂SO₄ at 1.6 M end concentration was used to effectively precipitate the tag-free S- α TSR protein at room temperature (20–22°C).

Sodium Dodecyl Sulfate Polyacrylamide Gel Electrophoresis (SDS-PAGE)

Purified recombinant proteins were analyzed by SDS-PAGE using 10% separating gels, except for the small α TSR-Hisx6 protein (~9.8 kDa) that needed a 16% separating gel. The gels were stained using commercial Coomassie Stain (Bio-Rad) as instructed by the manufacturer.

Gel Filtration Chromatography

Gel filtration was performed as described previously^{37–39} using an ÄKTA™ Fast Performance Liquid Chromatography system (FPLC, ÄKTA™ pure 25L, GE Healthcare Life Sciences) using a size exclusion column (Superdex 200, 10/300 GL, GE Healthcare Life Sciences). The size exclusion column was calibrated using gel filtration calibration kits (GE Healthcare Life Sciences), whereas the elution peaks of the proteins were defined using the formerly prepared S₆₀-VP8* nanoparticle (~3.4 mDa) [33] and GST-dimer (~52 kDa). Relative protein concentrations in the effluents were shown by absorbance at A₂₈₀.

Anion Exchange Chromatography

This was conducted to purify the tag-free S- α TSR protein as described previously.^{35,40} Briefly, the ÄKTA FPLC system equipped with a HiPrep Q HP 16/10 column (GE Healthcare Life Sciences) was used. First, the column was equilibrated with 7 column volumes (CVs) of 20 mM Tris buffer (pH 8.0, buffer A). After sample load, the column was washed by 7 CVs of buffer A, followed by elution of bound proteins by 8 CVs of buffer B (1 M NaCl in buffer A) via a linear gradient from 0% to 100%. Proteins in the effluent were indicated as UV absorbance at A₂₈₀, while the elution positions of the proteins were reported by percentages of buffer B.

CsCl Density Gradient Ultracentrifugation

This method was used to determine the density of the S₆₀- α TSR nanoparticle as described previously.³⁴ Briefly, 0.5 mL of the resin purified S- α TSR protein was blended with 10 mL of CsCl solution with a density of 1.3630 g/mL. After centrifugation for 45 hours at 288,000 x g using the Optima L-90K ultracentrifuge (Beckman Coulter), the CsCl gradient was fractionated by bottom puncture into 22 fractions (~0.5 mL each). After 200-fold dilution in PBS, the S- α TSR

protein in fractions was detected by EIA (see below) using antibody against norovirus VLPs.⁴¹ The CsCl densities of fractions were determined based on the refractive index.

Negative Stain Transmission Electron Microscopy (TEM)

After staining with 1% ammonium molybdate, the purified S- α TSR nanoparticles were inspected using an EM10 C2 microscope (Zeiss, Germany) at 80 kV at magnifications between 15,000x to 30,000x as described previously.³⁴

Image Processing and Modelling

Acquired micrographs of negatively stained S₆₀- α TSR nanoparticles were initially processed in RELION v4.0.⁴² A total of 874 particles were selected using the particle autopicking tool in RELION. After 2 rounds of reference-free 2D classification, a total of 689 particles remained after removing junk particles. The particle stack was split into even and odd numbered half datasets. JSPR^{43,44} was then used to reconstruct 5 independent initial models using a random subset of 200 particles at each iteration from each half dataset for 8 iterations. All particles for each half dataset were globally searched and further refined against the initial models. The resulting orientation parameters of each particle were compared and those with the most stable orientations were kept using the calcImageParamVariance.py script from the JSPR software package. The final density map was produced after 2 additional iterations of refinement of the most stable particles. Fitting of the density map was performed through UCSF Chimera software⁴⁵ using the S₆₀ nanoparticle cryo-EM structure of norovirus (strain VA387) S domain (residue 46 to 218) and the crystal structure of the α TSR monomer of *P. falciparum* (PDB code: 3VDJ).¹⁶ The crystal structures of the inner shell of the 60-valent feline calicivirus (FCV) VLP (PDB code: 4PB6)⁴⁶ was used to show the exposed C-terminal ends of the S domain.

Enzyme Immunoassay (EIA)-Based Glycan Binding Assay

EIA assays were utilized to determine interaction between the α TSR domains and HS glycans. Heparin sodium salt from porcine intestinal mucosa (Sigma-Aldrich) at 25 μ g/mL was coated on 96-well microtiter plates (Thermo Scientific) at 4°C overnight. After blocking with 5% (w/v) skim milk, the plates were incubated with serially diluted α TSR-Hisx6, GST- α TSR, or S₆₀- α TSR proteins. The bound proteins were detected using three different antibodies: 1) our in house made mouse sera after immunization with S₆₀- α TSR (1:2000 dilution) for detections of the α TSR-Hisx6 and the GST- α TSR proteins, 2) rabbit antibody against norovirus VLP⁴¹ (1:3000 dilution) for detection of the S₆₀ protein, or 3) rabbit antibody against GST for detection of the GST protein. The bound antibody was measured by corresponding horse-radish peroxidase (HRP)-conjugated secondary goat anti-mouse or rabbit IgG (Thermo Fisher Scientific) at 1:5000 dilution. The signal intensity was shown by optical density at wavelength 450 nm after incubated with the HRP substrates as described previously.³⁵

Mass Spectrometry

Electrospray ionization mass spectrometry (ESI-MS) was carried out in positive ion mode using a Q Exactive Hybrid Quadrupole-Orbitrap mass spectrometer (Thermo Fisher Scientific) equipped with the Nanospray FlexTM ion source. The nanoflow ESI (nanoESI) was performed using tips pulled from a borosilicate capillary (1.0 mm o.d., 0.75 mm i.d.) by a micropipette puller (P-1000, Sutter Instruments). A voltage of 0.9 kV was applied to a platinum wire inserted into the nanoESI tip and in contact with the sample solution. The capillary temperature 160°C, maximum inject time 200 ms, microscans 10, S-lens RF level 100, and resolution 17,000 were applied, whereas other instrument parameters were set at default values. Data acquisition and processing were performed using Xcalibur (Thermo Fisher Scientific, version 4.4).

Deconvolution of ESI Mass Spectra

Deconvolution of the ESI mass spectra was performed using UniDec software (version 3.2.0)⁴⁷ to determine the molecular weights of the GST- α TSR, α TSR-Hisx6, and GST proteins. The following parameters were used: *m/z* range 2000–5000 (for GST- α TSR and GST), and 1000–3000 (for α TSR); bin every 0.5 (ie, the raw MS data points were resampled at every 0.5 *m/z* to UniDec); charge range 5–20; mass range 60,000–80,000 for GST- α TSR, 46,000–64,000 for GST, and 8000–12,000 for α TSR-Hisx6; sample mass every 1.0 Da; Peak FWHM (Th) 1.0; peak shape function

Gaussian; and maximum number of iterations 100. A full description of the UniDec deconvolution parameters can be found from the User Manual at <http://unidec.chem.ox.ac.uk/>.

ESI-MS Binding Measurements

The affinities for the interactions between the two α TSR proteins (GST- α TSR and α TSR-Hisx6) or GST and the glycans in our HS library were quantified using ESI-MS binding assay. The measurements were performed in 200 mM aqueous ammonium acetate solution (pH 6.8, 25°C), whereby each protein was incubated with each ligand (separately) for 10 min and then subjected to ESI-MS analysis. The apparent (ie, macroscopic) dissociation constant (K_D) was calculated (Equation 1), from the total abundance (Ab) ratio (R , Equation 2) of the ligand-bound (PL)-to-free protein (P) ions and the initial concentrations of P ($[P]_0$) and ligand ($[L]_0$):

$$K_D = \frac{[L]_0}{R} - \frac{[P]_0}{R + 1} \quad (1)$$

$$R = \frac{Ab(PL)}{Ab(P)} = \frac{[PL]}{[P]} \quad (2)$$

In all cases, a single chain fragment (scFv, MW 26,539 Da) of the monoclonal antibody Se155-4, served as the reference protein (P_{ref}) to correct for any nonspecific binding during the nanoESI process. A detailed description of the method can be found elsewhere.⁴⁸ The reported affinities are average values from replicate measurements performed at a minimum of three different L concentrations.

Mass Photometry

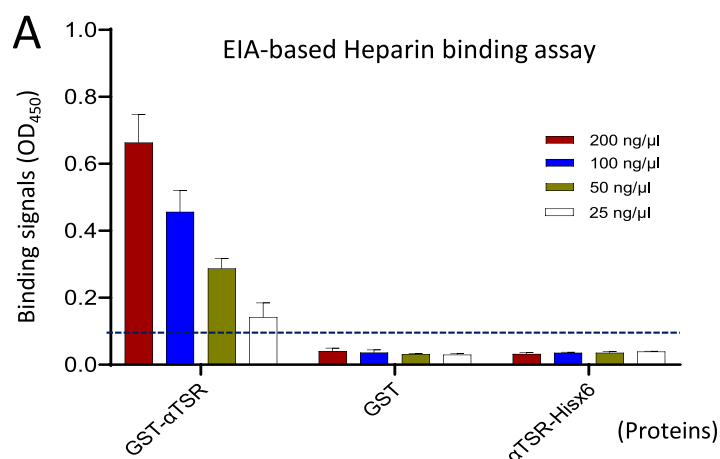
Mass photometry was carried out with a OneMP mass photometer (Refeyn Ltd, Oxford, UK) to determine the molecular weights of the S_{60} - α TSR nanoparticle. Briefly, the microscope coverslips (Thorlabs Inc) were pre-cleaned with isopropanol (LC-MS grade) and Milli-Q H₂O, followed by drying with a clean nitrogen stream. Clean coverslips were assembled for sample delivery using silicone CultureWell™ gaskets (Grace Bio-Labs, OR). For each acquisition, 15 μ L of sample diluted in 200 mM ammonium acetate (pH 6.8, 25°C) was introduced into a gasket. Following autofocus stabilization, a movie of 60s duration was recorded at 1 kHz, with exposure times varying between 0.6 ms and 0.9 ms. Each sample was measured at least three times independently. Data acquisition was performed using AcquireMP (Refeyn Ltd, version 2.5). The data were analyzed with DiscoverMP (Refeyn Ltd, version 2.5), which detects the events of proteins in the solution landing onto the glass coverslip surface and determines the respective interferometric scattering contrasts. The mean peak contrast of each sample solution was determined using Gaussian fitting, and converted to the mass value through contrast comparison with known mass standard calibrants (ie, contrast-to-mass calibration). The calibration curve was performed daily, including measuring bovine serum albumin (66, 132, 198 kDa), thyroglobulin (660 kDa) and apoferritin (480 and 960 kDa). The detailed description of the principles and data analysis method for the mass photometry can be found elsewhere.⁴⁹

HS Glycan Library

The 15 HS compounds in the HS glycan library, including HS DP2, HS DP4, HS DP6, HS15 to HS20, and HS22 to HS27 (see Figure 2, were provided by Prof. Geert-Jan Boons at Utrecht University and University of Georgia.

Immunization of Mice

Pathogen free BALB/c mice at age ~6 weeks from the Jackson Laboratory (Bar harbor, ME) were maintained at the Division of Veterinary Services of Cincinnati Children's Hospital Medical Center (CCHMC). Mice were divided randomly into four groups with 6 to 8 mice per group ($n = 6-8$) and the grouped mice were immunized with one of the following reagents: 1) S_{60} - α TSR nanoparticle (S_{60} - α TSR); 2) GST- α TSR fusion protein (GST-TSR); 3) α TSR-Hisx6 protein (TSR-Hisx6); and 4) S_{60} nanoparticle without the α TSR antigens (S_{60}). Immunogens that contained Alum adjuvant (Imject Alum, Thermo Fisher Scientific) at 25 μ L/dose (20 μ g/mouse/dose) were administered intramuscularly (IM) in the thigh muscle three times with 50 μ L injection volume at 2-week intervals. Blood samples were collected two



B Screening of a glycan library consisting of 15 heparan sulfate compounds

#	Name	Composition	Max # Na ⁺	Formular without Na ⁺	Monoisotopic mass (Da)	GST-αTSR <i>K_D</i> (μM)	αTSR-His <i>K_D</i> (μM)	GST <i>K_D</i> (μM)
1	HS DP2	α-L-IdoA(2S)-α-D-GlcNS(6S)-O-(CH ₂) ₅ NH ₂	4	C ₁₇ H ₃₂ N ₂ O ₂₀ S ₃	680.0711	380±130	3930±560	NB
2	HS DP4	[α-L-IdoA(2S)-α-D-GlcNS(6S)] ₂ -O-(CH ₂) ₅ NH ₂	8	C ₂₉ H ₅₁ N ₃ O ₃₉ S ₆	1257.0424	430±130	1000±100	NB
3	HS DP6	[α-L-IdoA(2S)-α-D-GlcNS(6S)] ₃ -O-(CH ₂) ₅ NH ₂	12	C ₄₁ H ₇₀ N ₄ O ₅₈ S ₉	1834.0137	330±70	2040±400	NB
15	HS15	β-D-GlcA(1-4)-α-D-GlcNAc(1-4)-α-L-IdoA(2S)(1-4)-α-D-GlcNAc-O-	3	C ₃₃ H ₅₅ N ₃ O ₂₆ S	941.2795	400±140	NB	NB
16	HS16	β-D-GlcA(1-4)-α-D-GlcNS(1-4)-α-L-IdoA(2S)(1-4)-α-D-GlcNS-O-	5	C ₂₉ H ₅₁ N ₃ O ₃₀ S ₃	1017.1720	390±70	NB	NB
17	HS17	β-D-GlcA(1-4)-α-D-GlcNAc(1-4)-β-D-GlcA(2S)(1-4)-α-D-GlcNAc(6S)-O-	4	C ₃₃ H ₅₅ N ₃ O ₂₉ S ₂	1021.2363	500±130	NB	NB
18	HS18	β-D-GlcA(1-4)-α-D-GlcNS(1-4)-β-D-GlcA(2S)(1-4)-α-D-GlcNS(6S)-O-	6	C ₂₉ H ₅₁ N ₃ O ₃₃ S ₄	1097.1288	540±210	1880±490	NB
19	HS19	β-D-GlcA(1-4)-α-D-GlcNAc(1-4)-β-D-GlcA(2S)(1-4)-α-D-GlcNAc-O-	3	C ₃₃ H ₅₅ N ₃ O ₂₆ S	941.2795	500±120	NB	NB
20	HS20	β-D-GlcA(1-4)-α-D-GlcNS(1-4)-β-D-GlcA(2S)(1-4)-α-D-GlcNS-O-	5	C ₂₉ H ₅₁ N ₃ O ₃₀ S ₃	1017.1720	390±120	2670±450	NB
22	HS22	β-D-GlcA(1-4)-α-D-GlcNAc(1-4)-α-L-IdoA(2S)(1-4)-α-D-GlcNS-O-	4	C ₃₁ H ₅₃ N ₃ O ₂₈ S ₂	979.2257	1110±20	NB	NB
23	HS23	β-D-GlcA(1-4)-α-D-GlcNS(1-4)-α-L-IdoA(2S)(1-4)-α-D-GlcNS(6S)-O-	6	C ₂₉ H ₅₁ N ₃ O ₃₃ S ₄	1097.1288	430±60	NB	NB
24	HS24	β-D-GlcA(1-4)-α-D-GlcNAc(1-4)-α-L-IdoA(2S)(1-4)-α-D-GlcNAc(6S)-O-	4	C ₃₃ H ₅₅ N ₃ O ₂₉ S ₂	1021.2363	1000±390	2750±580	NB
25	HS25	β-D-GlcA(1-4)-α-D-GlcNAc(1-4)-α-L-IdoA(2S)(1-4)-α-D-GlcNS(6S)-O-	5	C ₃₁ H ₅₃ N ₃ O ₃₁ S ₃	1059.1825	600±60	NB	NB
26	HS26	β-D-GlcA(1-4)-α-D-GlcNAc(1-4)-α-L-IdoA(1-4)-α-D-GlcNAc(6S)-O-	3	C ₃₃ H ₅₅ N ₃ O ₂₆ S	941.2795	320±50	NB	NB
27	HS27	β-D-GlcA(1-4)-α-D-GlcNS(1-4)-α-L-IdoA(1-4)-α-D-GlcNS(6S)-O-	5	C ₂₉ H ₅₁ N ₃ O ₃₀ S ₃	1017.1720	550±140	NB	NB

Figure 2 Binding of the αTSR proteins to heparan sulfate (HS) glycans. **(A)** Binding of the αTSR containing proteins to heparin sodium salt by EIA based binding assays. Y-axis shows binding signals in optical density (OD₄₅₀), while X-axis denotes the proteins at indicated concentrations: red columns, 200 ng/μL; blue columns, 100 ng/μL; brown columns, 50 ng/μL; and while columns, 25 ng/μL. The cutoff signal value (OD₄₅₀=0.1) is shown by a dashed line. **(B)** Screening of a HS glycan library consisting of 15 HS glycans using the αTSR-Hisx6, the GST-αTSR, and GST (indicated by blue fonts) as probes, respectively, through electrospray ionization mass spectrometry (ESI-MS). The order (#), name, composition, maximum number of sodium (Max # Na⁺), formula without sodium (Na⁺), and monoisotopic mass (Da) of each HS glycan, as well as their binding affinities in dissociation constants (*K_D*, μM) to each of the GST-αTSR, the αTSR-Hisx6, or GST protein are shown. In the composition column, the numbers of sulfates are shown in red fonts. The *K_D* is an average value (μM) ± standard deviation. NB ≡ no binding.

weeks after the second immunization through the tail vein and two weeks after the final vaccination via heart puncture and sera were processed via an established protocol.⁵⁰

EIAs for α TSR-Specific IgG Titer Determination

EIAs were applied to define α TSR specific IgG antibody titers as described previously.^{51,52} Briefly, gel-filtration purified GST- α TSR (for S₆₀- α TSR, α TSR-Hisx6, or S₆₀ protein immunized mouse sera) or α TSR-Hisx6 (for GST- α TSR protein immunized mouse sera) proteins at 5 μ g/mL were coated to plates as capture antigens. After blocked with nonfat milk, plates were incubated with serially diluted mouse sera. The bound IgG was detected by goat-anti-mouse IgG-HRP conjugate (1:5000, MP Biomedicals). The α TSR-specific antibody titers were defined as maximum dilutions of sera with positive signals (OD₄₅₀ \geq 0.2).

Immunofluorescence Assays (IFAs)

Air dried *P. falciparum* sporozoites were stained using the mouse sera after immunization of the α TSR containing proteins as described previously.^{53,54} Briefly, sporozoites of *P. falciparum* from mosquito salivary glands on slides (kindly provided by Dr. Photini Sinnis at Johns Hopkins University) were warmed to ambient temperature. After blocked with 1% BSA in 1x TBS (pH 7.4), the slide was incubated with 2-fold serially (2000x to 256,000x) diluted mouse sera after immunization of the S₆₀- α TSR nanoparticle or the α TSR-Hisx6 protein (see above) in a humid chamber. After washing, fluorophore-conjugated secondary antibody (Millipore-Sigma) was added. The slide was washed and mounted using Citifluor Mountant Media by placing a cover glass on the slide and seal with nail polish. The slide was viewed with fluorescence microscope at 20x to 40x magnifications. Stain titers were defined as the maximum dilutions of the mouse sera that showed recognizable staining of *Plasmodium* sporozoites.

Statistical Analysis

Statistical differences between two data groups were calculated by GraphPad Prism 9.0 (GraphPad Software, Inc.) using unpaired *t* test. Differences were considered non-significant when *P*-values are >0.05 (marked as “ns”), significant when *P*-values are <0.05 (marked as “*”), highly significant when *P*-values are <0.01 (marked as “**”), and extremely significant when *P*-values are <0.001 (marked as “***”), respectively.

Ethics Statement

All animal studies were completed in compliance with the recommendations in the Guide for the Care and Use of Laboratory Animals (23a) of the National Institute of Health (NIH). The protocols were approved by the Institutional Animal Care and Use Committee (IACUC) of the Cincinnati Children’s Hospital Research Foundation (Animal Welfare Assurance No. A3108-01). No human subjects were involved in this study.

Results

Generation and Characterization of the α TSR-Hisx6 and GST- α TSR Proteins

After expression in *E. coli*, soluble α TSR-Hisx6 protein in the bacteria lysate was purified using the Cobalt resin at a yield of ~15 mg/L bacteria culture. SDS-PAGE using a 16% gel showed the protein band at ~10 kDa (Figure 1B). Similarly, soluble GST- α TSR protein was isolated using the Glutathione Sepharose 4 Fast Flow medium (GE Healthcare Life Sciences) at a yield of ~40 mg/L bacteria culture. SDS-PAGE revealed the GST- α TSR protein at ~35 kDa (Figure 1C). Soluble GST with a ~26 kDa was also produced as a control (Figure 1D). The purified proteins were further analyzed by MS, revealing a molecular weight of 9801 Da for the α TSR-Hisx6 protein (Figure 1E) as monomers, 69,909 Da for the GST- α TSR protein (Figure 1F) as dimers, and 54,281 Da for GST protein (Figure 1G) as dimers.

Binding of the α TSR Fusion Proteins to HS Glycans

The CSP is known to interact with HSPGs as receptors to facilitate *Plasmodium* infection in the liver,^{15,18–20} likely using the α TSR region as the HSPG binding domain,^{20–23} although this has been contradicted in some studies.^{16,24} To clarify

this issue, the two α TSR proteins generated above were tested for their binding to heparin glycans by EIA-based binding assays. The results (Figure 2A) showed that the GST- α TSR fusion proteins bound well heparin, but the α TSR-Hisx6 did not bind. As negative controls, both GST without the α TSR domain did not bind heparin (Figure 2A), indicating the binding signals were from the α TSR protein. It was noted that the α TSR-Hisx6 protein did not bind heparin, consistent with a previous observation.¹⁶

To further verify these observations, the GST- α TSR, α TSR-Hisx6, and GST proteins were used as probes to screen a HS glycan library consisting of 15 HS compounds, respectively, by highly sensitive ESI-MS binding assays. The results (Figure 2B) demonstrated that the GST- α TSR bound all 15 HS glycans in the library, revealing dissociation constants (K_D values) in the μ M range to 13 HS glycans (Figure 2B, GST- α TSR column), which are in typical affinity range for protein-glycan interactions. Noteworthy, the α TSR-Hisx6 protein also bound weakly six HS glycans with K_D values in mM range between 1.0 and 3.9 mM (Figure 2B, α TSR-His column) which were substantially higher than those of the GST- α TSR protein. The remaining nine glycans in the library did not reveal detectable binding to the α TSR-Hisx6 protein. As the negative control, GST did not bound any of the 15 HS compounds (Figure 2B, GST column). While the screening data verified the EIA-based binding results, they also showed that α TSR-Hisx6 protein bound some HS glycans with low affinity. These data strongly suggested that, as part of the CSP, the α TSR may need a fused protein as a foundation to support its proper conformations for high binding affinity to HS glycans, a scenario resembling the VP8* domain of rotavirus VP4 in binding to its glycan receptors^{55–58} (see discussion).

Production of the S₆₀- α TSR Nanoparticle

Soluble S- α TSR protein with a C-terminal Hisx6 tag (Figure 3A) was produced via the *E. coli* system at yields of ~20 mg/L bacterial culture using the Cobalt resin. SDS-PAGE analysis revealed the target protein at expected ~35 kDa (Figure 3B). Gel-filtration chromatography of the protein showed a major (P1) and two minor (P2 and P3) elution peaks (Figure 3C). The major peak (P1) at high molecular weights represented the self-assembled S₆₀- α TSR nanoparticle (see below), while the minor P2 and P3 were at similar positions like the previously observed S-VP8* dimer and S-VP8* monomer,³⁴ suggesting that they were the S- α TSR dimers and monomers, respectively. Negative stain TEM inspection of the P1 sample showed nanoparticles at ~25 nm in diameter (Figure 3D), confirming that most S- α TSR protein self-assembled into S₆₀- α TSR nanoparticle. Interestingly, TEM examination of the P2 and P3 samples also revealed similar nanoparticles (Figure 3E and F) at less numbers compared with that of the P1. This observation suggested that the nanoparticle assembly may be dynamic, in which the S- α TSR dimers and monomers can further assemble into S₆₀- α TSR nanoparticles.

Validation of the S₆₀- α TSR Nanoparticle Formation

The assembly of the S₆₀- α TSR nanoparticle was further demonstrated using mass photometry analysis of the resin-purified S- α TSR protein. The results (Figure 3G) revealed only one form of large complexes with the molecular weights of 1967 ± 84 kDa. This matches well with the expected 1942 kDa of the S₆₀- α TSR nanoparticle as a 60mer (T = 1 icosahedron) of S- α TSR protein with each copy having a mass of 32.37 kDa, which is calculated based on the amino acid sequences. No signals corresponding to a T = 3 (180-mer = 5826 kDa) or a T = 4 (240-mer = 7769 kDa) icosahedral particles of the S- α TSR protein were detected. The remaining signals from the mass photometry analysis did not match with the fold numbers of the S- α TSR protein, thus may represent the co-purified bacterial proteins (Figure 3B) or degraded forms of the S₆₀- α TSR nanoparticle. The formation of the S₆₀- α TSR nanoparticle was further demonstrated by 3D structural reconstruction (see below).

Density Analysis of the S₆₀- α TSR Nanoparticle

The resin purified S- α TSR protein was analyzed by CsCl density gradient ultracentrifugation. After centrifugation, the S- α TSR protein formed a visible band in the top quarter of the gradient (data not shown). EIA assays to detect the S- α TSR protein in the 22 fractions of the gradient confirmed that the S- α TSR protein formed a peak around fraction 17 (Figure 4A, F17) with a density of 1.305 g/mL. Negative stain TEM inspection of fraction 17 showed uniform S₆₀- α TSR nanoparticles in ring-like images (Figure 4B and C). It was noted that, unlike the S₆₀ nanoparticle with smooth surface,³⁴

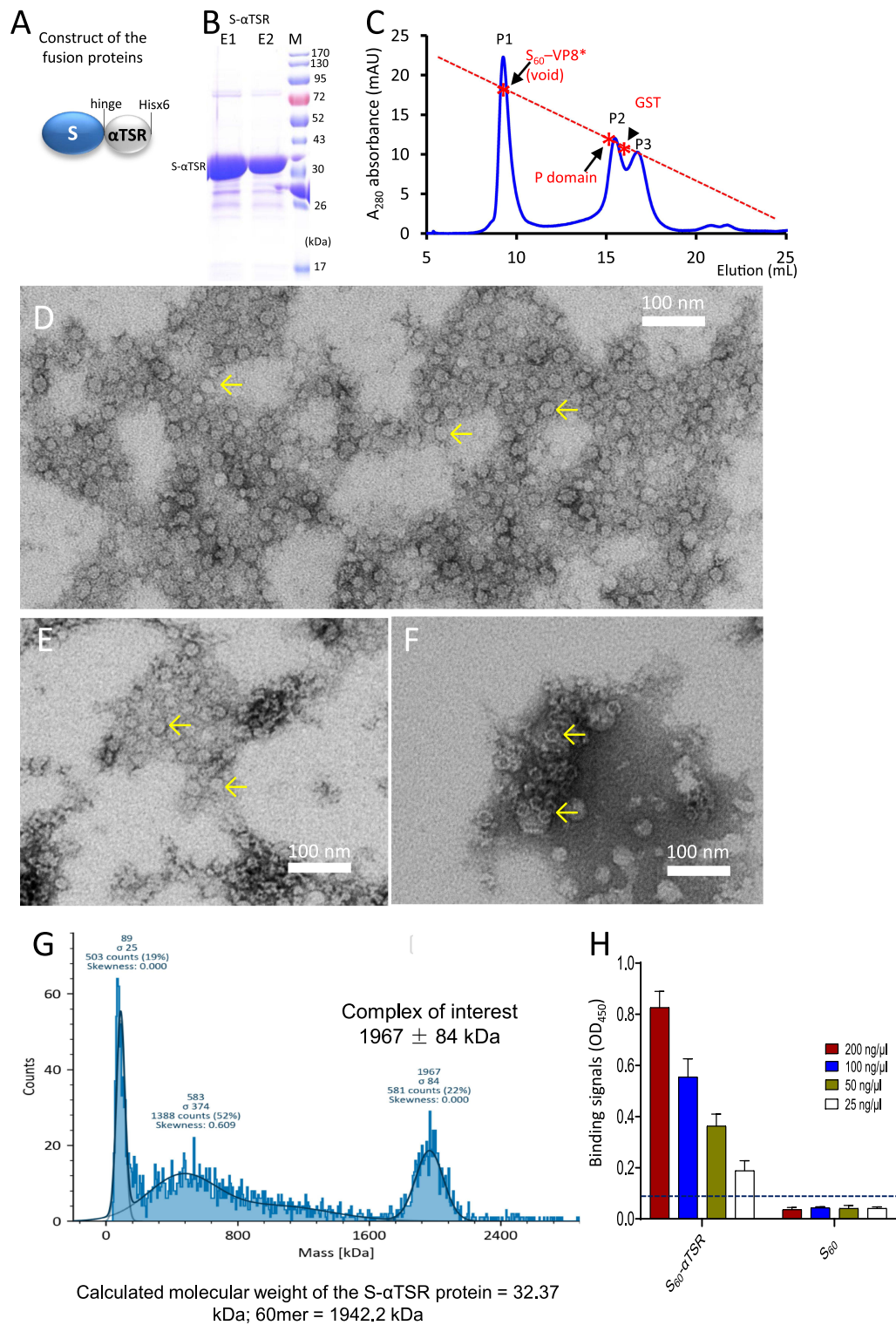


Figure 3 Production of the S₆₀-αTSR nanoparticles. **(A)** Schematic construct of the S-αTSR fusion protein with a C-terminal Hisx6 tag. S, modified norovirus S domain; hinge, the hinge region of norovirus VP1. **(B)** SDS-PAGE analysis of the resin purified S-αTSR fusion protein showing a band at ~32 kDa. M, protein standards with indicated molecular sizes. **(C)** A gel-filtration elution curve of the S-αTSR protein through a size exclusion column, showing a major peak (P1, void volume) representing the S₆₀-αTSR nanoparticle and two minor peaks (P2 and P3) representing the dimers and monomers of the S-αTSR protein. The elution peaks were calibrated using the S₆₀-VP8* nanoparticle (~3.4 mDa) [20], norovirus P dimer (~69 kDa),³⁸ and GST dimers (54 kDa) with their elution positions being shown by star symbols and arrows on a red dashed line. Y-axis shows UV absorbances at A₂₈₀ (mAU), while X-axis indicates elution volume (mL). **(D–F)** Micrographs of transmission electron microscopy (TEM) of protein samples from the P1, P2, and P3, respectively, showing typical S₆₀-αTSR nanoparticles (arrows). **(G)** Molecular weight determinations of the S₆₀-αTSR nanoparticle by mass photometry. The complex of interest showed a molecular weight of 1967 ± 84 kDa, matching well with calculated molecular weight (1942.2 kDa) of the S₆₀-αTSR nanoparticle. **(H)** Binding of the S₆₀-αTSR proteins to heparin sodium salt by EIA based binding assays. Y-axis shows binding signals (optical density (OD)₄₅₀), while X-axis denotes the S₆₀-αTSR proteins and its control proteins without the αTSR domain at indicated concentrations. The cutoff signal value (OD₄₅₀=0.1) is shown by a dashed line.

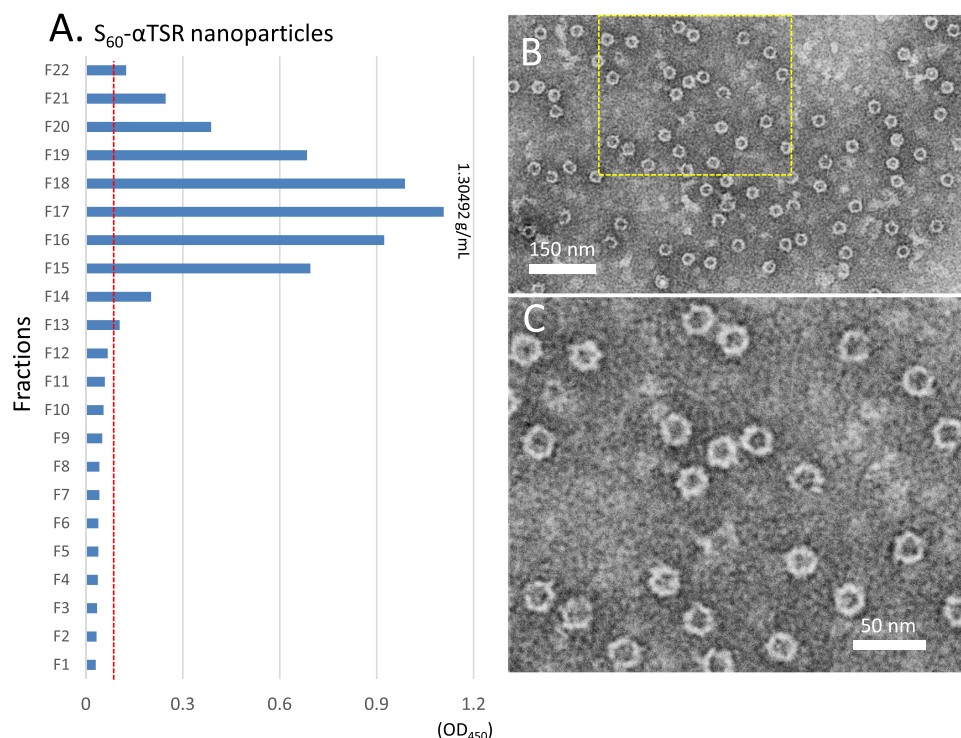


Figure 4 Analysis of the S₆₀-αTSR nanoparticles after purification by cesium chloride (CsCl) density gradient centrifugation. **(A)** Following centrifugation, the CsCl density gradient containing the S₆₀-αTSR nanoparticles was fractionated into 22 portions. The relative concentrations of the S₆₀-αTSR protein in the fractions were measured by EIA assays using antibody against norovirus VLP. Y-axis indicates the fraction numbers, while X-axis shows signal intensities in optical density (OD₄₅₀) with a red dashed line showing the cut-off signal at OD₄₅₀=0.1. **(B and C)** A TEM micrograph of the S-αTSR protein of fraction 17 showing uniform S₆₀-αTSR nanoparticles **(B)** with an enlargement of the framed area in **(C)** showing recognizable protrusions of αTSR antigens on the surface.

the S₆₀-αTSR nanoparticles revealed recognizable protrusions on the surface representing the displayed αTSR antigens (Figure 4C). These nanoparticle images appeared to be more uniform compared with those of gel-filtration purified ones (compared Figure 3D with Figure 4B and C).

Production of Tag-Free S₆₀-αTSR Nanoparticles

A scalable method was also developed for production of tag-free S₆₀-αTSR nanoparticles, which contains two major steps. First, after expression the tag-free S-αTSR protein (Figure 5A) was precipitated from the bacterial lysate using 1.6 M (NH₄)₂SO₄ (Figure 5B). Second, an anion exchange chromatography was performed to separate the (NH₄)₂SO₄ precipitated proteins, resulting in several elution peaks (Figure 5C). The S-αTSR protein was eluted in peak 6 (P6), corresponding to a NaCl concentration of 284 mM (28.4% buffer B), reaching to a purity of >90% (Figure 5D). TEM inspection confirmed the assembly of S₆₀-αTSR nanoparticles by the tag-free S-αTSR protein (Figure 5E).

Structure of the S₆₀-αTSR Nanoparticle

The 3D structure of the Hisx6-tagged S₆₀-αTSR nanoparticle was reconstructed at a resolution of ~20 Å based on the negative stain TEM images using JSPR. The final structure indicated that each particle contains 60 S-αTSR proteins, organizing into a T = 1 icosahedral symmetry with a diameter of ~23 nm (Figure 6A). Sixty surface protrusions were noted, representing the displayed αTSR domains. The section view shows the outer αTSR domains, the middle S₆₀ shell, and a central lumen (Figure 6B), respectively. Fitting of the S₆₀ particle model into the density map confirms S₆₀ in the interior shell region of the S₆₀-αTSR nanoparticle (Figure 6D–F). This fitting also left the outer protrusions unoccupied, confirming the αTSR regions. The crystal structures of *P. falciparum* αTSR (PDB code: 3VDJ)¹⁶ fitted into the density maps of the protrusions well (Figure 6G and H). A zoom-in view of the fitting indicated that the N-terminal ends of the

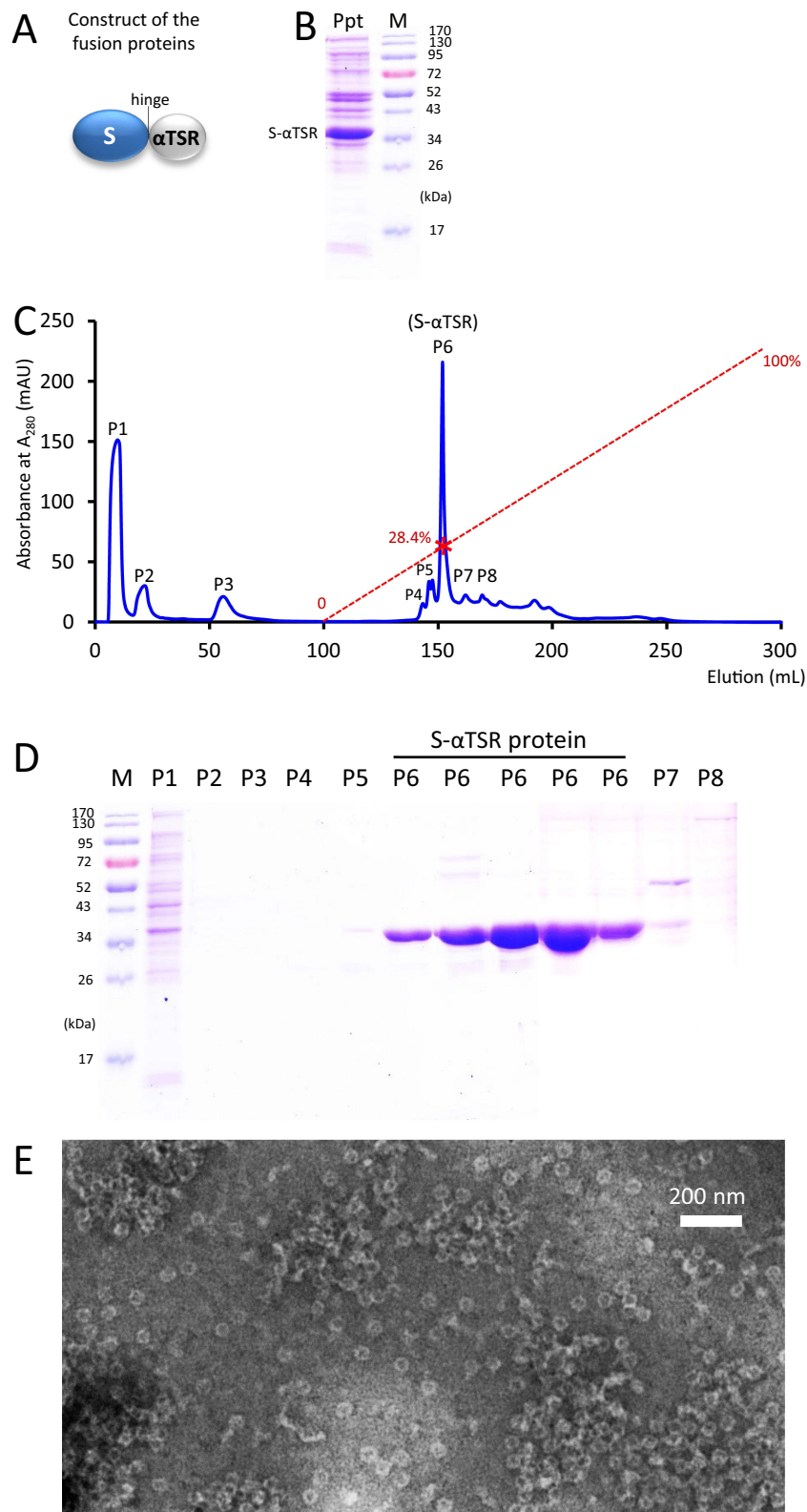


Figure 5 Production of tag-free S_{60} - α TSR nanoparticles. **(A)** The schematic construct of the tag-free S - α TSR proteins. **(B)** SDS-PAGE showing the ammonium sulfate precipitated S - α TSR protein (Ppt) from the bacterial lysis. **(C)** An anion exchange elution curve of the ammonium sulfate precipitated proteins from **(B)**. Y-axis shows UV absorbances at A_{280} (mAU), whereas X-axis indicates elution volume (mL). The red dashed line indicates the linear increase of the elution buffer B (0–100%) with a red star symbol indicating the percentage of buffer B at the elution position of the S - α TSR protein (28.4%). Eight elution peaks (P1 to P8) that were analyzed by SDS-PAGE are indicated. **(D)** SDS-PAGE analyses of the eight elution peaks from the ion exchange **(C)**. Five fractions from P6 were analyzed. M in **(B)** and **(D)** is protein standard with the molecular weights as indicated. The S - α TSR protein is eluted in P6. **(E)** A micrograph of transmission electron microscopy (TEM) of the protein sample from P6 showing typical S_{60} - α TSR nanoparticles.

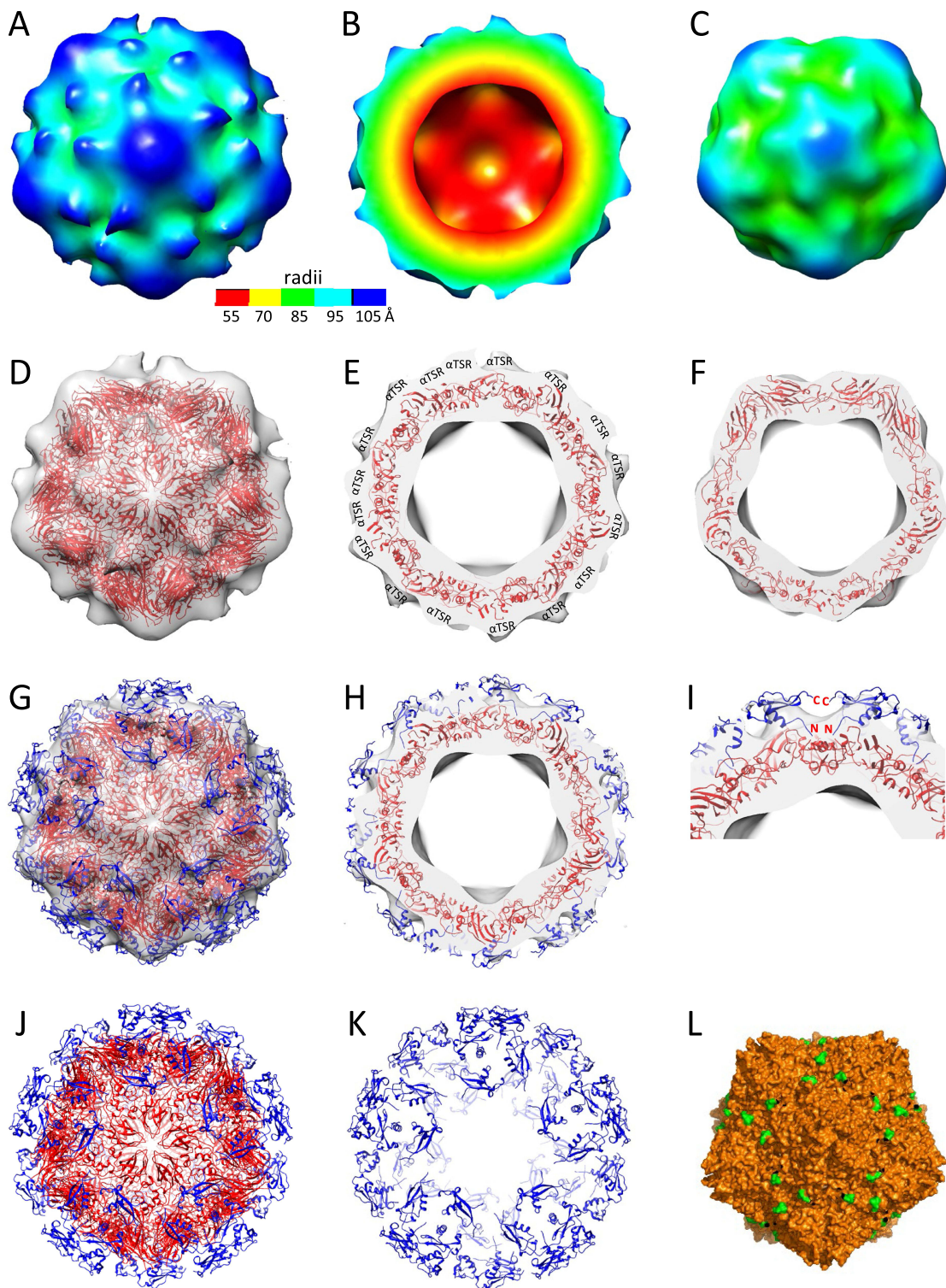


Figure 6 3D structures of the S_{60} - α TSR nanoparticle. (A to C) The S_{60} - α TSR nanoparticle in surface view (A) showing its external structure; section view (B) showing its cross section, internal lumen, and interior structure, as well as surface view (C) of the S_{60} inner shell, respectively, revealing a $T = 1$ icosahedral symmetry. The colored bar shows the radii of the structures in different color schemes. (D–F) Fitting of a GII.4 norovirus S_{60} nanoparticle model (strain VA387, red cartoon representation) into the electron density map of the S_{60} shell region, showing in full (D) and slice (E) views, as well as slice view of the S_{60} shell (F), respectively. The blank protrusions representing the displayed α TSR antigens are indicated in (E). (G and H) Fitting of the crystal structures of 60 α TSR antigens (blue cartoon representation) into the electron density maps of the protrusion regions, showing in full (G) and slice (H) views respectively. (I) A zoom-in view of the fitting region of (H) showing the N- and C-terminal ends of two α TSR antigens. (J and K) Structural model of the S_{60} - α TSR nanoparticle (J), with its surface displayed α TSR antigens (K). (L) Crystal structure of the inner shell of the 60-valent feline calicivirus (FCV) VLP (PDB code: 4PB6, Orange surface representation) showing the surface exposed C-terminal ends of the S domain. All images are shown at five-fold axes.

α TSRs go inward connecting to the inner shell (Figure 6I). This fitting further justified the S₆₀ nanoparticle shell and the displayed α TSR domains, the two components of the S₆₀- α TSR nanoparticle.

While the locations of the protrusions match well the C-termini of the S₆₀ nanoparticle (Figure 6, compared A with L), the sizes and shapes of the protrusion's distal heads fit less well with the α TSR crystal structures. This may be due to the flexibility of the displayed α TSR domains relative to the S₆₀ shell, attributing most likely to the flexible hinge and linker at the C terminal end of the S domains, as well as the flexible loop at the N terminus of the α TSR antigen. This phenomenon is commonly observed in structural biology. Based on the fitting, a structural model of the S₆₀- α TSR nanoparticle in PDB format was made (Figure 6J), showing its 60-surface displayed α TSR antigens, providing a structural basis for the high α TSR specific immune response in mice (see below).

The Nanoparticle-Displayed α TSRs Retained HS Glycan Binding Function

The S₆₀- α TSR nanoparticle was examined for their ability to bind HS glycan receptors. EIA-based binding assays showed that the resin purified S₆₀- α TSR nanoparticle bound heparin (Figure 3H); as a negative control the S₆₀ nanoparticle did not bind, indicating that the nanoparticle displayed α TSR antigens retained receptor binding function. The data further suggest that the nanoparticle-presented α TSR domains maintained their original structure and conformations, supporting the notion that the S₆₀- α TSR nanoparticle can be used as a vaccine candidate.

The S₆₀- α TSR Nanoparticle Enhanced Immune Response Toward the Displayed α TSR Antigens

Literatures showed that antigens with pathogen-associated molecular patterns (PAMPs) in polyvalent nature show higher immunogenicity compared with ones in mono- or low-valence.^{33,59-61} To test if this is true, mice were immunized with the S₆₀- α TSR nanoparticle using the dimeric GST- α TSR (Figure 1F) and the monomeric α TSR-Hisx6 (Figure 1E) proteins for comparisons. After two and three immunizations, the nanoparticle elicited high α TSR-specific IgG titers, reaching to 10,667 and 307,200, respectively (Figure 7A). These titers were at least 3.3 folds (after two immunizations) or 17.1 folds (for three immunizations) higher than those induced by the dimeric GST- α TSR protein or those elicited by the monomeric α TSR-Hisx6 protein (P s < 0.01). Thus, the nanoparticle displayed α TSR antigens improved their immune responses significantly.

The Nanoparticle Immunized Mouse Sera Stain Sporozoite at High Titer

To further show that the α TSR-specific antibody bound the CSPs on *Plasmodium* sporozoites as a potential indication of the antibody to inhibit the receptor binding function of the CSPs and thus block the infection of the parasites, IFAs were performed to stain the *P. falciparum* sporozoites using the above mouse sera in serial dilutions between 1000x and 256,000x. The results showed that sporozoites were stained specifically by the mouse sera (Figures 7B and 8). Specifically, mouse sera after immunization with the S₆₀- α TSR nanoparticle stained the *P. falciparum* sporozoites at a high titer of 66,560 that was 14.8 folds higher than the staining titer (4480) of sera after immunization with the α TSR-Hisx6 (Figure 7B). Several typical IFA micrographs, representing 16,000x (Figure 8A and B) and 128,000x (Figure 8C and D) dilutions of the sera after immunization with the S₆₀- α TSR nanoparticle, respectively, and 4000x dilution of the sera after immunization with the α TSR-Hisx6 protein (Figure 8E and F) are shown as examples. As a negative control, mouse sera after immunization with the S₆₀ nanoparticle did not stain the *P. falciparum* sporozoites even at much lower dilutions (Figure 8G and H). It was noted that, although many junks on the slides were seen under optical field using transmitted light, only the sporozoites are stained by the sera after immunization with the α TSR proteins specifically, supporting the potential of the S₆₀- α TSR nanoparticle as a malaria vaccine candidate.

Discussion

In this study, we first characterized two α TSR fusion proteins and provided solid evidence for the first time that the α TSR domain interacts with HS glycans and thus is a key functional domain of the CSP and an important vaccine target. Secondly, we generated a protein nanoparticle to display the α TSR antigens for enhanced immune response toward the

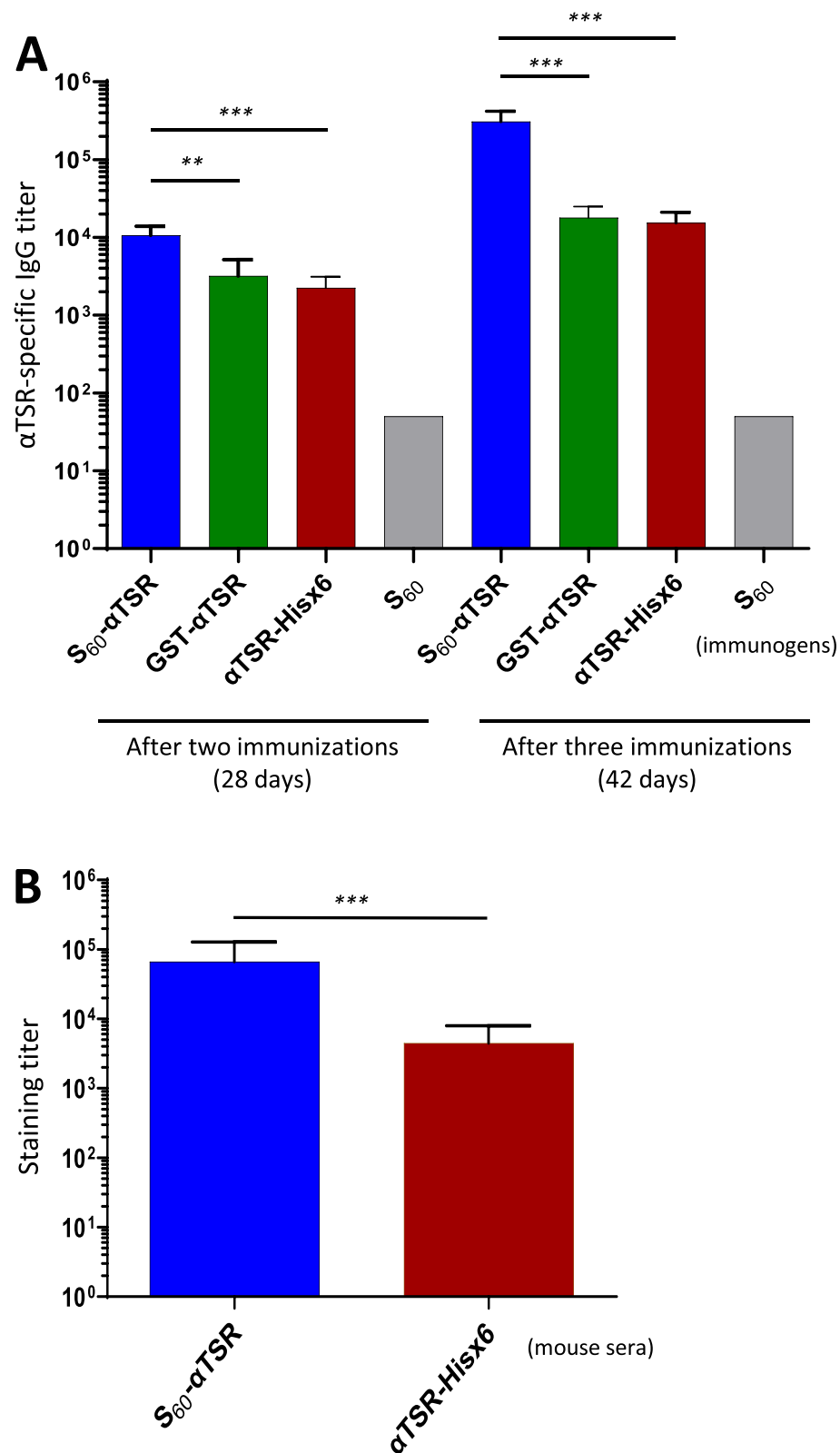


Figure 7 Immune response of the S_{60} - α TSR nanoparticle in mice. **(A)** α TSR-specific IgG titers induced by the polyvalent S_{60} - α TSR nanoparticle (S_{60} - α TSR), the dimeric GST- α TSR protein (GST- α TSR), and the monomeric α TSR-Hisx6 protein (α TSR-Hisx6) after two (left four columns) and three (right four columns) immunizations are shown. Y-axes indicate α TSR-specific IgG titers, while X-axes indicate various immunogens. The S_{60} nanoparticle is a negative control. **(B)** Staining titers of the mouse sera after three immunizations of the S_{60} - α TSR nanoparticle (S_{60} - α TSR), and the α TSR-Hisx6 protein (α TSR-Hisx6) to air-dried *P. falciparum* sporozoites. Y-axis indicates staining titers, while X-axes indicate the two mouse sera. Statistic differences between data groups are shown as “***” for highly significant with P -values < 0.01 or “****” for extremely significant with P -values < 0.001.

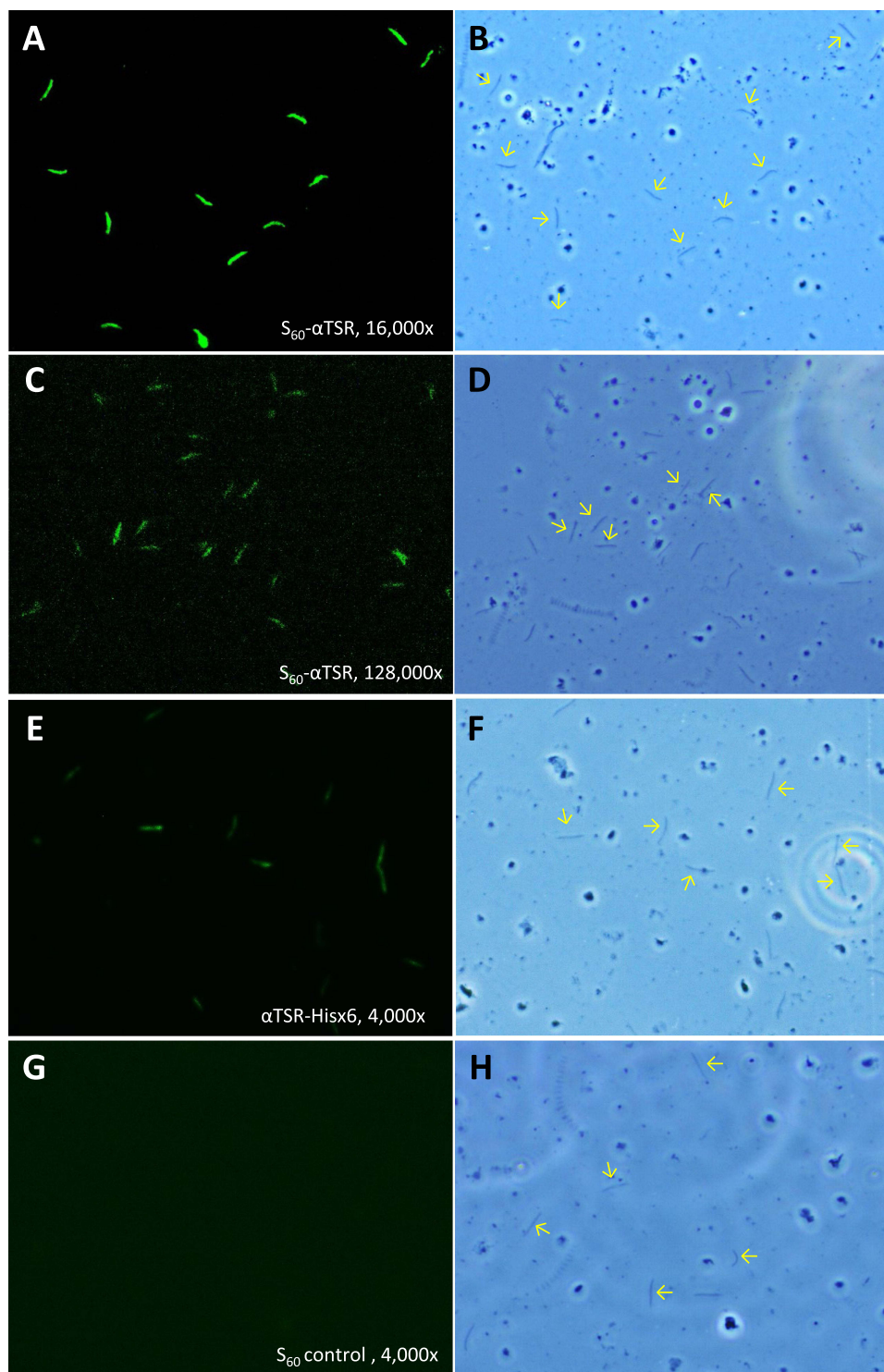


Figure 8 Representative micrographs of immunofluorescence assays (IFAs) of *P. falciparum* sporozoites by the mouse sera after immunization with the S_{60} - α TSR nanoparticle and controls. **(A–D)** Typical IFA micrographs representing 16,000x **(A and B)** and 128,000x **(C and D)** dilutions, respectively, of the sera after immunization with the S_{60} - α TSR nanoparticle. **(E and F)** Typical IFA micrographs representing 4000x dilution of the sera after immunization with the α TSR-Hisx6 protein. **(G and H)** Typical IFA micrographs representing 4000x dilution of the sera after immunization with the S_{60} nanoparticle without α TSR antigens. Each pair of micrographs consists of an IFA staining result **(A, C, E and G)** and the optical view of the same field **(B, D, F and H)**. Arrows in the optical field of view indicate some sporozoites showed in the corresponding IFA views.

displayed α TSR antigens, with a long-term goal to develop it into an effective malaria vaccine. The S_{60} - α TSR nanoparticle with or without a tag can be produced via the *E. coli* system. Both structural and functional analyses revealed that 60 α TSR antigens are properly presented on the surface of the nanoparticle with authentic receptor binding function. Immunological evidence showed that the polyvalent S_{60} - α TSR nanoparticle significantly improved the immunogenicity of the displayed α TSR antigens, eliciting high titer of α TSR-specific antibody that bound the CSPs of *P. falciparum* sporozoites at high titer. Therefore, the S_{60} - α TSR nanoparticle may serve as a promising malaria candidate vaccine.

The CSP is known to recognize HSPG receptors to allow sporozoites being arrested in the liver, which is an important step of *Plasmodium* infection to humans.^{15,18–20} The TSR (region II plus) of the CSP was suggested to be the HSPG-binding domain,^{20–23} although discrepancy was also reported.^{16,24} A critical limitation of the previous studies aiming to map the HSPG-binding domain of the CSP was their usage of short synthetic peptides, mostly being smaller than 25 amino acids in length. Based on the known crystal structure,¹⁶ the α TSR domain assembles into a highly specific 3D structure (Figure 9), in which amino acid residues that are distant in linear sequence may be close in 3D structure. Therefore, the HSPG binding site on the α TSR domain is most likely conformational, which is unlikely to be well represented by a short synthetic peptide. Particularly, the crystal structure reveals two disulfide bonds that are required for the proper folding of the α TSR domain,¹⁶ but none of the tested peptides contains these two pairs of cysteines (Figure 9E).

It is noted that, among the four peptides (Pf3 to Pf6) within the α TSR domain that were tested for HS glycan binding, only the longest one with 32 residues (Pf6, 341-IQVRIKPGSANKPKDELTDYANDIEKKICKMEK-372) bound heparin/heparan sulfate.²⁴ Analysis of the four peptides on the known α TSR crystal structure¹⁶ (Figure 9A–D) showed that only Pf6 contains full sequences to form the intact top structure of the α TSR domain (Figure 9D, purple portion), suggesting that the top moiety formed by sequences from I346 to I364 might contain the HSPG binding site. Noteworthy, the relative closeness between the N- and C-termini at the basis of the α TSR domain¹⁶ implies that the top region may form a distal head extending to the surface of parasite sheath. However, as discussed above, whether the synthetic peptides can fold into properly functional conformations remains in question. Particularly, none of these peptides contain the four cysteines for the disulfide bonds that are necessary for proper α TSR structures (Figure 9E).¹⁶

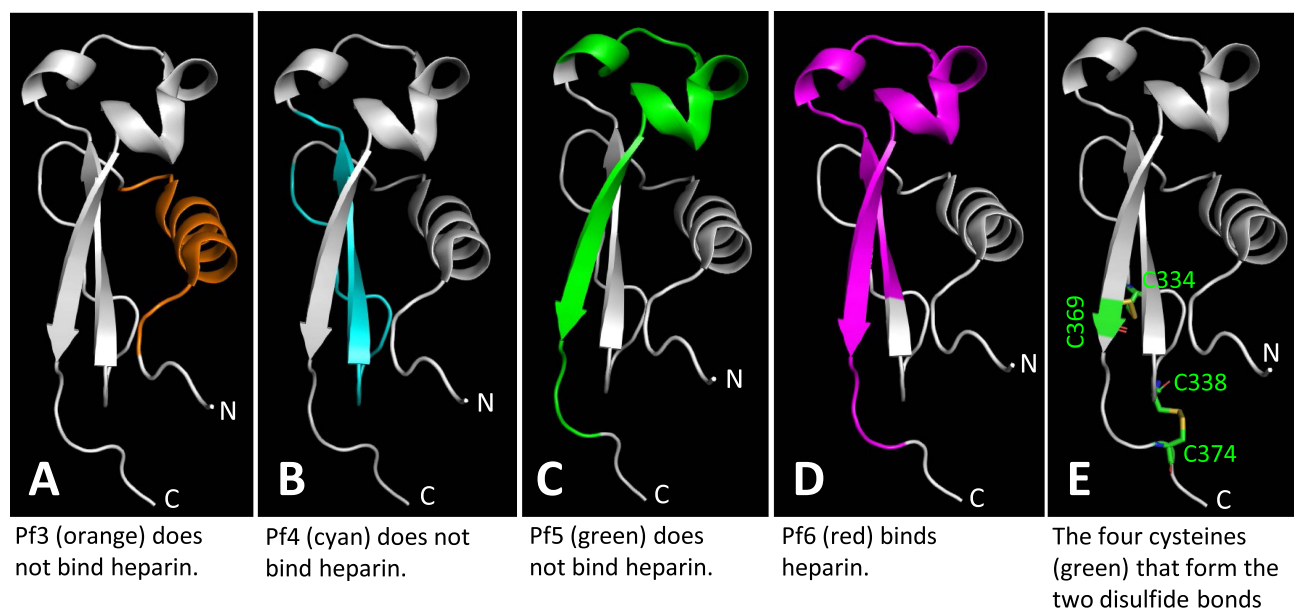


Figure 9 Structural analyses of four tested α TSR related peptides for binding to heparan sulfates and the two disulfide bonds reported in literature. The regions and structures of the four α TSR peptides, Pf3 (A, Orange), Pf4 (B, cyan), Pf5 (C, green), and Pf6 (D, purple),²⁴ corresponding to the known crystal structure of the α TSR¹⁶ (in grey) are shown. The four cysteines forming the two disulfide bonds are shown in green in (E).

In a study focusing on the α TSR structure by crystallography, Doud et al showed that the C-terminally Hisx6-tagged α TSR did not reveal binding signals using a heparin-sepharose column.¹⁶ Similarly, we did not observe binding signals of the α TSR-Hisx6 protein to heparin in an EIA-based binding assay in this study. However, employing highly sensitive ESI-MS binding assays, we detected weak binding of the α TSR-Hisx6 to six different HS compounds in a HS glycan library with K_D values in the mM range (Figure 2). Thus, the α TSR-Hisx6 does bind HS glycans at low affinity, which cannot be detected by an EIA- or column-based binding assays. Strikingly, unlike the α TSR-Hisx6 protein, two other α TSR fusion proteins, the GST- α TSR and S₆₀- α TSR, bound both heparin and many other HS glycans in the HS glycan library strongly with K_D values in the μ M range (Figure 2). Since the Hisx6 tag is small, the α TSR-Hisx6 is basically a free protein (Figure 1E), whereas the α TSR in the GST- α TSR and the S₆₀- α TSR proteins was supported by large (>21 kDa) protein. This scenario reminds us of the rotavirus VP8* domains. As the distal head of rotavirus spike protein VP4, the recombinant VP8* domain alone does not show detectable binding to its glycan receptors in EIA-based glycan binding assays, but GST-VP8* fusion proteins,^{55–58} S₆₀-VP8*^{34,62} and P₂₄-VP8*⁴⁰ bound the same glycans well.

While the mechanism behind this phenomenon remains elusive, one explanation is that the α TSR domain, like rotavirus VP8* domain, is a part of a large protein and may need a structural support from a big protein to confer proper conformations for optimal binding affinity to the HS glycans. Structural support by a big protein may mimic the scenario that α TSR is a part of the CSP with ~400 amino acids, like the case of VP8* domain as a part of rotavirus VP4. As a next step, we plan to identify the HS glycan binding site of the α TSR domain using nuclear magnetic resonance (NMR) and crystallography, following the footsteps of our previous studies on rotavirus VP8* domain interacting with its glycan receptors.^{63,64}

The CSP has been shown as an immunodominant, protective antigen.²⁵ We have now shown that the α TSR is a key functional domain of the CSP and the α TSR domain has been demonstrated to contain polymorphic T-cell epitopes eliciting CD4⁺ T-cell response with protection,³⁰ strongly supporting that the α TSR is an excellent vaccine target. However, since the α TSR domain is small (~9.8 kDa) with apparently low immunogenicity in its free, monomeric form (Figure 7), a polyvalent platform will help to enhance its immune response. The 60 valent S₆₀ nanoparticle has been shown to be a potent platform to display antigens for improved immunogenicity.^{34,35,65,66} Our results in this study showed that the S₆₀- α TSR nanoparticle elicited significantly higher α TSR-specific antibody titers than those induced by the monomeric α TSR-Hisx6 or the dimeric GST- α TSR proteins. The mouse antisera after immunization with the S₆₀- α TSR nanoparticle bound the CSPs of the *P. falciparum* sporozoites at a very high titer. We assume that such antibody- α TSR binding would block the HSPG binding function of the CSPs and thus would be able to prevent the liver invasion by the *P. falciparum* sporozoites, making the S₆₀- α TSR nanoparticle a promising malaria vaccine candidate. Due to the lack of a suitable facility, we are reaching out to potential collaborators to evaluate the neutralization and/or protective efficacy of our S₆₀- α TSR nanoparticle vaccine.

Another advantage of our technology is that the S₆₀- α TSR nanoparticle can be produced using the *E. coli* system, providing a low-cost method for future production of our vaccine. This is particularly important because malaria occurs mostly in the source-deprived developing countries. The known crystal structure of the α TSR protein expressed by yeast *Pichia* did not reveal any glycosylation,¹⁶ indicating that production of our S₆₀- α TSR nanoparticle via the *E. coli* system is appropriate. In fact, the data that the *E. coli* expressed S₆₀- α TSR nanoparticle bound HS glycans and that the α TSR-specific antibody elicited by the S₆₀- α TSR nanoparticle bound authentic CSPs on *P. falciparum* sporozoites supported our hypothesis.

Conclusion

In summary of the data presented in this study, we conclude that the α TSR is an important receptor-binding domain of the CSP and serves as an ideal vaccine target. The S₆₀- α TSR nanoparticle displaying multiple α TSR antigens may be a promising vaccine candidate against attachment and infection of *Plasmodium* parasites.

Abbreviations

TSR, thrombospondin type-I repeat; CSP, circumsporozoite protein; HSPG, heparan sulfate proteoglycan; HS, heparan sulfate; 3D, three-dimensional; WHO, the World Health Organization; GPI, glycosylphosphatidylinositol; GSK, GlaxoSmithKline; VLP,

virus-like particle; BSA, bovine serum albumin; SDS-PAGE, Sodium dodecyl sulfate polyacrylamide gel electrophoresis; FPLC, fast performance liquid chromatography system; TEM, transmission electron microscopy; FCV, feline calicivirus; EIA, enzyme immunoassay; ESI-MS, electrospray ionization mass spectrometry; HRP, horse-radish peroxidase; IFA, immunofluorescence assay; NIH, National Institute of Health; IACUC, Institutional Animal Care and Use Committee; Ns, non-significant.

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Disclosure

Ming Tan has a financial interest in the S particle vaccine platform technology that Cincinnati Children's Hospital Medical Center licensed to Blue Water Vaccines, Inc. The authors report no other conflicts of interest in this work.

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