



Bioinformatics-based Design of a Multi-epitope Chimeric Vaccine Candidate Targeting *Toxoplasma gondii*

Ebrahim Nayeri¹, Tooran Nayeri^{2,*}, Mahdi Fasihi-Ramandi³ and Ezatollah Ghasemi^{2,4}

¹Taleghani Hospital, Kermanshah University of Medical Sciences, Kermanshah, Iran

²Infectious and Tropical Diseases Research Center, Dezful University of Medical Sciences, Dezful, Iran

³Molecular Biology Research Center, Baqiyatallah University of Medical Sciences, Tehran, Iran

⁴Department of Medical Parasitology, School of Medicine, Dezful University of Medical Sciences, Dezful, Iran

Abstract:

Introduction: The development of an effective vaccine against *Toxoplasma gondii* (*T. gondii*) remains complex and challenging due to the biological complexity and antigenic diversity of this parasite and requires further study. Therefore, this study aimed to design a novel chimeric protein as a vaccine candidate containing four immunogenic antigens (GRA1, SAG1, ROP18, and MIC13).

Methods: Amino acid positions from GRA1 (81-180), SAG1 (201-300), ROP18 (321-420), and MIC13 (201-300) were selected and linked using A(EAAAK)A. The secondary and tertiary structures, antigenicity, allergenicity, physicochemical properties, codon optimization, and the secondary structure of the mRNA of the GSRM (GRA1-SAG1-ROP18-MIC13) protein were analyzed using various bioinformatics tools.

Results: The GSRM protein had 442 amino acids with a Molecular Weight (MW) of 47,953.46 Da and a theoretical Isoelectric Point (pI) of 5.61. An aliphatic index of 80.11 was obtained for the construct. With an instability index of 43.07, the protein appears to exhibit moderate stability. The GRAVY of the protein was estimated to be -0.316, indicating that the protein is hydrophilic. The results also showed that this protein is antigenic and non-allergenic. Based on the evaluation of tertiary structures, GSRM was selected as the best structure for the vaccine candidate. Furthermore, analysis of the secondary structure of mRNA ($\Delta G = -387.70$ kcal/mol) revealed the absence of stable hairpin structures at the 5' end, and therefore translation is possible.

Discussion: *In silico* analyses indicate that GSRM possesses favorable immunological and physicochemical properties. Combining antigenic fragments from different life stages of *T. gondii* may enhance the effectiveness of the immune response.

Conclusion: The accuracy and precision of the *in silico* analyses should be confirmed by subsequent experimental studies *in vitro* and *in vivo*.

Keywords: *Toxoplasma gondii*, *In silico*, GRA1, SAG1, ROP18, MIC13.

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*Address correspondence to this author at the Infectious and Tropical Diseases Research Center, Dezful University of Medical Sciences, Dezful, PC 6461643993, Iran; Tel: 06142429731; E-mail: tooran.nayeri@gmail.com

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1. INTRODUCTION

Toxoplasma gondii (*T. gondii*) is an obligate intracellular protozoan parasite that belongs to the phylum Apicomplexa. This parasite has been reported in a wide range of hosts

(warm-blooded and cold-blooded animals) [1, 2]. Tachyzoites, bradyzoites (found in tissue cysts), and sporozoites (found in oocysts) are the three main stages in the parasite's life cycle. Tachyzoites are involved in the

acute phase of infection, and bradyzoites are associated with the chronic phase [3]. Infection in humans occurs through consumption of raw or undercooked meat containing cysts, ingestion of food or water contaminated with oocysts, or through the placenta during pregnancy [1, 4]. In healthy people, the infection is usually asymptomatic. However, in pregnant women, the parasite can cross the placenta and be transmitted to the fetus [5]. Infection can have dangerous consequences in patients with human immunodeficiency virus, organ transplant recipients, and cancer patients [6]. Moreover, this parasite causes significant economic losses in livestock, especially sheep and goats, worldwide [7].

The drugs used to treat toxoplasmosis are a combination of pyrimethamine and sulfonamides such as sulfadiazine [8]. However, these drugs are only effective on tachyzoites and often have various side effects such as hypersensitivity, hematotoxicity, teratogenicity, and allergic reactions [9, 10]. Given these limitations, the best option is to develop an effective vaccine to control human and animal toxoplasmosis. Recent advances in immunoinformatics have enabled the prediction of T and B-cell epitopes and the identification of potential vaccine targets. Compared to traditional vaccine design, *in silico* approaches offer notable advantages, including enhanced efficiency, reduced cost, and shortened time of experimental stages. Vaccine efficacy remains limited due to the complex life cycle of *T. gondii*, which includes tachyzoite, bradyzoite, and sporozoite stages and multiple antigens. Therefore, multi-epitope vaccine approaches that target antigens from different stages of parasite development may provide greater and more potent protection against toxoplasmosis [11]. Among the *T. gondii* antigens involved in protective immunity, some of the Surface Antigens (SAGs), Microneme Proteins (MICs), Rhoptry Proteins (ROPs), and Dense Granule Antigens (GRAs) are considered potential antigen targets. GRA1 is secreted by tachyzoites, bradyzoites, and sporozoites and contributes to parasite survival within the parasitophorous vacuole [12]. SAG1, a dominant surface antigen of tachyzoites, elicits both humoral and cellular immune responses [13]. ROP18, a rhoptry kinase, plays a key role in virulence and intracellular survival [14, 15], while MIC13 is involved in host cell adhesion and parasite dissemination through the intestinal epithelium [16]. Given these roles, the present study sought to design a chimeric protein vaccine candidate containing immunodominant fragments of the GRA1, SAG1, ROP18, and MIC13 antigens.

2. MATERIALS AND METHODS

2.1. Retrieval of Protein Sequences

Amino acid sequences of the four *T. gondii* antigens (GRA1, SAG1, ROP18, and MIC13) from the RH strain were retrieved in FASTA format from the publicly available sequence database, including the National Center for Biotechnology Information (NCBI; <https://www.ncbi.nlm.nih.gov/>) and Universal Protein (UniProt; <http://www.uniprot.org/>). These sequences served as the basis for subsequent immunoinformatics analyses.

2.2. Prediction of Transmembrane Domains and Signal Peptides

TMHMM Server (version 2.0) (<http://www.cbs.dtu.dk/services/TMHMM-2.0/>) was used to identify transmembrane domains of the four proteins (GRA1, SAG1, ROP18, and MIC13) [14]. The UniProt database (<http://www.uniprot.org/>) and SignalP 4.1 server (<http://www.cbs.dtu.dk/services/SignalP/>) were used to evaluate the peptide signal of these proteins [17]. These analyses were performed to exclude protein fragments located within membrane-spanning regions and to ensure that selected antigenic sequences are accessible to the host immune system.

2.3. B-Cell Epitope Identification

The Immune Epitope Database (IEDB) online service (<http://tools.iedb.org/bcell/>) was used to predict the B-cell epitopes using various parameters, including antigenicity, flexibility, surface accessibility, hydrophilicity, and beta-turn propensity [14].

2.4. Prediction of MHC Class I and II Binding Epitopes

To determine peptides capable of binding mouse Major Histocompatibility Complex (MHC) alleles, predictions were conducted using the PRED (BALB/c) web server (<http://cvc.dfci.harvard.edu/balbc/>). For class II molecules, alleles H2-IA_d and H2-IE_d were analyzed; for class I, H2-K_d, H2-L_d, and H2-D_d were included [18].

2.5. Construction of the Chimeric Protein

To achieve the best placement of epitopes derived from proteins GRA1, SAG1, ROP18, and MIC13 in the structure of the chimeric antigen, four immunodominant peptide domains with different arrangements were fused by a rigid linker A(EAAAK)A.

2.6. Secondary and Tertiary Structure Prediction

The secondary structure of the protein was predicted using online services, namely Garnier-Osguthorpe-Robson (GOR) IV (https://npsa-prabi.ibcp.fr/cgi-bin/npsa_automat.pl?page=npsa_gor4.html) [17]. Moreover, the tertiary structure of the chimeric protein was predicted by the ITASSER server (<https://zhanglab.ccmb.med.umich.edu/I-TASSER/>) [19].

2.7. Structural Validation

Validation of the tertiary structure of the chimeric protein was performed by the analysis of the Ramachandran plot (<https://swissmodel.expasy.org/assess>) [18].

2.8. Prediction of Antigenicity, Allergenicity, and Solubility

VaxiJen (version 2.0) (<http://www.ddgpharmfac.net/vaxijen/VaxiJen/VaxiJen.html>) estimated the antigenicity of this protein [20]. Allergenicity was assessed with the AlgPred server (<http://www.imtech.res.in/raghava/algpred/>) [18]. Protein solubility was estimated after

expression in *Escherichia coli* (*E. coli*) using the SOLpro tool (<http://scratch.proteomics.ics.uci.edu/>) [21].

2.9. Physicochemical Characterization

The physicochemical features of the designed chimeric protein, including Molecular Weight (MW), theoretical Isoelectric Point (pI), instability index, aliphatic index, and Grand Average of Hydropathicity (GRAVY), were predicted by the ProtParam server (http://web.expasy.org/prot_param/) [22].

2.10. Codon Optimization for Expression

Codon optimization was performed using the European Bioinformatics Institute (EBI) database (https://www.ebi.ac.uk/Tools/st/emboss_backtranseq/) to increase protein expression efficiency [23].

2.11. Prediction of mRNA Secondary Structure

The mfold tool was used to predict the secondary structure of mRNA (<http://unafold.rna.albany.edu/?q=mfold>). This tool determines the Minimum Free Energy (MFE) associated with different structures of mRNA, such as loops and hairpins [24].

3. RESULTS

3.1. Information About Genes

Among available sequences, the longest sequence for each antigen was selected [GRA1(UniProt: B9PHR1), SAG1 (UniProt: C7E5T3), ROP18 (UniProt: Q2PAY2), and MIC13 (UniProt: H9BC62)]. Fragments with 100, 100, 100, and 100 amino acids, respectively, from each antigen were selected to be used as chimeric antigens.

3.2. Prediction of Transmembrane Domains and Signal Peptides

Analysis of the TMHMM server determined that only GRA1 has a transmembrane domain, and this region was not used to construct the chimeric protein. SignalP 4.1 predicted signal peptides at positions 1-24 (GRA1), 1-25 (SAG1), 1-47 (ROP18), and 1-22 (MIC13).

3.3. B-cell and T-cell Epitope Prediction

B-cell epitopes for each antigen are summarized in Table 1. Epitopes predicted to bind MHC-I and MHC-II molecules with scores > 9 are presented in Table 2.

Table 1. Predicted linear B-cell epitopes of GRA1, SAG1, ROP18, and MIC13 proteins identified using the IEDB analysis resource.

	GRA1	SAG1	ROP18	MIC13
Bepipred Linear Epitope	67-78, 87-98 , 122-145 , 170-180	20-80, 80-145, 145-245 , 245-300 , 300-330	20-110, 125-140, 180-220, 240-340 , 355-410 , 515-550	70-95, 172-185, 208-222 , 240-260 , 305-320, 330-405, 417-445
Beta-Turn	65-80, 86-98 , 108-112 , 120-125 , 135-145 , 173-186	45-65, 90-115, 125-140, 160-170, 180-265 , 285-295 , 305-315	40-145, 165-265, 285-380 , 400-520	60-90, 105-133, 150-165, 200-220 , 305-320, 330-345, 395-405, 420-445
Accessibility	65-75, 87-90 , 105-114 , 119-125 , 138-147 , 167-180	60-100, 120-155, 165-210 , 220-295	50-190, 220-320, 445-550	175-200, 218-235 , 270-278 , 285-318 , 360-381
Flexibility	70-80, 85-95 , 105-115 , 120-128 , 137-145 , 165-182	60-115, 120-170, 190-290 , 305-315	40-340 , 370-450 , 480-550	70-97, 175-185, 205-245 , 255-278 , 305-318, 360-380, 400-410, 420-445
Antigenicity	78-86 , 90-103 , 115-120 , 132-137 , 145-170	25-120, 140-220 , 280-300 , 310-336	20-40, 60-155, 190-285, 300-420 , 470-540	100-130, 150-170, 235-270 , 280-305 , 330-345, 440-453
Hydrophilicity	67-75, 85-100 , 106-113 , 120-145 , 155-180	55-115, 125-140, 160-240 , 250-320	40-145, 165-340 , 360-460 , 480-550	70-95, 173-185, 207-222 , 240-260 , 305-320, 330-345, 395-407, 415-445

Table 2. Predicted MHC class I and II T-cell epitopes of GRA1, SAG1, ROP18, and MIC13 proteins based on IEDB analysis.

Protein	Amino Acid Start Position ¹			-	-	Number of Binding Epitopes ²		Total ³
Name	H2-Kd	H2-Ld	H2-Dd	I-Ad	I-Ed	MHC- I	MHC- II	
GRA1	--	--	122, 137	81, 94, 114, 142, 144, 148, 151, 155	110, 115, 155, 165, 171	2	13	15
SAG1	242, 273	--	206, 221, 227, 259, 261, 275, 288	217, 247, 260, 268, 278, 296, 300	214, 221, 241, 267, 271, 279, 292, 298	9	15	24
ROP18	321, 362	--	323, 371, 381, 386, 410, 420	328, 344, 352, 354, 356, 363, 380, 383, 388, 391, 393, 397	339, 344, 378, 383, 394, 405, 411, 416, 418	8	21	29
MIC13	--	--	222, 228, 229	222, 227, 234, 237, 244, 248, 257, 265, 271, 284, 299	297, 298, 300	3	14	17

Note: 1: Epitopes with a score higher than 9 identified using the IEDB analysis resource.

2: The number of epitopes with a score higher than 9 in the selected antigen regions.

3: Total number of predicted epitopes with a score higher than 9 in the selected antigen regions.

3.4. Segment Selection

Based on epitope prediction analyses, the fragments GRA1 (81-180), SAG1 (201-300), ROP18 (321-420), and MIC13 (201-300) were selected and joined using rigid linkers. A total of 24 chimeric arrangements were created from all possible permutations of the four antigenic fragments. Among them, eight chimeric proteins (RSGM, RGSM, GRSM, GSRM, MSGR, SRGM, SGRM, and RMSG) showed the best stability and predicted folding based on secondary and tertiary structure analysis. Of these eight chimeric proteins, the GSRM construct (GRA1-SAG1-ROP18-MIC13) with the A(EAAAK)A linker showed the highest predicted antigenicity, the most

favorable structural features, and the highest confidence score (C-score) and was therefore selected as the final vaccine candidate.

3.5. Prediction and Analysis of Secondary and Tertiary Structures

The 442-residue GSRM protein contains 41.18% alpha-helices, 42.53% random coils, and 16.29% extended strands (Fig. 1A and B). Typically, a high proportion of α -helices (> 40%) and balanced coil content support proper folding and structural stability. Tertiary structure modeling yielded a C-score of -2.21 (Fig. 2A and B), which falls within the acceptable range of -5 to 2 for I-TASSER models.



Fig. (1). Secondary structure prediction of the GSRM chimeric protein using the GOR IV server: (A) predicted secondary structure composition (α -helix, β -strand, and coil) and (B) graphical representation of the predicted secondary structure.

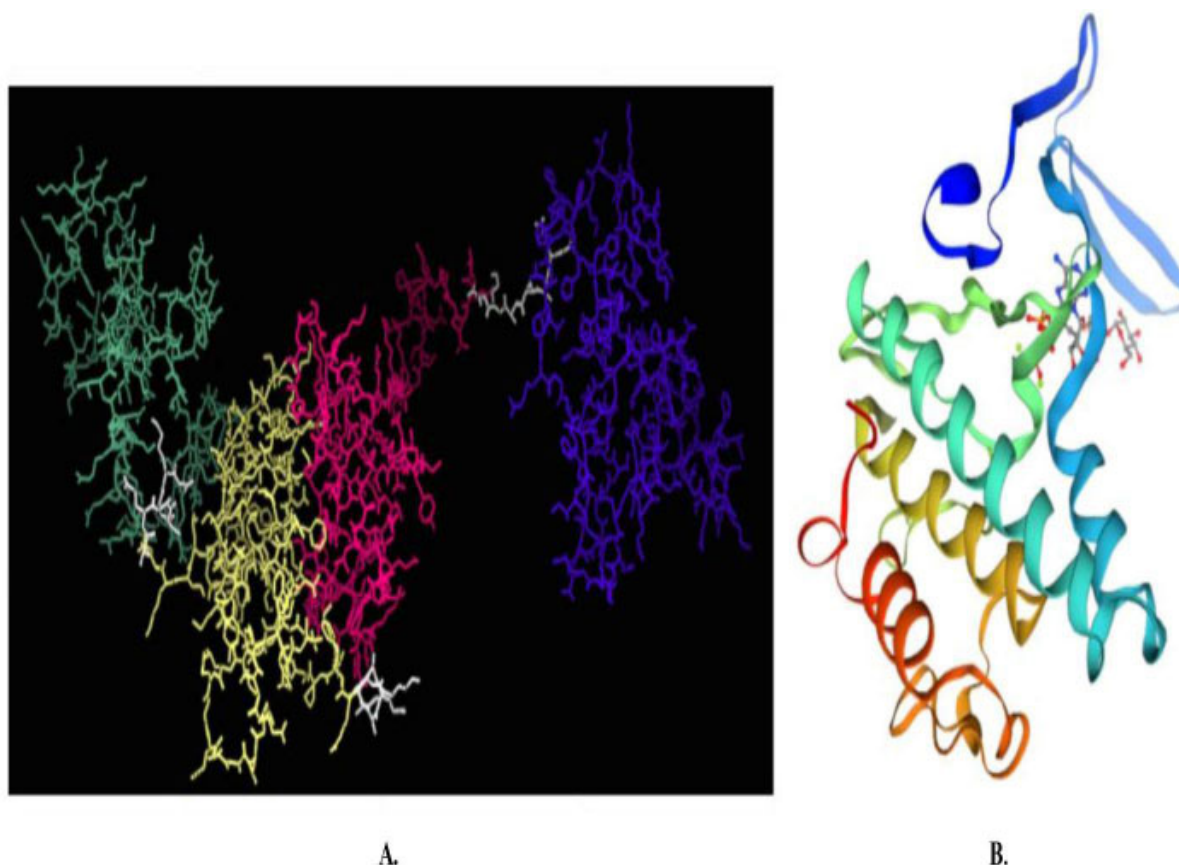


Fig. (2). Predicted 3D structure of the GSRM protein: (A) I-TASSER model showing domains (GRA1: blue, SAG1: pink, ROP18: yellow, MIC13: green, and linker: white) and (B) 3D model visualization generated by the SWISS-MODEL server.

3.6. Tertiary Structure Validation

Ramachandran plot analysis *via* SWISS-MODEL showed that 84.90% of the residues were located in the desired regions, 10.41% in the allowed regions, and 4.69% in the outlier regions (Fig. 3). When more than 90% of the residues are located in the favorable and allowed regions and less than 5% are in the outlier regions, the GSRM has acceptable stereochemical quality.

3.7. Antigenicity, Allergenicity, and Solubility Assessment

The Vaxijen score was estimated to be 0.6701. A common threshold of 0.5 is considered for potential protective antigens, and antigenicity above this threshold indicates strong antigenic potential. AlgPred analysis confirmed that the GSRM protein is nonallergenic. SOLpro predicted a solubility probability of 0.656861, and since values above 0.5 indicate that a protein is likely to be soluble when expressed in *E. coli*, this result suggests good expression potential in a prokaryotic system.

3.8. Prediction of Physicochemical Properties

The protein has a MW of 47,953.46 Da and a theoretical pI of 5.61. Negatively charged residues (Asp +

Glu) total 61, and positively charged residues (Arg + Lys) total 49. Predicted half-life: 30 hours in mammalian reticulocytes (*in vitro*), > 20 hours in yeast (*in vivo*), and > 10 hours in *E. coli* (*in vivo*). The instability index is 43.07, and since values < 40 indicate stable proteins, this protein is considered unstable. The aliphatic index of the protein is 80.11, indicating good thermal stability. The GRAVY score of -0.316 indicates hydrophilicity.

3.9. Optimization of the Codon

Reverse translation and codon adaptation of the designed GSRM protein were performed using the EMBOSS Backtranambig tool on the EBI server, with *E. coli* as the host organism. This optimization replaces rare codons with synonymous codons preferred by *E. coli*, enhancing translational efficiency and minimizing ribosomal stalling.

3.10. Prediction of mRNA Secondary Structure

The best-predicted structure had $\Delta G = -387.70$ kcal/mol, and the first 10 nucleotides at the 5' end did not form a stable hairpin or pseudoknot, supporting efficient ribosome binding and translation initiation (Fig. 4). These results indicate that the optimized GSRM transcript is suitable for robust expression in a prokaryotic system.

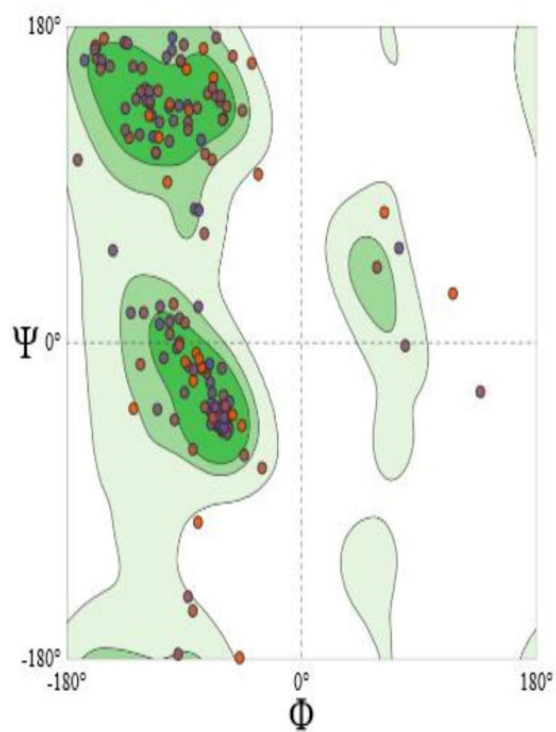


Fig. (3). Validation of the predicted GSRM tertiary structure using Ramachandran plot analysis. The plot shows 84.90% of residues in favored regions, 10.41% in allowed regions, and 4.69% as outliers.

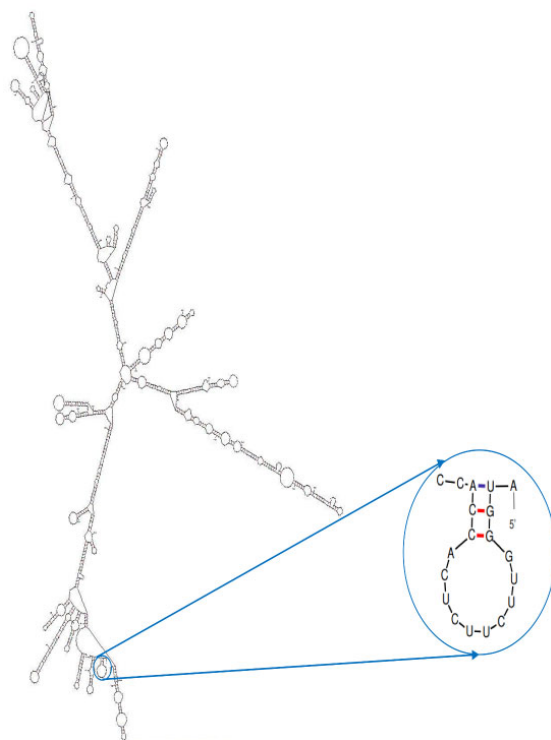


Fig. (4). mFold-predicted secondary structure of the optimized GSRM sequence ($\Delta G = -387.70$ kcal/mol).

4. DISCUSSION

The ability of *T. gondii* to evade the host immune response, its complex life cycle, and its antigenic diversity remain a challenge in the development of vaccines against this parasite. Therefore, this study aimed to develop a chimeric protein vaccine containing B- and T-cell epitopes of GRA1, SAG1, ROP18, and MIC13 antigens of *T. gondii* and analyze various aspects of the chimeric protein using different bioinformatics tools.

In the *in silico* approach, any antigen can be a vaccine candidate. In this study, a variety of antigens (surface antigens such as SAGs and secretory antigens such as GRAs, ROPs, and MICs) were used for vaccine design. GRA1, SAG1, ROP18, and MIC13 were selected based on their roles in parasite survival, invasion, and pathogenicity. In addition, the aim was to select antigens that are expressed in all three stages of the parasite life cycle to elicit a more comprehensive immune response, as stage-specific antigens typically provide limited protection. On the one hand, the immune response generated against the combination of epitopes of these antigens is much more effective than the immune response against a vaccine that consists of only one antigen and stimulates an immune response against that single antigen [11, 14, 24]. Recent advances in the field support the rationale behind our design. For example, Nayeri *et al.* (2025) performed an *in silico* analysis of *T. gondii* surface and secretory proteins. They designed a novel chimeric vaccine and demonstrated a stronger immune response elicited against a combination of multiple antigenic epitopes [25]. Ghaffari and Rahimi (2024) constructed a multi-epitope peptide vaccine based on calcium-dependent protein kinases using immunoinformatics; their analyses suggested favorable antigenicity, solubility, and safety. Therefore, it becomes necessary to conduct experimental tests [26]. Similarly, another recent *in silico* study used tachyzoite-specific SAG1-Related Sequence (SRS) proteins to design a peptide vaccine containing T-cell and B-cell epitopes that showed high antigenicity, non-allergenicity, and predicted interaction with human TLR4 [27]. Moreover, a genome-wide epitope vaccine study by Li *et al.* (2022) evaluated *T. gondii* membrane proteins and identified candidate epitopes for broad coverage, underscoring the power of comprehensive bioinformatics strategies [28]. In contrast, the present study introduces several methodological innovations. Rather than stitching together minimal epitopes, we selected antigenic fragments of approximately 100 amino acids from four antigens (GRA1, SAG1, ROP18, and MIC13), generated all 24 possible permutations of these fragments using the rigid A(EAAAK)A linker, and carefully screened the resulting structures for stability, folding quality, antigenicity, Ramachandran analysis, and physicochemical profile. This vaccine design strategy preserves the structural elements of the original antigens and could create a more stable and immunogenic vaccine candidate.

The selection of antigenic fragments was performed using rigorous epitope prediction analyses targeting B and

T cell epitopes, ensuring that the designed construct could effectively stimulate an immune response. This is particularly important given the intracellular nature of *T. gondii* and the need for strong cellular immunity alongside humoral immunity. Notably, the A(EAAAK)A linker sequence used between the antigenic fragments has previously been shown to promote proper folding and functional independence of the individual domains, minimizing potential steric hindrance or epitope masking [29]. These design considerations are essential to ensure that the epitopes remain structurally accessible and immunologically active.

Our structural predictions showed that the GSRM protein has a balanced secondary structure, with a significant proportion of alpha-helices and random coils, which are favorable for stable folding and antigen presentation. Tertiary structure modeling revealed a moderate C-score (-2.21), indicating a relatively reliable Three-Dimensional (3D) conformation. Ramachandran plots confirmed the tertiary structure of the protein. The results showed that more than 90% of the residues were located in the favorable and allowed regions.

Moreover, immunoinformatic analyses predicted that the GSRM is antigenic (Vaxijen score: 0.6701) and nonallergenic, addressing two primary prerequisites for a safe and effective vaccine candidate. The pI is also one of the important physicochemical parameters that can be used to estimate the solubility of a protein at a certain pH, meaning that the protein precipitates in a solution close to the pI [24]. The pI is useful for establishing a suitable buffering system for protein purification based on pH-focused methods [24]. The presence of an acidic isoelectric point and a negative GRAVY score (-0.316) further indicates that the protein is likely to be hydrophilic and accessible to the host immune system, which is advantageous for immunogenicity. Although the instability index slightly exceeded the threshold for stability, the predicted solubility and favorable antigenicity scores suggest the protein remains a viable candidate for vaccine development. A MW (47953.46 Da) indicates that GSRM could be a suitable antigen (it has been indicated that antigens with a MW of < 5-15 kDa do not provide good immunity) [18] and an aliphatic index (80.11) indicative of good thermostability.

Codon optimization and mRNA secondary structure analysis are important steps in establishing a link between *in silico* vaccine design and practical protein production. High-throughput expression of antigen in *E. coli* facilitates purification and experimental testing. Codon optimization reduces the likelihood of ribosomal stalling, and stable mRNA folding, and the absence of inhibitory structures at the 5' end ensures translation of the chimeric protein.

This study is based exclusively on *in silico* analyses, which, although powerful for screening candidates, cannot fully simulate the complexity of immune responses in living systems. Structural predictions, antigenicity scores, allergenicity assessments, and solubility estimates are computational approximations that require experimental validation. Furthermore, epitope prediction tools vary in

accuracy, and the selected fragments may behave differently when expressed as a recombinant protein *in vitro* or *in vivo*. The instability index suggests that the final construct may require stabilization strategies during expression and purification. Additionally, the immunogenic performance of the chimeric protein cannot be inferred without testing in animal models, including evaluation of cellular immunity, antibody responses, and assessing survival rates in mice infected with different strains. Therefore, the findings here should be considered preliminary and require experimental testing.

CONCLUSION

In conclusion, the bioinformatics-driven design of the GSRM protein represents a promising step towards the development of an effective multiepitope vaccine against *T. gondii*. By combining multiple antigens from different stages of the parasite's life cycle, this construct aims to overcome the limitations of single-antigen vaccines and provide broad, cross-stage protection. This work will provide a solid foundation for further experimental studies and help control toxoplasmosis in humans and animals.

AUTHORS' CONTRIBUTIONS

The authors confirm their contributions to the paper as follows: T.N.: conceived the study and designed the study protocol; T.N.: Was the supervisor of this research; E.N.: Performed the bioinformatics analysis; T.N.: Drafted the manuscript; E.N., M.F. and E.G.: Critically revised the manuscript; and All authors read and approved the final version of the manuscript.

LIST OF ABBREVIATIONS

SAGs	= Surface Antigens
MICs	= Microneme Proteins
ROPs	= Rhoptry Proteins
GRAs	= Dense Granule Proteins
NCBI	= National Center for Biotechnology Information
UniProt	= Universal Protein Resource
IEDB	= Immune Epitope Database
MHC	= Major Histocompatibility Complex
GOR IV	= Garnier-Osguthorpe-Robson IV
MW	= Molecular Weight
pI	= Isoelectric Point
GRAVY	= Grand Average of Hydropathicity
EBI	= European Bioinformatics Institute
MFE	= Minimum Free Energy
C-score	= Confidence Score

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

This study did not involve human participants, animals, patient data, or biological specimens. However, the

research project was registered and approved by the institutional ethics committee in accordance with national regulations governing research projects in medical universities. Therefore, an ethics approval code was assigned to the study (Ethics Code: IR.DUMS.REC.1404.014). The ethics approval was administrative and regulatory in nature and did not reflect the involvement of human or animal subjects.

HUMAN AND ANIMAL RIGHTS

No animals/humans were used for studies that are the basis of this research.

CONSENT FOR PUBLICATION

Not applicable.

AVAILABILITY OF DATA AND MATERIALS

The data and supportive information is available within the article.

FUNDING

None.

CONFLICT OF INTEREST

The authors declare no conflict of interest, financial or otherwise.

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