Imaging of High and Low Resolution Ebola Envelope GP Structures Composited with in silico Models of Difficult-to-Resolve Sections

Garry W Lynch1,*, Stefanie S Portelli1, Siu Wai Wong1, Maria Rocha Costa1, Ana Paula Mello Lemgruber2, Peter Williamson3, John S Sullivan1,6, Robert Booy1,4 and W Bret Church1

1Faculty of Medicine, The University of Sydney, Camperdown, NSW, 2006, Australia
2Faculty of Veterinary Science, The University of Sydney, Camperdown, NSW, 2006, Australia
3Faculty of Pharmacy, The University of Sydney, Camperdown, NSW, 2006, Australia
4The National Centre for Immunossurveillance Research, The Children’s Hospital Westmead, Westmead, NSW, 2114, Australia
5School of Medical Sciences, UNSW, Kensington, NSW 2052, Australia

Abstract

The Ebola surface glycoprotein, GP, facilitates receptor binding, cell infection and cytopathology, and is a principal target for immune protection. The structure of the GP central core structure and antibody target for the 1976 Ebola Zaire Mayinga strain of Ebola has been resolved by X-ray crystallography (e.g., pdb-ID: 3CSY). But other important GP regions have defied crystal structure determination. These include the apical mucin-like domain (MLD) that contains immuno-protective sites and is most distal to the membrane, and the membrane proximal C terminus region (meD) where GP is tethered to the Ebola surface. A molecular structure-based strategy in vaccine design necessitates detailed fine-structure knowledge of exposed sites for immuno-targeting. To address the lack of MLD and meD structures we have performed computational modeling of those regions using the programs iTasser, Phyre2 and CABS-flex. Candidate MLD model structures for the 1976 Mayinga strain were screened for continuity and fit with the 3CSY core and tested further for 3D compliance for its fit within the spatial boundaries of a low-resolution GP trimer cryoelectron tomograph (EMDB-ID: EMD-6003). Under these constraints only 1 of 6 initial MLD models generated by iTasser or Phyre2 met the required structural criteria of continuity, orientation and spatial fit. That best-fit structure (MLDm1p1) was additionally evaluated using the MolProbity geometric analysis, as well by Phyre2 and CABS-flex program. Notably, documented protective antibody sites of the MLD mapped to the MLDm1p1 apex surface, as anticipated for domain functionality or exposure to antibody attack. These studies describe best fit proximate in silico models of the MLD and meD regions of GP to assist molecular understanding and therapy design. Biochemical and immunologic validation and refinements of the models are continuing, to establish to their biостructural compatibilities and value to inform on GP structure-function and immuno-target potential.

Introduction

The 2014 Ebola Virus (EBOV) outbreak in West Africa had up to the end of March 2015 claimed more than 20,236 lives from 24,753 infections [1]. Now more than a year on from its emergence the crisis appears to be contained and finally a couple of vaccine constructs have been in field clinical trials, and show some evidence of protection for the EBOV 2014 strain [2,3].

The wider protective potential of vaccines in the pipeline to protect against any other Ebola strains that might emerge remains a significant unknown, and whether they could provide any protection against the more distant and highly lethal Marburg filovirus. There is therefore a high need for a variety of efficacious drugs and vaccines to pre-empt any such future outbreaks of Ebola or Marburg. Structure based drug and vaccine design has of recent times matured into being of value in the development of therapeutics for infectious disease agents. Our structure based studies of the Ebola GP protein follows from an approach developed in the study of Collateral Immunity® for broad-strain seasonal antibody responses and protection of Influenza [4-7]. In this present article we elaborate on an invited medical image published by this journal earlier this year on EBOV GP structure [8].

Ebola Envelope GP: The envelope surface glycoprotein, GP, protrudes from the Ebola virus surface as a homo-trimer that contains the receptor binding and membrane fusion determinants that define cell tropism, entry and infection [9,10]. Consequently a detailed understanding of GP structure and interactions as the principle target for immune defense and for vaccine development is paramount. Indeed, detailed knowledge of the high-resolution, 3.4 Ångstrom, crystal-struture determination of the central core (pdb structure identifier: 3CSY), that contains receptor binding and fusion sites [9], has been invaluable, and particularly so for the realization of specific target sites for immune antibodies. However, that resolved structure, the most comprehensive of all performed to date, represents only about half of the whole pre-fusion protein sequence of GP. Unfortunately a number of other GP sections are without detailed structural information as they have been recalcitrant to their determination in the crystal. Within those undefined regions are important structural and functional features and for which high resolution information would be invaluable. The focus of this article is on two of those unresolved sections, namely the mucin-like domain (MLD that spans aa 313-501) of GP1 that contains immunodominant antibody sites for host immune protection [11-13], and a principal focus of this study, and the C-terminal section of GP2 (meD aa 658-676) at the GP-virion surface envelope interface.

*Corresponding author: Garry Lynch, Faculties of Medicine and Veterinary Science, Room 551 Gunn Building (B19), University of Sydney, Camperdown, NSW, 2006, Australia, Tel: 61-286270258; E-mail: garry.lynnch@sydney.edu.au

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Defining protein structure

X-ray crystallography has emerged over time as the principal technology for the description of high-resolution protein structures and remains the most validated procedure. Despite its considerable value it has limitations and can provide variants of the structure not found in nature. Furthermore X-ray crystallography cannot capture all of the possible structures a protein may achieve, and many of these may be temporal and subject to their biologic context and environment (e.g., pH). With crystallography likely capturing only one of many possible structural configurations. Many proteins or sections of proteins can be recalcitrant to X-ray evaluation, as they are either refractory to crystallization or do not contribute to ordered diffraction patterns, due to being heterogeneous in composition, hydrophobic, have mobile elements, or post-translationally modified. The most notable and substantial section that has remained unresolved in the GP 3CSY structure, despite the presence of its sequence in the protein construct used for crystallisation, is the mucin-like domain (MLD). In the attached native state the MLD is the most membrane-distal segment of the envelope GP and projects out above the central, chalice-like, 3CSY core [9,10]. Characteristically this highly glycosylated mucin-like domain would provide a surface environment compatible for EBOV to mix and circulate in mucosal surfaces, by virtue of its similarity with the heterogeneous mucin family of proteins [13,14], that populate the mucosa, and with which it likely shares a degree of structural similarity. Notably the entire mucin-family is poorly represented in the structural database. The high degree of glycosylation of this glycoprotein family is in itself problematic for crystal structure refinement.

Another region of GP for which there is currently minimal structural knowledge is the membrane-associating region, which anchors GP into the EBOV envelope. Hence, we asked the question, whether useful structural information on the MLD and the membrane-associating meD sections could be attained from computer-based in silico modeling [16]. To assist in evaluating plausible in silico simulations, a low-resolution cryoelectron tomograph structure determination of the GP trimer of the EBOV surface has recently been published [16], and for our studies provides a necessary framework to which acceptable models need to be compatible.

Rational of the study approach

Success in correctly assigning and defining protein structures and as evaluated biennially with the CASP (Critical Assessment of protein Structure Prediction) challenges [18] has realistically improved the potential and value for determining structural information for proteins that defy description by other means. The GP of MLD is considered to fit within the ‘highly motile’, ‘structurally heterogeneous’ or ‘inherently disordered’ category of proteins, and for which no structures have been obtained. The quest is to visualize and determine the structures the GP MLD domain that are exposed for antibody binding, and that might present to cell interactions and to the immune system as a potential target or for avoidance thereof. Before technologies arise that are capable of determining the basic structure(s) of such regions the point of this study was aimed at attempting to generate feasible models of the MLD site of EBOV GP.

Templates, programs and procedures

EBOV Study Strain and sequence: For the modeled structures in this study the Ebola Mayinga 1976 strain Zaire subtype (UniProt Taxon ID: 128952) was the template reference strain sequence used for analysis to ensure direct comparison and consistency with the Ebola strain sequence upon which the resolved core GP 3CSY structure was generated. That isolate was: Ebola virus/H.sapiens-tc/COD/1976/ Yambuku-Mayinga, NCBI: NC_002549. With the specific protein studied; the 676 amino acid (aa) envelope protein (UniprotKB ID: Q65320; and Proteome ID: UP000007209).

Molecular Modelling Platforms: The molecular modelling of the MLD and meD segments was performed using the Phyre2 [18,19] (http://www.sbg.bio.ic.ac.uk/phyre2); I-Tasser [20,21] (http://zhanglab.csb.med.umich.edu/I-TASSER/); or CABS-flex [22,23] (http://biocomp.chem.uw.edu.pl/CABSflex/) software tools. Our model structure of the mucin-like region (MLD) was achieved using Phyre2 [19] and covered amino acids (aa) 313 to 501 across GP1 and included a 313-335 aa overlap with the 3CSY structure [9]. Modelling, rendering or alignment analyses of the GP structures was performed using RASMOl [24], PyMOL [25], and Swiss-PdbViewer, SPDBV_4.1.0_OSX computational tools. Further, our simulations for the membrane-proximal/associating meD region of GP2 (aa 658-676), was achieved using Phyre2. That short C-terminus, 13 aa, model structure was manually aligned with the 3CSY and EMD-9003 structures, and rendered together in PyMOL.

EBOV GP Reference Structures: The template scaffold structures used in this study were the high resolution 3.4 Å 3CSY derived by x-ray crystallography [9] for a truncated and modified GP sequence of the Mayinga 76 strain, and a low-resolution CET produced from Ebola virus-like particles expressing full-length EBOG GP deposited into the Electron Microscopy Data Bank (EMDB accession number 6003 (EMD-6003) (http://www.emdbanken.org/) [16] (note1).

Structure Visualisation and Alignment: Best-fit simulations from the in silico structure predictions were mapped against the pre-cuplar protein crystal 3CSY rendered in PyMOL. Successful model candidates were further aligned and rendered into the low-resolution, trimeric cryo-electron tomogaphy (CET) structure [16], to assess additional 3D-orientation and spatial compliance.

Structure Geometry Profiling: The coordinates in pdb format of best-fit GP protein Phyre2-models experimentally obtained for the mucin-like (Phyre2 and CABS-Flex structures) and C-terminus regions (Phyre2 structure) were submitted for MolProbity analysis (http://molprobity.biochem.duke.edu). Assessments of molecular conflict, and rotamer and Ramachandran characteristics obtained using the MolProbity profiling tool [26,27], were compared against comparable properties generated for several other well determined protein structures. Proteins used for comparison were; the A3 collagen binding domain of von Willebrand Factor (pdb: 1AO3), Human Serum Albumin (: 1BM0), Bovine Serum Albumin (:4FSS), EBOV GP core (: 3CSY)(1976 Mayinga Strain), and EBOV GP core (: 3VE0)(1976 Boniface Strain) protein structures.

Immunomapping: Established sites of protective antibodies for the MLD region of GP were mapped onto the Phyre2 MLD model (MLDm1p1). Sites assessed were; i) the IEDB epitope ID: 156605 by the neutralising antibody CAG7 [11,28-30] and ii) a protective site of the Zaire Mayinga strain for the non-neutralising antibody H13F6, but not protective for other EBOV strains [11,12,31]. These were rendered in PyMOL onto the composite image shown in Figure 1.

Structure Refined: For assessment and profiling of structure flexibility the MLDm1p1 model coordinates were submitted to the CABS-flex program server [22,23] for analysis (http://biocomp.chem.uw.edu.pl/CABSflex).
Results

Composite imaging of resolved and in silico modeled GP domains and structures

In the absence of defined high resolution structures for the mucin-like (amino acids 313-501) and membrane spanning (amino acids 658-676) domains of GP we have performed in silico modeling of these regions and composited those models together with the published low EMD-6003 and high 3CSY structures of Ebola GP. Under the imposed structural requirements only 1 of 6 MLD computer simulated models generated by either iTasser or Phyre2 satisfied the spatial fit criteria set above; for continuity with 3CSY, and compatibility and 3D position within the EMD-6003 trimer footprint. The successful model, ID: MLDm1p1, was modeled using Phyre2. The central composite of Figure 1 shows the X-ray crystal resolved structure, and the in silico MLD and meD models mapped together and within the recent cryo-electron spatial EBOV GP trimer tomograph [16]. Shown on the periphery of that image are the space-filled models of the complete low-resolution cryoelectron tomograph (Cryo-ET) trimer structure of GP (EMD-6003) and the high-resolution X-ray crystal structure of the GP trimer core (pdb 3CSY) [9]. On the right of the composite are the in silico space-filled models from Phyre2 simulations of a GP1 mucin-like domain monomer and the GP2 C-terminus membrane stalk domain trimer. In the centre image each
of these structural elements has been assembled as a composite, built in ribbon display, and within the Cryo ET global framework. Together these structures represent a majority (~85%) of the GP protein, with the 3 separate mucin-like domains at the very top of the composite (yellow), a central core trimer (white) and at the bottom the membrane domain (yellow). Structural visualizations of the refined and the in silico-modeled structures are from pdbviewer, Rasmol [24] or PyMOL [25] rendering. Each of the different GP domain sequence determinations were generated for the 1976 Ebola Zaire strain, and imaged to represent a single, prefusion, GP trimer, knob that projects out from the envelope surface. Noting as mentioned this composite of most of the GP is without proposed structures for the GP regions of amino acid sequences: 1-30, 190-213, 238-299 or 599-658). For direct reference of the respective regions already resolved or modeled, a linear map of the complete GP protein sequence has been included in bar format in Figure 1.

Model Validation

Attempts to validate computationally derived molecular models, without experimental or biological evidence and using only computational tools can have considerable limitations. Therefore we have taken 2 different approaches for validation of the MLD model, one based on biological data 1) using antibody binding- site mapping, and alternately 2) by computational assessment.

Relevance to known functional, structural or biologically identified sites

A necessary requirement for any proposed MLD structure is that sites for biologically protective antibodies to the viral envelope GP must be surface exposed and oriented away from the MLD-3CSY interface, which by definition would physically preclude antibody attack at the EBOV envelope surface. When tested for positioning of the sites for the binding of the protective antibodies C4G734 (marked in green) and H13F6f13 (marked in red) these were mapped onto the proximate MLD model (Figure 1), As shown those sites were found to map to the exterior surface and namely to the top and side surfaces of the model, as would be predicted. Whilst this does not definitively confirm this exact conformation is correct it is appropriate. Importantly, if the converse had been observed, such a structure would be considered an irrelevant configuration and excluded from further consideration.

Computational assessment, validation and molecular refinement

Molecular flexibility: Test algorithms were applied to evaluate the proximate structures modeled here. Due to the inherent characteristics of the MLD and failure to be resolved by crystallography it was deemed useful to computationally test the flexibility of our Phyre2 MLDm1p1 model, using the CAB-Flex web tool. That assessment revealed 12 related and similar structures, indicating a reasonable degree in flexibility for the MLD-m1p1. The model MLDm2c2 was selected for further study and compared with the parent Phyre2 MLDm1p1 structure (Table 1). The MLDm2c2 is highly similar, but has notable differences as evident on alignment.

Geometry profiles: The MLD models (MLDm1p1 and MLDm1c2) co-ordinates were submitted to the MolProbity program to assess their geometric rotameric and Ramachandran properties [26,27]. To put the measured geometries for the best-fit MLD and meD models in perspective, these were matched and compared with the equivalent parameters obtained in MolProbity for a number of other protein structures considered relevant for comparison (Table 1). These were the A3 collagen binding domain of von Willebrand Factor, human albumin, bovine serum albumin, the GP core 3CSY structure the MLD (MLD-m1p1, MLD-m2c2) and meD. The respective protein structure properties and MolProbity Rotamer and Ramachandran Plot analyses are presented in Table 1.

Comparisons of the acceptable rotamer and Ramachandran plot profiles for the MLD and meD models were made against the profiles obtained for the likely semi-rigid (e.g., A3 collagen-binding domain of vwf) and flexible (e.g., albumin) proteins and x-ray crystallography defined 3CSY and 3VE0 structures. As predicted the A3 domain of vwf gave the best adherence for both rotamer and Ramachandran plot assessments. In general the in silico models gave geometry profiles within the range of established and defined structures. However, the MLDm1p1 had the second poorest rotamer score of 12.25% and by far the worst Ramachandran value showing 20.19% of outliers. Notably the CABS-Flex MLDn2c2 derivative of the MLDm1p1, on geometry profiling gave remarkably improved scores of 1.20% poor Rotamers and 4.84% Ramachandran outliers, comparable to the geometries of highly validated structures. This reveals a similar, but even more refined MLD structure structure in the MLDm2c2 model and better compatibility with accepted structure parameters than the MLDm1p1 model.

Conclusions

The essential boundaries and constraints placed for the modeling of MLD and meD regions of GP was that they must be compatible with the resolved low (CEM) and high (crystal) resolution structures. Fitting this basic criteria, the in silico model MLD model structure obtained in these studies was shown to map with i) continuity to the resolved 3CSY structure, and ii) within the 3D-boundaries of the EMD-6003 trimer structure [16]. In immunology based testing of the model for biologic relevance, the G4G7 and H13F6f protective antibody sites were found to map onto the MLD1p1, at the most exposed surface and apex of the MLD region, as would be appropriate for functionality and immune attack. Additional confidence in the Phyre2 generated MLDm1p1 and MLDm1c2 models, was also achieved using computational geometric validation. This included examination of the structural flexibility of the MLDm1p1 using the program CABS-Flex, which returned 12 additional similar but varied model structures and is indicative of considerable structural flexibility for the MLD. A representative MLDm2c2 model was revealed as highly similar, but with notable structural differences to the MLDm1p1, when the two structures were aligned (Table 1). Furthermore, the geometry profiles for the in silico models based on MolProbity analysis, fitted within the range of values for the established and defined structures. Albeit the MLDm1p1 model had the second weakest score for both Rotamer and Ramachandran values. However, it was found that the CABS-Flex MLDn2c2 derivative of the MLDm1p1, on geometry profiling gave scores of 98.71% acceptable rotamers and 95.16% acceptable Ramachandran plotting, and was far more comparable to the geometric characteristics of highly validated structures. This has revealed a similar but valuable refinement of the MLD (i.e., the MLDm2c2), for future evaluation, and comparisons of the biological feasibility of both models. Additional structure validations are being pursued, such as compatibility with carbohydrate mapping. Whilst these current studies have mainly focussed on the MLD domain, additional models for the membrane-associating C-terminus have also been built, including a short 13 aa region. Further studies and more extended models that overlap with the 3CSY and 3VE0 core structures are now needed.
**Table 1A**

<table>
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<th>Protein</th>
<th>VWF A3 Domain (Coll Type I/III bind region)</th>
<th>Human Serum Albumin 1BM0</th>
<th>Bovine Serum Albumin (BSA)</th>
<th>3CSY 3CSYFH</th>
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<tr>
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<td>N/A</td>
<td>N/A</td>
<td>Mayinga Zaire 1976</td>
</tr>
<tr>
<td>Pdb ID</td>
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<td>1BM0</td>
<td>4F55</td>
<td>3CSY</td>
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<tr>
<td>Protein size:</td>
<td>186 aa</td>
<td>581 aa</td>
<td>581 aa</td>
<td>187 aa</td>
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**Table 1B**

<table>
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<tbody>
<tr>
<td>Strain</td>
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<td>N/A</td>
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<tr>
<td>Pdb ID</td>
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<td>13 aa</td>
<td>188 aa</td>
<td>188 aa</td>
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**Table 1:** Computational Geometric Comparative MolProbity analyses of GP Resolved Core structures & the Proximate MLD and meD Models. Table 1A and 1B Show respectively the Rotamer and Ramachandran Geometry Comparison Molprobity Profiles of Table 1A) the resolved X-ray crystallography structures for the A3 collagen-binding domain of von Willebrand Factor (vWF, pdb: 1AO4), human (pdb: 1BM0) and bovine (pdb: 4F55), and the central the Ebola GP protein core structures of pdb: 3CSY (1976 EBOV Zaire strain) and in 2A) the 3VE0 (1976 EBOV Sudan strain), and their geometries compared to computational models of the 1976 EBOV Zaire sequence membrane proximal GP 658-676 aa C terminus mpcDm1p1 (modeled in Phyre2), and the 313-501 aa Mucin-like-domain MLDm1p1 (modeled in Phyre2) and the MLDm2c12 (modeled in CABS). For this figure each of the crystal or modeled protein structures were visually prepared in PyMol. The very last figure in the sequence showed a combined alignment of the structurally similar Phyre2 and CABS models for the MLD. Note: The displayed structures differ in relative size between panel 1A (at 100%), and panel 1B (at 150%), to enhance visibility of the shorter protein constructs.
The GP in silico models shown here fit a range of structural and biomapping criteria, but it needs to be appreciated that without additional qualification, they remain as models or proxies of what those domains might be, and serve at present as interim simulations to provoke new approaches for biotesting, en route to the development and determination of more substantiated structures. Certainly, related similar structures generated by the CABS-Flex program and with even stronger geometric validation scores immediately offers new refinements upon which the different models can be tested for, in biological assays and further quality checks. These structures herein provide a starting point upon which experiments will be designed for the biotesting and identification of new antibody target sites. It will therefore also be valuable to know if such simulations of difficult to resolve domains can for the future be beneficial for vaccine and drug design. And whether synergistic use of molecular modeling and cryoelectron tomography (e.g., as used here) may offer a viable alternative for the fine structure determination of proteins that are resistant to crystal structure refinement. An enticing prospect for the future may be that sub Aångstrom level cryoelectron tomograph determinations [34,35] may overcome this technical obstacle and be used together with ab initio modeling, to give correct and fully verifiable, high-definition assignments of true native structures. To achieve this would be a holygrail in the realization of difficult-to-resolve protein structures of biological and therapeutic importance.

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Disclosure

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