

1 Title:

2 **Glycan shield of the ebolavirus envelope glycoprotein GP**

3

4 Peng W.¹, Rayaprolu V.^{2,4}, Parvate A.D.^{2,5}, Pronker M.F.¹, Hui S.^{2,6}, Parekh D.², Shaffer K.^{2,7}, Yu
5 X.², Saphire E.O.^{2,3}, Snijder J.^{1*}

6

7 ¹ Biomolecular Mass Spectrometry and Proteomics, Bijvoet Center for Biomolecular Research
8 and Utrecht Institute of Pharmaceutical Sciences, Utrecht University, Padualaan 8, 3584 CH
9 Utrecht, The Netherlands

10 ² Center for Infectious Disease and Vaccine Discovery, La Jolla Institute for Immunology, La Jolla,
11 CA 92037, USA.

12 ³ Department of Medicine, University of California, San Diego, La Jolla, CA 92039 USA

13 ⁴ Current affiliation - Pacific Northwest Center for CryoEM, Portland, OR 97225

14 ⁵ Current affiliation - Environmental Molecular Sciences Laboratory, Pacific Northwest National
15 Laboratory, Richland, WA 99354

16 ⁶ Current affiliation – Molecular Microbiology and Microbial Pathogenesis Program, Washington
17 University School of Medicine, Saint Louis, MO 63108, USA

18 ⁷ Current affiliation – Division of Biology and Biological Engineering, California Institute of
19 Technology, Pasadena, CA 91125, USA

20

21 *corresponding author: j.snijder@uu.nl

22

23

24 **Abstract**

25 The envelope glycoprotein GP of the ebolaviruses is essential for host cell attachment and entry.
26 It is also the primary target of the protective and neutralizing antibody response in both natural
27 infection and vaccination. GP is heavily glycosylated with up to 17 predicted N-linked sites,
28 numerous O-linked glycans in its disordered mucin-like domain (MLD), and three predicted C-
29 linked mannosylation sites. Glycosylation of GP is important for host cell attachment to cell-
30 surface lectins, as well as GP stability and fusion activity. Moreover, it has been shown to shield
31 GP from neutralizing activity of serum antibodies. Here, we use mass spectrometry-based
32 glycoproteomics to profile the site-specific glycosylation patterns of ebolavirus GP. We detect up
33 to 16 unique O-linked glycosylation sites in the mucin-like domain, as well as two O-linked sites
34 in the head and glycan cap domains of the receptor-binding GP1 subunit. Multiple O-linked
35 glycans are observed at the S/T residues of N-linked glycosylation sequons, suggesting possible
36 crosstalk between the two types of modifications. We also confirmed the presence of C-
37 mannosylation at W288 in the context of trimeric GP. We find heterogenous, complex N-linked
38 glycosylation at the majority of predicted sites as expected. By contrast, the two conserved sites
39 N257 and N563 are enriched in unprocessed high-mannose and hybrid glycans, suggesting a
40 role in host-cell attachment via DC-SIGN/L-SIGN. We discuss our findings in the context of
41 antibody recognition to show how glycans contribute to and restrict neutralization epitopes. This
42 information on how N-, O-, and C-linked glycans together build the heterogeneous glycan shield
43 of GP can guide future immunological studies and functional interpretation of ebolavirus GP-
44 antibody interactions.

45 Introduction

46 Ebola virus is a member of the *Filoviridae* family [1, 2]. Since its initial discovery in 1976, it has
47 caused recurring outbreaks of disease in Central and West Africa upon spillover into the human
48 population from an as-yet unidentified animal host reservoir, or recrudescence from convalescent
49 humans [3, 4]. Detection of viral RNA and isolation of a new ebolavirus species (Bombali) from
50 bats have pointed to these animals as a likely reservoir [3-5], similar to the related Marburg virus
51 (MARV) for which the evidence is more established [6-8]. Outbreaks of ebolaviruses have typically
52 been limited to the order of 10-1000 cases by contact tracing and isolation, but in 2013-2015 an
53 outbreak with over 28000 confirmed cases and over 11000 deaths occurred in Sierra Leone,
54 Liberia and Guinea [2, 9]. This outbreak accelerated the development of an effective vaccine and
55 improved therapies against Ebola virus disease [10]. Still, clinical manifestation of Ebola virus
56 infection has historically been associated with mortality rates ranging from 30% to 90% and even
57 the most successful therapies to date provide only a modest improvement of mortality rates and
58 don't offer a cure for advanced disease [2, 10]. Six species of ebolavirus have currently been
59 discovered, including Ebola (a.k.a. Zaire; EBOV), Sudan (SUDV), Bundibugyo (BDBV), Tai Forest
60 (TAFV), Reston (RESTV) and Bombali (BOMV), of which all but the latter two are known to cause
61 severe disease in humans [2].

62 The ebolaviruses are enveloped and contain an 18kb genome of non-segmented, negative sense,
63 single-stranded RNA that encodes seven genes: *NP*, *VP35*, *VP40*, *GP*, *VP30*, *VP24* and *L* [2].
64 The *GP* gene encodes the full-length envelope glycoprotein (GP) as well as two truncated
65 secreted versions (sGP and ssGP) by transcriptional editing [11-13]. The full-length envelope GP
66 is a trimeric class I viral fusion protein and plays an important role in host cell attachment and
67 entry [11]. Following virus internalization by macropinocytosis [14-17], GP binds the Niemann-
68 Pick C1 (NPC1) receptor, triggering fusion of the viral envelope with the host membrane, thereby
69 delivering the ribonucleoprotein complexes in the cytosol where replication will take place [18-20].
70 GP is also the primary target of antibodies produced upon natural infection or vaccination and of
71 monoclonal antibodies developed as antiviral therapeutics [21]. Full-length GP is translated as a
72 ~670 amino acid precursor and is cleaved by host furin into two disulfide-linked subunits: GP1
73 and GP2 [22, 23]. GP1 is responsible for receptor binding and consists of 4 domains: base, head,
74 glycan cap and mucin-like domain (MLD) [11]. The GP2 subunit contains the fusion peptide and
75 has the strongest conservation between different members of the filoviruses [24-26].

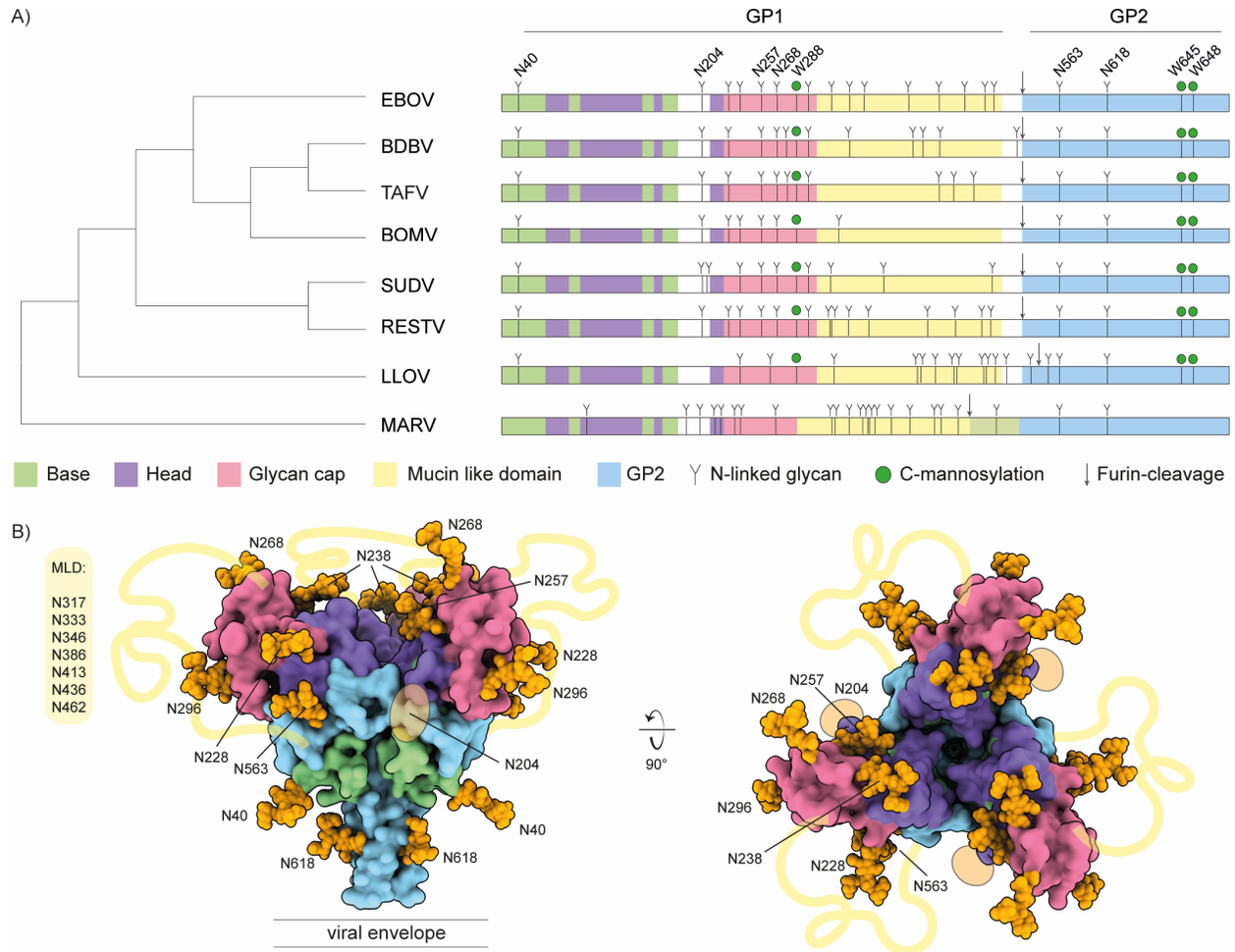
76 There are up to 17 N-linked glycosylation sites in ebolavirus GP, 15 of which are in the GP1
77 subunit, primarily in the glycan cap and MLD (see Figure 1A). The N-linked glycans mediate host-

78 cell attachment through C-type lectins DC-SIGN/L-SIGN [27-29] and have been implicated in
79 shielding GP from binding by neutralizing antibodies [28, 30-36]. Whereas the overall sequences
80 and especially the N-linked glycosylation sites in the base, head and glycan cap are relatively well
81 conserved among ebolavirus species, those in the MLD are highly variable. Further, the MLD is
82 also modified with numerous O-linked glycans. The GP2 subunit contains two N-linked
83 glycosylation sites that are conserved in all known mammalian filoviruses and play important roles
84 in GP expression, stability and cell entry [36, 37]. Besides the numerous N- and O-linked glycans,
85 there are also two predicted tryptophan C-mannosylation motifs in GP. These motifs consist of a
86 WXXW sequence near the glycan cap, and a tandem WXXWXXW sequence in the membrane-
87 proximal region of GP, where a mannose residue may be linked to the C2 atom of the first
88 tryptophan's indole group. The biological function of C-mannosylation is generally not well
89 understood, but known to play a role in the folding, stability and trafficking of secreted
90 glycoproteins, including components of the complement system and gel-forming mucins [38, 39].
91 The C-mannosylation motifs in GP are conserved in all ebolavirus species and the related Lloviu
92 virus (LLOV), but not Marburg virus (MARV). So far, C-mannosylation in the glycan cap has been
93 confirmed in the secreted version sGP [40], but its presence in full-length GP and role in the
94 infection cycle remain unclear.

95 Glycomics studies have confirmed the presence of both complex N- and O-linked glycans in GP
96 [30, 35], but little is known about the site-specific patterns of glycan processing. As glycans play
97 a crucial role in host cell attachment and immune evasion, a better understanding of these
98 patterns in the context of GP structure may help understand mechanisms of infectivity and epitope
99 shielding. Here, we present an in-depth glycoproteomics study of the N-, O- and C-linked glycans
100 of ebolavirus GP. We compare recombinant soluble GP ectodomains of EBOV and BDBV from
101 both human HEK293 and insect S2 cells (material from S2 cells was included as it is often used
102 as a diagnostic and research tool). We demonstrate that the conserved N-linked glycans at N257
103 and N563 are enriched in under-processed high-mannose and hybrid structures in both viral
104 species and cellular expression platforms, suggesting a specific role in host cell attachment
105 through binding of the cell surface lectins DC-SIGN/L-SIGN (which have a markedly higher affinity
106 for high-mannose glycans). We observe that the MLD is modified by numerous O-glycans,
107 comprising a mixture of truncated Tn-antigen and extended, sialylated core 1 and 2 structures,
108 depending on the expression platform. Moreover, we find several O-linked glycosylation sites
109 within the serine/threonine residues of N-linked glycosylation sequons, as well as evidence for O-
110 linked glycans outside the MLD in both EBOV and BDBV GP. We also confirm C-mannosylation
111 in the glycan cap of both ebolavirus species, which only occurs in the HEK293 expression

112 platform. These key findings were confirmed in glycoproteomics experiments on virus-like
113 particles formed by co-expression of full-length GP and VP40. We discuss the observed
114 glycosylation profile in the context of known structures of GP in complex with neutralizing
115 antibodies. Our findings provide a framework to understand the contributions and restrictions of
116 GP glycosylation to the neutralization epitopes of antiviral antibodies.

117



118

119 *Figure 1. A) Schematic of filovirus GP domain structure with annotated N-linked glycosylation and*
120 *C-mannosylation. B) Pseudomodel of EBOV GP with core pentasaccharide of N-linked glycans*
121 *shown as orange spheres.*

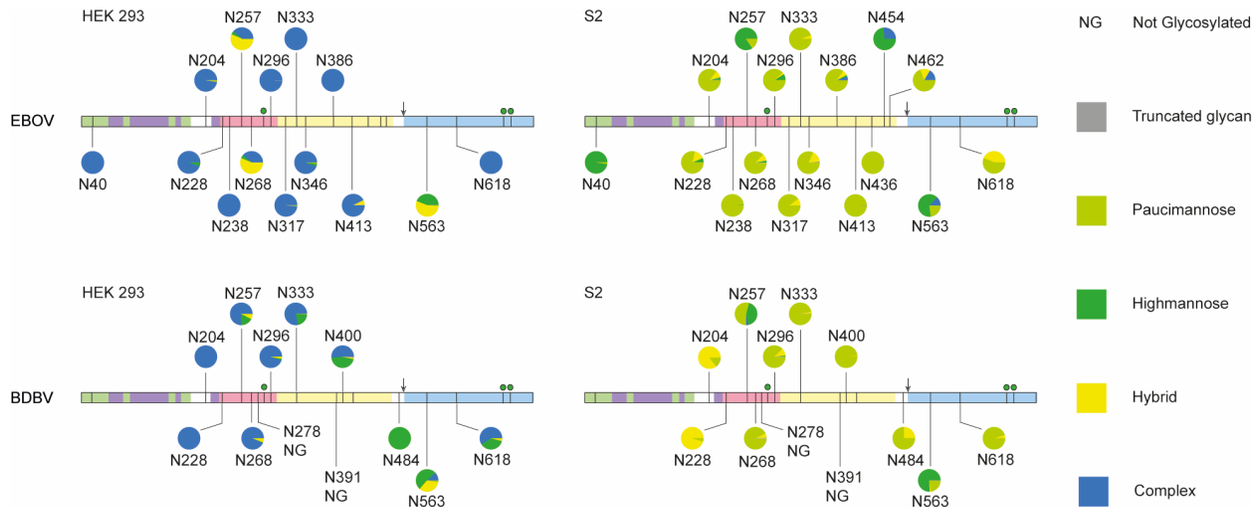
122

123 **Results**

124 We compared the pattern of predicted N-linked glycosylation sites (NXS/T) and C-mannosylation
125 sites (WXXW) of all known ebolavirus species and the related filoviruses MARV and LLOV (see
126 Figure 1A and the sequence alignment in Supplementary Figure S1). The number of predicted N-
127 linked glycosylation sites varies from 9 in BOMV to 17 in EBOV and RESTV. Two of these
128 glycosylation sites are situated in the GP2 subunit (N563 and N618) and they are conserved in
129 all ebolavirus species, MARV and LLOV. All remaining sites are located within the GP1 subunit,
130 especially the glycan cap and MLD. Only 4 sites in GP1 are fully conserved in all ebolavirus
131 species: N40 in the base (also present in LLOV), N204 in a flexible loop between the base and
132 head domains, and N257 and N268 in the glycan cap. All other N-linked glycosylation sites in the
133 glycan cap are shared between a smaller set of ebolavirus species, but virtually all N-linked sites
134 within the MLD are unique, in line with the disordered nature and high overall sequence variability
135 of this region. There are 3 predicted C-mannosylation sites conserved in all ebolavirus species
136 and LLOV, but conspicuously missing in MARV. The first WXXW motif is situated in the glycan
137 cap, at W288 (in EBOV), close to the junction with the MLD. In addition, a tandem WXXWXXW
138 motif is situated at W645/W648 in the membrane proximal region of the GP2 subunit.

139 To visualize the N-glycan shield, we built a pseudomodel of EBOV GP with the core
140 pentasaccharide of each site linked to the corresponding residue of GP1/GP2 (see Figure 1B).
141 The GP trimer forms a chalice-shaped structure with GP2 as the stem, and GP1 as the bowl on
142 top. The conserved sites N40, N204, N257, N268, N563 and N618 are distributed evenly across
143 the structure, whereas the remaining sites are situated primarily at the rim of the bowl extending
144 outwards from the glycan cap. The glycans occupy much of the available surface of GP.
145 Moreover, the disordered MLD connects the tip of the glycan cap with the lower base of the cup
146 and can be expected to further shield the surface of GP. This pseudo model includes only the
147 common core pentasaccharide of the N-linked glycans and it is not known how the glycans are
148 processed in the context of folded GP, as predicted sites are not always glycosylated and the
149 processing from high-mannose precursors to hybrid and mature complex glycans may depend on
150 many unpredictable factors, including local structural constraints. We investigated the patterns of
151 site-specific glycosylation of ebolavirus GP with LC-MS/MS based glycoproteomics experiments,
152 using recombinant soluble ectodomains (GP Δ TM) of EBOV and BDBV, as well as the
153 corresponding full-length GP from virus-like particles produced by co-expression with VP40. We
154 compared GP Δ TM from human HEK293 and insect S2 cells, both commonly used for structural
155 biology studies, experimental immunizations, antibody selection and serological tests.

156



157

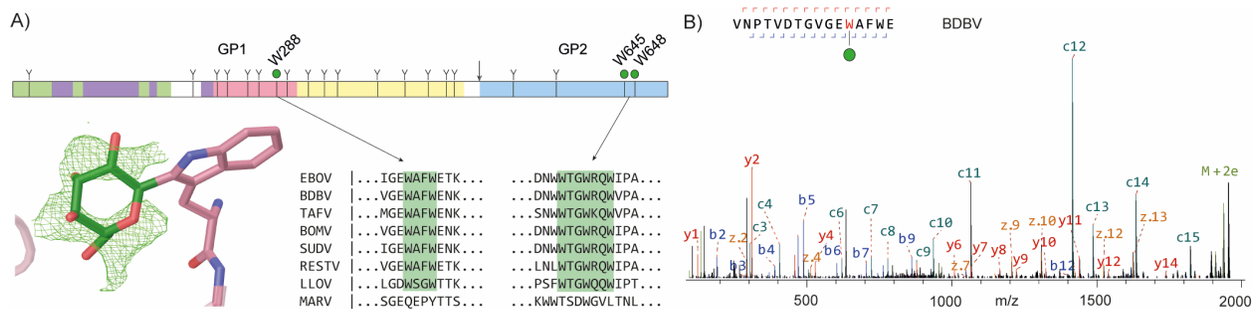
158 **Figure 2. Overview of site-specific N-linked glycan processing in ebolavirus GP Δ TM from HEK293**
159 **and S2 cells as determined by LC-MS/MS. The glycans were classified by HexNAc content as**
160 **truncated, paucimannose, high-mannose, hybrid or complex. Shown is the average of a duplicate**
161 **experiment.**

162

163 Our results cover 14/17 and 17/17 predicted sites in EBOV GP from HEK293 and S2 cells,
164 respectively, as well as 12/14 predicted sites of BDBV GP from both expression platforms (see
165 Figure 2 and Supplementary Table S1). As expected, the N-linked glycosylation patterns of GP
166 from HEK293 and S2 cells are dominated by complex and paucimannose/hybrid glycans,
167 respectively. The glycosylation of GP from especially HEK293 cells is extremely heterogeneous,
168 with some sites carrying over 40 unique glycan compositions. Predicted sites N278 and N391 in
169 BDBV were only detected as unglycosylated asparagines. Most detected glycan compositions
170 are compatible with di-, tri- and tetra-antennary, galactosylated complex glycans with or without
171 a single (core) fucose residue and a variable number of terminal sialic acids, as previously
172 described in glycomics analyses [30, 35]. While complex glycans dominate the overall picture,
173 selected sites show clear and robust enrichment of unprocessed glycans, (*i.e.* high-mannose and
174 hybrid structures in the HEK293-derived samples). These include particularly the conserved N257
175 and N563 sites, in both EBOV and BDBV GP. In good agreement with these observations in the
176 HEK293-derived samples, N257 and N563 are also enriched in unprocessed high-mannose
177 glycans in the S2-derived samples of both EBOV and BDBV GP, indicating that processing of
178 these sites is somehow structurally restricted. Our pseudomodel of EBOV GP indicates that N257

179 may be partially buried between the head domain and glycan cap (*i.e.* its first asparagine-linked
180 GlcNAc residue), and N563 similarly between the head domain and GP2. Whereas sites
181 N40/N268/N454 in EBOV GP, and N400/N454 in BDBV GP also show elevated levels of
182 unprocessed glycans in selected samples, we refrain from any conclusions on these sites due to
183 a relatively shallow coverage in the underlying mass spec data and the lack of agreement between
184 HEK293/S2 or EBOV/BDBV samples. Nevertheless, the data clearly indicate a lack of processing
185 at the conserved N257 and N563 sites in both tested ebolavirus species and expression
186 platforms. These findings are confirmed in LC-MS/MS experiments of EBOV and BDBV virus-like
187 particles derived from HEK293 cells, where we also detected a large fraction of high-mannose
188 and hybrid glycans at N257/N563 against a background of highly processed complex glycans at
189 the remaining covered sites (see Supplementary Figures S2 and S3). The abundance of
190 unprocessed glycans is most prominent at site N563, where the vast majority of glycans consists
191 of hybrid and high-mannose forms in both full-length GP and GPΔTM. At site N257, the
192 abundance of unprocessed glycans is markedly lower in the full-length EBOV GP from VLPs.

193



194

195 **Figure 3. C-mannosylation in ebolavirus GP. A) schematic domain structure ebolavirus GP with**
196 **highlighted C-mannosylation sites. The modelled structure of C-mannosylated W288 was based**
197 **on average Fo-Fc density (shown in green) of twelve isomorphous GP crystal structures. B) LC-**
198 **MS/MS spectrum of C-mannosylated W288 from BDBV GPΔTM. Note the prominent c11-c12**
199 **peaks provide direct evidence for the presence and localization of the Hex (+162 Da) modification**
200 **of W288.**

201

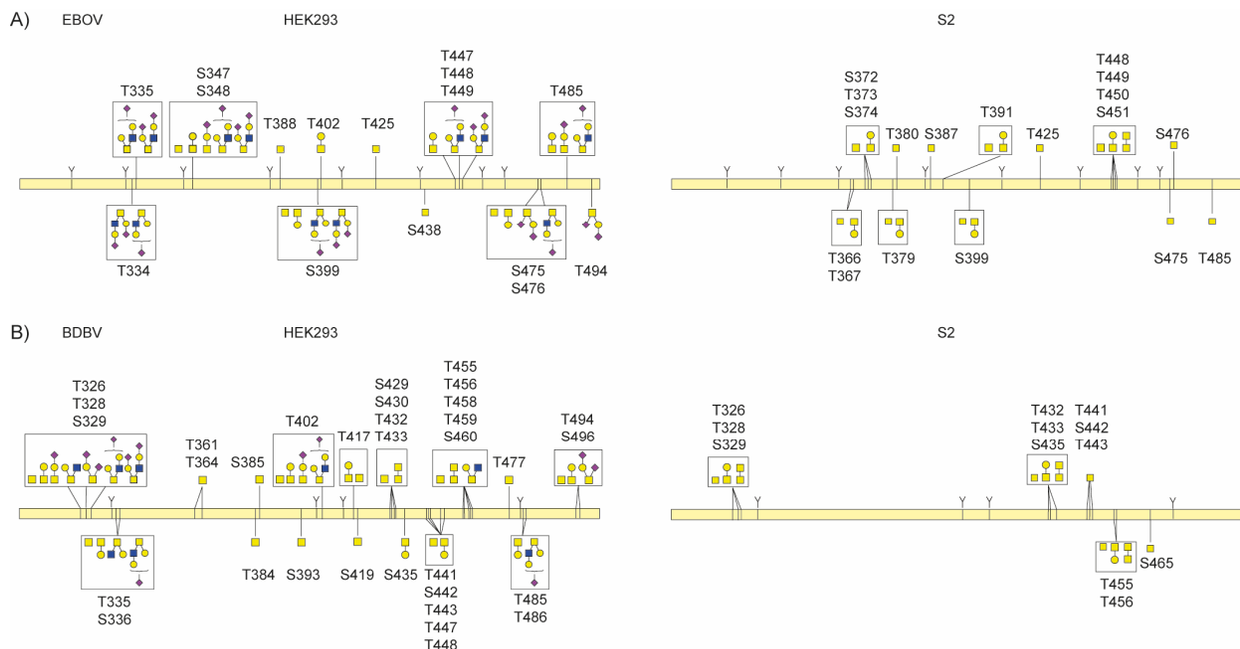
202 Our experiments also cover the C-mannosylation site at W288 in the glycan cap (see Figure 3).
203 The GPΔTM constructs used here are truncated before the second C-mannosylation motif at
204 W645/W648 and therefore not covered in these experiments. The GP samples derived from
205 HEK293 cells both contain a mixture of C-mannosylated and unmodified W288, with an estimated

206 occupancy of 1-10%. In contrast, this modification is completely absent in both samples derived
207 from S2 cells. The presence of C-mannosylated W288 was confirmed in the glycoproteomics
208 experiments on full-length GP in the virus-like particles formed by co-expression with VP40 (see
209 Supplementary Figure S4). Unfortunately, we could not detect any peptides that cover the second
210 C-mannosylation motif at W645/W648 in these samples.

211 The W288 site is situated in a lesser ordered region of the glycan cap (the β 17- β 18 loop) just
212 before the start of the MLD, which is deleted in the constructs of most structural studies. While
213 most available GP structures do not model the corresponding region, we identified a set of 12
214 deposited isomorphous GP crystal structures of HEK293-derived material with electron density for
215 W288 and its adjacent residues [41-45]. The individual crystal structures did not show a clear F_o -
216 F_c density corresponding to the C2-linked mannose, but after averaging all available electron
217 density maps, a clear ring structure did appear. The weak observed electron density is consistent
218 with the low occupancy of the modification observed in our glycoproteomics experiments. The
219 C2-linked mannose residue was modelled in the extra density, positioning it at the exposed
220 surface of the glycan cap, pointing towards the center of the β 17- β 18 loop.

221 We also mapped out the patterns of O-linked glycosylation in ebolavirus GP (see Figure 4). In
222 contrast to N- and C-linked glycosylation, there is no clear sequence motif to predict O-linked
223 glycosylation sites. The modification generally occurs in serine/threonine-rich disordered regions,
224 such as the MLD of filovirus GPs. Whereas the presence of O-glycans in the MLD is well-known,
225 the precise localization of these modifications remains unclear (the MLD contains more than 50
226 possible S/T residues). For these experiments, we first removed all N-linked glycans by PNGase
227 F digestion. This reduces the complexity of the glycopeptide mixture to facilitate O-linked
228 glycopeptide identification and site localization, while leaving a clear mark at the digested N-
229 glycan site by deamidation of the asparagine residue (resulting in a +1 Da mass shift).

230 In EBOV GP, we detected 12 unique O-linked glycosylation sites in the MLD of GP Δ TM from
231 HEK293 cells versus 12 in S2 cells, with 5 sites in common. In BDBV GP we detected 16 unique
232 O-linked glycosylation sites in the MLD of GP Δ TM from HEK293 cells versus 5 in S2 cells, with 4
233 sites in common. Whereas O-linked glycosylation was dominated by simple Tn antigen and core
234 1 structures in samples from S2 cells, samples from HEK293 cells also contained extended and
235 sialylated core 1 and core 2 structures, especially in EBOV GP. Multiple unique glycan
236 compositions were often detected for a given site, further adding to the extreme heterogeneity of
237 GP due to its glycosylation.



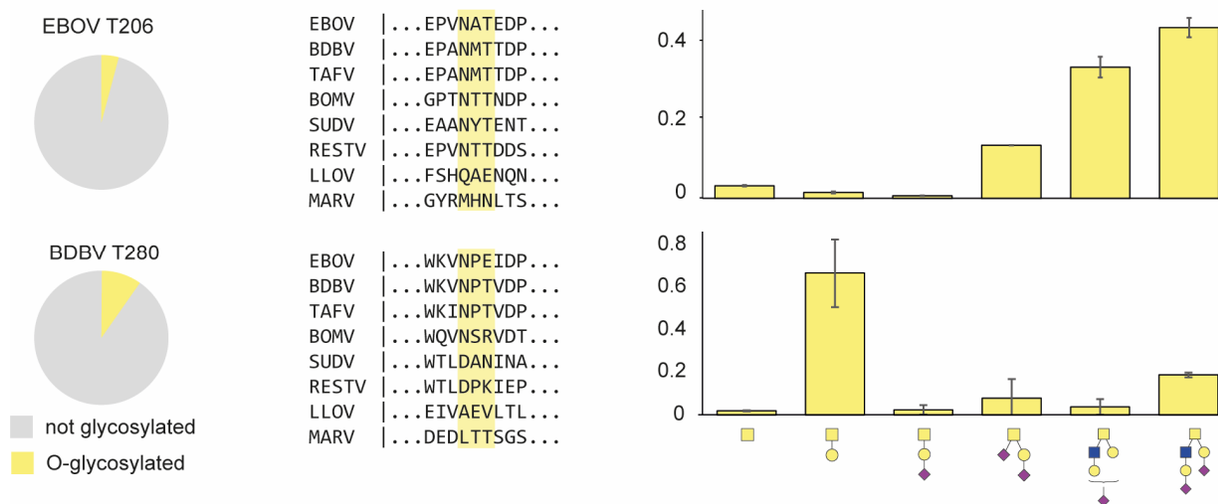
238

239 **Figure 4. O-linked glycosylation in the MLD of ebolavirus GP Δ TM from EBOV (A) and BDBV (B).**
 240 **Glycans drawn within a box represent multiple detected compositions per site. Glycans connected**
 241 **to multiple indicated sites could not be unambiguously localized from the LC-MS/MS data.**

242

243 We also detected several O-glycosylation sites outside the MLD of both EBOV and BDBV GP
 244 (see Figure 5). In BDBV GP we detected O-linked glycosylation at T280, with 6 unique glycan
 245 compositions amounting to an estimated total occupancy of ~10%. This threonine residue is part
 246 of a putative NPT glycosylation sequon, but we only detect the unmodified asparagine, which
 247 remains unprocessed presumably because of the following proline residue. The modified
 248 threonine is shared only by BDBV and TAFV GP, but absent in EBOV, SUDV, BOMV, RESTV,
 249 MARV and LLOV. We also detected O-linked glycosylation at T206 in EBOV GP, with 6 unique
 250 glycan compositions and an estimated occupancy of ~5%. This threonine is part of the
 251 glycosylation sequon of N204, which is fully occupied by N-glycans as evidenced by the
 252 deamidated asparagine and the N-linked glycoproteomics data discussed earlier. The N-linked
 253 glycosylation sequon including the modified threonine is conserved among all ebolavirus species,
 254 but does not exist in MARV and LLOV. Residues adjacent to this sequon show substantial
 255 variation between ebolavirus species and modified T206 was not detected in the BDBV GP
 256 samples. The close juxtaposition of N- and O-glycans is also observed in the MLD of both EBOV
 257 and BDBV GP, where T335 is part of the N-linked glycosylation sequon of N333 and detected in
 258 GP Δ TM from both species as an O-linked glycosylation site. Similarly, S348/T388/S438 in EBOV

259 GP and T402/T488 in BDBV GP are all part of N-glycosylation sequons. Finally, we also observed
 260 O-linked glycosylation within the strep-tag of the constructs (see Supplementary Figure S5). The
 261 presence of the O-linked glycans at T206 (EBOV) and T280 (BDBV) outside the MLD could be
 262 confirmed in our glycoproteomics measurements of full-length GP from virus-like particles (see
 263 Supplementary Figure S4).
 264

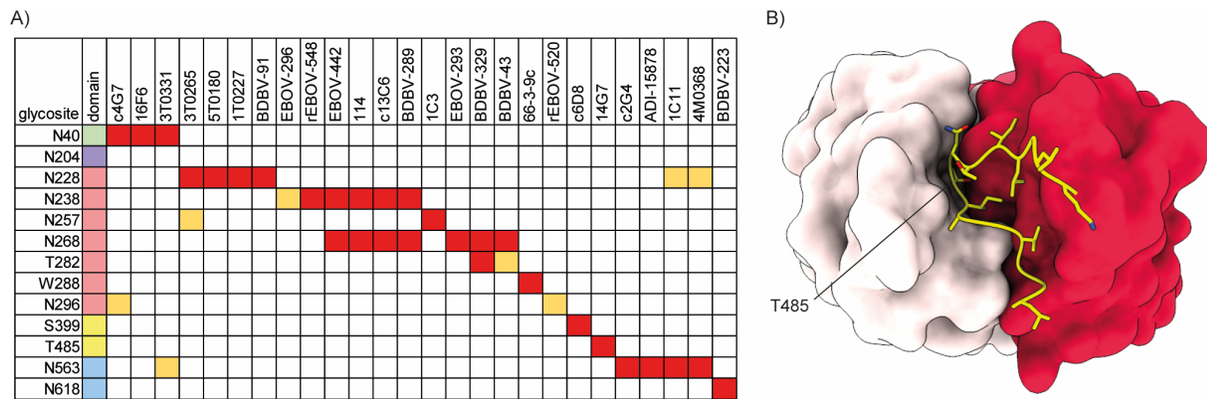


265
 266 **Figure 5. O-linked glycosylation outside the MLD in ebolavirus GP. (left) pie charts represent the**
 267 **occupancy of the O-linked modification. (middle) sequence conservation of the detected O-linked**
 268 **glycosylation sites. (right) distribution of glycan types at the indicated sites (average +/- stdev from**
 269 **duplicate experiments)**

270
 271 This high extent of glycosylation must be accommodated by antibodies against ebolavirus GP.
 272 To understand the contribution of glycans to neutralization epitopes (and restrictions they impose)
 273 we screened the Protein Data Bank for structural models of ebolavirus GP in complex with
 274 neutralizing antibodies and also looked for linear B-cell epitopes reported in literature that span
 275 the glycosylation sites we detected in our experiments (see Figure 6A) [46-56]. The glycan-rich
 276 epitopes we report in this overview include both cases of direct contacts between modelled GP
 277 glycans and CDR residues, as well as brushing interactions of adjacent glycans with the
 278 framework regions of the variable domains, which may sterically restrict binding. It should be
 279 noted that glycans are typically incompletely modelled in GP-antibody structures and that
 280 inference of these brushing interactions is not an exact determination. This overview indicates

281 that neutralizing antibodies span a broad range of epitopes that cover essentially all N-linked
 282 glycosylation sites (N204 and the entire MLD have not yet been modelled in structural studies
 283 and are therefore not represented in this analysis). The conserved GP1 glycans (N40, N257, and
 284 N268) all contribute to the epitopes of neutralizing antibodies, with possible glycan-antibody
 285 interactions for N268 reported in the epitopes of as many as 7 unique monoclonal antibodies.
 286 This includes the therapeutic monoclonal antibody Mab114, which makes additional contacts with
 287 N238. The components of the therapeutic ZMapp mixture (c2G4, c4G7 and c13C6) also interact
 288 with N40, N238, N268 and N563. The epitopes of the three components in the therapeutic REGN-
 289 EB3 mixture are not defined to atomic detail (and therefore not included in the overview), but
 290 published negative stain EM reconstructions suggest possible interaction with N563 and the
 291 glycan cap [57].

292



293

294 *Figure 6. Glycan-containing epitopes of monoclonal antibodies against ebolavirus GP. A)*
 295 *overview of glycosylation sites associated with indicated epitopes. Direct contacts and clashes*
 296 *with CDRs are indicated in red, brushing interactions with framework regions in light orange. B)*
 297 *highlighted structure of 14G7 in complex with a linear epitope from the MLD (PDB ID:2Y6S),*
 298 *showing that the O-linked glycosylation site T485 is deeply buried in the cleft between heavy (dark*
 299 *red) and light chain (light pink).*

300

301 Besides N-glycans, we also noted several putative interactions with C-mannosylated W288 and
 302 O-linked glycosylation sites. The monoclonal antibodies BDBV-329 and BDBV-43 are in close
 303 proximity to T280 with their CDRH3 and framework 3 regions, respectively. The monoclonal
 304 antibody 66-3-9c binds a linear epitope that spans the C-mannosylation site W288. Similarly,
 305 c6D8 binds a linear epitope that spans the O-linked glycosylation site S399. The monoclonal

306 antibody 14G7 binds to a linear epitope in the MLD that spans the O-linked glycosylation site
307 T485. A crystal structure of the 14G7 Fab in complex with its unglycosylated epitope reveals that
308 T485 is buried deep within the cleft between heavy and light chains, where it is in direct contact
309 with CDRH3 residues (see Figure 6B). It is therefore unlikely to accommodate the bulky O-glycans
310 detected in our experiments, further highlighting the potential epitope shielding effects of not just
311 N-glycans, but also O-glycans in the MLD.

312

313 **Discussion**

314 Here we have presented a comprehensive overview of glycosylation in ebolavirus GP, using
315 glycoproteomics to resolve the patterns of site-specific N-, O- and C-linked glycans. In the GP
316 samples derived from HEK293 cells, we observed heterogeneous, complex N-glycosylation
317 overall, but noted enrichment of unprocessed glycans at two conserved sites N257 and N563. It
318 is known that ebolavirus GP interacts with cell surface lectins DC-SIGN/L-SIGN in a high-
319 mannose glycan dependent manner [27-29]. Our results indicate that the unprocessed glycans
320 present at N257 and N563 may be primarily responsible for the interaction, thereby facilitating
321 host cell attachment and infectivity.

322

323 We also detected up to 16 unique site-specific O-glycans in the MLD of GP, revealing a
324 heterogeneous mixture of not only simple Tn-antigen (*i.e.* a single GalNAc), but also extended
325 sialylated core 1 and core 2 structures. The sites detected in our experiments are likely just the
326 tip of the iceberg, as the dense decoration of the MLD with O-glycans may make proteolytic
327 digestion of the MLD especially difficult and the presence of multiple glycans in the same peptide
328 makes it exponentially more challenging to confidently make assignments from the raw LC-
329 MS/MS data.

330

331 We further confirmed the presence of the C-mannosylation site in GP at W288, which is
332 completely conserved in all ebolavirus species and LLOV, but not MARV. The second motif at
333 W645/W658 is also missing in MARV, but whereas the MARV sequence at W288 completely
334 diverges from the ebolavirus species, the MARV region corresponding to W645/W648 is similarly
335 rich in tryptophan residues (see Figure 3A). The MARV GP sequence in this region is thereby
336 primed to acquire a C-mannosylation motif through a single deletion or a tryptophan substitution
337 at any of four adjacent positions. Conversely, this could also indicate that the C-mannosylation

338 motifs were present in a common ancestor with ebolaviruses and LLOV but lost in MARV.
339 Whereas C-mannosylation is known to be important for the stability and folding of secreted human
340 glycoproteins [38, 39], its role in ebolavirus replication and pathogenesis remains unclear.

341
342 The presence of heterogeneous N-, O- and C-linked glycosylation add up to a staggering
343 complexity of GP composition. In the case of EBOV GP, the 17 N-glycosylation sites alone, each
344 linked to on average a dozen unique glycan compositions, already give rise to an enormous
345 number of permutations. Add to this the heterogeneity of O-linked glycosylation and a picture
346 emerges where no two copies of GP on a virion are strictly identical. Meanwhile, the glycans
347 represent a major component of the overall GP structure. The 17 N-glycans already contribute
348 approximately one quarter of the molecular weight of GP, all situated at its exposed surface
349 (counting ~74 kDa of polypeptide and on average 1.5 kDa per N-glycan). Although antibodies
350 evidently mount a neutralizing response to infection or vaccination, the high variability of the
351 exposed GP surface due to heterogeneous glycosylation must frustrate the overall binding
352 efficiency of antibodies that accommodate glycans in their epitope and restrict good binders to
353 only the core elements of glycans that are common between the countless variations of GP
354 present on the surface of mature virions. From this perspective the glycans contribute to immune
355 evasion not only by sterically shielding neutralization epitopes, but also by blurring the molecular
356 identity of the envelope glycoprotein.

357
358 Glycoproteomics studies on other envelope glycoproteins from divergent virus species, such as
359 HIV-1, Lassa virus, MERS-CoV, SARS-CoV-2 and the herpesviruses, all show a similar trend of
360 heterogeneous complex glycosylation with unprocessed glycans at selected sites [58-64]. From
361 studies on HIV-1 gp120 it has been shown that the lack of processing of certain N-glycans is
362 caused by local crowding and reduced accessibility to processing enzymes, resulting in
363 enrichment of high-mannose glycans [65-68]. Neither glycan at N257 and N563 in ebolavirus GP
364 fits this description, but both have their first N-linked GlcNAc residue partially buried in interactions
365 with surrounding side chains. An intriguing possibility is that these interactions at the base of the
366 glycan limit its conformational degrees of freedom to negatively impact processing at the
367 antennae. Whereas N-linked glycosylation is universally known to play a role in the replication
368 cycles of enveloped viruses, O-linked glycosylation is less well-studied and perhaps less
369 common. Recent studies on a range of herpesvirus glycoproteins, SARS-CoV-2 Spike, the
370 attachment proteins G of paramyxoviruses, and hepatitis C virus E2 point to a role in envelope
371 glycoprotein processing, trafficking, host cell attachment, and immune evasion [58, 59, 69-72].

372

373 All three types of glycosylation observed in ebolavirus GP will alter its antigenic surface. The
374 overview provided in Figure 6 illustrates how the site-specific glycosylation observed in our
375 experiments contributes to and restricts the epitopes of currently known neutralizing antibodies.
376 Whereas bulkier types of glycans at specific sites may indeed modulate the binding affinity of the
377 indicated antibodies, the presented overview shows quite the opposite of the glycans' shielding
378 effects in ebolavirus GP. That interpretation would be a kind of survivorship bias, as the
379 monoclonal antibodies have been selected for binding and neutralization. The effective shielding
380 would perhaps be better illustrated by the antibodies that don't bind or neutralize ebolavirus
381 infection because of the steric clashes with glycans. Several studies have illustrated this by
382 showing greatly enhanced sensitivity of (pseudo) ebolavirus to serum neutralization after glycan
383 removal by mutagenesis [31, 32].

384

385 Future studies may shed light on how the heterogenous glycan composition of GP may modulate
386 antibody binding. Similarly, the close juxtaposition of N- and O-linked glycan raises the intriguing
387 possibility of an interplay between the two types of modifications. Furthermore, the exact role of
388 C-mannosylation and O-linked glycans outside the MLD also remain to be investigated.
389 Nevertheless, we have presented a comprehensive overview of ebolavirus GP glycosylation that
390 may provide a useful framework for future immunological and structural studies on GP-antibody
391 interactions.

392

393 **Acknowledgements**

394 The authors would like to thank everyone in the Biomolecular Mass Spectrometry and Proteomics
395 group at Utrecht University for support and helpful discussions. This research was funded by the
396 Dutch Research Council NWO Gravitation 2013 BOO, Institute for Chemical Immunology (ICI;
397 024.002.009) to JS, and NIAID U19 AI142790 to EOS.

398

399 **Materials and Methods**

400 **Ebolavirus GP sequence analysis.**

401 The indicated full-length GP reference sequences were downloaded from UniProt. The LLOV-GP
402 sequence had to be reconstructed from the two separate GP1 and GP2 entries in UniProt.
403 Sequence alignment was performed with ClustalX 2.1 [73]. The sequence IDs and resulting
404 alignment are provided in the Supplementary Information. The cladogram was generated with
405 FigTree (version 1.4.4). N-linked glycosylation sites were predicted by identifying all NXS/T
406 sequences with NetNGlyc-1.0. C-mannosylation sites were predicted by identifying all WXXW
407 sequences by manual inspection.

408

409 **Pseudomodel building of glycosylated EBOV GP**

410 A homology model EBOV GP (strain Mayinga '76) was generated with SWISS-MODEL to fill in
411 missing loops using PDB ID 5jq3 as a template [45, 74]. The core pentasaccharides were added
412 to GP1 and GP2 subunits separately with GLYCAM Glycoprotein Builder (GLYCAM Web, Woods
413 group 2021). The full trimer was reconstructed by alignment with the biological assembly of 5jq3.
414 The loop containing N204 was manually removed because it produced clashes with neighboring
415 subunits in the full trimer. The figures were generated with ChimeraX 1.2.5 [75].

416

417 **Glycan-containing epitopes of monoclonal antibodies**

418 Structures of monoclonal antibodies in complex with ebola virus GP were retrieved from the PDB
419 (with PDB IDs: 2y6s, 3s88, 5fhc, 5kel, 5kem, 5ken, 6ea7, 6n7j, 6pci, 6qd7, 6qd8, 6s8d, 7kej,
420 7kew, 7kex, 7kfe, 7kf9 and 7kfb) [46-56]. The structures were aligned with the glycosylated
421 ZEBOV GP pseudomodel described above, using the MatchMaker function of ChimeraX 1.2.5,
422 and glycans within 6 Å of the CDRs or framework regions of the modelled antibodies were
423 included in the overview.

424

425 **Modelling of C-mannosylated W288**

426 identified 12 isomorphous published crystal structures of HEK293-derived EBOV GP samples in
427 the PDB (PDB codes 6f6n, 6f6i, 6f54, 6nae, 5jqb, 5jq7, 5jq3, 6g9b, 6g9i, 6g95, 6hro and 6hs4)
428 [41-45]. To obtain higher signal-to-noise ratios from these maps, the $2F_o-F_c$ and F_o-F_c difference
429 maps were averaged using COOT [76]. This averaged map showed clear ring-shaped electron
430 density next to W288 in the F_o-F_c difference map at a contour level of 2.7 root mean square
431 deviation. A C-mannosyl group was modelled next to W288 using the EBOV GP structure 6hs4
432 as a template in COOT. To accommodate realistic geometry, the tryptophan had to be
433 repositioned slightly, albeit still in agreement with the local electron density. Care was taken to
434 model the mannose with a ring-flipped 1C_4 chair conformation [77-79].

435

436 **Production and purification of ebola virus GP ectodomains**

437 EBOV and BDBV GP were produced by both transient transfection of HEK293T cells and stable
438 transfection of *Drosophila melanogaster* S2 cells. Lipofectamine 3000 (Invitrogen) was used to
439 transfect HEK293T cells, and Effectene (Qiagen) was used to produce stable S2 cells with a
440 modified pMT-puro vector plasmid containing the GP gene of interest and stable selection of
441 transfected cells with 6 μ g/mL puromycin. HEK293T cells were grown at 37°C with 5% CO₂ in
442 DMEM media (Gibco) supplemented with 10% FBS in T75 flasks and expanded into 10-stack
443 flasks (Corning) for transfection. S2 cells were selected at 27°C in complete Schneider's medium
444 and then transferred to Insect Xpress medium (Lonza) for large-scale expression in 2-liter
445 Erlenmeyer flasks. Secreted GP ectodomain expression was induced with 500 mM CuSO₄, and
446 supernatant harvested after 4 days. Ebola virus GP was engineered with a double Strep-tag at
447 the C terminus to facilitate purification using Strep-trap HP 5mL column (GE) and then further
448 purified by Superdex 200 size exclusion chromatography (SEC) in 25 mM Tris-buffered saline
449 (Tris-HCl, pH 7.5, 150 mM NaCl [TBS]).

450

451 **Production and purification of ebolavirus-like particles**

452 EBOV and BDBV virus-like particles were produced by transfecting HEK293T cells.
453 Polyethylenimine (PEI) was used to transfect HEK293T cells with a modified phCMV plasmid
454 containing the full-length GP gene of interest and a modified pTriEx plasmid containing the full-
455 length EBOV VP40 gene at a 2:5 ratio (w:w), respectively. The VLP supernatant was clarified by

456 centrifugation after 48 hours. The clarified supernatant was further purified using a 20% sucrose
457 cushion ultra-centrifuge spin at 106,800xg for 3hrs. The cushion and supernatant was carefully
458 decanted and the pellet washed with sterile PBS 2 times. Following the wash, the pellet was
459 incubated overnight in 0.75mL of PBS and resuspended.

460

461 **Glycoproteomics sample preparation**

462 For N-linked glycan analysis, the recombinant GP was denatured at 95 °C in a final concentration
463 of 2% sodium deoxycholate (SDC), 200 mM Tris/HCl, 10 mM tris(2-carboxyethyl)phosphine, pH
464 8.0 for 10 min followed with 30 min reduction at 37 °C for 30 min. Samples were next alkylated
465 by adding 40 mM iodoacetamide and incubated in the dark at room temperature for 45 min. 3 µg
466 recombinant GP was used for each protease digestion. Samples were split in three for parallel
467 digestion with trypsin (Promega), alpha lytic protease (Sigma), and gluC (Sigma)-trypsin. For each
468 protease digestion, 18 µL of the denatured, reduced, and alkylated samples was diluted in a total
469 volume of 100 µL 50 mM ammonium bicarbonate, adding proteases in a 1:15 ratio (w:w) for
470 incubation overnight at 37 °C. For the gluC-trypsin digestion, gluC was added first for two hours,
471 followed by incubation with trypsin overnight. After overnight digestion SDC was removed through
472 precipitation by adding 2 µL formic acid (FA) and centrifugation at 14,000 rpm for 20 min.
473 Following centrifugation, the supernatant containing the peptides was collected for desalting on a
474 30 µm Oasis HLB 96-well plate (Waters). The Oasis HLB sorbent was activated with 100%
475 acetonitrile and subsequently equilibrated with 10% formic acid in water. Next, peptides were
476 bound to the sorbent, washed twice with 10% formic acid in water and eluted with 100 µL of 50%
477 acetonitrile/5% formic acid in water (v/v). The eluted peptides were vacuum-dried and
478 resuspended in 100 µL of 2% formic acid in water. For O-linked glycan analysis, the recombinant
479 GP was first treated with PNGase F (Sigma) to remove N-glycans. 4 µL PNGase F was added to
480 the sample in PBS and incubated at 37 °C overnight. Following N-glycan removal, GPs were
481 digested following the same protocol as for N-linked glycan analysis, using parallel digestion with
482 trypsin and aLP. Both N- and O-linked analyses were performed in duplicate.

483

484 **Glycoproteomics LC-MS/MS measurements**

485 For each sample and protease digestion, approximately 0.15 µg of peptides were run by online
486 reversed phase chromatography on an Agilent 1290 UHPLC or Dionex UltiMate 3000 (Thermo

487 Fisher Scientific) coupled to a Thermo Scientific Orbitrap Fusion mass spectrometer.
488 A Poroshell 120 EC C18 (50 cm x 75 μ m, 2.7 μ m, Agilent Technologies) analytical column and
489 a ReproSil-Pur C18 (2 cm x 100 μ m, 3 μ m, Dr. Maisch) trap column were used for peptide
490 separation. The duplicate samples were analyzed with two different mass spectrometry methods,
491 using identical LC-MS parameters and distinct fragmentation schemes. In one method, peptides
492 were subjected to Electron Transfer/Higher-Energy Collision Dissociation fragmentation. In the
493 other method, all precursors were subjected to HCD fragmentation, with additional EThcD
494 fragmentation triggered by the presence of glycan reporter oxonium ions. A 90-min LC gradient
495 from 0% to 44% acetonitrile was used to separate peptides at a flow rate of 300 nl/min. Data was
496 acquired in data-dependent mode. Orbitrap Fusion parameters for the full scan MS spectra were
497 as follows: a standard AGC target at 60 000 resolution, scan range 350-2000 m/z, Orbitrap
498 maximum injection time 50 ms. The ten most intense ions (2+ to 8+ ions) were subjected to
499 fragmentation. For the EThcD fragmentation scheme, the supplemental higher energy collision
500 dissociation energy was set at 27%. MS2 spectra were acquired at a resolution of 30,000 with an
501 AGC target of 800%, maximum injection time 250 ms, scan range 120-4000 m/z and dynamic
502 exclusion of 16 s. For the triggered HCD-EThcD method, the LC gradient and MS1 scan
503 parameters were identical. The ten most intense ions (2+ to 8+) were subjected to HCD
504 fragmentation with 30% normalized collision energy from 120-4000 m/z at 30,000 resolution with
505 an AGC target of 100% and a dynamic exclusion window of 16 s. Scans containing any of the
506 following oxonium ions within 20 ppm were followed up with additional EThcD fragmentation with
507 27% supplemental HCD fragmentation. The triggering reporter ions were: Hex(1) (129.039;
508 145.0495; 163.0601), PHex(1) (243.0264; 405.0793), HexNAc(1) (138.055; 168.0655; 186.0761),
509 Neu5Ac(1) (274.0921; 292.1027), Hex(1)HexNAc(1) (366.1395), HexNAc(2) (407.166),
510 dHex(1)Hex(1)HexNAc(1) (512.1974), and Hex(1)HexNAc(1)Neu5Ac(1) (657.2349). EThcD
511 spectra were acquired at a resolution of 30,000 with a normalized AGC target of 400%, maximum
512 injection time 250 ms, and scan range 120-4000 m/z.

513

514 **Glycoproteomics data analysis**

515 The acquired data was analysed using Byonic (v3.9.6 [80]) against a custom database of
516 recombinant ebola virus GP protein sequences and the proteases used in the experiment,
517 searching for glycan modifications with 12/24 ppm search windows for MS1/MS2, respectively.
518 Up to six missed cleavages were permitted using C-terminal cleavage at R/K for trypsin, R/K/E/D
519 for gluC-trypsin, or T/A/S/V for alpha lytic protease. For N-linked analysis, carbamidomethylation

520 of cysteine was set as fixed modification, oxidation of methionine/tryptophan as variable common
521 1, and hexose on tryptophan as variable rare 1. N-glycan modifications were set as variable
522 common 2, allowing up to max. 2 variable common and 1 rare modification per peptide. All N-
523 linked glycan databases from Byonic were merged into a single non-redundant list to be included
524 in the database search. All reported glycopeptides in the Byonic result files were manually
525 inspected for quality of fragment assignments. All glycopeptide identifications were merged into
526 a single non-redundant list per sequon. Glycans were classified based on HexNAc content as
527 truncated (≤ 2 HexNAc; < 3 Hex), paucimannose (2 HexNAc, 3 Hex), high-mannose (2 HexNAc;
528 > 3 Hex), hybrid (3 HexNAc) or complex (> 3 HexNAc). Byonic search results were exported to
529 mzIdentML format to build a spectral library in Skyline (v20.1.0.31 [81]) and extract peak areas
530 for individual glycoforms from MS1 scans. The full database of variable N-linked glycan
531 modifications from Byonic was manually added to the Skyline project file in XML format. Reported
532 peak areas were pooled based on the number of HexNAc, Fuc or NeuAc residues to distinguish
533 truncated, paucimannose, high-mannose, hybrid, and complex glycosylation, or the degree of
534 fucosylation and sialylation, respectively. For O-linked analysis, all the same protease digestion
535 parameters and peptide modifications were used, with the addition of deamidation at
536 asparagine/glutamine as variable rare 1. O-glycan modifications were set as variable common 6,
537 allowing a maximum of 6 variable common and 2 rare modifications per peptide.

538

539 **Data Availability**

540 The raw LC-MS/MS files and glycopeptide identifications have been deposited to the
541 ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier
542 PXD031459. All reagents and relevant data are available from the authors upon request.

543

544

545 **References**

546

- 547 1. Burk, R., L. Bollinger, J.C. Johnson, J. Wada, S.R. Radoshitzky, G. Palacios, S. Bavari,
548 P.B. Jahrling and J.H. Kuhn, *Neglected filoviruses*. FEMS microbiology reviews, 2016.
549 **40**(4): p. 494-519.
- 550 2. Jacob, S.T., I. Crozier, W.A. Fischer, A. Hewlett, C.S. Kraft, M.-A. de La Vega, M.J. Soka,
551 V. Wahl, A. Griffiths and L. Bollinger, *Ebola virus disease*. Nature reviews Disease
552 primers, 2020. **6**(1): p. 1-31.
- 553 3. Leroy, E.M., B. Kumulungui, X. Pourrut, P. Rouquet, A. Hassanin, P. Yaba, A. Délicat, J.T.
554 Paweska, J.-P. Gonzalez and R. Swanepoel, *Fruit bats as reservoirs of Ebola virus*.
555 Nature, 2005. **438**(7068): p. 575-576.
- 556 4. Marí Saéz, A., S. Weiss, K. Nowak, V. Lapeyre, F. Zimmermann, A. Düx, H.S. Kühl, M.
557 Kaba, S. Regnaut and K. Merkel, *Investigating the zoonotic origin of the West African*
558 *Ebola epidemic*. EMBO molecular medicine, 2015. **7**(1): p. 17-23.
- 559 5. Goldstein, T., S.J. Anthony, A. Gbakima, B.H. Bird, J. Bangura, A. Tremeau-Bravard, M.N.
560 Belaganahalli, H.L. Wells, J.K. Dhanota and E. Liang, *The discovery of Bombali virus adds*
561 *further support for bats as hosts of ebolaviruses*. Nature microbiology, 2018. **3**(10): p.
562 1084-1089.
- 563 6. Amman, B.R., S.A. Carroll, Z.D. Reed, T.K. Sealy, S. Balinandi, R. Swanepoel, A. Kemp,
564 B.R. Erickson, J.A. Comer and S. Campbell, *Seasonal pulses of Marburg virus circulation*
565 *in juvenile Rousettus aegyptiacus bats coincide with periods of increased risk of human*
566 *infection*. 2012.
- 567 7. Swanepoel, R., S.B. Smit, P.E. Rollin, P. Formenty, P.A. Leman, A. Kemp, F.J. Burt, A.A.
568 Grobbelaar, J. Croft and D.G. Bausch, *Studies of reservoir hosts for Marburg virus*.
569 Emerging infectious diseases, 2007. **13**(12): p. 1847.
- 570 8. Towner, J.S., X. Pourrut, C.G. Albariño, C.N. Nkogue, B.H. Bird, G. Grard, T.G. Ksiazek,
571 J.-P. Gonzalez, S.T. Nichol and E.M. Leroy, *Marburg virus infection detected in a common*
572 *African bat*. PloS one, 2007. **2**(8): p. e764.
- 573 9. Team, W.E.R., *Ebola virus disease in West Africa—the first 9 months of the epidemic and*
574 *forward projections*. New England Journal of Medicine, 2014. **371**(16): p. 1481-1495.
- 575 10. Hoenen, T., A. Groseth and H. Feldmann, *Therapeutic strategies to target the Ebola virus*
576 *life cycle*. Nature Reviews Microbiology, 2019. **17**(10): p. 593-606.
- 577 11. Lee, J.E. and E.O. Saphire, *Ebolavirus glycoprotein structure and mechanism of entry*.
578 Future virology, 2009. **4**(6): p. 621-635.

- 579 12. Mehedi, M., D. Falzarano, J. Seebach, X. Hu, M.S. Carpenter, H.-J. Schnittler and H.
580 Feldmann, *A new Ebola virus nonstructural glycoprotein expressed through RNA editing*.
581 Journal of virology, 2011. **85**(11): p. 5406-5414.
- 582 13. Volchkova, V.A., H. Feldmann, H.-D. Klenk and V.E. Volchkov, *The nonstructural small*
583 *glycoprotein sGP of Ebola virus is secreted as an antiparallel-orientated homodimer*.
584 Virology, 1998. **250**(2): p. 408-414.
- 585 14. Aleksandrowicz, P., A. Marzi, N. Biedenkopf, N. Beimforde, S. Becker, T. Hoenen, H.
586 Feldmann and H.-J. Schnittler, *Ebola virus enters host cells by macropinocytosis and*
587 *clathrin-mediated endocytosis*. The Journal of infectious diseases, 2011. **204**(suppl_3): p.
588 S957-S967.
- 589 15. Mulherkar, N., M. Raaben, J.C. de la Torre, S.P. Whelan and K. Chandran, *The Ebola*
590 *virus glycoprotein mediates entry via a non-classical dynamin-dependent macropinocytic*
591 *pathway*. Virology, 2011. **419**(2): p. 72-83.
- 592 16. Nanbo, A., M. Imai, S. Watanabe, T. Noda, K. Takahashi, G. Neumann, P. Halfmann and
593 Y. Kawaoka, *Ebolavirus is internalized into host cells via macropinocytosis in a viral*
594 *glycoprotein-dependent manner*. PLoS pathogens, 2010. **6**(9): p. e1001121.
- 595 17. Saeed, M.F., A.A. Kolokoltsov, T. Albrecht and R.A. Davey, *Cellular entry of ebola virus*
596 *involves uptake by a macropinocytosis-like mechanism and subsequent trafficking*
597 *through early and late endosomes*. PLoS pathogens, 2010. **6**(9): p. e1001110.
- 598 18. Gong, X., H. Qian, X. Zhou, J. Wu, T. Wan, P. Cao, W. Huang, X. Zhao, X. Wang and P.
599 Wang, *Structural insights into the Niemann-Pick C1 (NPC1)-mediated cholesterol transfer*
600 *and Ebola infection*. Cell, 2016. **165**(6): p. 1467-1478.
- 601 19. Miller, E.H., G. Obernosterer, M. Raaben, A.S. Herbert, M.S. Deffieu, A. Krishnan, E.
602 Ndungo, R.G. Sandesara, J.E. Carette and A.I. Kuehne, *Ebola virus entry requires the*
603 *host-programmed recognition of an intracellular receptor*. The EMBO journal, 2012. **31**(8):
604 p. 1947-1960.
- 605 20. Wang, H., Y. Shi, J. Song, J. Qi, G. Lu, J. Yan and G.F. Gao, *Ebola viral glycoprotein*
606 *bound to its endosomal receptor Niemann-Pick C1*. Cell, 2016. **164**(1-2): p. 258-268.
- 607 21. Saphire, E.O., S.L. Schendel, B.M. Gunn, J.C. Milligan and G. Alter, *Antibody-mediated*
608 *protection against Ebola virus*. Nature immunology, 2018. **19**(11): p. 1169-1178.
- 609 22. Volchkov, V.E., H. Feldmann, V.A. Volchkova and H.-D. Klenk, *Processing of the Ebola*
610 *virus glycoprotein by the proprotein convertase furin*. Proceedings of the National
611 Academy of Sciences, 1998. **95**(10): p. 5762-5767.

- 612 23. Sanchez, A., Z.-Y. Yang, L. Xu, G.J. Nabel, T. Crews and C.J. Peters, *Biochemical*
613 *analysis of the secreted and virion glycoproteins of Ebola virus*. Journal of virology, 1998.
614 **72**(8): p. 6442-6447.
- 615 24. Ito, H., S. Watanabe, A. Sanchez, M.A. Whitt and Y. Kawaoka, *Mutational analysis of the*
616 *putative fusion domain of Ebola virus glycoprotein*. Journal of virology, 1999. **73**(10): p.
617 8907-8912.
- 618 25. Malashkevich, V.N., B.J. Schneider, M.L. McNally, M.A. Milhollen, J.X. Pang and P.S. Kim,
619 *Core structure of the envelope glycoprotein GP2 from Ebola virus at 1.9-Å resolution*.
620 Proceedings of the National Academy of Sciences, 1999. **96**(6): p. 2662-2667.
- 621 26. Weissenhorn, W., A. Carfi, K.-H. Lee, J.J. Skehel and D.C. Wiley, *Crystal structure of the*
622 *Ebola virus membrane fusion subunit, GP2, from the envelope glycoprotein ectodomain*.
623 Molecular cell, 1998. **2**(5): p. 605-616.
- 624 27. Alvarez, C.P., F. Lasala, J. Carrillo, O. Muñiz, A.L. Corbí and R. Delgado, *C-type lectins*
625 *DC-SIGN and L-SIGN mediate cellular entry by Ebola virus in cis and in trans*. Journal of
626 virology, 2002. **76**(13): p. 6841-6844.
- 627 28. Lin, G., G. Simmons, S. Pöhlmann, F. Baribaud, H. Ni, G.J. Leslie, B.S. Haggarty, P.
628 Bates, D. Weissman and J.A. Hoxie, *Differential N-linked glycosylation of human*
629 *immunodeficiency virus and Ebola virus envelope glycoproteins modulates interactions*
630 *with DC-SIGN and DC-SIGNR*. Journal of virology, 2003. **77**(2): p. 1337-1346.
- 631 29. Simmons, G., J.D. Reeves, C.C. Grogan, L.H. Vandenberghe, F. Baribaud, J.C. Whitbeck,
632 E. Burke, M.J. Buchmeier, E.J. Soilleux and J.L. Riley, *DC-SIGN and DC-SIGNR bind*
633 *ebola glycoproteins and enhance infection of macrophages and endothelial cells*. Virology,
634 2003. **305**(1): p. 115-123.
- 635 30. Collar, A.L., E.C. Clarke, E. Anaya, D. Merrill, S. Yarborough, S.M. Anthony, J.H. Kuhn,
636 C. Merle, M. Theisen and S.B. Bradfute, *Comparison of N-and O-linked glycosylation*
637 *patterns of ebolavirus glycoproteins*. Virology, 2017. **502**: p. 39-47.
- 638 31. Dowling, W., E. Thompson, C. Badger, J.L. Mellquist, A.R. Garrison, J.M. Smith, J.
639 Paragas, R.J. Hogan and C. Schmaljohn, *Influences of glycosylation on antigenicity,*
640 *immunogenicity, and protective efficacy of ebola virus GP DNA vaccines*. Journal of
641 virology, 2007. **81**(4): p. 1821-1837.
- 642 32. Iraqi, M., A. Edri, Y. Greenspan, K. Kundu, P. Bolel, A. Cahana, A. Ottolenghi, R. Gazit,
643 L. Lobel and A. Braiman, *N-Glycans mediate the ebola virus-GP1 shielding of ligands to*
644 *immune receptors and immune evasion*. Frontiers in cellular and infection microbiology,
645 2020. **10**: p. 48.

- 646 33. Jeffers, S.A., D.A. Sanders and A. Sanchez, *Covalent modifications of the Ebola virus*
647 *glycoprotein*. Journal of virology, 2002. **76**(24): p. 12463-12472.
- 648 34. Lennemann, N.J., B.A. Rhein, E. Ndungo, K. Chandran, X. Qiu and W. Maury,
649 *Comprehensive functional analysis of N-linked glycans on Ebola virus GP1*. MBio, 2014.
650 **5**(1): p. e00862-13.
- 651 35. Ritchie, G., D.J. Harvey, U. Stroehrer, F. Feldmann, H. Feldmann, V. Wahl-Jensen, L.
652 Royle, R.A. Dwek and P.M. Rudd, *Identification of N-glycans from Ebola virus*
653 *glycoproteins by matrix-assisted laser desorption/ionisation time-of-flight and negative ion*
654 *electrospray tandem mass spectrometry*. Rapid Communications in Mass Spectrometry:
655 An International Journal Devoted to the Rapid Dissemination of Up-to-the-Minute
656 Research in Mass Spectrometry, 2010. **24**(5): p. 571-585.
- 657 36. Wang, B., Y. Wang, D.A. Frabutt, X. Zhang, X. Yao, D. Hu, Z. Zhang, C. Liu, S. Zheng
658 and S.-H. Xiang, *Mechanistic understanding of N-glycosylation in Ebola virus glycoprotein*
659 *maturation and function*. Journal of Biological Chemistry, 2017. **292**(14): p. 5860-5870.
- 660 37. Lennemann, N.J., M. Walkner, A.R. Berkebile, N. Patel and W. Maury, *The role of*
661 *conserved N-linked glycans on Ebola virus glycoprotein 2*. The Journal of infectious
662 diseases, 2015. **212**(suppl_2): p. S204-S209.
- 663 38. Furmanek, A. and J. Hofsteenge, *Protein C-mannosylation: facts and questions*. Acta
664 biochimica polonica, 2000. **47**(3): p. 781-789.
- 665 39. Julenius, K., *NetCGlyc 1.0: prediction of mammalian C-mannosylation sites*. Glycobiology,
666 2007. **17**(8): p. 868-876.
- 667 40. Falzarano, D., O. Krokhin, G. Van Domselaar, K. Wolf, J. Seebach, H.-J. Schnittler and
668 H. Feldmann, *Ebola sGP—the first viral glycoprotein shown to be C-mannosylated*.
669 Virology, 2007. **368**(1): p. 83-90.
- 670 41. Plewe, M.B., N.V. Sokolova, V.R. Gantla, E.R. Brown, S. Naik, A. Fetsko, D.D. Lorimer,
671 D.M. Dranow, H. Smutney and J. Bullen, *Discovery of Adamantane Carboxamides as*
672 *Ebola Virus Cell Entry and Glycoprotein Inhibitors*. ACS medicinal chemistry letters, 2020.
673 **11**(6): p. 1160-1167.
- 674 42. Ren, J., Y. Zhao, E.E. Fry and D.I. Stuart, *Target identification and mode of action of four*
675 *chemically divergent drugs against ebolavirus infection*. Journal of medicinal chemistry,
676 2018. **61**(3): p. 724-733.
- 677 43. Shaikh, F., Y. Zhao, L. Alvarez, M. Iliopoulou, C. Lohans, C.J. Schofield, S. Padilla-Parra,
678 S.W. Siu, E.E. Fry and J. Ren, *Structure-based in silico screening identifies a potent*

- 679 *ebolavirus inhibitor from a traditional Chinese medicine library*. Journal of medicinal
680 chemistry, 2019. **62**(6): p. 2928-2937.
- 681 44. Zhao, Y., J. Ren, E.E. Fry, J. Xiao, A.R. Townsend and D.I. Stuart, *Structures of Ebola*
682 *virus glycoprotein complexes with tricyclic antidepressant and antipsychotic drugs*.
683 Journal of medicinal chemistry, 2018. **61**(11): p. 4938-4945.
- 684 45. Zhao, Y., J. Ren, K. Harlos, D.M. Jones, A. Zeltina, T.A. Bowden, S. Padilla-Parra, E.E.
685 Fry and D.I. Stuart, *Toremifene interacts with and destabilizes the Ebola virus*
686 *glycoprotein*. Nature, 2016. **535**(7610): p. 169-172.
- 687 46. Cohen-Dvashi, H., M. Zehner, S. Ehrhardt, M. Katz, N. Elad, F. Klein and R. Diskin,
688 *Structural basis for a convergent immune response against Ebola Virus*. Cell host &
689 microbe, 2020. **27**(3): p. 418-427. e4.
- 690 47. Dias, J.M., A.I. Kuehne, D.M. Abelson, S. Bale, A.C. Wong, P. Halfmann, M.A.
691 Muhammad, M.L. Fusco, S.E. Zak and E. Kang, *A shared structural solution for*
692 *neutralizing ebolaviruses*. Nature structural & molecular biology, 2011. **18**(12): p. 1424-
693 1427.
- 694 48. Ehrhardt, S.A., M. Zehner, V. Krähling, H. Cohen-Dvashi, C. Kreer, N. Elad, H. Gruell,
695 M.S. Ercanoglu, P. Schommers and L. Gieselmann, *Polyclonal and convergent antibody*
696 *response to Ebola virus vaccine rVSV-ZEBOV*. Nature medicine, 2019. **25**(10): p. 1589-
697 1600.
- 698 49. King, L.B., B.R. West, C.L. Moyer, P. Gilchuk, A. Flyak, P.A. Ilinykh, R. Bombardi, S. Hui,
699 K. Huang and A. Bukreyev, *Cross-reactive neutralizing human survivor monoclonal*
700 *antibody BDBV223 targets the ebolavirus stalk*. Nature communications, 2019. **10**(1): p.
701 1-8.
- 702 50. Milligan, J.C., C.W. Davis, P.A. Ilinykh, K. Huang, P. Halfmann, R.W. Cross, V. Borisevich,
703 K.N. Agans, J.B. Geisbert and C. Chennareddy, *Asymmetric and Non-Stoichiometric*
704 *Recognition Results in Broad Protection Against Ebolaviruses by a Two-Antibody Cocktail*.
- 705 51. Misasi, J., M.S. Gilman, M. Kanekiyo, M. Gui, A. Cagigi, S. Mulangu, D. Corti, J.E.
706 Ledgerwood, A. Lanzavecchia and J. Cunningham, *Structural and molecular basis for*
707 *Ebola virus neutralization by protective human antibodies*. Science, 2016. **351**(6279): p.
708 1343-1346.
- 709 52. Murin, C.D., P. Gilchuk, P.A. Ilinykh, K. Huang, N. Kuzmina, X. Shen, J.F. Bruhn, A.L.
710 Bryan, E. Davidson and B.J. Doranz, *Convergence of a common solution for broad*
711 *ebolavirus neutralization by glycan cap-directed human antibodies*. Cell reports, 2021.
712 **35**(2): p. 108984.

- 713 53. Olal, D., A. Kuehne, S. Bale, P. Halfmann, T. Hashiguchi, M.L. Fusco, J.E. Lee, L.B. King,
714 Y. Kawaoka and J.M. Dye, *Structure of an Ebola virus-protective antibody in complex with*
715 *its mucin-domain linear epitope*. Journal of Virology, 2011.
- 716 54. Pallesen, J., C.D. Murin, N. De Val, C.A. Cottrell, K.M. Hastie, H.L. Turner, M.L. Fusco,
717 A.I. Flyak, L. Zeitlin and J.E. Crowe, *Structures of Ebola virus GP and sGP in complex*
718 *with therapeutic antibodies*. Nature microbiology, 2016. **1**(9): p. 1-9.
- 719 55. West, B.R., C.L. Moyer, L.B. King, M.L. Fusco, J.C. Milligan, S. Hui and E.O. Saphire,
720 *Structural basis of pan-ebolavirus neutralization by a human antibody against a*
721 *conserved, yet cryptic epitope*. MBio, 2018. **9**(5): p. e01674-18.
- 722 56. Wilson, J.A., M. Hevey, R. Bakken, S. Guest, M. Bray, A.L. Schmaljohn and M.K. Hart,
723 *Epitopes involved in antibody-mediated protection from Ebola virus*. Science, 2000.
724 **287**(5458): p. 1664-1666.
- 725 57. Pascal, K.E., D. Dudgeon, J.C. Trefry, M. Anantpadma, Y. Sakurai, C.D. Murin, H.L.
726 Turner, J. Fairhurst, M. Torres and A. Rafique, *Development of clinical-stage human*
727 *monoclonal antibodies that treat advanced Ebola virus disease in nonhuman primates*.
728 The Journal of infectious diseases, 2018. **218**(suppl_5): p. S612-S626.
- 729 58. Bagdonaite, I. and H.H. Wandall, *Global aspects of viral glycosylation*. Glycobiology, 2018.
730 **28**(7): p. 443-467.
- 731 59. Hargett, A.A. and M.B. Renfrow, *Glycosylation of viral surface proteins probed by mass*
732 *spectrometry*. Current opinion in virology, 2019. **36**: p. 56-66.
- 733 60. Snijder, J., M.S. Ortego, C. Weidle, A.B. Stuart, M.D. Gray, M.J. McElrath, M. Pancera, D.
734 Veessler and A.T. McGuire, *An antibody targeting the fusion machinery neutralizes dual-*
735 *tropic infection and defines a site of vulnerability on Epstein-Barr virus*. Immunity, 2018.
736 **48**(4): p. 799-811. e9.
- 737 61. Walls, A.C., X. Xiong, Y.-J. Park, M.A. Tortorici, J. Snijder, J. Quispe, E. Cameroni, R.
738 Gopal, M. Dai and A. Lanzavecchia, *Unexpected receptor functional mimicry elucidates*
739 *activation of coronavirus fusion*. Cell, 2019. **176**(5): p. 1026-1039. e15.
- 740 62. Watanabe, Y., J.D. Allen, D. Wrapp, J.S. McLellan and M. Crispin, *Site-specific glycan*
741 *analysis of the SARS-CoV-2 spike*. Science, 2020. **369**(6501): p. 330-333.
- 742 63. Wörner, T.P., T.M. Shamorkina, J. Snijder and A.J. Heck, *Mass spectrometry-Based*
743 *structural virology*. Analytical Chemistry, 2020. **93**(1): p. 620-640.
- 744 64. Yao, H., Y. Song, Y. Chen, N. Wu, J. Xu, C. Sun, J. Zhang, T. Weng, Z. Zhang and Z. Wu,
745 *Molecular architecture of the SARS-CoV-2 virus*. Cell, 2020. **183**(3): p. 730-738. e13.

- 746 65. Behrens, A.-J. and M. Crispin, *Structural principles controlling HIV envelope glycosylation*.
747 Current opinion in structural biology, 2017. **44**: p. 125-133.
- 748 66. Go, E.P., A. Herschhorn, C. Gu, L. Castillo-Menendez, S. Zhang, Y. Mao, H. Chen, H.
749 Ding, J.K. Wakefield and D. Hua, *Comparative analysis of the glycosylation profiles of*
750 *membrane-anchored HIV-1 envelope glycoprotein trimers and soluble gp140*. Journal of
751 virology, 2015. **89**(16): p. 8245-8257.
- 752 67. Go, E.P., G. Hewawasam, H.-X. Liao, H. Chen, L.-H. Ping, J.A. Anderson, D.C. Hua, B.F.
753 Haynes and H. Desaire, *Characterization of glycosylation profiles of HIV-1*
754 *transmitted/founder envelopes by mass spectrometry*. Journal of virology, 2011. **85**(16):
755 p. 8270-8284.
- 756 68. Raska, M., K. Takahashi, L. Czernekova, K. Zachova, S. Hall, Z. Moldoveanu, M.C. Elliott,
757 L. Wilson, R. Brown and D. Jancova, *Glycosylation patterns of HIV-1 gp120 depend on*
758 *the type of expressing cells and affect antibody recognition*. Journal of Biological
759 Chemistry, 2010. **285**(27): p. 20860-20869.
- 760 69. Bagdonaite, I., R. Nordén, H.J. Joshi, S.L. King, S.Y. Vakhrushev, S. Olofsson and H.H.
761 Wandall, *Global mapping of O-glycosylation of varicella zoster virus, human*
762 *cytomegalovirus, and Epstein-Barr virus*. Journal of Biological Chemistry, 2016. **291**(23):
763 p. 12014-12028.
- 764 70. Bräutigam, J., A.J. Scheidig and W. Egge-Jacobsen, *Mass spectrometric analysis of*
765 *hepatitis C viral envelope protein E2 reveals extended microheterogeneity of mucin-type*
766 *O-linked glycosylation*. Glycobiology, 2013. **23**(4): p. 453-474.
- 767 71. Brun, J., S.a. Vasiljevic, B. Gangadharan, M. Hensen, A. V. Chandran, M.L. Hill, J.
768 Kiappes, R.A. Dwek, D.S. Alonzi and W.B. Struwe, *Assessing Antigen Structural Integrity*
769 *through Glycosylation Analysis of the SARS-CoV-2 Viral Spike*. ACS central science,
770 2021. **7**(4): p. 586-593.
- 771 72. Watanabe, Y., T.A. Bowden, I.A. Wilson and M. Crispin, *Exploitation of glycosylation in*
772 *enveloped virus pathobiology*. Biochimica et Biophysica Acta (BBA)-General Subjects,
773 2019. **1863**(10): p. 1480-1497.
- 774 73. Larkin, M.A., G. Blackshields, N.P. Brown, R. Chenna, P.A. McGettigan, H. McWilliam, F.
775 Valentin, I.M. Wallace, A. Wilm and R. Lopez, *Clustal W and Clustal X version 2.0*.
776 bioinformatics, 2007. **23**(21): p. 2947-2948.
- 777 74. Waterhouse, A., M. Bertoni, S. Bienert, G. Studer, G. Tauriello, R. Gumienny, F.T. Heer,
778 T.A.P. de Beer, C. Rempfer and L. Bordoli, *SWISS-MODEL: homology modelling of*
779 *protein structures and complexes*. Nucleic acids research, 2018. **46**(W1): p. W296-W303.

- 780 75. Goddard, T.D., C.C. Huang, E.C. Meng, E.F. Pettersen, G.S. Couch, J.H. Morris and T.E.
781 Ferrin, *UCSF ChimeraX: Meeting modern challenges in visualization and analysis*. Protein
782 Science, 2018. **27**(1): p. 14-25.
- 783 76. Emsley, P. and K. Cowtan, *Coot: model-building tools for molecular graphics*. Acta
784 crystallographica section D: biological crystallography, 2004. **60**(12): p. 2126-2132.
- 785 77. de Beer, T., J.F. Vliegthart, A. Loeffler and J. Hofsteenge, *The Hexopyranosyl Residue*
786 *That Is C-Glycosidically Linked to the Side Chain of Tryptophan-7 in Human RNase Us Is.*
787 *alpha.-Mannopyranose*. Biochemistry, 1995. **34**(37): p. 11785-11789.
- 788 78. Frank, M., D. Beccati, B.R. Leeftang and J.F. Vliegthart, *C-Mannosylation Enhances*
789 *the Structural Stability of Human RNase 2*. Iscience, 2020. **23**(8): p. 101371.
- 790 79. Jonker, H.R., K. Saxena, A. Shcherbakova, B. Tiemann, H. Bakker and H. Schwalbe,
791 *NMR Spectroscopic Characterization of the C-Mannose Conformation in a*
792 *Thrombospondin Repeat Using a Selective Labeling Approach*. Angewandte Chemie,
793 2020. **132**(46): p. 20840-20846.
- 794 80. Bern, M., Y.J. Kil and C. Becker, *Byonic: advanced peptide and protein identification*
795 *software*. Current protocols in bioinformatics, 2012. **40**(1): p. 13.20. 1-13.20. 14.
- 796 81. Pino, L.K., B.C. Searle, J.G. Bollinger, B. Nunn, B. MacLean and M.J. MacCoss, *The*
797 *Skyline ecosystem: Informatics for quantitative mass spectrometry proteomics*. Mass
798 spectrometry reviews, 2020. **39**(3): p. 229-244.

799