Review

Hidden Relationships between \(\text{N}\)-Glycosylation and Disulfide Bonds in Individual Proteins

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Abstract: \(\text{N}\)-Glycosylation (NG) and disulfide bonds (DBs) are two prevalent co/post-translational modifications (PTMs) that are often conserved and coexist in membrane and secreted proteins involved in a large number of diseases. Both in the past and in recent times, the enzymes and chaperones regulating these PTMs have been constantly discovered to directly interact with each other or colocalize in the ER. However, beyond a few model proteins, how such cooperation affects \(\text{N}\)-glycan modification and disulfide bonding at selective sites in individual proteins is largely unknown. Here, we reviewed the literature to discover the current status in understanding the relationships between NG and DBs in individual proteins. Our results showed that more than 2700 human proteins carry both PTMs, and fewer than 2\% of them have been investigated in the associations between NG and DBs. We summarized both these proteins with the reported relationships in the two PTMs and the tools used to discover the relationships. We hope that, by exposing this largely understudied field, more investigations can be encouraged to unveil the hidden relationships of NG and DBs in the majority of membranes and secreted proteins for pathophysiological understanding and biotherapeutic development.

Keywords: posttranslational modifications; \(\text{N}\)-glycosylation; disulfide bonds; membrane and secreted proteins; endoplasmic reticulum quality control; protein structure and function

1. Introduction

Both \(\text{N}\)-glycosylation (NG) and disulfide bonds (DBs) can form co- and post-translational modifications (PTMs) [1] on proteins in the endoplasmic reticulum (ER) while they pass through the secretory pathway. These two modifications are not only common but often evolutionarily conserved in membrane and secreted proteins from prokaryotes to eukaryotes. As two critical modifications, they facilitate protein folding and regulate protein structure, function, stability, and cellular localization. Defects in either one can fail protein ER quality control, trigger unfolded protein response, and cause pathological conditions ranging from heritable congenital disorders of glycosylation as an example to acquired disorders such as cancers, dementia, diabetes, autoimmune, infectious, and cardiovascular diseases [2,3].

Owing to their importance, both NG and DBs have been frequently investigated in numerous proteins. In UniProt [4], over 2700 human proteins have both PTMs annotated. However, the associations between NG and DBs have only been well examined in a few model proteins, such as influenza hemagglutinin (HA) [5–7]. Both past and recent discoveries have shown that the enzymes responsible for adding, processing, and degrading \(\text{N}\)-glycans are closely related to proteins with the abilities of adding, isomerizing, and breaking disulfide bonds. Beyond some well-studied model proteins, how NG and DB
formation affect each other in the remaining individual proteins is the focus of this review. By reviewing research articles, we generally cataloged the observed relationships from the host proteins into three kinds, including inhibition, promotion, and no relationship. More importantly, we noticed that the majority of the studies did not investigate the interactions between the two PTMs but studied them separately. As a result, some of the observed structural and functional changes in proteins can be a synergistic effect of both instead of one PTM. The reasons are likely that the tools required for these studies were not widely recognized and/or the importance of the associations between them were not commonly appreciated. To raise awareness, this review not only summarizes molecular foundations to support a potentially complex relationship and the current known relationships between NG and DBs in individual proteins but also enlists the relevant tools that can be used for addressing these relationships and provides our recommendations for future high-throughput analysis.

2. Molecular Foundations

When nascent proteins are translocated into the ER, NG and DB formation are two closely associated events. NG is a complex process that refers to a dolichol pyrophosphate donor transferring a tetradecasaccharide precursor en bloc to the nitrogen atom of the asparagine side chain in a consensus sequence (namely, sequon), i.e., N–X–S/T, in which X can be any amino acid but not proline [8]. Newly added N-glycans are further processed while glycosylated proteins fold and mature in the ER. Glycoproteins that fail the quality control in the ER are subjected to the ER-associated degradation process (ERAD) [9]. N-Glycans are bulky. The addition of an N-glycan can shape the local conformation of a polypeptide chain through the first GlcNAc residue, the triose core, and the outer sugar moieties [10–12]. Either intra- or extramolecular N-glycans can stabilize a loose polypeptide conformation via both hydrophilic and hydrophobic interactions with the peptide backbone [13]. The stabilized conformation can bring two free cysteine side-chains in close proximity to promote an otherwise slow or energetically costly DB formation, as experimentally proven in model peptides and whole proteins [12–15].

The covalent disulfide bond is formed by oxidizing two free thiol groups in two cysteine residues in the vicinity. In the ER, DBs can be formed, isomerized, or broken by a number of oxidoreductases. Over 20 ER resident oxidoreductases, also called protein disulfide isomerase (PDI) family enzymes, have been discovered [16]. The characteristic feature of PDI family proteins is the thioredoxin fold, C–X–X–C. This motif is usually the catalytic domain of oxidoreductases. These enzymes can use the cysteines in the motif to exchange electrons in the thiols of the substrate to either form or break DBs [17]. In addition, the mixed DBs formed between the oxidoreductase and the substrate through the thioredoxin fold can often stabilize the substrate and allow further modifications, such as N-glycosylation, by other cooperating enzymes in the same complex or nearby [18]. Overall, PDIs can carry out a number of functions, including oxidoreduction, isomerization, and chaperone activity [19].

Due to the spatial requirement of DBs, the protein structure can be largely perturbed or stabilized after the formation of novel DBs or the reshuffling of existing DBs. For glycoproteins carrying bulky N-glycans to fold and mature in the ER, the formation of DBs can block or restrain the access of glycoenzymes or chaperones to N-glycans. Interestingly, glycoenzymes and glycochaperones are often accompanied by oxidoreductases in the ER, or the glycoenzyme itself has a thioredoxin fold that can interact with thiols in substrates, as recently reviewed by Patel et al. [9]. For example, the oxidoreductases of MAGT1 or TUSC3 are part of STT3B of OST, the enzyme responsive to adding N-glycans to proteins; the oxidoreductases of ERP57 and ERP72 form complexes with CNX/CRT, the ER chaperones of N-glycoproteins; in the ER N-glycosylation quality control system, UGGT1 itself has thioredoxin-like motifs and interacts with SELENOF (Sep15), a selenocysteine-containing oxidoreductase [20]; further along the thread in the ERAD, BIP binds PDI, P5, and ERdj5,
whereas EDEMs bind TXNDC11, P4HB, ERp46, etc. The intriguing roles of NG and DBs are just beginning to be realized.

Beyond the ER, both NG and DBs can be further edited. For example, after exiting the ER, N-glycans can be processed in the Golgi [21]. For DBs, a number of enzymes both inside and outside of cells can break and reform these covalent bonds, which often function as switches to activate or suppress host protein functions [22]. How NG and DBs relate to each other beyond the ER remains a question [9]. A complete elucidation of these interactions requires kinetic information of the modification in the time domain, as well as spatial information on their cellular localization. As introduced later, kinetic measurements of PTMs can be performed through traditional pulse-chase or mass spectrometry (MS). The former has been well developed, while the latter is emerging [23]. Spatial information can be delineated through inhibitors specific to an enzymatic step or stressors to a particular organelle.

Nevertheless, not all DBs are linked to every NG in a protein. The functions of both NG and DBs are pleiotropic. The Weerapana group has cataloged the functions of reactive cysteines into five types, including structural, regulatory, redox catalytic, catalytic nucleophile, and metal-binding cysteines [24]. Similarly, the functions of NG range from facilitating local folding to promoting secondary structures, increasing protein stability, protecting hydrophobic surfaces, reducing aggregation, guiding trafficking, and functioning as epitopes for molecular recognition [25].

Given the close molecular associations between enzymes functioning in the formation and processing of the two PTMs in the ER, as well as the broad availability of both PTMs in membrane and secreted proteins, we asked what the current observed relationships of these two PTMs are in individual proteins. Given that one-third of the mammalian proteome is produced through the secretory pathway [26], most proteins have annotated DBs and NG, as shown in UniProt. We anticipated that rich information can be obtained from such exercises. Below, we summarize our findings on both the relationships in individual proteins and the tools used to study them.

3. Overview of the Studies

To obtain the relevant articles, we searched PubMed using various keywords, including “N-glycosylation and disulfide bonds”, “N-linked glycosylation and disulfide bonds”, “N-linked glycans and disulfide bonds”, “carbohydrate and disulfide bond”, or “glycans and disulfide”. Furthermore, we included the associated studies cited in the selected papers. The combined list had nearly 600 articles.

After establishing the list, we first examined the types of proteins that were studied. Most studied proteins were membrane and secreted proteins of humans, model organisms, and viruses, as shown in Figure 1. Others examined proteins from venomous species such as snakes and wasps, parasites such as hookworms, and agricultural species such as vegetables, fishes, and silkworms. Most of these proteins were surface receptors, adhesion molecules, secreted or membrane-tethered enzymes, cytokines, hormones, signaling molecules, and extracellular matrix proteins. For viruses, the most studied proteins were surface glycoproteins involved in recognition and fusion with host cells. All these results suggested the importance of these two PTMs in diseases originating from cell–protein interactions and cell–cell communications.

Second, we broadly classified all the reports into four categories. For most studies that included both PTMs but ignored their relationships, we cataloged them as “no studies”. We further divided the investigated relationships into three categories: promotion, inhibition, and independent relationships, as summarized in Figure 2 and detailed in Table 1. In Figure 2, the promoting relationship dominated current studies. This relationship, however, was often observed passively by the diminished or inhibited formation of one type of PTM when the other type of PTM was experimentally removed. To a much lesser extent, an inhibitory relationship was observed, in which the formation of one PTM blocked the development of the other. Only in very limited reports did proteins show independent
modifications of these two PTMs, in which the removal of one PTM did not affect the formation or the processing of the other. In the four sections below, we use detailed examples to illustrate these observed relationships.

Figure 1. Distribution of species hosting proteins studied for N-glycosylation and disulfide bonds.

Figure 2. Distribution of different relationships between the N-glycosylation and disulfide bonds in the literature examined.
Table 1. Summary of the proteins showing interactions between N-glycosylation and disulfide bonds.

| UniProt Accession | Protein Name                          | Position of DBs                          | Position of N-Glycans | Class      | Reference |
|-------------------|---------------------------------------|------------------------------------------|-----------------------|------------|-----------|
| A0A346IHA8        | CD4-binding region of the envelope glycoproteins | 50–70 a, 219–248 a, 229–240 a, 369–402 a, 362–429, 597–603 a | 390, 447             | No relation | [27]      |
| Q9Y6R1            | SLC4 Na⁺-coupled transporter (NBCe1-A) | 583–585, 630–642 c                         | 597, 617              | Inhibition  | [28]      |
| Q99062            | Granulocyte colony-stimulating factor receptor | 107–118, 153–162, 224–271, 364–371      | 27, 69, 104, 110, 365, 450, 547, 555, 586 | Promotion  | [29]      |
| Q62635            | Gastric mucin (rat)                    | 160 a, 420 a, 667 a, 767 a, 837 a, 892 a, 1136 a, 1151 a, 1212 a, 1227 a, 1243 a, 1350 a | Promotion  | [30]      |
| P49018            | GPI-anchor transamidase (GPI8) yeast   | 85 (interchain with C-194 in GPI16) b      | 256 a, 346 a          | Promotion  | [31]      |
| P00750            | Tissue-type plasminogen activator (t-PA) | 41–71, 69–78, 86–97, 91–108, 110–119, 127–208 b, 148–190 b, 179–203 b, 215–296, 236–278, 267–291, 299–430, 342–358 b, 350–419 b, 444–519 b, 476–492 b, 509–537 b | 117, 184, 448 | Inhibition | [32,33]  |
| P04275            | Von Willebrand factor (VWF)            | 35–162 d, 57–200 d, 388–524 d, 410–559 d, 432–440 d, 509–695, 767–808, 776–804, 810–821, 867–996 d, 889–1031 d, 898–993 d, 914–921 d, 1060–1084, 1071–1111, 1089–1091, 1126–1130, 1149–1169, 1153–1165, 1196–1199, 1234–1237, 1272–1458, 1669–1670, 1686–1872, 1879–1904, 1899–1940 d, 1927–2088, 1950–2085 d, 1972–2123 d, 1993–2001 d, 2724–2774 b, 2739–2788 b, 2750–2804 b, 2754–2806 b, | 99 a, 156 a, 211 a, 666 a, 857 e, 1147 (atypical) e, 1231 e, 1515, 1574 e, 2223 e, 2290 e, 2357 e, 2400 e, 2546, 2585 e, 2790 e | Promotion | [34–36]  |
| P0DN86            | Human chorionic gonadotropin beta-subunit | 9–57, 23–72, 26–110, 34–88, 38–90, 93–100 | 13, 30              | Promotion  | [37–39]  |
| P32906            | Yeast-alpha1,2 mannosidase             | 340–385, 468–471                         | 96, 155, 224          | Promotion  | [41]      |
| O60896            | Receptor activity-modifying protein 3 (RAMP3) | 40–72 b, 57–104 b                         | 28, 57, 70, 102       | Inhibition  | [42]      |
| UniProt Accession | Protein Name                          | Position of DBs                      | Position of N-Glycans | Class      | Reference |
|-------------------|--------------------------------------|--------------------------------------|-----------------------|------------|-----------|
| Q13087            | PDIA2                                | 18 (interchain)                      | 127, 284, 516         | Inhibition | [43]      |
| P04853            | Hemagglutinin-neuraminidase (Sendai virus) | 129 (interchain)                  | 77, 499, 511          | Promotion  | [21,44]  |
| Q91UL0            | Hemagglutinin-neuraminidase (NDV)    | 123 (interchain), 172–196, 186–247, 238–251, 344–461, 455–465, 531–542 | 119, 341, 433, 481    | Inhibition | [45,46]  |
|                   | Hemagglutinin (Influenza A virus)    | 14–466, 52–277, 64–76, 97–139, 281–305, 473–477 | 8, 22, 38, 81, 165, 285, 483 | Promotion  | [47,48]  |
| P01229            | Human lutropin subunit beta          | 29–77 b, 43–92 b, 46–130 b, 54–108 b, 58–110 b, 113–120 b | 30                    | Promotion  | [49]      |
| P35625            | Metalloproteinase inhibitor 3 (TIMP3) | 24–91, 26–118, 36–143, 145–192 d, 150–155 d, 163–184 d |                      | Inhibition  | [50]      |
| Q16820            | Meprin A                             | 103–255, 124–144, 265–427, 273 a (interchain), 305 (interchain), 492 (interchain) d, 608–619 d, 613–628 d, 630–643 d | 41, 152, 234, 270, 330, 426, 452, 546, 553 | Promotion  | [51,52]  |
| P01130            | Low-density lipoprotein (LDL) receptor | 27–39 e, 34–52 e, 46–63 e, 68–82 e, 75–95 e, 89–104 e, 109–121 b, 116–134 e, 128–143 e, 148–160 e, 155–173 e, 167–184 e, 197–209 e, 204–222 e, 216–231 e, 236–248 e, 243–261 e, 255–270 e, 276–289 e, 284–302 e, 296–313 e, 318–329 e, 325–338 e, 340–352 e, 358–368 e, 364–377 e, 379–392 e, 667–681 e, 677–696 e, 698–711 e | 97 a, 156, 272, 515 a, 657 | Inhibition  | [53–55]  |
| Q02817            | MUC2 mucin                           | 59–67 b, 37–169 d, 59–206 d, 391–528 d, 413–563 d, 435–443 d, 680–992 d, 882–1027 d, 891–989 d, 909–916 d, 4481–4622 d, 4503–4661 d, 4527–4535 d, 5075–5122 b, 5089–5136 b, 5098–5152 b, 5102–5154 b | 163 a, 423 a, 670 a, 770 a, 894 a, 1139 a, 1154 a, 1215 a, 1230 a, 1246 a, 1787 a, 1820 a, 4339 a, 4351 a, 4362 a, 4373 a, 4422 a, 4438 a, 4502 a, 4616 a, 4627 a, 4752 a, 4787 a, 4881 a, 4955 a, 4970 a, 5019 a, 5038 a, 5069 a | Promotion  | [56,57]  |
| P12476            | VP7                                  | 82–135, 165–249, 191–244, 196–207 | 69                    | Promotion  | [22,54]  |
| P04156            | Major prion protein                  | 179–214                             |                       | Promotion  | [58]      |
| P05026            | Sodium/potassium-transporting ATPase subunit beta-1 | 126–149, 159–175, 213–276 | 158, 193, 265 | Promotion  | [59,60]  |
| P05231            | Interleukin-6 (IL6)                   | 45–51, 74–84                         | 46                    | Inhibition  | [61–63]  |
### Table 1. Cont.

| UniProt Accession | Protein Name                  | Position of DBs       | Position of N-Glycans | Class       | Reference |
|-------------------|-------------------------------|-----------------------|-----------------------|-------------|-----------|
| H2AM12            | Glycoprotein Gc               | 523–550, 580–589, 591–598, 471–487 | 493, 686              | Promotion   | [64]      |
| P22146            | 1,3-beta-Glucanosyltransferase| 74–103, 216–348, 234–265, 370–421, 372–462, 379–445, 398–403 | 40 a, 57, 95 a, 149 a, 165 a, 253, 283 a, 321 a, 409 a, 495 a | Promotion   | [65,66]   |
| P32623            | Probable glycosidase CRH2     | 493, 686              |                       | Promotion   | [66]      |
| Q9UMF0            | Intercellular adhesion molecule-5 (ICAM5) | 55–99, 59–103, 142–198, 249–302 | 54,74,137,195,214,274,316,371,397,582, 636,645,762,793,794 | Promotion   | [67,68]   |
| O75829            | Chondromodulin-I              | 131–193 b, 282–286 b, 283–323 b, 293–317 b, 297–313 b | 243 a            | No relation | [69]      |
| P40225            | Thrombopoietin                | 7–151, 29–85          | 197, 206, 234, 255, 340 a, 348 a | No relation | [70,71]   |
| Q9UNQ0            | ABCG2 protein                 | 592–608, 603 (interchain) | 596                   | Promotion   | [8,72,73] |
| P56817            | beta-Site APP-cleaving enzyme (BACE) | 216–420, 278–443, 330–380 | 153 a, 172 a, 223 a, 354 a | Promotion   | [74]      |
| Q9H1U4            | Multiple epidermal growth factor-like domains protein 9 (MEGF9) | 204–217 a, 206–224 a, 226–235 a, 238–251 a, 254–266 a, 256–272 a, 274–283 a, 286–298 a, 301–310 a, 303–317 a, 320–329 a, 332–346 a, 349–360 a, 351–371 a, 374–383 a, 386–397 a, 400–415 a, 402–422 a, 425–434 a, 437–449 a | 40 a, 182 a, 205 a, 218 a, 245 a, 267 a, | Inhibition   | [75]      |
| P35634            | Cathepsin                     | 30–118, 54–136 c, 255–298 c, 291–331 c, 321–337 c | 29               | Inhibition   | [76,77]   |
| P08709            | Coagulation factor VII        | 348–367               | 360                   | Inhibition   | [76]      |
| Q07837            | rBAT                          | 242–273, 571–666, 673–685 | 214, 261, 332, 495, 513, 575 | Promotion   | [78]      |
| O75355            | Nucleoside triphosphate diphosphohydrolase 3 (NTPDase3) | 92–116, 261–308, 289–334, 347–353, 399–422 | 81 a, 149 a, 238 a, 381 a, 392 a, 402 a, 454 a | Inhibition   | [79]      |
| P08563            | E2 glycoprotein (rubella virus) | 6 (interchain)                 | 53, 71, 115 | Promotion   | [80]      |
| Q9H955            | Fukutin-related protein (FKRP) | 6 (interchain)                 | 172, 209 | No relation | [81]      |
| O14773            | Tripeptidyl-peptidase I       | 111–122, 365–526, 522–537 | 210, 222, 286, 313, 443 | Promotion   | [82–85]   |
| P15813            | CD1d                          | 120–184, 224–279 | 20, 42, 108, 163 | Promotion   | [86–89]   |
| O14656            | TorsinA                       | 44–162 a, 280–319 | 143, 158 | Promotion   | [90,91]   |
### Table 1. Cont.

| UniProt Accession | Protein Name | Position of DBs | Position of N-Glycans | Class     | Reference     |
|-------------------|--------------|-----------------|-----------------------|-----------|---------------|
| P21825            | Translocation protein SEC62 | 153, 62 e | Promotion [92] |
| P04070            | Protein C    | 331–345         | 97, 248, 313, 329 (atypical) | Promotion [93] |
| K7WJ21            | PtrMAN6      | 448, 452, 456   | 23, 194, 227, 375, 392 | No relation [94] |
| P01857            | Immunoglobulin G1 Fc | 27–83 d, 103 e, 109 e, 112 e, 144–204 d, 250–308 d | 180, 297 | Promotion [84,96–98] |
|                   | Nicotinic acetylcholine receptor fragment | 128–142 | Promotion [12] |
| Q07108            | CD69         | 68 (interchain) e, 85–96 e, 113–194 e, 173–186 e | 111 (atypical), 166 | Inhibition [99,100] |
| P15813            | Antigen-presenting glycoprotein CD1d | 120–184 e, 224–279 e | 38 e, 60 e, 126 e, 181 e | Promotion [101,102] |
| P04439            | MHC class I heavy chain | 125–188 e, 227–283 e | 110 e | Promotion [103,104] |

*—predicted by sequence analysis (accessed from UniProt); †—by similarity (accessed from UniProt); ‡—a discrepancy between NG sites in the abstract and main text was found in [28]. We used the sites mentioned in the main text as they correspond with UniProt; d—PROSITE-ProRule annotation; e—UniProt.
4. Promoting Relationship

Feng et al. [37] demonstrated through kinetic studies of intracellular folding of the human chorionic gonadotropin (hCG)-β subunit that NG facilitated the rapid formation of DBs and the folding of the hCG-β subunit, which harbors six DBs [39]. The relative positions of NG and DBs are shown in Figure 3. Lacking the two NG sites slowed down the folding of the β subunit more than fourfold from 7 min to 33 min in CHO cells, and the slow formation of DBs retained the misfolded proteins up to 5 h in the ER before degradation [37]. The co-expression of the α subunit could assist the appropriate folding and secretion of the β subunit of the hormone lacking the NG. Among the six DBs in the hCG-β subunit shown in Figure 3, the formation of Cys34–Cys88 occurred earlier than that of Cys9–Cys57/Cys38–Cys90, while the remaining three pairs occurred later [39]. The first three pairs are important in protein folding and secretion and N-glycan processing. Eliminating these early formed DBs rendered part of the N-glycans to be high mannose instead of complex glycans, which were sensitive to ER quality control and degradation.

**Figure 3.** Schematic of the relative position between N-glycosylation and disulfide bonds and their relationship in the selected proteins discussed here. S4A4, UniProt ID of human SLC4 Na⁺-coupled transporter, NBCe1; CGB3, UniProt ID of human chorionic gonadotropin beta-subunit; 9HIV1, UniProt ID of human immunodeficiency virus envelope glycoprotein; CNMD, UniProt ID of human chondromodulin-1. Colors indicate different disulfide bond pairs. The N-glycan sign does not represent the actual sugar structure.
Another study was conducted in the β subunit of Na, K-ATPase. Na, K-ATPase is a plasma membrane transporter that is responsible for the maintenance of potassium and sodium homeostasis in animal cells [59]. The functional β subunit is a type II glycoprotein composed of a large C-terminal ectodomain with three NG sites (Asn\textsubscript{158}, Asn\textsubscript{193}, Asn\textsubscript{265}) and three conserved DBs (Cys\textsubscript{126}–Cys\textsubscript{149}, Cys\textsubscript{159}–Cys\textsubscript{175}, and Cys\textsubscript{213}–Cys\textsubscript{276}) [59]. The mutation of Cys\textsubscript{126}–Cys\textsubscript{149} increased the non-glycosylated proportion of the protein compared to the wildtype from the Western blot [60], suggesting a promoting relationship. Mutating each of the three glycosylation sites indicated their involvement in initial folding [59]. The acquisition of at least one sugar moiety was necessary for the β subunit to ensure its association with the α subunit through pulse chase. Interestingly, when all three N-glycans were removed, the protein did not form aggregates through DBs but permanently associated with BIP from degradation [59].

In addition to hCG and Na, K-ATPase, multiple other examples also indicate a “strengthening” relationship between NG and DB. Mirazimi and Svensson [105] showed that the chief role of NG on rotavirus VP7 is to facilitate correct intermolecular DB formation in dimerization. Removal of NG induced VP7 misfolding through random intermolecular DBs. Similar effects were also observed for MUC2 [56], vWF [36], meprin A [52], and hemagglutinin [47], as shown in Table 1.

5. Inhibitory Relationship

The hemagglutinin-neuraminidase (HN) glycoprotein of Newcastle disease virus (NDV) is responsible for virus attachment to host cell receptors, thereby initiating infection [4]. The HN protein is a type II membrane protein containing six potential NG sites: Asn\textsubscript{119}, Asn\textsubscript{341}, Asn\textsubscript{433}, Asn\textsubscript{481}, Asn\textsubscript{508}, and Asn\textsubscript{538} [45]. Among them, only four (Asn\textsubscript{119}, Asn\textsubscript{341}, Asn\textsubscript{433}, Asn\textsubscript{481}) are utilized for NG [106]. The protein also has 13 cysteine residues in the ectodomain [106], as summarized in Table 1. The cysteine residue closest to the membrane anchor (Cys\textsubscript{122}) is involved in an intermolecular DB [107,108], while the other 12 cysteine residues form intramolecular DBs [107].

McGinnes and Morrison found that intramolecular DBs might play a critical role in the usage of glycosylation sites [107]. They explored whether DB formation could be a determinant of the two unused glycosylation sites, Asn\textsubscript{508} (site 5) and Asn\textsubscript{538} (site 6), in HN protein [45]. Removing Cys\textsubscript{531}–Cys\textsubscript{542} flanking the unused glycosylation site Asn\textsubscript{538} by mutation or DTT promoted the NG of Asn\textsubscript{538} for an efficiency of 39–59% and 26–27%, respectively [45]. The successful NG was supported by the deglycosylation analysis with endo H. Under similar conditions, the usage of the non-glycosylated site 5, Asn\textsubscript{508}, which is far from any DB, was not improved. Together, these results suggest that the glycosylation of Asn\textsubscript{538} is under steric hindrance by the DB in the vicinity [45], whereas the non-glycosylated Asn\textsubscript{508} could be caused by other factors not related to DBs [45].

Another study investigating the lack of sequon utilization in tissue-type plasminogen activator (t-PA) reported similar findings that folding and DB formation of t-PA negatively impact the extent of core N-glycosylation [33]. As a result, they suggested that variable usage of glycosylation sites could be caused by the transient accessibility and appropriate orientation of the sequon relative to the transferase or dolichol-linked donor in a folding event [33].

Human sodium bicarbonate cotransporter 1, NBCe1 (SLC4A4 gene), is an electrogenic sodium/bicarbonate cotransporter localized in the plasma membrane [4]. The malfunction of this gene is related to a series of diseases in the kidney, eye, ear, brain, and tooth. All SLC4 Na\textsuperscript{+}–coupled transporters are multipass transmembrane proteins containing a large extracellular loop (EL-3) with multiple NG consensus sites and four highly conserved cysteines [109]. NBCe1-A, one of the three variants, is a homodimer, and its two EL-3 loops form unique conformations that are potentially critical to the function of the protein [28]. In the EL-3 loop of NBCe1-A, two sequons are glycosylated (Asn\textsubscript{597} and Asn\textsubscript{617}) but not Asn\textsubscript{592}, and four conserved cysteines form two intramolecular DBs (Cys\textsubscript{583}–Cys\textsubscript{585} and Cys\textsubscript{617}–Cys\textsubscript{642}), as shown in Figure 3 [28].
In a detailed study evaluating the interplay between DBs and NG to define the EL-3 loop topology in NBCe1, it was found that the two EL-3 loops of the dimer formed a unique clove conformation [28]. This conformation was “finely tuned” by glycosylation [28]. In the absence of Cys\(^{583}\)–Cys\(^{585}\) or the two NG sites, the third NG site, Asn\(^{592}\), became glycosylated. With glycans, the two DBs were deeply buried from the external surface of the EL-3 loop, which can sustain DDT-induced denaturation and enzymatic digestion under basic conditions. Losing both DBs and NG made the loop adopt an extended structure that could not be recognized by the designated antibody and was susceptible to chymotrypsin digestion [28]. Instead of considering the steric hindrance between DBs and NG at Asn\(^{592}\), the authors hypothesized that Asn\(^{592}\) was originally glycosylated at the nascent polypeptide chain of NBCe1-A and later removed when DBs and other glycosylation sites were formed [28]. Removing NG did not affect the formation of two DBs; however, an additional removal of one cysteine in the DBs by mutagenesis promoted the formation of intermolecular DBs in the homodimer, which maintained transport function [28]. A complex relationship must exist between NG and DBs in determining the final folding of EL-3 during NBCe1-A protein maturation; however, no kinetic experiments were performed to monitor protein folding, and ER maturation was not specifically probed. It is, therefore, difficult to elucidate how ER resident enzymes facilitate these processes. The study, on the other hand, had systematic structural delineation by a combinatorial mutation of all four cysteines in two DBs for a total of 12 mutants.

6. Independent Relationship

Envelope glycoprotein 160 (gp160) on human immunodeficiency virus (HIV) is critical for viral binding to the CD4 receptor and fusion with CD4\(^+\) cells. The precursor gp160 needs to be cleaved to gp120 and gp41 to activate the binding domain on gp120 with CD4. A study on the linkage region between gp120 and gp41, which is also the future binding site of gp120 to CD4, suggested that DBs and NG in this region function independently [27]. The relative position between NG and DBs in this region is shown in Figure 3. Cys\(^{402}\) and Cys\(^{429}\) are both located in the linker region but form separate DBs, of which Cys\(^{402}\) is critical for cleavage. Mutation of Cys\(^{402}\) not only prevented the cleavage but also affected the transport of gp160 and the future binding of gp120 to CD4\(^+\) cells [110,111]. Around Cys\(^{402}\), there are two occupied NG sites Asn\(^{390}\) and Asn\(^{447}\). Mutating these NG sites did not affect disulfide bonding through Cys\(^{402}\) or the relevant functions, suggesting an independent relationship between DBs and NG.

In another study of the 25 kD extracellular matrix protein chondromodulin-I (ChM-I), NG was critical in its solubility but had no effect on DBs [69]. ChM-I is a secreted protein and has two separate domains, in which the hydrophilic N-terminal domain is heavily glycosylated by one O-glycan and two N-glycans, whereas the hydrophobic C-terminal domain harbors four DBs. As the two domains are separated, the removal of either the NG or the N-terminal domain seems to have no effect on the formation of DBs in the C-terminal domain, as shown in Figure 3.

7. Unknown Relation

For most of our searched studies that concerned both DBs and NG, the exact relationship between the two PTMs was not experimentally examined. Half of these studies focused on experimental mapping of their sites without functional studies. One-third of the remaining studies only predicted the potential DBs and NG by sequence alignment or computational modeling without experimental data. For the articles that did examine the functions of both modifications, many of them did not study or discuss their interactions but rather examined them separately. For a very small number of papers, the potential interactions were hypothesized but not experimentally verified.

For example, a very nice study investigated the role of NG and DB in the rat G protein-coupled receptor class C, group 6, member A (GPRC6A) [112], a widely expressed GPCR that functions importantly in many diseases ranging from metabolic syndrome...
to cancer [113,114]. This protein is a class C GPCR with a large N-terminal extracellular domain (ECD), which contains a Venus-flytrap (VFT) domain and a cysteine-rich domain (CRD) [4]. The VFT domain is for ligand binding, and the CRD domain is for signal transfer [112].

It was observed that the ECD domain of GPRC6A consists of nine sequons [112]. Only seven asparagine residues carry N-glycans. Five of them are in the VFT domain, and Asn\textsuperscript{555} and Asn\textsuperscript{567} are located in the CRD [112]. The VFT domain also has two conserved cysteines (Cys\textsuperscript{122} and Cys\textsuperscript{131}), whereas the CRD domain has nine conserved cysteines, eight of which form intra-CRD DBs [113]. Through analysis of different mutants by SDS-PAGE, it was found that Asn\textsuperscript{555} is important for protein surface expression and that Asn\textsuperscript{567} regulates receptor function. Furthermore, from the studies of two cysteines, Cys\textsuperscript{122} and Cys\textsuperscript{131}, C\textsuperscript{131} contributed to the formation of a homodimer through an intermolecular disulfide bridge [112], and Cys\textsuperscript{122} contributed to the interdomain DB between VFT and CRD [115]. Mutation of C131A abolished the intermolecular DB and homodimer formation but did not impair receptor surface expression and its function, whereas mutation of C122A was responsible for the lowered signal response (40%) and higher (50%) surface expression [112]. This result suggested that the C122A mutation causes certain conformational changes. Not only is Cys\textsuperscript{122} next to Asn\textsuperscript{121}, but the DB between CRD and VFT domains can also be largely shaped by the seven N-glycans carried by these two domains. It is likely that NG plays a role in the potential conformation changes or intermolecular DB formation; however, no experiments or discussion were presented on the relationships between NG and cysteine disulfide bridges or cysteines in the paper.

Another study explored the role of the conserved cysteines and NG sites among all alphaherpesviruses such as herpes simplex virus 1 (HSV-1) in virus production and membrane fusion by single- and double-site directed mutagenesis [116]. Glycoprotein K (gK) is a conserved virion protein in all alphaherpesviruses [117]. The N-terminal extracellular domain of gK is important for HSV-1 to enter neurons via axonal termini. This domain contains two conserved NG sites at Asn\textsuperscript{48} and Asn\textsuperscript{58} and four conserved cysteines for two potential disulfide pairs of Cys\textsuperscript{37}–Cys\textsuperscript{114} and Cys\textsuperscript{82}–Cys\textsuperscript{243} according to single-cysteine mutation and computational modeling [116]. It was found that viruses lacking Asn\textsuperscript{58} or lacking both sites (Asn\textsuperscript{48}, Asn\textsuperscript{58}) had enhanced fusion [116]. Interestingly, deletion of Cys\textsuperscript{37} or Cys\textsuperscript{114} led to a gK-null phenotype of few plaques, whereas mutation of Cys\textsuperscript{58} or Cys\textsuperscript{243} caused enhanced cell fusion. The authors provided an extensive discussion on the potential interactions between NG and DB on the basis of the known studies and hypothesized that the removal of NG at Asn\textsuperscript{58} could displace the DB formation, as the deletion of the Cys\textsuperscript{82}–Cys\textsuperscript{243} disulfide recapitulated a similar fusion phenotype of Asn58A. However, the authors did not further verify this hypothesis, such as examining the presence of DBs through gel shift assays, labeling assays for free thiols, or MS characterization. Therefore, the authors in the end did not entail the specific relationship more than stating the presence of “a potentially important relationship” [116].

Related to the ongoing COVID-19 pandemic, the immunogen SARS-CoV-2 spike protein and its endogenous binding target ACE2 are both heavily glycosylated with numerous DBs. The SARS-CoV-2 S protein has a total of 22 sequons that to various extents are all glycosylated [118–125], seven O-glycosylations [118,125], and 40 cysteines with 15 DBs [118]. Similarly, ACE2 was mapped to have seven NG sites [36,125], one O-glycosylation [125], and four DBs [126,127]. According to mutations, molecular dynamics simulations of protein structures, and sequence alignment studies, eliminating certain DBs or NG on both ACE2 and SARS-CoV-2 S proteins can alter binding affinity to each other and change virus infectivity. For example, Cys\textsuperscript{480}–Cys\textsuperscript{488} is considered the most important pair in the receptor-binding domain (RBD) of SARS-CoV-2 S proteins, and this pair participates in binding to the N-terminal of the host receptor that forms a stable SARS-COV-2 and ACE2 complex [126–128]. In addition, Cys\textsuperscript{133}–Cys\textsuperscript{141} of ACE2 is responsible for making the loop at dimer interference [76,128] and is predicted to be crucial for making interactions with
the spike protein of SARS-CoV-2 [127]. Deletion of Asn⁹⁰ glycosylation of ACE2 increased the binding to S proteins, while removal of Asn³⁲ of ACE2 decreased virus binding and infection [125,129]. From the perspective of SARS-CoV-2 S protein, the Ser³⁰⁹ neutralizing antibody binds Asn³⁴ glycosylated RBD [32], and double mutations of N165A/N234A [35] and N331Q/N343Q [53] in S protein both reduced the binding between the immunogen and the receptor. Despite the extensive and rapid studies of NG and DBs in S proteins and ACE2 in the past 2 years, no studies have examined the relationship between DBs and NG in these two proteins. This phenomenon clearly indicates the severe understudy in this important field.

8. Common Methods

Successful studies of the relationships between DB and NG relied on suitable tools. Tables 2 and 3 summarize common methods used in studies of NG and DB, respectively.

Table 2. Summary of methodologies for the identification and structural analysis of N-glycosylation in proteins.

| NG detection          | Staining procedures [130] |
|-----------------------|---------------------------|
|                       | Resolve protein on SDS-PAGE and stain the gel for glycoproteins |
| Affinity-based methods [130] | Saccharide-binding protein (lectin-based) |
|                       | Enzyme-based methods |
|                       | Antibody-based methods |
| NMR spectroscopy [131,132] | X-ray crystallography [133] |
| Circular dichroism (CD) spectroscopy [64] | |
| NG structural analysis | Requires N-glycan removal prior to further analysis which can be achieved by |
|                       | Enzymatic removal [130,134] |
|                       | PNGase F |
|                       | Endo-H and peptide: N-glycanase [67] |
|                       | Chemical removal [130] |
|                       | β-Elimination |
|                       | Alkaline borohydride [135] |
|                       | Hydrazinolysis |
| Chromatography         | Weak anion exchange (WAX) [136] |
|                       | Gel filtration |
|                       | High-performance anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD) [137] |
|                       | Normal-phase high-performance liquid chromatography (NP-HPLC) [138] |
|                       | Reverse-phase HPLC (RP-HPLC) [139] |
|                       | Mass spectrometry [140–146] |
|                       | MALDI-MS, ESI-MS, or LC–ESI-MS [147] |
|                       | LC–MS/MS [148] |
|                       | Targeted MS/MS [149] |

NG functional analysis

Chemical tools: inhibitors of glycosyltransferases and glycosidases (in vivo) [150]

- Tunicamycin
- Plant alkaloids: australine, castanospermine [151], deoxyxojirimycin [151], deoxymannojirimycin, kifunensine, swainsonine, and mannostatin A

Physical tools: adjust temperature, ATP, pH, etc.

Site-directed mutagenesis

Genetic glyco-engineering [152]

- Involves introduction of heterologous glycosylation machinery or inactivation of endogenous enzymes.
Table 3. Summary of the methodologies for the detection and analysis of disulfide bonds in proteins.

| Quantify the amount of unbound cysteine residues | Chemical labeling and spectroscopic detection [153] |
|-------------------------------------------------|---------------------------------------------------|
| • Ellman assay                                   |                                                   |
| • N-(1-pyrenyl)maleimide (NPM) labeling          |                                                   |

| Reduction methods                                 | Reducing enzymes [153]                             |
|--------------------------------------------------|---------------------------------------------------|
| • 2-Mercaptoethanol (ME)                          |                                                   |
| • Dithiothreitol (DTT)                            |                                                   |
| • Tris(2-carboxyethyl)phosphine (TCEP)            |                                                   |
| • Tris(2-hydroxyethyl)phosphine (THP)             |                                                   |

| Detection/Analysis of DB                         | Edman degradation sequencing [154]                |
|-------------------------------------------------|---------------------------------------------------|
| • Electrophoretic methods [153,160]              |                                                   |
| • Capillary electrophoresis sodium dodecyl sulfate (CE-SDS) |   |
| • Nonreducing SDS polyacrylamide gel electrophoresis (SDS-PAGE) [160,161] |       |
| • Diagonal gel electrophoresis [162]             |                                                   |

| Mass spectrometry approaches: Front-end separation [153,163–169] | Mass spectrometry approaches: Fragmentation types |
|-----------------------------------------------------------------|---------------------------------------------------|
| • LC–MS/MS                                                      | • Activated ion ETD (AI-ETD) [170,171]            |
| • LC–ESI-MS/MS                                                  | • CID [172,173]                                  |
| • ETD-MS/MS                                                     | • HCD [174]                                     |
| • ESI-MS/MS                                                     | • EThcD [175,176]                                |
| • MS/MS/MS                                                      | • Ultraviolet photodissociation (UVPD)-MS [177]  |

| Partial proteolysis by enzymes such as trypsin and pepsin       |                                                   |
|----------------------------------------------------------------|---------------------------------------------------|
| • Epitope tags such as haemagglutinin, hexahistidine, V5, FLAG (visualized using antibodies) |               |
| • Biotin tags                                                   |                                                   |

| Detect structural changes                                      |                                                   |
|----------------------------------------------------------------|---------------------------------------------------|
| • Site-directed mutagenesis                                    |                                                   |

X-ray analysis of N-glycosylation can be challenging due to the heterogenous glycoforms that impede diffraction-quality crystallization. Nevertheless, the 3D structures of N-glycans in glycoproteins have been growing in PDB [178]. Various methods have been explored, including engineering the host cell glycosylation machinery to produce homogenous N-glycan-modified proteins for X-ray crystallography [133]. Many NMR methods have been developed to study the structure of N-glycans, intact N-glycoproteins, N-glycoprotein complexes, and model N-glycosylated peptides [131,132]. Synthetic model glycopeptides have unique advantages in terms of forming well-defined sequences and structures to interrogate their conformational effect in great detail under NMR [132]. In particular, the relations between NG and DBs have been characterized using synthetic model peptides, such as those derived from nicotinic acetylcholine receptor and prion protein [12,58]. In addition, MS has also recently been developed to decipher glycoprotein complex interactions through N-glycans, in which various glycoforms can be examined individually [179].

For molecular engineering, mutagenesis is another common method that is widely used to accurately pinpoint the site of modifications and to study the structural and functional consequence upon complete, permanent, and selective removal of some or all of these PTMs. The most frequently used mutagenesis is single-amino-acid substitution, even though deletion of one or a chain of amino acids exists. It is worth mentioning the choice of amino-acid replacements. Commonly, cysteines are mutated to alanines (A) or serines (S),
whereas asparagines are replaced by aspartic (D) and glutamic acids (E) or glutamines (Q), even though some studies replaced threonine (T)/serine (S) in the sequons to abolish NG.

Chemical removal or tagging of DBs and NG is also frequently employed to identify their presence and functions. For DBs, reducing agents can be used to disrupt the covalent bond, and thiol-reactive chemical groups can modify the free thiols to distinguish them from those that are occupied by DBs. Either a gel shift assay or MS can be employed downstream to identify changes in these chemical perturbations. Chemical removal of NG can be catalyzed by enzymes. Using enzymatic selectivity, different glycans can be readily distinguished. For example, according to the degree of processing, N-glycans have three types, i.e., high mannose, complex, and hybrid N-glycans. Endoglycosidase (endo H) cannot cleave complex N-glycans, yet N-glycosidase F (PNGase F) can [134]. Therefore, the two enzymes are frequently used to delineate the type of N-glycans. Chemical removal can also be facilitated by small molecules, such as base-assisted beta-elimination and hydrazinolysis [130], which are less selective than enzymatic reactions.

For structural characterization upon changes by DBs and NG, in addition to the instrumental approaches mentioned above, several biochemical approaches have also been developed. First, enzymes, particularly peptidase/protease, have been used to examine the overall structures of proteins. Both trypsin and pepsin were used to assess the compactness of the protein folding according to digestion efficiency. Second, for specific epitopes of a protein, antibodies were developed for rapid and specific recognition.

In addition to the above in vitro analysis methods, in vivo tools to interrogate the pathways in the formation and processing of the two PTMs were made available. A selective perturbation to the pathways can be achieved through molecularly engineered knockout, knockdown, or knock-in of a particular enzyme or chaperone. Targeted changes can also be elicited through pharmacological inhibition by small molecules. Less selective conditions, such as dithiothreitol (DTT) and 2-mercaptoethanol (2-ME) treatments, were also used to induce a global reduction of all DBs in vivo.

One important aspect of characterizing the relationships between NG and DBs is to examine the kinetics of protein maturation in the secretory pathway, as well as the kinetics of enzymatic reactions that regulate NG and DB formation and processing. Due to the migration shift in the gel after the formation of N-glycans and DBs, the protein maturation kinetics have often been examined by pulse-chase gel-shift assays. Regarding the kinetics of glycoenzymes with oxidoreductase activities, available studies are very limited. Historically, radioisotope labeling was used to probe enzymatic reactions in vitro to build predictive models for N-glycosylation [180,181]. Later, the MS characterization of glycans with and without stable isotope labeling was adopted for safer measurements [182–184]. Recently, targeted MS was developed to quantify the kinetics of stable isotope-labeled glycopeptides for more accurate modeling, in which not only the rates of glycan synthesis/processing but also the amino-acid sequences around glycosylation sites were monitored for “cell, enzyme, and glycosylation” site-specific analysis [23].

Among all the techniques, we would like to highlight MS in the characterization of the two modifications. With the advent of modern instrumentation, the method measures the mass of peptides down to sub-ppm accuracy and attomole sensitivity. The instrument holds promise for sequencing, i.e., structurally resolved, complex biological samples including all proteins, peptides, nucleic acids, carbohydrates, lipids, and metabolites within [151]. It has been playing ever increasingly important roles in deconvoluting the structures of proteins, including their DBs and glycosylation. In recent years, the application of the technique has been moved from studying one protein at a time to studying a sub-proteome by enrichment through a native moiety or chemical tagging. For example, glycoproteins and glycopeptides can be enriched by lectins or hydrophilic metals, charcoal, organic sorbents, or chemical bonding to sorbents through hydrazone and boronic acid diesters and subsequently analyzed by MS for identification, quantification, or structural interpretation [185–187]. Similarly, the thiol group has been a conventional substrate for MS analysis either at the individual protein level or at the proteome level using methods such
as ICAT [188]. With both label- and label-free quantitation rapidly developing, MS characterization holds the potential both for steady-state analyses and for kinetic interrogations in modeling biological processes. Several reviews are available that summarize the field of MS characterization in either DBs [166,188] or NG [185–187].

9. Final Remarks

It is widely accepted that both NG and BDs can affect the folding, maturation, trafficking, and degradation of host proteins; however, how to effectively control and engineer these PTMs in individual proteins for disease prevention and treatments, respectively, remains in its infancy. Unlike existing reviews on the mechanisms of enzymes functioning in NG and DB formation pathways, our study focused on the relationships of the two PTMs discovered in individual proteins. After reviewing more than 500 papers that investigated both modifications in one protein, we noticed that most studies only mapped their positions or studied their functions separately. Fewer than 100 articles have experimentally addressed the relationships between the two PTMs. We summarized the studied proteins in Table 1.

From the intriguing cooperation observed between N-glycoenzymes and oxidoreductases in the ER, it is envisaged that close relationships between NG and DBs are anticipated to widely exist in membrane and secreted proteins. Compared to the total human proteins annotated in UniProt with DBs and NGs, the proteins in Table 1 comprise less than 2%. As a result, for the most studied STT3B and STT3A substrate preferences, their complete responsive sites in all substrate proteins are still elusive. A recent discovery-based study carried out by proteomics on STT3A and STT3B substrate pools uncovered some interesting proteins [75] that could not be explained by the known mechanisms. During CNX/CRT-chaperoned N-glycoprotein folding, oxidoreductases are involved. It is known that there are ERp57 obligate and ERp57 facultative substrates [189]. ERp72 was discovered to be the alternative enzyme that acts on facultative substrates in the absence of ERp57; however, it is unclear how the obligate and facultative substrates are determined in vivo. For other complexes, such as EDEMs in ERAD, their protein substrates are just starting to emerge.

Even though relatively few proteins have been studied on the mutual relations between NG and DBs as exemplified in Figure 4, they are important disease biomarkers and therapeutic targets. Understanding the function and regulation of these PTMs is, therefore, critically important in disease treatment and prevention. Most of the studies used mutagenesis to remove one or both PTMs for structural and functional effects; however, a few studies introduced novel PTMs into proteins to engineer protein-based drugs/vaccines for better stability and efficacy [48,190,191]. In addition, knowledge obtained from studying these modifications can help researchers gain insights into early disease diagnosis and prevention [50,58].
Figure 4. Molecular pathophysiology of extracellular loop 3 of NBCe1-A caused by mutual relationships between N-glycosylation and disulfide bonds.

Due to the diverse structures possessed by individual proteins and the complex relationships, it is important to examine the entire sub-proteome to gain comprehensive knowledge. The current intriguing interactions between the two PTMs derived from model proteins should serve as an encouraging start for an exciting field. We believe that the MS-led new generation of high-throughput high-accuracy analyses can quickly move this field forward. Lastly, we hope that this review will encourage future studies to investigate the relationship between NG and DBs and to better disclose their hidden linkages in the remaining 98% of the proteins for novel insights into their structural and functional roles.

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References
1. Adams, B.M.; Oster, M.E.; Hebert, D.N. Protein Quality Control in the Endoplasmic Reticulum. *Protein J.* 2019, 38, 317–329. [CrossRef] [PubMed]
2. Grek, C.; Townsend, D.M. Protein Disulfide Isomerase Superfamily in Disease and the Regulation of Apoptosis. *Endoplasmic Reticulum Stress Dis.* 2014, 1, 4–17. [CrossRef] [PubMed]
32. Vlahos, C.J.; Wilhelm, O.G.; Hassell, T.; Jaskunas, S.R.; Bang, N.U. Disulfide pairing of the recombinant kringle-2 domain of tissue plasminogen activator produced in Escherichia coli. J. Biol. Chem. 1991, 266, 10070–10072. [CrossRef]
33. Allen, S.; Naim, H.Y.; Bulleid, N.J. Intracellular folding of tissue-type plasminogen activator. Effects of disulfide bond formation on N-linked glycosylation and secretion. J. Biol. Chem. 1995, 270, 4797–4804. [CrossRef] [PubMed]
34. Cruz, M.A.; Handin, R.I.; Wise, R.J. The interaction of the von Willebrand factor-A1 domain with platelet glycoprotein Ib/IX. The role of glycosylation and disulfide bonding in a monomeric recombinant A1 domain protein. J. Biol. Chem. 1993, 268, 21238–21245. [CrossRef]
35. Marti, T.; Rosselet, S.J.; Titani, K.; Walsh, K.A. Identification of disulfide-bridged substructures within human von Willebrand factor. Biochemistry 1987, 26, 8099–8109. [CrossRef]
36. McKinney, T.A.; Goode, E.C.; Birdsey, G.M.; Nowak, A.A.; Chan, A.C.; Lane, D.A.; Laffan, M.A. Specific N-linked glycosylation sites modulate synthesis and secretion of von Willebrand factor. Blood 2010, 116, 640–648. [CrossRef]
37. Feng, W.; Huth, J.R.; Norton, S.E.; Rudder, R.W. Asparagine-linked oligosaccharides facilitate human chorionic gonadotropin beta-subunit folding but not assembly of prefolded beta with alpha. Endocrinology 1995, 136, 52–61. [CrossRef]
38. Feng, W.; Matzuk, M.M.; Mountjoy, K.; Bedows, E.; Rudder, R.W.; Boime, I. The asparagine-linked oligosaccharides of the human chorionic gonadotropin beta subunit facilitate correct disulfide bond pairing. J. Biol. Chem. 1995, 270, 11851–11859. [CrossRef]
39. Moriwaki, T.; Suganuma, N.; Furuhashi, M.; Kikkawa, F.; Tomoda, Y.; Boime, I.; Nakata, M.; Mizuochi, T. Alteration of N-linked oligosaccharide structures of human chorionic gonadotropin beta-subunit by disruption of disulfide bonds. Glycoconj. J. 1997, 14, 225–229. [CrossRef]
40. Kuglstatter, A.; Stühle, M.; Neumann, C.; Muller, C.; Schaefer, W.; Klein, C.; Benz, J. Structural differences between glycosylated, disulfide-linked heterodimeric Knob-into-Hole Fc fragment and its homodimeric Knob-Knob and Hole-Hole side products. Protein Eng. Des. Sel. PEDS 2017, 30, 649–656. [CrossRef]
41. Lipari, F.; Herscovics, A. Role of the cysteine residues in the alpha1,2-mannosidase involved in N-glycan biosynthesis in Saccharomyces cerevisiae. The conserved Cys340 and Cys385 residues form an essential disulfide bond. J. Biol. Chem. 1996, 271, 27615–27622. [CrossRef] [PubMed]
42. Flahaut, M.; Pfister, C.; Rossier, B.C.; Firsov, D. N-Glycosylation and conserved cysteine residues in RAMP3 play a critical role for the functional expression of CRLR/RAMP3 adrenomedullin receptor. Biochemistry 2003, 42, 10333–10341. [CrossRef]
43. Walker, A.K.; Soo, K.Y.; Levina, V.; Talbo, G.H.; Atkin, J.D. N-linked glycosylation modulates dimerization of protein disulfide isomerase family A member 2 (PDI2). FEBS J. 2013, 280, 233–243. [CrossRef] [PubMed]
44. Vidal, S.; Mottet, G.; Kolakofsky, D.; Roux, L. Addition of high-mannose sugars must precede disulfide bond formation for proper folding of Sendai virus glycoproteins. J. Virol. 1989, 63, 892–900. [CrossRef] [PubMed]
45. McGinnis, L.W.; Morrison, T.G. Disulfide bond formation is a determinant of glycosylation site usage in the hemagglutinin-neuraminidase glycoprotein of Newcastle disease virus. J. Virol. 1997, 71, 3083–3089. [CrossRef]
46. Pitt, J.J.; Da Silva, E.; Gorman, J.J. Determination of the disulfide bond arrangement of Newcastle disease virus hemagglutinin-neuraminidase glycoprotein of Newcastle disease virus. J. Biol. Chem. 2000, 275, 6469–6478. [CrossRef]
47. Daniels, R.; Kurowski, B.; Johnson, A.E.; Hebert, D.N. N-linked glycan structures of human chorionic gonadotropin beta subunit facilitate correct disulfide bond pairing. J. Biol. Chem. 1995, 270, 14651–14658. [CrossRef] [PubMed]
48. Thornlow, D.N.; Macintyre, A.N.; Oguin, T.H.; Karlsson, A.B.; Stover, E.L.; Lynch, H.E.; Sempowski, G.D.; Schmidt, A.G. Altering the Immunogenicity of Hemagglutinin Immunogens by Hyperglycosylation and Disulfide Stabilization. Front. Immunol. 2021, 12, 737973. [CrossRef]
49. Suzuki, S.; Furuhashi, M.; Suganuma, N. Additional N-glycosylation at Asn13 rescues the human LHbeta-subunit from disulfide-linked aggregation. Mol. Cell. Endocrinol. 2000, 160, 157–163. [CrossRef]
50. Naessens, S.; De Zeytijd, J.; Suy, D.; Vandenbroucke, R.E.; Smeets, F.; Van Cauwenbergh, C.; Leroy, B.P.; Coppieters, F. The N-terminal p.(Ser38Cys) TIMP3 mutation underlying Sorsby fundus dystrophy is a founder mutation disrupting an intramolecular disulfide bond. Hum. Mutat. 2019, 40, 539–551. [CrossRef]
51. Arolas, J.L.; Broder, C.; Jefferson, T.; Guevara, T.; Sterchi, E.E.; Bode, W.; Stocker, W.; Becker-Pauly, C.; Gomis-Ruth, F.X. Structural basis for the sheddase function of human meprin beta metallocproteinase at the plasma membrane. Proc. Natl. Acad. Sci. USA 2012, 109, 16131–16136. [CrossRef] [PubMed]
52. Ishmael, S.S.; Ishmael, F.T.; Jones, A.D.; Bond, J.S. Protease domain glycans affect oligomerization, disulfide bond formation, and stability of the meprin A metalloprotease homo-oligomer. J. Biol. Chem. 2006, 281, 37404–37415. [CrossRef] [PubMed]
53. Liu, T.; Qian, W.J.; Gritsenko, M.A.; Camp, D.G.; 2nd; Monroe, M.E.; Moore, R.J.; Smith, R.D. Human plasma N-glycoproteome analysis by immunoaffinity subtraction, hydrazide chemistry, and mass spectrometry. J. Proteome Res. 2005, 4, 2070–2080. [CrossRef]
54. McCormick, L.M.; Urade, R.; Arakaki, Y.; Schwartz, A.L.; Bu, G. Independent and cooperative roles of N-glycans and molecular chaperones in the folding and disulfide bond formation of the low-density lipoprotein (LDL) receptor-related protein. Biochemistry 2005, 44, 5794–5803. [CrossRef] [PubMed]
55. Zelcer, N.; Hong, C.; Boyadjian, R.; Tontonoz, P. LXRx regulates cholesterol uptake through Idol-dependent ubiquitination of the LDL receptor. Science 2009, 325, 100–104. [CrossRef]
56. Asker, N.; Axelsson, M.A.; Olofsson, S.O.; Hansson, G.C. Dimerization of the human MUC2 mucin in the endoplasmic reticulum is followed by a N-glycosylation-dependent transfer of the mono- and dimers to the Golgi apparatus. J. Biol. Chem. 1998, 273, 18857–18863. [CrossRef]

57. Bell, S.L.; Xu, G.; Khatri, I.A.; Wang, R.; Rahman, S.; Forstner, J.F. N-linked oligosaccharides play a role in disulphide-dependent dimerization of intestinal mucin Muc2. Biochem. J. 2003, 373, 893–900. [CrossRef]

58. Bosques, C.J.; Imperiali, B. The interplay of glycosylation and disulphide formation influences fibrillization in a prion protein fragment. Proc. Natl. Acad. Sci. USA 2003, 100, 7593–7598. [CrossRef]

59. Beggah, A.T.; Jaunin, P.; Geering, K. Role of glycosylation and disulphide bond formation in the beta subunit in the folding and functional expression of Na, K-ATPase. J. Biol. Chem. 1997, 272, 10318–10326. [CrossRef]

60. Laughery, M.D.; Todd, M.L.; Kaplan, J.H. Mutation analysis of alpha-beta subunit interactions in the delivery of Na, K-ATPase heterodimers to the plasma membrane. J. Biol. Chem. 2003, 278, 34794–34803. [CrossRef]

61. Clogston, C.L.; Boone, T.C.; Crandall, B.C.; Mendiz, E.A.; Lu, H.S. Disulfide structures of human interleukin-6 are similar to those of human granulocyte colony stimulating factor. Arch. Biochem. Biophys. 1989, 272, 144–151. [CrossRef]

62. May, L.T.; Shaw, J.E.; Khanna, A.K.; Zabriskie, J.B.; Sehgal, P.B. Marked cell-type-specific differences in glycosylation of human interleukin-6. Cytokine 1991, 3, 204–211. [CrossRef]

63. Orita, T.; Oh-edah, M.; Hasegawa, M.; Kubon1wa, H.; Esaki, K.; Ochi, N. Polypeptide and carbohydrate structure of recombinant human interleukin-6. J. Biol. Chem. 1991, 266, 18857–18863. [CrossRef] [PubMed]

64. Roman-Sosa, G.; Karger, A.; Kraatz, F.; Aeber, A.; Wernike, K.; Maksimov, P.; Lillig, C.H.; Reimann, I.; Brocchi, E.; Keller, M.; et al. The amino terminal subdomain of glycoprotein Ge of Schmallemborg virus: Disulfide bonding and structural determinants of neutralization. J. Gen. Virol. 2017, 98, 1259–1273. [CrossRef]

65. Mohd Yusuf, S.N.; Bailey, U.M.; Tan, N.Y.; Jamaluddin, M.F.; Schulz, B.L. Mixed disulfide formation in vitro between a glycoprotein substrate and yeast oligosaccharyltransferase subunits Ost3p and Ost6p. Biochem. Biophys. Res. Commun. 2013, 432, 438–443. [CrossRef]

66. Schulz, B.L.; Stirnimann, C.U.; Grisham, J.P.; Brozzo, M.S.; Fritsch, F.; Mohorko, E.; Capitani, G.; Glockshuber, R.; Grutter, M.G.; Aebi, M. Oxidoreductase activity of the large subunit of yeast oligosaccharyltransferase. J. Biol. Chem. 2009, 284, 475–485. [CrossRef]

67. Ohgumori, T.; Nanao, T.; Morita, A.; Ikekita, M. Asn54-linked glycan is critical for functional folding of intercellular adhesion molecule-5. Glycoconj. J. 2012, 29, 47–55. [CrossRef]

68. Zhang, H.; Gasnasnovas, J.M.; Jin, M.; Liu, J.H.; Gahnberg, C.G.; Springer, T.A.; Wang, J.H. An unusual allosteric mobility of the C-terminal helix of a high-affinity alphal integrin I domain variant bound to ICAM-5. Mol. Cell 2008, 31, 432–437. [CrossRef]

69. Kondo, J.; Shibata, H.; Miura, S.; Yamakawa, A.; Sato, K.; Higuchi, Y.; Shukummi, C.; Hiraki, Y. A functional role of the glycosylated N-terminal domain of chondromodulin-I. J. Bone Miner. Metab. 2011, 29, 23–30. [CrossRef]

70. Hoffman, R.C.; Andersen, H.; Walker, K.; Krakover, J.D.; Patel, S.; Stamm, M.R.; Osborn, S.G. Peptide, disulfide, and glycosylation structures of human interleukin-6 produced in Chinese hamster ovary cells. J. Biol. Chem. 2009, 284, 3976–3984. [CrossRef] [PubMed]

71. Kato, T. Protein characteristics of thrombopoietin. J. Gen. Virol. 2002, 83, 14849–14861. [CrossRef]

72. May, L.T.; Shaw, J.E.; Khanna, A.K.; Zabriskie, J.B.; Sehgal, P.B. Peptide, disulfide, and glycosylation mapping of recombinant human thrombopoietin from ser1 to Arg246. Biochemistry 1996, 35, 14849–14861. [CrossRef]

73. Kato, T. Protein characteristics of thrombopoietin. Stem Cells 1996, 14 (Suppl. S1), 139–147. [CrossRef] [PubMed]

74. Diop, N.K.; Hrycyna, C.A. N-Linked glycosylation of the human ABC transporter ABCG2 on asparagine 596 is not essential for expression, transport activity, or trafficking to the plasma membrane. Biochemistry 2005, 44, 5420–5429. [CrossRef] [PubMed]

75. Wakabayashi-Nakao, K.; Tamura, A.; Furukawa, T.; Nakagawa, H.; Ishikawa, T. Quality control of human ABCG2 protein in the endoplasmic reticulum: Ubiquitination and proteasomal degradation. Adv. Drug Deliv. Rev. 2009, 61, 66–72. [CrossRef]

76. Fischer, F.; Molinari, M.; Bodendorf, U.; Paganetti, P. The disulphide bonds in the catalytic domain of BACE are critical but not essential for amyloid precursor protein processing activity. J. Neurochem. 2002, 80, 1079–1088. [CrossRef] [PubMed]

77. Cherepanova, N.A.; Venev, S.V.; Leszyk, J.D.; Shaffer, S.A.; Gilmore, R. Quantitative glycoproteomics reveals new classes of oxidoreductase activity necessary for N-glycosylation of cysteine-proximal acceptor sites in glycoproteins. J. Cell Biol. 2014, 206, 525–539. [CrossRef]

78. Ruiz-Canada, C.; Kelleher, D.J.; Gilmore, R. Cottranslational and posttranslational N-glycosylation of polypeptides by distinct mammalian OST isoforms. Cell 2009, 137, 273–283. [CrossRef]

79. Rius, M.; Sala, L.; Chillaron, J. The role of N-glycans and the C-terminal loop of the subunit rBAT in the biogenesis of the cystinuria-associated transporter. Biochem. J. 2016, 473, 233–244. [CrossRef]

80. Ivanenkov, V.V.; Meller, J.; Kirley, T.L. Characterization of disulphide bonds in human nucleoside triphosphate diphosphohydrolase 3 (NTPDase3): Implications for NTPDase structural modeling. Biochemistry 2005, 44, 8998–9012. [CrossRef]

81. Qiu, Z.; Hobman, T.C.; McDonald, H.L.; Seto, N.O.; Gillam, S. Role of N-linked oligosaccharides in processing and intracellular transport of E2 glycoprotein of rubella virus. J. Virol. 1992, 66, 3514–3521. [CrossRef]

82. Alhamidi, M.; Kjeldsen Buvang, E.; Fagerheim, T.; Brox, V.; Lindal, S.; Van Ghelue, M.; Nilssen, O. Fukutin-related protein resides in the Golgi cisternae of skeletal muscle fibres and forms disulphide-linked homodimers via an N-terminal interaction. PLoS ONE 2011, 6, e22968. [CrossRef] [PubMed]

83. Pal, A.; Kraetzner, R.; Gruene, T.; Grapp, M.; Schreiber, K.; Gronborg, M.; Urlaub, H.; Becker, S.; Asif, A.R.; Gartner, J.; et al. Structure of tripeptidyl-peptidase I provides insight into the molecular basis of late infantile neuronal ceroid lipofuscinosis. J. Biol. Chem. 2009, 284, 3976–3984. [CrossRef] [PubMed]
109. Kao, L.; Sassani, P.; Azimov, R.; Pushkin, A.; Abuladze, N.; Peti-Peterdi, J.; Liu, W.; Newman, I.; Kurtz, I. Oligomeric structure and minimal functional unit of the electrogenic sodium bicarbonate cotransporter NBCe1-A. *J. Biol. Chem.* 2008, 283, 26782–26794. [CrossRef]

110. Hemmeng, A.; Bolmstedt, A.; Flodby, P.; Lundberg, L.; Gidlund, M.; Wigzell, H.; Olafsson, S.O. Cysteine 402 of HIV gp120 is essential for CD4-binding and resistance of gp120 to intracellular degradation. *Arch. Virol.* 1999, 109, 269–276. [CrossRef]

111. Rider, P.J.F.; Naderi, M.; Bergeron, S.; Chouljenko, V.N.; Brylinski, M.; Kousoulas, K.G. Cysteines and N-Glycosylation Sites Derived from the GlycoDelete HEK293 cell line enable diffraction-quality crystallogenesis. *FEBS Lett.* 2015, 588, 588–597. [CrossRef] [PubMed]

112. Norskov-Lauritsen, L.; Jorgensen, S.; Brauner-Osborne, H. N-glycosylation and disulfide bonding affects GPRC6A receptor expression, function, and dimerization. *FEBS Lett.* 2015, 588, 588–597. [CrossRef] [PubMed]

113. Zhang, S.; Go, E.P.; Ding, H.; Anang, S.; Kappes, J.C.; Desaire, H.; Sodroski, J. Analysis of glycosylation and disulfide bonding of SARS-CoV-2 spike protein. *J. Virol.* 1990, 64, 2250–2259. [CrossRef] [PubMed]

114. Sanda, M.; Morrison, L.; Goldman, R. N and O glycosylation of the SARS-CoV-2 spike protein. *Anal. Chem.* 2020, 92, 14730–14739. [CrossRef]

115. Muto, T.; Tsuchiya, D.; Morikawa, K.; Jingami, H. Structures of the extracellular regions of the group II/III metabotropic glutamate receptors. *Proc. Natl. Acad. Sci. USA* 2007, 104, 3759–3764. [CrossRef]

116. Rider, P.J.F.; Naderi, M.; Bergeron, S.; Chouljenko, V.N.; Brylinski, M.; Kousoulas, K.G. Cysteines and N-Glycosylation Sites Derived from the GlycoDelete HEK293 cell line enable diffraction-quality crystallogenesis. *FEBS Lett.* 2015, 588, 588–597. [CrossRef] [PubMed]

117. Go, E.P.; Ding, H.; Anang, S.; Kappes, J.C.; Desaire, H.; Sodroski, J. Analysis of glycosylation and disulfide bonding of wild-type SARS-CoV-2 spike glycoprotein. *J. Virol.* 2022, 96, e01626-21. [CrossRef]

118. Brun, J.; Vasiljevic, S.; Gangadharan, B.; Hensen, M.; Chandran, A.V.; Hill, M.L.; Kappes, J.L.; Dwek, R.A.; Alonzi, D.S.; Struve, W.B.; et al. Analysis of SARS-CoV-2 spike glycosylation reveals shedding of a vaccine candidate. *bioRxiv* 2020. [CrossRef]

119. Yao, H.; Song, Y.; Chen, Y.; Wu, N.; Xu, J.; Sun, C.; Zhang, J.; Weng, T.; Zhang, Z.; Wu, Z.; et al. Molecular Architecture of the SARS-CoV-2 Spike Proteins Using Signature Ions-Triggered Electron-Transfer/Higher-Energy Collisional Dissociation (EThcD) Mass Spectrometry. *Anal. Chem.* 2020, 92, 14730–14739. [CrossRef]

120. Wang, D.; Baudys, J.; Bundy, J.L.; Barr, J.R. Comprehensive Analysis of the Glycan Complement of SARS-CoV-2 Spike Proteins Using Signature Ions-Triggered Electron-Transfer/Higher-Energy Collisional Dissociation (EThcD) Mass Spectrometry. *Anal. Chem.* 2020, 92, 14730–14739. [CrossRef]

121. Hori, Y.; Kozaki, K.; Kato, T.; Yamada, H.; Kato, Y.; Ishi, K.; Kato, T.; Kuroda, K.; Hori, Y.; et al. The crystal structure of the extracellular region of the group II metabotropic glutamate receptor. *ACS Omega* 2019, 4, 3301–3309. [CrossRef] [PubMed]

122. Sanda, M.; Morrison, L.; Goldman, R. N and O glycosylation of the SARS-CoV-2 spike protein. *Anal. Chem.* 2020, 92, 14730–14739. [CrossRef]

123. Watanabe, Y.; Allen, J.D.; Wrapp, D.; McLellan, J.S.; Crispin, M. Site-specific glycan analysis of the SARS-CoV-2 spike glycoprotein from an advanced vaccine candidate. *Cell Host Microbe* 2020, 28, 586–601.e586. [CrossRef]

124. Bangaru, S.; Ozorowski, G.; Turner, H.L.; Antanasijevic, A.; Huang, D.; Wang, X.; Torres, J.L.; Diedrich, J.K.; Tian, J.H.; Portnoff, A.D.; et al. Structural analysis of full-length SARS-CoV-2 spike protein from a vaccine candidate. *Nature* 2020, 589, 1089–1094. [CrossRef]

125. Zhao, P.; Praisman, J.L.; Grant, O.C.; Cai, Y.; Xiao, T.; Rosenbaum, K.E.; Aoki, K.; Kellman, B.P.; Bridger, R.; Barouch, D.H.; et al. Virus-Receptor Interactions of Glycosylated SARS-CoV-2 Spike and Human ACE2 Receptor. *Cell Host Microbe* 2020, 28, 586–601.e586. [CrossRef]

126. Hori, Y.; Kozaki, K.; Kato, T.; Yamada, H.; Kato, Y.; Ishi, K.; Kato, T.; Kuroda, K.; Hori, Y.; et al. The crystal structure of the extracellular region of the group II metabotropic glutamate receptor. *ACS Omega* 2019, 4, 3301–3309. [CrossRef] [PubMed]

127. Singh, J.; Dhindsa, R.S.; Misra, V.; Singh, B. SARS-CoV2 infectivity is potentially modulated by host redox status. *Comput. Struct. Biotechnol. J.* 2020, 18, 3705–3711. [CrossRef]

128. Lan, J.; Ge, J.; Yu, J.; Shan, S.; Zhou, H.; Fan, S.; Zhang, Q.; Shi, X.; Wang, Q.; Zhang, L.; et al. Structure of the SARS-CoV-2 spike receptor-binding domain bound to the ACE2 receptor. *Nature* 2020, 581, 215–220. [CrossRef]

129. Mehdipour, A.R.; Hummer, G. Dual nature of human ACE2 glycosylation in binding to SARS-CoV-2 spike. *Comput. Struct. Biotechnol. J.* 2020, 62, 22–30. [CrossRef]

130. Roth, Z.; Yeheskel, G.; Khaila, I. Identification and Quantification of Protein Glycosylation. *Int. J. Carbohydr. Chem.* 2012, 2012, 640923. [CrossRef]

131. Gimeno, A.; Valverde, P.; Arda, A.; Jimenez-Barbero, J. Glycan structures and their interactions with proteins. A NMR view. *Curr. Opin. Struct. Biol.* 2020, 62, 22–30. [CrossRef]

132. Engineer, S.; Bloch, Y.; De Munck, S.; Mikula, A.; Bento, I.; Savvides, S.N.; Meijers, R. Homogeneously N-glycosylated proteins derived from the GlycoDelete HEK293 cell line enable diffraction-quality crystallogenesis. *Acta Crystallogr. Sect. D Struct. Biol.* 2020, 76, 1244–1255. [CrossRef]

133. Freeze, H.H.; Kranz, C. Endoglycosidase and glycoamidase release of N-linked glycans. *Curr. Protoc. Mol. Biol.* 2010, 89, 17.13A.1–17.13A.25. [CrossRef]
135. Kinsella, M.G.; Wight, T.N. Isolation and characterization of dermran sulfate proteoglycans synthesized by cultured bovine aortic endothelial cells. J. Biol. Chem. 1988, 263, 19222–19231. [CrossRef]

136. Guile, G.R.; Wong, S.Y.; Dwek, R.A. Analytical and preparative separation of anionic oligosaccharides by weak anion-exchange high-performance liquid chromatography on an inert polymer column. Anal. Biochem. 1994, 222, 231–235. [CrossRef]

137. Townsend, R.R.; Hardy, M.R. Analysis of glycoprotein oligosaccharides using high-pH anion exchange chromatography. Glycobiology 1991, 1, 139–147. [CrossRef]

138. Ilan, A.; Pinto, G.; Melchiorre, C.; Carpentieri, A.; Faraco, V.; Amoresano, A. Protein Glycosylation Investigated by Mass Spectrometry: An Overview. Cells 2020, 9, 9. [CrossRef] [PubMed]

139. Meachref, Y.; Novotny, M.V. Structural investigations of glycoconjugates at high sensitivity. Chem. Rev. 2002, 102, 321–369. [CrossRef]

140. Sagi, D.; Kienz, P.; Denecke, J.; Marquardt, T.; Peter-Katalinic, J. Glycoproteomics of N-glycosylation by in-gel deglycosylation and matrix-assisted laser desorption/ionisation-time of flight mass spectrometry mapping: Application to congenital disorders of glycosylation. Proteomics 2005, 5, 2689–2701. [CrossRef] [PubMed]

141. Zhang, H.; Li, X.J.; Martin, D.B.; Aebisher, R. Identification and quantification of N-linked glycoproteins using hydrazide chemistry, stable isotope labeling and mass spectrometry. Nat. Biotechnol. 2003, 21, 660–666. [CrossRef] [PubMed]

142. Zhu, F.; Trinidad, J.C.; Clemmer, D.E. Glycopeptide Site Heterogeneity and Structural Diversity Determined by Combined Lectin Affinity Chromatography/IMS/CID/MS Techniques. J. Am. Soc. Mass Spectrom. 2015, 26, 1092–1102. [CrossRef] [PubMed]

143. Bloom, J.W.; Madanat, M.S.; Ray, M.K. Cell line and site specific comparative analysis of the N-linked oligosaccharides on human ICAM-1des454-532 by electrospray ionization mass spectrometry. Biochemistry 1996, 35, 1856–1864. [CrossRef]

144. Han, L.; Costello, C.E. Mass spectrometry of glycans. Anal. Biochem. 1991, 204, 44–66. [CrossRef]

145. Ford, K.L.; Zeng, W.; Heazlewood, J.L.; Bacic, A. Characterization of protein N-glycosylation by tandem mass spectrometry using complementary fragmentation techniques. Front Plant Sci 2015, 6, 674. [CrossRef]

146. Escobedo, J.; Bertozzi, C.; Schnaar, R.L. Chemical Tools for Inhibiting Glycosylation. In Essentials of Glycobiology, 3rd ed.; Varki, A., Cummings, R.D., Esko, J.D., Stanley, P., Hart, G.W., Aebi, M., Darvill, A.G., Kinoshita, T., Packer, N.H., Prestegard, J.H., et al., Eds.; Cold Spring Harbor: Huntington, NY, USA, 2015; pp. 701–712. [CrossRef]

147. Wainwright, L.J.; Field, M.C. Quality control of glycosylphosphatidylinositol anchor attachment in mammalian cells: A biochemical study. Biochem. J. 1997, 321 Pt 3, 655–664. [CrossRef]

148. Barolo, L.; Abbriano, R.M.; Commault, A.S.; George, J.; Kahlke, T.; Fabris, M.; Padula, M.P.; Lopez, A.; Ralph, P.J.; Pernice, M. Oxidative Folding of Proteins: Basic Principles, Cellular Regulation and Engineering; Royal Society of Chemistry: London, UK, 2018; pp. 81–98. [CrossRef]

149. Haniu, M.; Acklin, C.; Kenney, W.C.; Rohde, M.F. Direct assignment of disulfide bonds by Edman degradation of selected peptide polypeptide ions. Anal. Biochem. 1985, 145, 200–204. [CrossRef]

150. Thannhauser, T.W.; Konishi, Y.; Scheraga, H.A. Analysis for disulfide bonds by electrophoresis. In The Protein Protocols Handbook; Walker, J.M., Ed.; Humana Press: Totowa, NJ, USA, 2009; pp. 1057–1062. [CrossRef]

151. Barolo, L.; Abbriano, R.M.; Commault, A.S.; George, J.; Kahlke, T.; Fabris, M.; Padula, M.P.; Lopez, A.; Ralph, P.J.; Pernice, M. Oxidative Folding of Proteins: Basic Principles, Cellular Regulation and Engineering; Royal Society of Chemistry: London, UK, 2018; pp. 81–98. [CrossRef]

152. Haniu, M.; Acklin, C.; Kenney, W.C.; Rohde, M.F. Direct assignment of disulfide bonds by Edman degradation of selected peptide polypeptide ions. Anal. Biochem. 1985, 145, 200–204. [CrossRef]

153. Aitken, A.; Learmonth, M. Quantitation of cysteine residues and disulfide bonds by electrophoresis. In The Protein Protocols Handbook; Walker, J.M., Ed.; Humana Press: Totowa, NJ, USA, 2009; pp. 1057–1062. [CrossRef]

154. Aitken, A.; Learmonth, M. Quantitation of cysteine residues and disulfide bonds by electrophoresis. In The Protein Protocols Handbook; Walker, J.M., Ed.; Humana Press: Totowa, NJ, USA, 2009; pp. 1057–1062. [CrossRef]

155. Haniu, M.; Acklin, C.; Kenney, W.C.; Rohde, M.F. Direct assignment of disulfide bonds by Edman degradation of selected peptide polypeptide ions. Anal. Biochem. 1985, 145, 200–204. [CrossRef]

156. Haniu, M.; Acklin, C.; Kenney, W.C.; Rohde, M.F. Direct assignment of disulfide bonds by Edman degradation of selected peptide polypeptide ions. Anal. Biochem. 1985, 145, 200–204. [CrossRef]

157. Haniu, M.; Acklin, C.; Kenney, W.C.; Rohde, M.F. Direct assignment of disulfide bonds by Edman degradation of selected peptide polypeptide ions. Anal. Biochem. 1985, 145, 200–204. [CrossRef]

158. Haniu, M.; Acklin, C.; Kenney, W.C.; Rohde, M.F. Direct assignment of disulfide bonds by Edman degradation of selected peptide polypeptide ions. Anal. Biochem. 1985, 145, 200–204. [CrossRef]

159. Haniu, M.; Acklin, C.; Kenney, W.C.; Rohde, M.F. Direct assignment of disulfide bonds by Edman degradation of selected peptide polypeptide ions. Anal. Biochem. 1985, 145, 200–204. [CrossRef]

160. Haniu, M.; Acklin, C.; Kenney, W.C.; Rohde, M.F. Direct assignment of disulfide bonds by Edman degradation of selected peptide polypeptide ions. Anal. Biochem. 1985, 145, 200–204. [CrossRef]
164. Cramer, C.N.; Haselmann, K.F.; Olsen, J.V.; Nielsen, P.K. Disulfide Linkage Characterization of Disulfide Bond-Containing Proteins and Peptides by Reducing Electrochemistry and Mass Spectrometry. *Anal. Chem.* 2016, 88, 1585–1592. [CrossRef] [PubMed]

165. Gorman, J.J.; Wallis, T.P.; Pitt, J.J. Protein disulfide bond determination by mass spectrometry. *Mass Spectrom. Rev.* 2002, 21, 183–216. [CrossRef] [PubMed]

166. Lakhub, J.C.; Shipman, J.T.; Desaire, H. Recent mass spectrometry-based techniques and considerations for disulfide bond characterization in proteins. *Anal. Bioanal. Chem.* 2018, 410, 2467–2484. [CrossRef]

167. Tsai, P.; Chen, S.; Huang, S.Y. Mass spectrometry-based strategies for protein disulfide bond identification. *Rev. Anal. Chem.* 2013, 32, 257–268. [CrossRef]

168. Wiesner, J.; Reesemann, A.; Evans, C.; Suckau, D.; Jabs, W. Advanced mass spectrometry workflows for analyzing disulfide bonds in biologics. *Expert Rev. Proteom.* 2015, 12, 115–123. [CrossRef]

169. Zhou, Y.; Dong, J.; Vachet, R.W. Electron transfer dissociation of modified peptides and proteins. *Curr. Pharm. Biotechnol.* 2011, 12, 1558–1567. [CrossRef]

170. Sarbu, M.; Ghislai, R.M.; Zamfir, A.D. Recent developments and applications of electron transfer dissociation mass spectrometry in proteomics. *Amino Acids* 2014, 46, 1625–1638. [CrossRef]

171. Wu, S.L.; Jiang, H.; Hancock, W.S.; Karger, B.L. Identification of the unpaired cysteine status and complete mapping of the 17 disulfides of recombinant tissue plasminogen activator using LC-MS with electron transfer dissociation/collision induced dissociation. *Anal. Chem.* 2010, 82, 5296–5303. [CrossRef]

172. Lakhub, J.C.; Clark, D.F.; Shah, I.S.; Zhu, Z.; Go, E.P.; Tolbert, T.J.; Desaire, H. Disulfide Bond Characterization of Endogenous IgG3 Monoclonal Antibodies Using LC-MS: An Investigation of IgG3 Disulfide-mediated Isoforms. *Anal. Methods* 2016, 8, 6046–6056. [CrossRef]

173. Clark, D.F.; Go, E.P.; Toumi, M.L.; Desaire, H. Collision induced dissociation products of disulfide-bonded peptides: Ions result from the cleavage of more than one bond. *J. Am. Soc. Mass Spectrom.* 2011, 22, 492–498. [CrossRef]

174. Olsen, J.V.; Macek, B.; Lange, O.; Makarov, A.; Horning, S.; Mann, M. Higher-energy C-trap dissociation for peptide modification analysis. *Nat. Methods* 2007, 4, 709–712. [CrossRef] [PubMed]

175. Frese, C.K.; Altelaar, A.F.; van den Toorn, H.; Nolting, D.; Griepe-Raming, J.; Heck, A.J.; Mohammed, S. Toward full peptide sequence coverage by dual fragmentation combining electron-transfer and higher-energy collision dissociation tandem mass spectrometry. *Anal. Chem.* 2012, 84, 9668–9673. [CrossRef] [PubMed]

176. Liu, F.; van Breukelen, B.; Heck, A.J. Facilitating protein disulfide mapping by a combination of pepsin digestion, electron transfer higher energy dissociation (EThC-D), and a dedicated search algorithm SlinkS. *Mol. Cell Proteom.* 2014, 13, 2776–2786. [CrossRef] [PubMed]

177. Agarwal, A.; Diedrich, J.K.; Julian, R.R. Direct elucidation of disulfide bond partners using ultraviolet photodissociation mass spectrometry. *Anal. Chem.* 2011, 83, 6455–6458. [CrossRef]

178. Suga, A.; Nagae, M.; Yamaguchi, Y. Analysis of protein landscapes around N-glycosylation sites from the PDB repository for understanding the structural basis of N-glycoprotein processing and maturation. *Glycobiology* 2018, 28, 774–785. [CrossRef]

179. Wu, D.; Struve, W.B.; Harvey, D.J.; Ferguson, M.A.J.; Robinson, C.V. N-glycan microheterogeneity regulates interactions of plasma proteins. *Proc. Natl. Acad. Sci. USA* 2018, 115, 8763–8768. [CrossRef]

180. Bendia, B.; Ward, L.D.; Simpson, R.J. Proteins of the Golgi apparatus. Purification to homogeneity, N-terminal sequence, and enrichment of redox-modified proteins and their applications. *Proteomics* 2014, 14, 197–213. [CrossRef] [PubMed]
189. Solda, T.; Garbi, N.; Hammerling, G.J.; Molinari, M. Consequences of ERp57 deletion on oxidative folding of obligate and facultative clients of the calnexin cycle. *J. Biol. Chem.* **2006**, *281*, 6219–6226. [CrossRef]

190. Watt, G.M.; Lund, J.; Levens, M.; Kolli, V.S.; Jefferis, R.; Boons, G.J. Site-specific glycosylation of an aglycosylated human IgG1-Fc antibody protein generates neoglycoproteins with enhanced function. *Chem. Biol.* **2003**, *10*, 807–814. [CrossRef]

191. Mathys, L.; Balzarini, J. Several N-Glycans on the HIV Envelope Glycoprotein gp120 Preferentially Locate Near Disulphide Bridges and Are Required for Efficient Infectivity and Virus Transmission. *PLoS ONE* **2015**, *10*, e0130621. [CrossRef]