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9-O acetylated gangliosides in health and disease ²

Luis Vicente Herrera-Marcos ¹ , Dil Sahali 1,2 and Mario Ollero 1,* 3

1. Univ Paris Est Créteil, INSERM, IMRB, F-94010 Créteil, France 4

2. AP-HP, Hôpitaux universitaires Henri Mondor, Service de Néphrologie, Créteil, F- 94010 France.Service 5 Néphrologie, AP-HP, Hôpital Henri Mondor, F-94010 Créteil, France 6

Review 1

 \blacksquare Correspondence: mario.ollero@inserm.fr 7

Abstract: Glycosphingolipids comprise a lipid class characterized by the presence of sugar moieties 8 attached to a ceramide backbone. The role of glycosphingolipids in pathophysiology has gained 9 relevance in the last years in parallel to the development of analytical technologies. Within this vast 10 family of molecules, gangliosides modified by acetylation represent a minority. Described for the 11 first time in the 80s, their relation to pathologies has resulted in an increased interest for their func- 12 tion in normal and diseased cells. This review presents the state of the art on 9-O acetylated gangli- 13 osides and their link to cellular disorders. 14

Keywords: glycosphingolipid; sphingolipid; acetylation; cancer; sialic acid 15

1. Discovery and chemistry 17

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Glycosphingolipids constitute a subcategory of sphingolipids in which a ceramide 18 backbone is linked to one or more sugar residues. Among glycosphingolipids, gangli- 19 osides contain at least one residue of sialic acid, anciently known as neuraminic acid (Fig- 20 ure 1). Gangliosides are subdivided according to the number of sialic acid residues, e.g. 21 monosialylated (GM), disialylated (GD), trisialylated (GT), and further classified accord- 22 ing to the number of neutral sugar residues subtracted from a maximum of 5 (e.g. GD1 23 contains 4 neutral residues, where "1" indicates 5-4=1) (Table 1). The sialic acid moiety 24 contained in the ganglioside molecule can present structural modifications, such as acet-
25 ylation. This modification can be present in other biomolecules containing sialic acid res- 26 idues, such as glycoproteins. 27

1.1. Types of acetylation and first findings in cells 30

Modifications of sialic acid were first discovered in the secreted products of subman- 31 dibular glands from cattle [1]. Those include O-glycoloyl, N-glycoloyl, O-acetyl and N- 32 acetyl forms, where glycoloyl and acetyl groups are formed by hydroxylation and acety- 33 lation of sialic acid, respectively (Figure 1). The acetyl and glycoloyl transferase activities 34 necessary to ensure these modifications were found in cytosolic and microsomal extracts 35 from these tissues [2, 3]. The O-acetyl transferase reaction conveying the acetyl group to 36 the sialic acid moiety (sialate O-acetyl transferase -SiAOAT- activity) has been recently 37 attributed to the enzyme CASD1 (CAS1 domain containing) by means of genome editing 38 approaches [4]. This acetylation can be reversed by the 9-O-acetylesterase or sialidase ac- 39 tivity (SIAE), found in several microorganisms and mammal brain tissue and resulting in 40 the release of acetyl residues (Figure 2) [5-8]. Interestingly, the presence of a 9-O acetyl 41 group in sialic acid can have an impact on the activity of sialidases, which remove sialic 42 acid from larger molecules [9]. 43

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Biomolecules **2022**, *12*, x FOR PEER REVIEW 2 of 25

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Figure 1. (Acetyl/Glycoloyl)-Ganglioside structure. A: Schematic representation of GM3 as an ex- 46 ample of ganglioside. * To note: gangliosides are a type of Glycosphingolipid but neither gangli- 47 osides nor glycosphigolipids are considered types of ceramides. Ceramide is a structural component 48 of all glycosphingolipids (including gangliosides). Sialic acid carbons are numbered as 1 to 9, start- 49 ing by the left side of the molecule. B: Different types of sialic acid modifications in mammalian 50 gangliosides mentioned in the text. N-acetylated (acetyl groups, in black, bound to the N atom) and 51 O-acetylated forms (bound to an O atom) are represented on the upper part. An N-glycoloylated (a 52 glycoloyl group, in black) bound to the N atom) and O-glycoloyl (bound to the O atom) are repre- 53 sented on the lower part. **54**

Biomolecules **2022**, *12*, x FOR PEER REVIEW 4 of 25

Figure 2. Enzymatic conversion of a ganglioside to its acetylated form and responsible enzymes 56 in human. CASD1: CAS1 domain containing (Uniprot ref. Q96PB1). SIAE: Sialate O-acetylesterase 57
(Uniprot ref. Q9HAT2). Green rectangle: sphingoid base. Blue rectangle: fatty acyl chain. Red hexa-(Uniprot ref. Q9HAT2). Green rectangle: sphingoid base. Blue rectangle: fatty acyl chain. Red hexa- 58 gons: neutral sugar residues. Purple double triangle: sialic acid residue. 59

Sialic acid O-acetylation can be present both in proteins and lipids. A membranebound acetyl-transferase activity was found associated with the modification of endogenous glycoprotein-bound sialic acids, while a soluble activity was linked to the modifica- 62 tion of exogenous, non-glycosidically bound sialic acids. This finding was further extended to brain tissue from pig and cow [10]. These first discoveries did not make the distinction between protein-bound and lipid-bound acetylated sialic acids. The first isolation of a ganglioside containing 9-O acetylation was obtained in the mouse brain, within 66 trisialo-ganglioside [11] and tetrasialo-ganglioside [12] structures. This was followed by 67 the guinea pig kidney [13], bovine buttermilk [14, 15], codfish brain [16, 17], rat and equine erythrocytes [18, 19], as well as less common species, like feather starfis[h \[20\].](https://www.zotero.org/google-docs/?pdl2yG) In rat erythrocytes, a combination of thin layer chromatography, gas chromatography, and an enzymatic treatment with *Vibrio cholerae* sialidase could identify GD1a (GD1 of the "a" series, 71 bearing 1 sialic acids on the galactose in position II; 0-, b- and c-series bearing 0, 2 and 3 respectively) (Table 1) and not GM1 as the main ganglioside containing this modification. In equine erythrocytes, NMR and fast atom bombardment mass spectrometry (FABMS) could identify 9-O-acetyl-GM3 (9-O-acGM3) [19]. In human tissue, an analysis in normal thyroid gland, resulted in the identification of a potential presence of 9-O acetyl gangliosides, defined as containing alkali-labile sialic acid [21]. Also, an antibody claimed to 77 recognize 9-O acetylated GD3 ($9P$ -O-acGD3) was able to bind normal human melanocytes [22], and so did another one isolated from melanoma cells [23]. This newly detected form was characterized by NMR and FABMS and further found in other species and tissues, such as rainbow trout, where it accounts for 23% of total gangliosides [24, 25]. Finally, an acetylated trisialylated form, 9-O-acGT2, was first identified in cod brain [16].

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1.2. Chemical structure and interactions 87

In GD1a, the N-acetylated sialic acid is linked to the outer galactose residue [26]. Conformational studies have been performed by molecular dynamics modeling and NMR on 9-O-acGD1a, concluding that acetylation does not modify the overall conformation of the ganglioside [26]. Specific interaction with a purified IgG fraction from human serum was suggested by the same study. More recently, a study on GM3 indicated that neither 9-O-acetylation nor 9-N-acetylation induce significant conformational changes on dihedral angles or the secondary structure, those being limited to the sialic acid glycerol chain and confirming structural similarities between both forms [27].

Concerning the composition in terms of sphingoid bases and acyl chains, this varies among species and no particular association with 9-O acetylation can be inferred from the scarce data available. Studies made on bovine buttermilk O-acetylated gangliosides have revealed C18-sphingosine as the sphingoid base and C18:0, C22:0, C23:0 and C24:0 as the main fatty acyl chains [15]. In rainbow trout ovarian fluid the structure differs, as it contains 4-sphingenine as sphingoid base, and C24:1 among fatty acids [25]. In another fish, mullet milt, 9-O-acGM3 is majoritarian, containing mostly C18:1/C16:0 fatty acids [28]. In feather starfish, C16 sphingosine is accompanied by C22:0 or C24:0 as the most common acyl chains [20].

1.3. Enzyme regulation

Sialic acid O-acetylation appears as a cell specific and developmentally regulated process. This is based on a tightly regulated activity of 9-O-acetyltransferases. Pioneering 107 studies indicate that sialyltransferase action regulates the expression of O-acyltransferases [29]. Cloning of this sialyltransferase (sialate-O-acetyltransferase, CASD1) was an elusive task. In one of the attempts, an open reading frame corresponding to a truncated form of the GC Vitamin D binding protein (VDBP) was found specifically responsible for sialic acid 9-O-acetylation of glycoproteins, while a fusion protein between a bacterial tetracycline resistance gene repressor and a sequence of the \overrightarrow{P} 3 plasmid (Tetrfusion) was able to acetylate gangliosides [30]. An interesting observation is that the product of O-acetylation makes the sialic acid moiety resistant to sialidase [31], which could have functional implications. Also, the natural forms of acetylated GD3 -a disialylated ganglioside- present the modification at the terminal sialic acid moiety, as compared to synthetic forms [32]. In another study it was shown that O-acetyltransferases use preferentially di- and tri-sialo- 118

gangliosides as substrates rather than mono-sialogangliosides [33]. Acetyltransferase ac- 119 tivity on GD3 (9-O-acGD3) is unchanged by the endoplasmic reticulum-to-Golgi transfer 120 stimulator brefeldin A, suggesting that the activity resides in the same Golgi compartment 121 as GD3 synthase, which is not the case for 9-O-acGD2 synthesis [34]. This suggests differ- 122 ent compartments and potentially different enzymes for GD3 and GD2 modification. Nev- 123 ertheless, 9-O-acGD2 can be synthesized either from GD2 by acetylation or from 9-O- 124 acGD3 by glycosylation. It must be noted that biosynthesis of 9-O-acetylated gangliosides 125 requires a transfer of the acetyl group from Acetyl-CoA. The Acatn acetyl-CoA transporter 126 was identified in mice as intervening in this process, and being mainly expressed during 127 embryogenesis [35]. 128

9-O-acetylation of GD3 has been proposed to be induced in Chinese hamster ovary 129 (CHO) cells by stable expression of its precursor, GD3, through activation of the *Tis21* 130 gene [36]. Moreover, when cells are incubated in the presence of exogenous GD3, cellular 131 9-O-acGD3 is detected after 6h and a half-life of 24h is observed, suggesting the induction 132 of the biosynthetic enzymatic machinery. This process, also reported in human fibroblasts, 133 is inhibited by blocking clathrin-mediated internalization of GD3 [37]. Conversely, Tis21 134 does not seem to be involved in the upregulation of 9-O-acGD3 synthesis that occurs in a 135 GM2/GD2 synthase knockout mouse model to compensate for the lack of complex gan- 136 gliosides [38]. In this model Vitamin D receptor and acetyl CoA transporter are not up- 137 regulated, suggesting an alternative mechanism of synthesis. 138

Reports on pharmacological agents exerting an impact on these synthesis reactions 139 are scarce. In one of the few examples, it has been shown that salicylate leads to deacety- 140 lation of gangliosides [39]. Also, cytidinmonophosphate-sialic acid and acetyl-CoA inhibit 141 *in vitro* sialyl transferase activity [40]. 142

In addition to enzyme activity, the regulation of enzyme expression must be consid- 143 ered. To date, no precise regulatory mechanisms for CASD1 or SIAE expression based on 144 experimental evidence have been published. Nevertheless, their promoters are defined in 145 the Ensembl database and several transcription factor binding sites have been confirmed 146 in numerous cell lines by ChIP-seq within the ENCODE project (Tables S1 and S2). In 147 addition, both promoters contain a CpG island (108 CpG in the CASD1 promoter and 50 148 CpG in the SIAE promoter) (figure S1). Interestingly, SIAE mRNA transcriptional variant 149 2 sequence starts upstream its CpG island, maybe as part of a mechanism to avoid silenc- 150 ing by methylation. Although the regulatory landscape of these two genes currently re- 151 mains unknown, according to Protein Atlas endocrine tissues present the highest CASD1 152 mRNA expression, followed by eye and digestive tract, while the protein has been found 153 in high abundance also in brain, pancreas, reproductive tissues, bone marrow and lym- 154 phoid tissues (https://www.proteinatlas.org/ENSG00000127995-CASD1/tissue). SIAE 155 mRNA shows a highest expression level in the gastrointestinal tract, while the highest 156 protein expression corresponds to brain, endocrine tissue, urinary system, male tissues 157 bone marrow and lymphoid tissues (https://www.proteinat- 158 las.org/ENSG00000110013-SIAE/tissue). 159

1.4. Methodological points 162

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The early studies and many of the follow up works have been based on the detection 163 of this type of modified gangliosides by monoclonal antibodies in combination with thin 164 layer chromatography (TLC) or immunohistochemistry (IHC). The so-called JONES, VIM- 165 2 [41], 13A and 27A [42], UM4D4 [43], CDW60 [44] and MT6004 [45] antibodies have been 166 shown to detect 9-O-acGD3, while the SGR37 monoclonal antibody detects distinctly the 167 de-N-acetyl form of GD3 [46]. It must be pointed out, though, that targeting lipid antigens 168 in IHC can be seriously impacted by the use of organic solvents for fixation and deparaf- 169 fination, such as acetone and xylol respectively. Special care must be taken, as an incorrect 170 fixation protocol is likely to induce artifactual results [47]. 171

Specific binding of Influenza C virus has also been considered as the basis of detec- 172 tion methods. This microorganism presents a higher affinity for 9-O-ac and a lower affin- 173 ity for 7-O-ac glycoconjugates [48, 49], regardless of the nature of the core moiety (lipid or 174 protein). Virus binding is also able to discriminate monoacetylated sialic acids from poly- 175 acetylated [48]. As a consequence, recombinant soluble influenza C hemagglutinin has 176 been used to characterize 9-O-acetyl sialylation [50]. Other molecules recognizing 9-acet- 177 ylated sialic acid and displaying a specificity for gangliosides are monocyte ficolins, 178 highly conserved oligomeric lectins involved in innate immunity [51]. 179

As explained above, chemical characterization has been mainly based on NMR and 180 FABMS. Finally, the evaluation of sialyl transferase and SIAE enzymatic activities have 181 added a functional dimension to some studies [52]. 182

2.1. In cell physiology 184

2.1.1. Embryogenesis 185

Human embryonic stem cells present a high abundance of 9-O-acGD3 that generally 186 decreases along differentiation [53, 54]. A particular type of cancer cells (NTERA-2, a hu- 187 man embryonic carcinoma line) has been used to study the ontogeny of glycolipids in 188 association with cell differentiation during embryonic development. In this model, gan- 189 glio-series, including 9-O-ac forms, replaced globo-series (glycosphingolipids containing 190 at least two neutral sugar residues and no sialic acid) when differentiation was induced 191 with retinoic acid [55]. 192

These molecules have been mainly studied in the context of nervous system devel- 193 opment. In particular, the presence of 9-O-acGD3 has been shown in neuroepithelial pre- 194 cursor cells [56]. An antigen expressed during neural development was identified as 9-O- 195 acGD3 [57]. In rat developing retina the pattern of 9-O-acGD3 and that of its precursor 196 GD3 were determined by the reactivity to several monoclonal antibodies (JONES, R24). 197 The two patterns differed, in the case of the 9-O acetylated form a rise was found between 198 day E15 and postnatal day 2, with a pronounced drop between day 2 and day 4 PN [58]. 199 9-O-acGD3 has also been found in primary cultures of both neurons and glia [\(reviewed](https://www.zotero.org/google-docs/?RzmRwZ) 200 in [\[59\]\).](https://www.zotero.org/google-docs/?RzmRwZ) In freshly dissociated retinal cells 9-O-acGD3 was found present on amacrine pho- 201 toreceptor and in ganglion cells [58]. In chick embryo, a monoclonal antibody (8A2) al- 202 lowed detecting 9-O-ac gangliosides in the optic fiber layer of central retina [60]. Another 203 study based on a monoclonal antibody staining and on sialidase sensitivity concluded that 204 a 9-O-ac form of GT3 (ganglioside C series) is also increased in rat cerebral cortex at day 205 14 of gestation, then progressively decreased and absent in adult rats [61], along with its 206 9-O-acGD3 counterpart [62]. 207

In the developing rat nervous system, acetylated gangliosides have been associated 208 with regions characterized by cell migration [63], such as the olfactory epithelium, where 209 they are involved in the formation of the mature olfactory bulb [64] and the hippocampus 210 [65]. They were detected in relation to the cell stream migrating from the lateral ventricle 211 rostral subventricular zone to the olfactory bulb, suggesting a function in cell migration 212 [66]. These gangliosides were also isolated from 10-day embryonic chicken brain [67]. 213 Concerning their cellular function, there is evidence that 9-O acetylated gangliosides play 214 a role in the extension of growth cones in neurites [68], along with a regulation of micro- 215 filament and microtubular structure of their cytoskeleton, probably modulating cell mo- 216 tility [69]. The same authors found 9-O-acGD3 localized to contact points of neural growth 217 cones, associated with beta-1-integrin and vinculin [70]. 218

The functional relevance during embryogenesis of 9-O acetylation of sialic acid was 219 studied by the generation of a transgenic mouse model overexpressing the sialic acid-spe- 220 cific acetylesterase of Influenza C virus under the control of the metallothionein promoter 221 [71]. This resulted in an arrest of development at the 2-cell stage. Using the phenylethan- 222 olamine-N-methyltransferase promoter, the authors induced expression in retina and ad- 223 renal gland, leading to impaired morphology and function of these organs. 224

2.1.2. Post-natal nervous system 225

The nervous system is generally rich in gangliosides, including 9-O-acGD3. In a 226 mouse model constitutively knocked out for GM2/GD2 synthase, the lack of complex gan- 227 gliosides is compensated by an accumulation of the precursors, namely GM3 and GD3, in 228 nervous tissue [72]. This accumulation also includes 9-O-acGD3, suggesting that this mol- 229 ecule can take over some of the functions of the absent glycosphingolipids [38]. In post- 230 natal rat retina a dorsal-ventral gradient of 9-O-acGD3 has been reported, an observation 231 based on the JONES monoclonal antibody [58], as well as in the adult olfactory bulb, but 232 at lower levels than in the developing nervous system [66]. In the chicken, 9-O acetylated 233 gangliosides were no longer detected in the adult in the central optic fiber. In contrast, 234 they would remain in the inner and outer plexiform layer, and in the outer nuclear layer 235 [60]. Likewise, 9-O-ac gangliosides have been found absent in rat adult hippocampus [65]. 236 In primary cell cultures from retina, they are present in the retinal ganglion but not in 237 Muller cells [60]. In the rat subventricular zone the presence of 9-O-acGD3 has been 238 demonstrated from neural stem and progenitor cells to the adult brain [73]. To add insight 239 on the subcellular distribution of these molecules, in olfactory ensheathing glia from rat, 240 9-O-acGD3 has been identified in membrane rafts [74]. 241

With respect to the potential function of these molecules in the nervous system, in 242 cerebellar astroglia isolated from rats, JONES staining was found in the contact sites of 243 migrating granule cells and in radial glia when cultured in the presence of neurons [63] 244 [75]. Another study suggested a role in the regulation of both neuronophilic and gliophilic 245 migration [76]. The staining is also present in neurons and glia involved in axonal regen- 246 eration of sciatic nerve in adult rats [77], which is defective in GD3 synthase knockout 247 mice [78]. The same antibody blocks migration in a dose-dependent manner, adding up 248 evidence to the participation of 9-O-acetyl gangliosides in granule cell migration [75, 79] 249 through a calcium signaling mechanism involving PY2 receptors [80]. Anti-9-O-aAcGD3 250 antibody-based inhibition of olfactory ensheathing glia migration has been observed in 251 organotypical cultures [81]; inhibition of neuronal migration has been shown *in vivo* in 252 normal mice [82, 83], and confirmed by videomicroscopy [84], while migration was also 253 blocked by a broad inhibitor of ganglioside synthesis (D-threo-1-phenyl-2-pal- 254 mitoylamino-3-pyrrolidino-1-propanol, inhibitor of the ganglioside precursor gluco- 255 sylceramide) [84]. However, the fact that antibody-based inhibition also occurs in GD3 256 synthase knockout mice, which are not supposed to contain the acetylated derivative, sug- 257 gests that the antibody inhibits migration through an alternative mechanism, while it also 258 raises questions on its specificity [83]. Nevertheless, sciatic regeneration is perturbed in 259 this mouse model and rescued by administration of exogenous GD3, which supports a 260 genuine role for downstream generated gangliosides [78]. 261

9-O-acetylated glycolipids have been detected in mammalian cerebellar Purkinje 262 cells [85], where they occupy the rostral lobes in mice [86]. They mostly mark the late onset 263 sagital banding patterns [87]. Interestingly, in the so-called nervous mutation model of 264 mouse Purkinje cells, the surviving mutant cells in the cerebellum correspond to those 265 positive for 9-O-acetylated gangliosides [86], mainly corresponding to 9-O-acGD3 [88]. 266

2.1.3. Immune system 267

Some glycolipid antigens at the surface of T lymphocytes were initially recognized 268 by monoclonal antibodies and defined as CDw60. These molecules have been shown to 269

induce costimulatory signals. The CDw60 antigen, recognized also by Influenza C virus 270 glycoprotein, was characterized as 9-O-acGD3 [89]. T lymphocytes (mostly CD4⁺) and 271 granulocytes present high amounts of this CD60 antigen, in contrast to the low levels pre- 272 sent in B cells, thymus cells and monocytes [90]. It was estimated that about 25% of pe- 273 ripheral T cells present a surface localization of CD60, while roughly all T cells express 274 modest amounts intracellularly in Golgi vesicles [91]. In an early report, a subtype of CD8+ 275 T cells, expressing also CD60 A – so called a T helper CD8+ CD60+ subset- of T cells was 276 rized as T help CD8⁺₇ was claimed to provideing help to B cells, while CD8⁺ 277 CD60- suppressed B cell differentiation. Both populations produced equally IL-2, but 278 CD60⁺ would secrete more IL-4 and less interferon gamma [92]. In spite of the low levels 279 initially reported, CD60 has been proposed as an activation marker of human B cells, as 280 peripheral and tonsillar B cells become CD60⁺ when activated by phorbol esters [93]. It 281 must be pointed out that another acetylated form of GD3, 7-O-acGD3, was also found in 282 human leukocytes, recognized by a specific monoclonal antibody that induced cell prolif- 283 eration [94]. T-cell receptor (TCR) activation results in decreased presence of detectable 9- 284 O-acetyl sialic acid at the surface of T cells, but this is mostly due to decreased sialomucins, 285 which also contain this residue, and not necessarily to gangliosides [50]. In peripheral 286 blood mononuclear cells (PBMC), treatment with a monoclonal antibody targeting 9-O- 287 acGD3, but not with another one against non-acetylated GD3, was able to induce phos- 288 phorylation of the spleen tyrosine kinase (Syk, p72), involved in T and B cell receptor sig- 289 nal transduction, resulting in phosphoinositide mobilization and cell proliferation [95]. 290

Following subsequent studies, CD60 was subdivided into CD60a (GD3), CD60b (O- 291 acetylated form), and CD60c (N-acetylated form) [96]. The CD60b form was found present 292 in tonsillar B cells in the activated germinal center, colocalizing in lipid rafts with Syk and 293 Lyn, in line with previous results [93, 95]. Hence, B cells can be costimulated by anti- 294 CD60b and anti-IgM/IL-4. Extrafollicular T cells also present with CD60b and can be co- 295 stimulated with anti-CD60 and phytohemagglutinin (PHA). Conversely, anti CD60c -rec- 296 ognizing the N-acetylated form- has been found sufficient to induce proliferation [96]. In 297 a thorough study on the presence of the three CD60 forms during differentiation of T cells 298 and B cells, CD4+ cells showed the strongest and CD8+ cells the weakest presence of CD60b 299 at the surface in thymocytes. Both T and B cells presented a CD60b staining in a patchy 300 fashion as compared to the other forms. Interestingly, subcellular distribution studies fol- 301 lowing biochemical methods showed 9-O-acGD3 mainly localized to non-raft microdo- 302 mains in T cells and to raft microdomains in B cells [45]. 303

2.1.4. Hematopoiesis 304

In human bone marrow, erythroid progenitors are rich in 9-O-acGD3, but the mole- 305 cule is progressively lost during maturation, becoming proapoptotic in mature erythro- 306 cytes [97]. The presence of 9-O-acGD3 in lymphoid and erythroid cells is reviewed in [98]. 307

2.1.5. Kidney 308

Cultured visceral glomerular epithelial cells -podocytes- contain the specific epitope 309 9-O-acGD3 recognized by several monoclonal antibodies, such as 13A and 27A. The latter 310 could immunoprecipitate with a non-characterized podocyte protein [42]. This epitope 311 was found by the 27A antibody to colocalize in podocyte lipid rafts with nephrin, a protein 312 present in the slit diaphragm, a structure responsible for the podocyte intercellular inter- 313 action and a main constituent of the glomerular filtration barrier. These seminal works 314 indicate the importance of this modified ganglioside in the physiology and the function 315 of the glomerular barrier [99]. 316

2.2. In cell pathology – diseases 317 2.2.1. Cancer 318

9-O-acetylation of gangliosides has been extensively associated with cancer, and even 319 considered as a marker of cell and tissue growth [100]. Very early studies on melanoma 320 cells found in extracts a thin layer chromatography band comigrating with 9-O-acetylated 321 gangliosides [101]. It was estimated that 10% of gangliosides in melanoma cells presented 322 this modification. These modified sialic acids, independently of their associated moiety - 323 either protein or sphingolipid-, were recognized by a monoclonal antibody prepared 324 against the rat brain tumor cell line B49. In another study, chromatographic comigration 325 with GD3 was found in cell extracts after isolation with a monoclonal antibody derived 326 from immunization of mice with WM164 melanoma cells [23]. It was estimated that all 327 nevus cell lines and one third of melanoma cell lines were positive to an antibody detect- 328 ing this modification, which was also found in lymphocytes infiltrating 30% of tumors. 9- 329 O-acGD3 has been ever since considered as a melanoma antigen [57, 102-104], as was 9- 330 O-acGD2 [105]. When evaluating different stages of Bomirski melanomas, 9-O-acGD3 was 331 found increased in the amelanotic, fast growing stage, as compared with the slow grow- 332 ing, highly differentiated forms [106], suggesting a role for the molecule in cell growth. Its 333 presence in nodular melanoma has been found greater than in metastatic acral lentiginous 334 melanoma [107]. However, it has not been found present in uveal melanoma [108, 109], 335 which may indicate that the acetylated varieties are characteristic of metastatic forms (cu- 336 taneous) as compared with non-metastatic (uveal). Interestingly, while other gangliosides, 337 such as GD2 and GD3, have been found increased in the serum of melanoma patients, this 338 is not the case of 9-O-acGD3 [110]. 339

In hamster melanoma, the O-acetylated form of GD3 was characterized as 7-O in- 340 stead of the human 9-O. The structure of the former is not very different from that of 341 buttermilk ganglioside, as it contains C18:0 sphingosine and a slightly different fatty acid 342 composition: C16:0, C18:0, C20:0, C22:0 and C24:0 [111]. In human melanoma a quite high 343 presence of C24:1 has been reported in both the 9-O-acGD3 and the GD3 precursor [23, 344 112]. Melanoma cells also display de-N-acGD3, with an intracellular and non-lysosomal 345 distribution [46]. In this case the main esterifying fatty acids are C16:0 and C18:0 [112]. 346

In mouse erythroleukemia cells 9-O-acGD3 is also present, but not detectable at the 347 surface, where 9-O-acetyl sialic acid is associated with sialomucins [113]. In lymphoblasts 348 from acute lymphoblastic leukemia patients' 9-O-acGD3 levels are increased [114]. An in- 349 creased SIAOAT enzymatic activity was detected in the microsomes of these cells. The 350 activity was found higher at diagnosis and decreased in remission, whereas SIAE activity 351 is down in the cytosol and in lysosomes [40, 52]. In Sézary syndrome, a very aggressive 352 leukemic form of cutaneous T cell lymphoma, circulating levels of CD60b (9-O-acGD3) 353 positive T cells were found associated with a poor prognosis [115]. 354

9-O acGD3, along with other gangliosides, has been proposed as a marker of several 355 neuroectodermal cancers. For example, it was detected in basal cell carcinoma cells and 356 found dramatically increased as compared to normal epidermis or dermis [116, 117]. It 357 has been suggested as a marker of small cell lung cancer [118]. Studies in breast tissue 358 have demonstrated the presence of CD60 antigen in the Golgi apparatus of normal ductal 359 cells, and increased in atypical hyperplasia and other benign lesions, as well as in mam- 360 mary carcinoma cells [119]. In well differentiated and invasive duct carcinoma the antigen, 361 identified as 9-O-acGD3, was found mostly present at the surface, with decreased pres- 362 ence in non-differentiated carcinomas [119]. In some breast cancer cell lines (Hs 578T and 363 SUM159PT) 9-O-acGD2 and not 9-O-acGD3 has been identified [120], and CASD1 demon- 364 strated as the enzyme responsible for its synthesis [121]. Both GD3 and 9-O-acGD3 were 365 detected and increased in 13 neural tumor cell lines [122], and in glioblastoma, where a 366 critical ratio between the two forms promoting tumor survival was established [123]. As 367 a consequence of all these findings, the presence of acetylated gangliosides in blood as 368 cancer biomarkers has been considered and specific testing by liquid chromatography- 369 mass spectrometry on dry blood samples has been developed [124]. 370

The link between 9-O-acetylation of gangliosides and cancer is underlined by its ef- 371 fect on apoptosis. GD3 is considered as a proapoptotic agent, at least *in vitro*, while its 9- 372 O acetylated form is shown as antiapoptotic [39, 125, 126]. The presence of 9-O-acGD3 in 373 Jurkat and Molt-4 cells prevents cell death induced by proapoptotic agents such as N- 374 acetyl sphingosine and daunorubicin [39]. Lymphoblasts from lymphoblastic leukemia 375 patients accumulate 9-O-acGD3 in mitochondrial membranes [114]. Unlike GD3, exoge- 376 nous 9-O-acGD3 prevents mitochondrial membrane depolarization, cytochrome C release 377 and caspase activation in lymphoblasts [114]. Interestingly, 9-O-acGD1, also known as 378 neurostatin, has antiproliferative effects on astrocytoma cells [127] and synthetic forms 379 have been produced and approved as anticancer drugs [128]. The potential regulation of 380 apoptosis by acetylated gangliosides (CD60) has been addressed in lymphocytes [96]. 381 However, a hematopoiesis study conducted on human bone marrow revealed a proapop- 382 totic impact of 9-O-acGD3 on mature erythrocytes, in contrast to its effect on lymphoblasts 383 [97]. 384

9-O acetyl-GD3 was consequently proposed as a potential target for immunotherapy 385 [129, 130]. The antibody response to injection in melanoma patients of 9-O-acGD3 ex- 386 tracted from buttermilk was studied, but the reactivity was not found antigen specific 387 [131], which underlies the problem of the low immunogenicity of the molecule. This was 388 improved by combining the antigen with very low density lipoproteins and enhanced by 389 IL-2, which could be used as adjuvants [132]. 9-N-acGD2, used as a stable surrogate of 9- 390 O-acGD2, has been also used as antigen, in this case conjugated with the carrier bacteriophage Qbeta, eliciting a strong and long lasting immune response in dog [133]. Interest- 392 ingly, a high titer of anti-9-O-acGD3 antibodies has been found in the serum of medullo- 393 blastoma patients [122]. Finally, in glioblastoma cells, several strategies based on hemag- 394 glutinin esterase cleavage of the acetyl group have been explored [123]. 395

2.2.2. Infection 396

Influenza C virus is known to infect cells through binding to N-acetyl-9-O-acetyl si- 397 alic acid, an ability that is shared with bovine coronavirus [134-135]. Treatment of cells 398 with 9-O acetylesterase confer resistance to infection, which is reversed by treating cells 399 with ganglioside preparations from bovine brain containing 9-O acetylated forms, sug- 400 gesting 9-O-acetylated gangliosides as potential receptors for this pathogen [5]. Binding 401 to 9-O-acGD1a has been demonstrated [136]. Conversely, influenza C virus is able to 402 slowly hydrolyze in vitro 9-O-acGD1a [7] and 9-O-acGT3 [137], since the hemagglutinin 403 encoded by the viral genome possesses a 9-O-acetyl sialic acid-specific acetyl esterase ac- 404 tivity [71]. Another pathogen, *Mycobacterium leprae*, invades Schwann cells with the help 405 of endogenous 9-O-acGD3, which is also upregulated upon infection. Immunoblocking of 406 the ganglioside reduces the demyelinization effect of the bacterium [138]. 407

2.2.3. Autoimmune diseases 408

9-O-acGD1b has been associated with Guillain-Barré syndrome, an autoimmune dis- 409 order characterized by the presence of anti-glycolipid antibodies in blood. The serum of a 410 subset of patients reacts with this modified ganglioside, along with the non-acetylated 411 form and with GM1, as found by ELISA and thin layer chromatography immunostaining 412

[139]. 413 Psoriatic basal and suprabasal keratinocytes express 9-O-acGD3 at the surface, and 414 the extent of expression is increased when these cells are subjected to material secreted by 415 T cells isolated from the same lesions, suggesting that soluble factors secreted by T cells 416 are responsible for this effect. In the same context, IL-4 and IL-13 induced upregulation 417 and interferon-gamma downregulation of the ganglioside, while the upregulation effect 418 was reduced by an anti-IL-13 antibody [43]. 419

2.2.4. Toxicology 420

Lead exposure has been associated with increased detection of several gangliosides 421 in kidney, including 9-O-acGD3 in glomeruli, using monoclonal antibodies and confirmed 422 by thin layer chromatography [44]. This was suggested by the authors of the work to con- 423 stitute a marker of lead exposure and to be associated with a dysregulation of apoptosis, 424 in that high levels of 9-O-acGD3 in glomeruli were correlated with a lower number of 425 apoptotic cells in the kidney. 426

3. Concluding remarks: from controversy to future prospects 427

The fact that detection systems target the acetylated sialic acid moiety, present in both 428 gangliosides and glycoproteins, leads to ambiguous interpretation of many results in the 429 absence of further biochemical characterization. Thus, a thorough study on the expression 430 of CD60 antigen in T cells and melanoma cells led to the conclusion that it corresponds 431 mostly to a glycoprotein marker in the former and a glycolipid in the latter [140]. Another 432 example of this ambiguity is the reported recognition by the JONES antibody of β1-integ- 433 rin in mouse cerebellum [83], which compromises some conclusions based on this partic- 434 ular tool. Considering these constraints, mass spectrometry reveals as the most reliable 435 approach to search for the distribution and biological effects of 9-O-acetylated gangli- 436 osides. 437

Some points raised by previous works will need to be clarified, while others are yet 438 unexplored. For example, a basic question is the relationship between cell cycle and 9-O- 439 acetylated ganglioside synthesis. Another one is the subcellular distribution of these mol- 440 ecules. Previous studies have shown their presence in mitochondria, at the plasma mem- 441 brane surface in and out of raft-like membrane microdomains, yet to date little is known 442 about their function in these compartments. Conversely, their presence in the nucleus has 443 not been explored. 444

Regarding the likely abundance of 9-O-ac gangliosides in membrane raft-like micro- 445 domains, it could be hypothesized a potential function as entry points to viral particles. It
has been shown that the sialic acid moieties of gangliosides, by means of their negative has been shown that the sialic acid moieties of gangliosides, by means of their negative charge, determine the electrostatic potential and thereby impact the interaction of viruses, 448 such as SARS-CoV-2 with host cells [141, $\frac{1}{142}$]. Interestingly, SARS-CoV-2 spike protein 449 binds preferentially to 9-N-ac and 9-O-ac sialic acid [143]. It is tempting to hypothesize 450 that 9-O-acetylation of gangliosides changes the dynamics of virus-raft interaction and 451 eventually virus entry. Whether this is the case, and whether the mechanism involves a 452 receptor like or a change in electrostatic interaction remains to be clarified. 453

While a reasonable body of knowledge has been gathered for 9-O-acetylated gangli- 454 oside in the context of cancer, an aspect that has been insufficiently addressed is their 455 implications in other pathologies, especially those accounting with alterations in lipid me- 456 tabolism -i.e. cardiovascular disease, type 2 diabetes mellitus, non alcoholic fatty liver dis- 457 ease- or lipid storage disorders. Likewise, the presence of 9-O-acetylated gangliosides in 458 circulating macromolecular structures, such as lipoproteins or extracellular vesicles is cur- 459 rently unexplored (apart from the enhanced immunogenicity of 9-O-acGD3 when ad- 460 sorbed onto very low density lipoproteins [132]). 461

Finally, in light of the available data summarized in this review (Table $2+$), a question 462 arises on the levels of 9-O-ac gangliosides found in physiological and pathological condi- 463 tions. As suggested by several studies, these molecules play a key role in cell survival and 464 cell mobility. These two properties are relevant to cancer cells to avoid immune defense 465 mechanisms and to propagate throughout the body. This would explain why some 9-O-
ac gangliosides are overabundant in cancer cells, hereby displaying a potential as cancer ac gangliosides are overabundant in cancer cells, hereby displaying a potential as cancer biomarkers. Nevertheless, these roles are also important in other cells in physiological 468 conditions. Consequently, 9-O-ac gangliosides are not exclusive of cancer cells and their 469 role as cancer biomarkers can be contested. For example, melanocytes increase their 9-O- 470 acGD3 content during carcinogenesis. However, other cells in physiological conditions 471 (e.g. podocytes, neuroblast cells and lymphocytes) have been proven to contain the same 472 molecule, which represents somehow a paradox. It can be hypothesized that their physi- 473 ological/pathological role in cells depends on a combination of at least two parameters, 474 namely abundance (as shown in [104]) and subcellular location. An additional parameter 475

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would be the ratio between 9-O-ac and non-acetylated counterparts [39, 125, 126], or be- 476 tween different types of acetylated forms (i.e. 9-O-ac, 7-O-ac, and N-ac). Even the fatty 477 acyl chain esterifying the ceramide moiety play a part [20, 25, 28]. This requires a global 478 analysis of all ganglioside forms, and further underlines the importance of mass spectrom-479 analysis of all ganglioside forms, and further underlines the importance of mass spectrometry-based methods. 480

In conclusion, the results so far point towards a relevant role of 9-O-ac gangliosides 481 nany tissues and cellular mechanisms. Nevertheless, the available information is in many tissues and cellular mechanisms. Nevertheless, the available information is 482 highly fragmented and further systematic research will be necessary to pursue the under-483 highly fragmented and further systematic research will be necessary to pursue the understanding of this fascinating puzzle. 484

Table 21. Synthesis of reported observations involving different 9-O-acetyl gangliosides in physio-
logical and pathological conditions logical and pathological conditions.

Biomolecules **2022**, *12*, x FOR PEER REVIEW 14 of 25

istration of exogenous GD3

Biomolecules **2022**, *12*, x FOR PEER REVIEW 15 of 25

IHC/TLC + Antibody (P-path): 9- O-ac glycolipids (9-OacGD3 and 9-O-acLDI)

Biomolecules **2022**, *12*, x FOR PEER REVIEW 16 of 25

Biomolecules **2022**, *12*, x FOR PEER REVIEW 17 of 25

Biomolecules **2022**, *12*, x FOR PEER REVIEW 18 of 25

Biomolecules **2022**, *12*, x FOR PEER REVIEW 19 of 25

* CHE-FcD = Hemaglutinin Esterase of Influenzavirus C fused to the carboxyl end with human IgG1 Fc region treated with diiso- 487 propylfluorophosphate to eradicate its estetase activity. CM: Confocal microscopy. FAB-MS: Fast atom bombardment mass spec- 488 trometry. IEM:Immuno Electron Microscopy. IF: Immunofluorescence. IHC:Immunohistochimestry. IP:Immunoprecipitation. 489 TLC: Thin layer Chromatography. 490

Abbreviations 491

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861