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² Characterisation of *Porphyromonas gingivalis* sialidase ³ and disruption of its role in host-pathogen interactions

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20 Abstract

- 21 Key to onset and progression of periodontitis is a complex relationship between oral bacteria and
- 22 the host. The organisms most associated with severe periodontitis are the periodontal pathogens of
- 23 the red complex; Tannerella forsythia, Treponema denticola, and Porphyromonas gingivalis. These
- 24 organisms express sialidases, which cleave sialic acid from host glycoproteins, and contribute to
- 25 disease through various mechanisms. Here, we expressed and purified recombinant *P. gingivalis*
- 26 sialidase SiaPG (PG_0352) and characterised its activity on a number of substrates, including host
- 27 sialoglycoproteins and highlighting inability to cleave diacetylated sialic acids a phenomenon
- 28 overcome by the NanS sialate-esterase from *T. forsythia*. Indeed SiaPG required NanS to maximise
- 29 sialic acid harvesting from heavily O-acetylated substrates such as bovine salivary mucin, hinting at
- 30 the possibility of interspecies cooperation in sialic acid release from host sources by these members
- of the oral microbiota. Activity of SiaPG and *P. gingivalis* was inhibited using the commercially
- 32 available chemotherapeutic zanamivir, indicating its potential as a virulence inhibitor, which also
- inhibited sialic acid release from mucin, and was capable of inhibiting biofilm formation of *P*.
- 34 gingivalis on oral glycoprotein sources. Zanamivir also inhibited attachment and invasion of oral

- 35 epithelial cells by *P. gingivalis* and other periodontal pathogens, both in monospecies but also in
- 36 multispecies infection experiments, indicating potential to suppress host-pathogen interactions of a
- 37 mixed microbial community. This study broadens our understanding of the multifarious roles of
- 38 bacterial sialidases in virulence, and indicates that their inhibition with chemotherapeutics could be
- 39 a promising strategy for periodontitis therapy.

40 Introduction

41 The black-pigmented, Gram negative, asaccharolytic anaerobe *Porphyromonas gingivalis* has been 42 the focus of much research due to its role in periodontitis where it is often referred to as a 43 "keystone pathogen" [1]. P. gingivalis exists as a small part of the microbiome, which sees a 44 population shift to include a greater proportion of certain bacterial species, a state now referred to 45 as dysbiosis [2]. In periodontal disease this population shift was observed in the 1990's, with certain 46 organisms considered particularly important to periodontal disease. Tannerella forsythia, 47 Porphyromonas gingivalis, and Treponema denticola form the so-called red complex which is 48 strongly associated with disease [3], and with the advent of next generation sequencing their 49 presence as a signature of periodontal disease has been well established, along with other 50 organisms which seem to play a causative role in periodontitis [2, 4]. During treatment of 51 periodontitis systemic antibiotics such as azithromycin can be administered as an adjunct to 52 mechanical removal of the subgingival plaque [5]. Despite this a proportion of patients poorly 53 respond to periodontal therapy, and in these patients (post-treatment) the levels of periodontal 54 pathogens are higher than in patients who respond to treatment, or in healthy controls [6]. This 55 difficulty in treatment has led to increased interest in development of an "anti-virulence approach" 56 to target the virulence factors of specific pathogens or groups of pathogens, including those 57 responsible for periodontitis. This has been seen in the case of the gingipains of *P. gingivalis*, with 58 several plant extracts [7, 8], and short peptides [9] shown to reduce gingipain activity, with in vitro 59 studies showing reductions in *P. gingivalis* biofilm formation [8], and disruption of bacterial nutrient 60 acquisition leading to decreased *P. gingivalis* growth [9]. Plant extracts that inhibit gingipains have 61 also shown disruption of host-P. gingivalis interactions; attachment and invasion of host cells, and 62 cytokine production [7]. Moreover, a recent study showed that treatment with a potent small 63 molecule gingipain inhibitor in mice reduced *P. gingivalis* infection in the brain and further blocked 64 A β_{1-42} production and neurodegenerative Alzheimer disease pathology [10].

65 Another virulence factor of periodontal pathogens including *P. gingivalis* are sialidases (also called 66 neuraminidases), enzymes frequently expressed by host-dwelling organisms which cleave terminal 67 sialic acid-a family of 9-carbon sugars-from host sialoglycans. In humans the most common sialic acid is N-acetylneuraminic acid (Neu5Ac), and is usually linked by its second carbon to the third or sixth 68 69 carbon of the underlying sugar (α 2-3 or α 2-6 linked sialic acid). Sialidases are well established 70 virulence factors in the gut and nasopharyngeal pathogens Vibrio cholerae and Streptococcus 71 pneumoniae, where they contribute to colonisation, persistence, and ultimately disease [11, 12]. 72 Interestingly, all three periodontal red complex pathogens express a sialidase; PG_0352 (termed 73 SiaPG here) in P. gingivalis, NanH in T. forsythia, and TDE0471 in T. denticola. Other periodontal

74 pathogens also possess mechanisms to process sialic acid in the absence of sialidase expression, 75 such as some Fusobacterium nucleatum spp., which possess neuraminate lyase for catabolism of 76 sialic acid [13]. It is thought that acquisition of sialic acid by organisms that do not possess sialidases 77 is due to the overall action of the microbial community of which they are part of. Additionally, 78 periodontal organisms tentatively associated with health such as *Veillonella* spp. [14] and the 79 recently characterised Tannerella HOT (oral taxon BU063) [15] do not possess sialidases. The 80 apparent importance of sialidases for periodontal pathogens has been investigated in studies of 81 sialidase knockout. T. forsythia nanH sialidase mutants displayed decreased attachment to oral 82 epithelial cells compared to their parent strain [16]. Different strains of P. gingivalis have also been 83 made deficient in SiaPG [17, 18], although the exact implications for virulence are less clear because 84 the sialidase mutants display defects in capsule formation. Mutant strains do show decreased 85 association with host cells, while the virulence of *AsiaPG* mutants in a mouse abscess model is also 86 decreased relative to the parent strain, where subcutaneous injections of wild type P. gingivalis 87 (strain W83) resulted in the formation of abscesses and ultimately lethal infection that were not 88 seen in sialidase deficient *P. gingivalis* [17]. Sialidase deficient mutants of *T. denticola* have also been 89 studied, and these display reduced complement evasion and decreased virulence in a mouse abscess 90 model [19].

Given the importance of sialidases for periodontal pathogen-host interactions and subsequent
virulence, abrogation of sialidase activity using sialidase inhibitors might exert anti-virulence effects
or be detrimental to the lifecycle of these organisms. This is particularly pertinent given recent
evidence that levels of sialidase transcript (*T. forsythia nanH* gene) are elevated in periodontal
disease [20]. Similarly, sialidase enzyme activity is raised in the gingival crevicular fluid of patients
with periodontal disease and this high sialidase activity is an indicator of poor responsiveness to
standard treatment [21].

Sialidase inhibitors are undergoing extensive development and some are available as a treatment for 98 99 influenza, with the sialidase inhibitors oseltamivir and zanamivir licensed for use globally. Sialidase 100 inhibition has been investigated in T. forsythia, where the inhibitor oseltamivir was shown to disrupt 101 biofilm formation on host sialoglycan substrates [16]. Considering the above, we set out to further 102 characterise the sialidase of P. gingivalis, performing a biochemical characterisation and 103 estabilishing an activity profile while also examining the potential of sialidase inhibitors to modulate 104 pathogenesis of *P. gingivalis* alone and in combination with a range of other periodontal pathogens 105 with the commercially available sialidase inhibitor zanamivir.

106 Methods

107 Bacterial strains, human cell lines, and culture conditions Tannerella forsythia strains used here 108 were the WT type strain ATCC 43037, supplied by William Wade, Queen Mary University of London, 109 UK, and the sialidase deficient nanH mutant (TFAnanH) [22]. Porphyromonas gingivalis strains used here were the WT strains ATCC 33277, 381 (Pg381), and a sialidase deficient mutant of Pg381 110 (Pg381∆siaPG). Fusobacterium nucleatum strain NCTC 25586 (subspecies nucleatum) was also used 111 112 in this study, supplied by William Wade. All strains were cultured at 37 °C under anaerobic conditions (10 % CO2, 10 % H2, 80 % N2) in a Don-Whitley mini-macs anaerobic cabinet. P. gingivalis 113 114 and F. nucleatum strains were cultured on Fastidious Anaerobe Agar (Lab M, Lancashire, UK) supplemented with 5 % (v/v) oxalated horse blood (Thermo Fisher Scientific, UK), for 2-3 days before 115 116 harvesting for use in experiments. T. forsythia strains were cultured on Fastidious Anaerobe Agar (Lab M, UK) supplemented with 5 % (v/v) oxalated Horse Blood (Thermo Fisher Scientific), 10 μ g ml⁻¹ 117 118 N-acetylneuraminic acid (NAM, Sigma Aldrich, UK), and 25 µg ml⁻¹ Gentamicin (Sigma-Aldrich) for 3-7 119 days before harvesting for use in experiments. In addition, TFAnanH and Pg381ASiaPG were 120 intermittently subcultured in the presence of 10 μ g ml⁻¹ erythromycin to ensure maintenance of the 121 antibiotic resistance cassette marker responsible for sialidase inactivation. Immortalized human oral 122 keratinocytes (OKF6/Tert2) (Dickson et al. 2000) were kindly provided by Dr J. Rheinwald (Harvard 123 Medical School, Cambridge, MA) and were grown in keratinocyte-serum free media supplemented 124 with defined growth supplements (DKSFM, Fisher Scientific, Loughborough, UK). The Oral Squamous 125 Cell Carcinoma (OSCC) cell line H357 (Thomas et al. 2001; Sugiyama et al. 1993)-a generous gift from 126 Professor S. Prime, University of Bristol, UK-was grown in Dulbecco's Modified Eagle Medium 127 (DMEM) supplemented with 10 % (v/v) foetal bovine serum (FBS), 2mM L-glutamine, and 100 units 128 ml⁻¹ Penicillin-Streptomycin (Sigma Aldrich). Cells were incubated at 37 °C, 5 % CO2 when not 129 undergoing media changes or passaging. Cells were grown to 70-90 % confluence and media was changed every 2-4 days. OBA-9 cells were maintained in keratinocyte basal medium KGM-2 130 131 supplemented with epidermal growth factor (EGF), bovine pituitary extract, epinephrine, transferrin, hydrocortisone and insulin, as per the manufacturer's recommendations (Lonza). Cells were used 132 133 after reaching near confluence.

Cloning, expression, and purification of recombinant proteins NanH and NanS were cloned,
expressed, and purified as previously described [23, 24]. A nucleotide sequence encoding SiaPG
(PG_0352, based on the genome of strain ATCC 33277) without the secretion signal sequence
(supplemental text), was codon optimised for expression in *E. coli*, and synthesized commercially
(ThermoFisher). The sequence also contained restriction sites for *Ndel* and *Xhol* at its 5' and 3' ends,
enabling it to be ligated into pET vectors. Restriction digest and ligation was performed using NEB
buffers, restriction enzymes, and T4 ligase, according to manufacturer's protocols. SiaPG was ligated

- into pET21a, and ultimately transformed into chemically competent *E. coli* BL21 (origami B).
- 142 Expression and purification of recombinant SiaPG was performed as previously described for NanH
- 143 [24].

144 Construction of *P. gingivalis siaPG* deletion mutant and genetic complementation.

145 For generating a sialidase deficient mutant, the siaPG gene (PG0352) in P. gingivalis 381 strain was inactivated by an allelic replacement strategy. The primers used in the construction and 146 147 complementation strategy are listed in Table 1. Briefly, a DNA fragment containing PG0352 coding 148 region and flanking sequences was PCR amplified with Primers #1 and #2 using the P. gingivalis 381 149 gDNA. This product was then used as a template to amplify the upstream and downstream regions 150 of PG0352 with primer sets #1 and #3 (for '5 end) and #2 and #5 (for '3 end), respectively. In parallel, 151 the ermF gene (797 bp) was amplified from the plasmid pVA2198 with Primers #4 and #6. Primers #3, #4, #5, and #6 contain overlapping sequences for *ermF* and PG0352 to allow generation of a 152 153 fusion fragment via overlap PCR. Finally, PCR products of PG0352 with flanking regions and ermF gene were combined in an overlap PCR reaction using primers #1 and #2. The overlap PCR product 154 155 was sequenced to confirm the correct fusion of fragments and then transformed into P. gingivalis 156 381 by electroporation as previously described [25]. Transformants were plated on TSB-blood agar 157 plates containing 10 μ g ml⁻¹ of erythromycin and incubated at 37 °C anaerobically for 10 days. 158 Following incubation, erythromycin-resistant colonies were screened by PCR and Southern blotting. 159 One representative deletion mutant as confirmed by PCR and Southern blotting (data not shown) 160 and showing no sialidase activity was selected and designated as Pg381∆siaPG. For 161 complementation of Pg381AsiaPG, in trans complementation with the *siaPG* gene on the self-162 replicating shuttle plasmid pT-COW was employed as we have performed previously [25]. Briefly, the siaPG gene fragment amplified by PCR with the primers FWD BamHI and REV Sall (Table 1) was 163 164 digested with BamHI and Sall and ligated into BamHI and Sall digestd pT-COW plasmid. The resulting 165 plasmid, pT-PG352C was transformation into Pg381∆siaPG by tri-parental conjugation involving with 166 Pg381AsiaPG, E. coli carrying shuttle vector pT-PG352C and E. coli carrying helper plasmid pRK231 4,5. The transformants were selected on TSB-blood agar plate containing 100 μ g ml⁻¹ of gentamicin 167 168 (Invitrogen), 10 μ g ml⁻¹ of erythromycin (Sigma) and 1 μ g ml⁻¹ of tetracycline (Sigma). The completed 169 clones with sialidase expression were saved and one random clone designated $\Delta siaPG^+$ served as a 170 representative.

Inhibition of whole periodontal pathogens and purified sialidases by zanamivir. Zanamivir was
provided by GlaxoSmithKline (GSK, Weybridge, UK). Whole *T. forsythia* or *P. gingivalis*, or purified
sialidases 2.5 nM NanH or 5 nM rSiaPG were incubated in the presence of 0.1 mM MUNANA, in 50

174 mM Sodium Phosphate 200 mM NaCl, pH 7.4 (to mimic host physiological conditions), in the

- presence of different concentrations of zanamivir (0-10 mM). Sialidase inhibition was expressed as
- the percentage change in fluorescence at a given concentration of inhibitor, relative to fluorescence
- in the absence of inhibitor. Sialidase inhibition (%) was plotted against log[inhibitor], and the
- variable slope model applied to obtain the IC50 of zanamivir for SiaPG and NanH in Graphpad Prism
- 179 7, using the equation: Y=100/(1+10 (LogIC50-X) × HillSlope). Y = sialidase activity relative to no
- 180 inhibitor condition (%), X = log[inhibitor], and Hillslope = steepness of the curve.

181 Biochemistry of SiaPG: Determination of optimum pH, reaction kinetics with MUNANA and

182 sialyllactose, and sialic acid release from mucin in concert with NanS from *T. forsythia*. Whole *P*.

183 *gingivalis* and purified SiaPG were tested for pH optima in a similar manner to *T. forsythia* and its

sialidase [24]. For whole cell assays, bacteria were resuspended from agar plates, washed 3x in

- phosphate buffer saline, pH 7.4 (PBS, Sigma Aldrich) and centrifugation at 10000 g for 2 minutes,
- and resuspended to an OD600 of 0.05 in appropriate reaction buffers (20 mM sodium citrate-citric
- acid (pH 3.0–6.4), sodium phosphate mono-basic/dibasic (pH 6.8–8.8) or sodium carbonate–sodium

bicarbonate (pH 9.2–10.5)). For pH optimum derivation for purified SiaPG, enzyme was used at 2.5

- nM. In both cases, cells or enzyme were incubated with 100 μ M Mu-NANA and quenched with
- addition of 100 mM sodium carbonate buffer, pH 10.5, at a volume ratio of 1 : 1.5 (reaction: sodium
- 191 carbonate), and sialidase activity quantified by measuring 4-MU fluorescence ($\lambda ex = 350 \text{ nm}$; $\lambda em =$
- 192 450 nm). For whole *P. gingivalis* assays were performed over 1-3 hours at 37 °C, incubation with
- 193 SiaPG was performed over 1-3 minutes at room temperature (~20 °C).

194 For derivation of Michaelis-Menten kinetics for SiaPG, 2.5 nM enzyme was incubated with varying 195 concentrations of MU-NANA and again quenched. A standard curve of the fluorescence signal of 4-196 MU at defined concentrations was also obtained, enabling sialidase activity to be expressed as 4-MU 197 released µmol min⁻¹ mg⁻¹ SiaPG. Similar conditions were used for derivation of kinetic parameters 198 with 3/6-siallyl-lactose, except no quenching was required and the levels of released sialic acid using 199 a modified Thiobarbituric acid (TBA) assay. In this case reactions were stopped at time points over 200 1-5 minutes by commencement of the TBA assay: Fifty microlitres of these reactions were added to 201 25 μl of 25 mM sodium periodate (Sigma–Aldrich) in 60 mM H₂SO₄ (Thermo Fisher) and incubated 202 for 30 min at 37 °C. This oxidation step was stopped by the addition of 20 μ l of 2% (w/v) sodium 203 meta-arsenite (Sigma-Aldrich) in 500 mM HCl. Forty-seven microlitres of this reaction were added to 204 100 μ l of 100 mM TBA, pH 9.0, and incubated at 95 °C for 7.5 min, resulting in the thiol-labelling of 205 free sialic acid. Upon centrifugation at 1500 G for 5 min, the pink chromophore in the clarified 206 supernatant was spectrophotometrically quantified at A₅₄₉. A standard curve of known sialic acid

207 (Neu5Ac, Carbosynth) concentrations was used to calculate sialic acid release in µmolmin⁻¹mg⁻¹

SiaPG. Finally, release of Neu5Ac from mucin was assessed using BSM and the TBA assay. PBS + 6μ M

- 209 BSM was incubated with combinations of 100 nM rSiaPG, 0.5 mM zanamivir, and/or the sialate-O-
- 210 acetylesterase NanS from *T. forsythia* (100 nM), and incubated for 30 minutes at 37 °C, before the
- 211 reaction was stopped and the TBA assay commenced as described above.
- 212

213 Analysis of Procainamide labelled glycans. SLeX and FA2G2S2 (A2F) glycans (Ludger Ltd.) underwent 214 procainamide labelling (see below), as were glycans released from recombinant human EPO as 215 previously described [23]. Briefly, human EPO was expressed in Chinese hamster ovary (CHO) cells (a gift from Antonio Vallin, Center for Molecular Immunology, La Habana, Cuba) and released using 216 217 peptide N-glycosidase F (PNGase F, E-PNG01; Ludger Ltd). EPO (in 17.5 µl) was denatured at 100 °C 218 for 5 mins with the addition of 6.25 μ l 2 % (w/v) SDS, and 1 M 2-mercaptoethanol, then incubated at 219 37 °C for 16 h with 1 μ l PNGase F and 1.25 μ l 15 % (w/v) Triton X-100. Released N-glycans were 220 fluorescently labelled with 2-aminobenzamide (2-AB) as described previously [26] using a LT-KPROC-221 96 kit (Ludger Ltd.). The released glycans were incubated with labelling reagents for 3 h at 65 °C. The 222 2-AB- labelled glycans were cleaned up using LudgerClean S Cartridges (Ludger Ltd), then incubated 223 with 1 μ l of 1 mg ml⁻¹ SiaPG in a final volume of 10 μ l (50 mM sodium acetate buffer, pH 5.5) for 16 h 224 at 37 °C. Glycans were cleaned-up and SiaPG removed using a LC-PROC-96 kit (Ludger Ltd.) 225 Procainamide labelled glycans (Ludger) were analysed by UHPLC-MS/MS. Here, 25 μl of each sample 226 was injected into a Waters ACQUITY UPLC Glycan BEH Amide column (2.1 x 150 mm, 1.7 μm particle 227 size, 130 Å pore size) at 40°C on a Dionex Ultimate 3000 UHPLC instrument with a fluorescence 228 detector (λ ex = 310 nm, λ em = 370 nm) attached to a Bruker Amazon Speed ETD. Mobile phase A 229 was a 50 mM ammonium formate solution (pH 4.4) and mobile phase B was neat acetonitrile. 230 Analyte separation was accomplished by a gradient running from 76-51% mobile phase B over 70 minutes at a flow rate of 0.4 ml min⁻¹. The Amazon Speed was operated in the positive sensitivity 231 232 mode using following settings: source temperature, 180 C; gas flow, 4 L min⁻¹; capillary voltage, 4500 233 V; ICC target, 200,000; maximum accumulation time, 50.00 ms; rolling average, 2; number of 234 precursor ions selected, 3; scan mode, enhanced resolution; mass range scanned, 400 to 1700. Data 235 was analysed using Bruker Compass Data Analysis software v4.1 and glycan diagrams made using 236 GlycoWorkbench v2.0. Glycan compositions were elucidated based on MS2 fragmentation.

Lectin staining for cell surface sialic acid- immunofluorescence microscopy. H357 cells were seeded
 at a density of 1.5x10⁵ cells ml⁻¹ into the wells of a 24-well tissue culture plate which contained
 sterile glass coverslips (BDH). These were incubated at 37 °C, 5 % CO₂ for 18 hours. Treatment with

240 SiaPG and NanH was performed by washing 2 x with PBS, then incubated with 200 nM of each 241 sialidase for 30 minutes at 37 °C. Cells then underwent sialic acid staining with lectins from Sambucus nigra (FITC conjugated SNA, Vector Labs, specific for a2-6 Neu5Ac linkages), Maackia 242 243 amuriensis (Biotinylated MAA, Vector Labs, specific for α 2-3 Neu5Ac linkages). After treatment, cells 244 were washed twice with 500 μ I PBS followed by application of lectins: 4 μ g ml⁻¹ SNA or 8 μ g ml⁻¹ 245 MAA, for 30 minutes at 37 °C, 5 % CO2. Cells were washed twice with 500 μl PBS, and conditions containing MAA underwent a second incubation with 2 μ g ml⁻¹ Texas Red-Streptavidin, for 30 246 247 minutes at 37 °C. Stained cells were washed three times with 500 μ l PBS and fixed with 500 μ l 2 % (w/v) paraformaldehyde for 15 minutes at 37 °C. Coverslips with fixed cells were removed from wells 248 249 and mounted onto glass slides with ProLong Gold Antifade mount (containing DAPI, ThermoFisher 250 Scientific). Mounted cells were incubated for at least 18 hours and visualised within one week. 251 Imaging was performed with an Axiovert 200M fluorescence microscope (Zeiss) and associated 252 Axiovert software (Zeiss). Images were processed using Fiji-imageJ, Software [27]. Fluorescence 253 background subtraction was performed for all images using the same parameters for each 254 fluorescence colour channel in a given experiment.

255 Biofilm assays. P. gingivalis was cultured in the wells of a 96-well tissue culture plate (poly-lysine 256 coated, Greiner): Bacteria from an agar plate were resuspended to an OD600 of 0.05 in Tryptic Soy 257 Broth (TSB; Sigma-Aldrich) supplemented with 2 % (w/v) yeast extract, 1 μ g ml⁻¹ vitamin K, 5 μ g ml⁻¹ hemin, 50 µg ml⁻¹ gentamicin, and either 6 mM Neu5Ac or no additional supplements. In conditions 258 259 testing host glycoproteins, 100 μ l of 6 μ M BSM, 100 % (v/v) pooled human saliva, or fetal bovine 260 serum (FBS) was added to plate wells and left to coat the wells by incubation at 4 °C overnight, 261 removed, and wells washed with PBS once. In conditions testing the effect of sialidase inhibition, 10 262 mM zanamivir was included. All media was equilibrated overnight in culture conditions, namely, 37 263 $^{\circ}$ C under anaerobic atmosphere (10 % CO₂, 10 % H₂, 80 % N₂). Biofilms were cultured for 5 days. To 264 quantify planktonic or total growth, all conditions underwent OD₆₀₀ measurement using a Tecan Infinite M200 plate reader. Quantification was carried out by either crystal violet staining or manual 265 266 counting of organisms. Crystal violet solution (0.1 % w/v) was added to each well and incubated at 267 room temperature for 30 minutes. After incubation, crystal violet was removed and wells were 268 gently washed 3-4 times with PBS before visualisation using light microscopy or extraction with 80: 269 20 ethanol: acetone. For manual counting, biofilms were gently washed twice using PBS to remove 270 planktonic cells, and vigorously resuspended in PBS and serial diluted where appropriate. Bacteria 271 were enumerated by counting using a Helber chamber (Hawksley) under phase contrast microscopy, 272 400 x magnification.

273 Antibiotic protection assays H357 or OKF6 cells were seeded into the wells of a 24-well tissue culture plate at a density of 1.2×10^5 - 2×10^5 cells and incubated for 24-48h. Media was removed, 274 the cells in one well were detached by trypsinisation and counted by haemocytometry, to determine 275 276 the number of cells per well. Remaining cells were incubated for 1 hour in media supplemented with 277 2 % (w/v) bovine serum albumin (BSA), then washed twice with PBS. Bacteria were quantified using 278 a Helber chamber (Hawksley) under phase contrast microscopy, 400 x magnification. Bacterial 279 suspensions were diluted in culture media (with no FBS or antibiotics) to a ratio of 1:100 host cells: 280 bacteria, and for inhibitor testing conditions 10 mM zanamivir was included. These suspensions were 281 incubated with the host cells at 37 °C, 5 % CO2 for 1.5 hours, or in empty wells containing no host cells-constituting a bacterial "viability" control. After incubation, the wells were washed twice with 282 283 PBS, and either harvested (see below)-forming the "total association" condition, or media with 200 284 μ g ml⁻¹ metronidazole was added, then incubated at 37 °C, 5 % CO₂ for 1 hour, forming the 285 "invasion" conditions. To harvest bacteria, wells were washed three times with PBS, lysed with 286 deionised water and scraped with a pipette tip for one minute. Sufficient lysis was checked by microscopy (this method reliably resulted in >99 % lysis). Harvested bacterial suspensions 287 288 underwent serial dilutions and were plated onto agar by Miles-Misra methodology. After incubation 289 and colony counting, the number of bacteria associated with host cells was determined and 290 expressed as a percentage of viable bacteria that were associated with and had invaded the OKF6 291 cells. By subtracting invaded organisms from associated, the number of "attached" bacteria was 292 obtained.

293 Assessment of zanamivir cytotoxicity Quantifying metabolism as an indicator of cell viability was 294 performed using 3-(4,5-Dimethylthiazol-2-yl)-2,5-Diphenyltetrazolium Bromide (MTT). Twenty five 295 thousand OKF6 cells in DKSFM were seeded into the wells of a 96 well tissue culture plate (Greiner), 296 and incubated for 24 hours at 37 °C, 5 % CO2. Cells were then washed twice with PBS, and media supplemented with 1 mg ml⁻¹ MTT, and 0, 2.5, 5, 7.5, or 10 mM zanamivir was added. Cells were 297 298 incubated for 2.5 hours at 37 °C, 5 % CO₂. MTT-supplemented media was removed, and cells were 299 washed twice using PBS. Formazan crystals were solubilised using isopropanol + 0.125 % (v/v) HCl, 300 and quantified by measuring absorbance at 540 nm, reference at 630 nm using a Tecan Infinite 301 M200 plate reader. A commercially available lactate dehydrogenase (LDH) assay, Cytotox 96 302 (Promega) was also used to assess cell membrane permeability as a measure of viability, according 303 to manufacturers' instructions, with cells seeded as described for MTT assays.

304 Results

305 SiaPG is a highly active, broad specificity sialidase, and desialylates host glycans. The coding 306 sequence of SiaPG without secretion signal sequence was synthesized as a codon optimised fragment and cloned into a pET21a plasmid. Expression was performed in E. coli BL21 (origami B), 307 308 and purified via Nickel affinity chromatography to high homogeneity (supplemental figure S1). The 309 commonly used fluorogenic 4-methylumbeliferyl-N-acetyl neuraminic acid (MUNANA) substrate was 310 used to assess sialidase activity with liberated 4-methylumbelliferone (4-MU) measured with 311 fluorescence change per minute used to quantify sialidase activity (Figure 1 A). We observed that 312 SiaPG has optimal activity under acidic conditions, with highest activity observed at a pH of 5.2-5.6. 313 However, it retained 40-50 % activity between pH 4.4 and 7.2, and ~20-30 % retained even at pH 4 314 and 8. Whole bacterial sialidase activity of *P. gingivalis* was also assessed, finding that \geq ~70 % 315 maximal activity was retained in the pH range 4.8-8.0, with the highest activity observed at pH 6.8, 316 (Figure 1 B). The stronger activity of whole *P. gingivalis* sialidase in a wider pH range compared to 317 the sialidase protein may be due to a buffering or stabilising effect of cell membranes or other 318 components and is relevant given that subgingival GCF pH is in the neutral range.

319 We also performed kinetics experiments to enable comparison of SiaPG to other sialidases. 320 Given the pH variations in the pathogens' ecological niche, i.e. the conditions the sialidases would 321 encounter in the host, we evaluated sialidase activity at SiaPG-optimum (pH 5.6) as well as 322 physiological pH (7.4) that would be present in the mouth, and in the presence and absence of 323 additional salt (figure 2). SiaPG had the greatest catalytic efficiency (k_{cat}/K_M) at pH 5.6 in the 324 presence of salt (161.0 μ M min⁻¹ with 200 mM NaCl, 124.7 μ M min⁻without NaCl). At pH 7.4, salt had a similar impact on catalytic efficiency of SiaPG (49.9 µM min with NaCl, 36.4 µM min without NaCl). 325 326 These values are broadly in line with those previously reported for the NanH sialidase of T. forsythia 327 [24], though salt had the opposite effect on NanH catalytic efficiency.

In addition to MUNANA, we assessed the ability of SiaPG to cleave 2-3' and 2-6' linked sialic acid, initially with the trisaccharide substrate sialyllactose (3- and 6- sialyllactose), using a TBA assay as previously described for NanH. SiaPG was capable of cleaving both linkage types (Figure 3), and although it had a much higher affinity (lower K_M) for the 2,6 linkage than 2,3, its reaction velocity with the 2,3 version means that it has a ~50 % higher catalytic efficiencies (K_{cat}/K_M) for the 3sialyllactose linked sugars.

Desialylation of more complex and physiologically relevant glycans by SiaPG was also
 assessed using HPLC and mass spectrometry. SLeX is an oligosaccharide containing α2-3 linked sialic
 acid, and is present in numerous host niches-on the plasma membrane of various cell types and at
 the termini of secreted glycans. SLeX may also play a role in *P. gingivalis* pathogenesis [28]. We

338 exposed soluble SLeX to SiaPG, subjected the reaction to HPLC, and found that SLeX was efficiently 339 desialylated by SiaPG (Figure 4A). Cleavage of α 2-6 linked sialic acid by SiaPG was tested using the 340 host-relevant FA2G2S2 (also known as A2F) glycan, a two-branched polysaccharide chain where both 341 branches terminate with a sialic acid, which is present on a number of secreted glycoproteins 342 including IgG. SiaPG was again capable of desialylating FA2G2S2 (Figure 4B). Notably, other peaks 343 were observed in the chromatogram for FA2G2S2, which are likely to correspond to differently 344 sialylated versions of the FA2G2S2 glycan, and these also appeared to be desialylated - notably the 345 mammalian NeuGc versions. It would be interesting to test the activity of SiaPG against such NeuGc 346 linked glycoconjugates, which the human host does not synthesize but obtains from dietary sources.

347 Finally, we wanted to test desialylation of glycans derived from a complex sialoglycoprotein 348 by SiaPG. To this end, we chose the host glycoprotein erythropoietin (EPO) due to its possession of 349 multiple bi-, tri- and tetra- branched glycans, the presence of α 2-3 and α 2-6 sialic acid linkages, and 350 because some of its NeuAc is mono or di-O-acetylated (Neu5,9Ac, and perhaps tri-acetylated 351 Neu5,8,9Ac). EPO was digested with SiaPG, N-glycans were procainamide labelled, and these 352 underwent UHPLC-MS/MS (Figure 5). As expected, the UHPLC chromatogram of undigested EPO 353 showed the presence of multiple peaks corresponding to various EPO glycans, further details and 354 structures of these are shown in supplemental table 1. Following digestion with SiaPG, we observed 355 a marked change in the chromatogram. The remaining peaks were investigated by MS/MS, providing 356 evidence that SiaPG was able to cleave the sialic acid from a wide-range of glycan structures. 357 However, this analysis also revealed the inability of SiaPG to cleave diacetylated (or triacetylated) 358 sialic acids (Figure 5, lower 2 panels). These analyses showed that extracted ion chromatogram (EIC) 359 corresponding to the MS2 ion for Neu5Ac trisaccharide (GlcNAc-Gal-Neu5Ac, m/z = ~657.25) was not 360 detected above background levels, while EIC corresponding to di- and tri- acetylated sialic acid 361 trisaccharides (GlcNAc-Gal-Neu5,9Ac and GlcNAc-Gal-Neu5,8,9Ac, $m/z = \sim 699.25$ and ~ 741.26) were 362 detected in high abundance (high peak intensity)- i.e. had been left uncleaved by SiaPG; a result expected given our previous studies on NanH [23]. 363

364 SiaPG releases Neu5Ac from mucins, an activity enhanced by the sialate-O-acetylesterase NanS

from *T. forsythia*. To explore activity of SiaPG further we next tested its activity with a glycoprotein
substrate relevant in the oral cavity- salivary mucin. Notably the commonly used Bovine
Submaxillary mucin (BSM) used in these experiments contains many terminal sialic acids that have
the second O-acetyl group, i.e. contain Neu5,9Ac [29]. Our previous data on NanH [23] and SiaPG
(this study) indicate terminal Neu5,9Ac is resistant to cleavage by periodontal sialidases. To counter
this, some bacteria express sialate-O-acetylesterases, such as NanS from *T. forsythia*, which remove

the second acetylation (forming Neu5Ac), and rendering the sialic acid susceptible to sialidase

372 cleavage. Indeed, we previously demonstrated that *T. forsythia* NanS enhances NanH sialidase

- activity on Neu5,9Ac terminal sugars and may act to aid sialidases from other species in harversting
- sialic acid [23]. To test the hypothesis that *T. forsythia* may act to enhance SiaPG activity we
- 375 performed sialic acid release assays on BSM. The data showed that NanS enhanced sialic acid release
- by SiaPG from BSM by ~2.5 -fold (from 302 to 746 pmol/min) (Figure. 7).

377 Zanamivir more efficiently inhibits sialidase activity of SiaPG and whole *P. gingivalis* than sialidase

378 activity of T. forsythia NanH and whole T. forsythia. As outlined, one aim of this study was to assess 379 the ability of a safe and FDA-approved drug in its ability to potentially reduce virulence of the 380 keystone periodontal pathogen, P. gingivalis. Therefore we assessed the impact of zanamivir on the 381 activity of whole P. gingivalis and another periodontal pathogen, T. forsythia, as well as their purified 382 sialidases using the MUNANA assay as described above. Both pathogens and purified sialidases 383 displayed decreased sialidase activity as zanamivir concentration was increased (Figure 6 A). The 384 decrease in activity was far more drastic for P. gingivalis, with a decrease in activity of ~70 % in the 385 presence of 10 mM zanamivir, compared to T. forsythia which only showed a decrease of ~25 % 386 (Figure 6 B). This apparent difference in efficacy for the two sialidases was confirmed by establishing 387 the IC50 of zanamivir for purified SiaPG and NanH (Figure 6 B), where zanamivir displayed a greater 388 inhibitory effect on SiaPG (369 μ M) than NanH (6130 μ M). These data were further enhanced when 389 we observed that zanamivir also inhibited sialidase activity in the context of removal of sialic acid 390 from mucin (Figure 7). There was no effect on NanS sialate-O-acetylesterase activity alone (data not 391 shown).

392 *P. gingivalis* biofilms on host sialoglycans are sialidase-dependent and disrupted by zanamivir. *P.*

gingivalis was cultured in microtitre plates, where well surfaces were coated with mucin (BSM), 393 394 saliva (pooled human saliva), or serum (FCS) in the presence or absence of 10 mM zanamivir. Total 395 growth (Figure 8 A) of P. gingivalis in the mucin-coated condition was significantly reduced in the 396 presence of zanamivir, from OD600 0.21 to 0.11 (p = < 0.001) i.e. zanamivir reduced total growth of P. 397 gingivalis in mucin-coated conditions by approximately half. In the presence of the other 398 glycoprotein-coated conditions (saliva and serum) there was a trend towards small reductions in 399 total growth in the presence of zanamivir but these did not reach statistical significance. Biofilm 400 formation (normalised to total growth-optical density) of P. gingivalis (Figure 8 B) was also reduced 401 when grown on mucin coated surfaces (30-40 % reduction, p = < 0.01), indicating that biofilm 402 formation on these surfaces was inhibited by zanamivir. Notably we also saw a consistent small

reduction on serum, but total growth (OD600) standardised to biofilm formation (bacteria ml⁻¹) of
 cultures grown on human saliva-coated surfaces was too variable to make conclusions.

405 In support of our evidence that sialidase is important when P. gingivalis grows on mucin coated 406 surfaces (as would often be the case in vivo), we also tested the effect of sialidase activity in this 407 regard in a related P. gingivalis strain – 381 (Pg381). For this purpose, a siaPG deletion mutant of P. 408 gingivalis 381 and its complemented strain were constructed (Pg381AsiaPG and AsiaPG+). 409 Pg381AsiaPG showed no sialidase activity as judged from lack of fluorescence on incubation with 410 MUNANA, while the silaidase activity was restored in the comlemented strain Δ siaPG⁺ (supplemental 411 figure S2). In biofilm assays, we observed a 2-fold reduction in biofilm growth in Pg381∆siaPG 412 (relative to the WT) on mucin coated surfaces, with this phenotype complemented by provision of 413 the SiaPG gene in the mutant (Δ siaPG+, figure. 9). These data expand our view of the role of sialidase

414 to a range of *P. gingivalis* strains and suggest it may be true across this species.

415 SiaPG releases 3- and 6- linked sialic acid from oral epithelial cells and is inhibited by zanamivir.

- 416 Studies of mutant strains and purified enzymes have shown that the well-studied *T. forsythia* NanH
- 417 can desialate oral epithelial cells [22, 24], while SiaPG mutant strains have been used to highlight P.
- 418 *gingivalis* sialidase-mediated release of sialic acid from erythrocytes [18]. In order to test the effect
- of SiaPG on physiologically relevant oral epithelial cells, we stained the OSCC line H357 with the
- 420 lectins SNA and MAA enabling fluorescent imaging of cell surface α 2-3 and α 2-6-linked sialic acid.
- 421 Exposure of epithelial cells to SiaPG (and *T. forsythia* NanH) was successful in desialylating cell
- 422 surfaces (this had been shown previously for NanH [22]), while 10 mM zanamivir inhibited
- 423 desialylation in the case of SiaPG, but not NanH (figure 10, supplemental figure S3).

424 Zanamivir inhibits periodontal pathogen-host cell association during monospecies infection. One

425 established aspect of *P. gingivalis* virulence is its ability to invade and reside within oral epithelial

426 cells. To this end we performed antibiotic protection assays to establish whether zanamivir had any

- 427 potential effect on host cell interaction and invasion. However, the effects of zanamivir on bacterial
- and human cell viability had to be established. To this end, oral epithelial cells were subjected to 10
- 429 mM zanamivir under antibiotic protection assay conditions, illustrating no effect of zanamivir on
- either membrane integrity (assayed by LDH) or cellular activity/ viability (via MTT assay)
- 431 (supplemental figure S4). Similarly, we incubated *P. gingivalis*, *T. forsythia* and *F. nucleatum* with
- 432 zanamivir under antibiotic protection assay conditions (2.5 hours in tissue culture medium), before
- 433 obtaining cfu counts, and revealed no detrimental effect of zanamivir on viability of any of these
- 434 species (supplemental figure S5).

435 Next we performed antibiotic protection assays on two oral epithelial cell lines, namely the OSCC cell 436 line H357- a frequently used model for oral epithelium-bacteria interactions in our laboratory [30, 31] and normal immortalised oral epithelial cell line OKF6 [32, 33]. In addition to P. gingivalis we 437 438 also tested the effect of zanamivir on two other key periodontal pathogens- namely T. forsythia and F. nucleatum subsp. nucleatum. For all three pathogens, in both cell lines, zanamivir significantly 439 440 reduced one or more aspects of host-bacteria association in both cell lines (figure 11). For T. forsythia, attachment, invasion, and total association (the sum of attached and invaded bacteria) 441 442 were all significantly reduced in both cell lines; from 3.2 to 1.0 %, 2.5 to 0.3 %, and 5.8 to 1.4 % in 443 OKF6 cells, and 4.6 to 1.2 %, 3.4 to 1.6 %, and 8.0 to 2.8 % in H357 cells (figure 11 A). For P. 444 *qinqivalis*, attachment, invasion, and total association were significantly reduced in OKF6 cells; from 445 0.9 to 0.1 %, 0.7 to 0.1 %, and 1.7 to 0.2 %, however in the H357 cell line, only invasion was 446 significantly reduced, from 4.5 to 1.8 % (figure 11 B). Finally, for F. nucleatum, invasion and total 447 association were significantly reduced in OKF6 cells; from 4.6 % to 1.3 %, and 5.6 to 2.6 %. For H357 448 cells attachment and total association were significantly reduced 21.8 % to 7.2 %, and 36.6 % to 18.2 449 % (figure 11 C).

450 To support the notion that sialidase activity is important for bacteria-host cell association, rather 451 than disruption of other processes by the presence of zanamivir, the human gingival cell line OBA-9 452 was exposed to *P. gingivalis* 381 (Pg381) and a *siaPG* deficient mutant (Pg381 Δ *siaPG*). Exposure to 453 the latter strain was performed without or with exogenous SiaPG . Pg381*\DeltasiaPG* displayed 454 significantly decreased association with host cells relative to the parent Pg381, which was restored by the presence of exogenous SiaPG in a concentration dependent manner (supplemental figure S6): 455 456 While 100 µg ml⁻¹ exogenous recombinant SiaPG resulted in partial restoration of host-bacteria 457 association of Pg381ΔsiaPG (WT; 0.50 %, Pg381ΔsiaPG; 0.15 %, Pg381ΔsiaPG+100 μg SiaPG; 0.26), 458 $200 \ \mu g/ml$ SiaPG resulted in complete restoration of host-bacterial association Pg381 (0.58 %).

459 Zanamivir inhibits attachment and invasion of oral epithelial cells during infection with multiple 460 periodontal pathogens. To at least partially mimic the *in vivo* situation, where none of these species 461 exist in isolation, we performed antibiotic protection assays where multiple periodontal pathogens 462 were used to infect cells. This is particularly relevant during host cell association, since periodontal 463 pathogens have been shown to act in synergy during attachment and invasion [34–36]. MOI was maintained at 1:100 host cells: bacteria, so for infections using two species, the ratio was 1:50:50 464 (host cells: species 1: species 2), and for infections with all three species, the ratio was 1:33:33:33 465 (host cells: species 1: species 2: species 3). T. forsythia, P. gingivalis, and F. nucleatum were used in a 466 467 variety of combinations to infect epithelial cells. In addition to quantifying the numbers of all

468 bacteria-not distinguishing between species during an assay-the levels of each individual species in a 469 given assay could be enumerated by colony counting on agar plates, since the colony morphologies 470 of *T. forsythia*, *F. nucleatum*, and *P. gingivalis* are distinct from each other (supplemental figure S7). 471 All possible combinations were tested in this way. For ease of discussion, only the total number of 472 bacteria in each combination are described here, with complete results for each individual species 473 provided in supplemental figures S8-S11. For T. forsythia and F. nucleatum co-infection, zanamivir 474 significantly reduced both invasion and total association of bacteria with host cells, from 7.4 to 1.8 475 %, and 19.6 to 7.6 % (p<0.01 and p<0.05), respectively (Figure 12 A). For T. forsythia and P. gingivalis 476 co-infection, the variation between experiments meant that none of the reductions were statistically 477 significant (where p <0.05, Figure 12 B), i.e. attachment was reduced from 12.1 to 0.8 % (p=0.076), invasion from 2.8 to 1.4 % (p=0.25), and total association from 14.4 to 2.5 % (p=0.062). For F. 478 479 nucleatum and P. gingivalis, zanamivir significantly reduced attachment, invasion, and total association of bacteria with host cells, from 1.9 to 0.8 % (p=<0.05), 4.0 to 1.56 % (p=<0.05), and 5.9 480 481 to 2.4 % (p=<0.01), respectively (figure 12 C). Finally, all three species (i.e. T. forsythia, F. nucleatum 482 and P. gingivalis) were infected into the cells as a three-species community. Here, strikingly, zanamivir significantly reduced attachment, invasion, and total association of bacteria with host 483 484 cells, from 17.7 to 7.7 % (p = <0.001), 9.6 to 7.15 % (, p = <0.05), and 26.14 to 14.6 % (p = <0.001), 485 respectively (figure 12 D); indicating this approach might have impact on cellular interaction in vivo.

486 Discussion

487 In this study we characterised the activity of the sialidase (SiaPG) from P. gingivalis on a number of 488 glycan and proteinaceous substrates, including relevant host sialoglycans SleX, FA2G2S2 and EPO, 489 and establishing activity across a range of pH values. During these studies, it became apparent that 490 SiaPG could cleave both α 2-3 and α 2-6 linked Neu5Ac, but not diacetylated Neu5,9Ac. These data 491 also agreed with low sialic acid release from the heavily diacetylated BSM protein, and this activity 492 being enhanced by coincubation with an enzyme capable of removing O-acetyl groups on sialic acid 493 residues - namely the NanS sialate-O-acetylesterase from the fellow oral bacterium T. forsythia [23]. 494 We hypothesise that in vivo NanS enhances the ability of P. gingivalis to cleave sialic acid from 495 glycoproteins - although given the fact that P. gingivalis does not produce its own esterase or utilise 496 sialic acid for nutrition we can only postulate that somehow removal of sialic acid opens up sites for 497 adhesion to oral surfaces, or might modulate immune responses mediated by Siglecs (sialic acid-498 binding immunoglobulin-like lectins) or allow P. gingivalis to access underlying sugars or the protein 499 component of the glycoproteins themselves, and may explain synergies between these organisms 500 observed by others [37-39].

501 We tested the ability of zanamivir to inhibit SiaPG and the previously characterised NanH sialidase 502 from another red complex pathogen, T. forsythia. While the inhibition (IC50) of SiaPG activity on MUNANA was determined to be within the micromolar range (~350 μ M), this was not the case for *T*. 503 504 forsythia, which required >6 mM zanamivir to reduce its activity by half. A similar trend was also 505 seen for whole bacteria. The relatively low efficacy of zanamivir for some bacterial sialidases is 506 observed in other studies where sialidases from the pathogens S. pneumoniae and Gardnerella 507 vaginalis are apparently only inhibited to limited extents [40, 41]. Nevertheless, here we showed 508 that P. gingivalis sialidase was effectively inhibited by zanamivir. At present we have no mechanistic 509 explanation for the variation between the two sialidases, but postulate it may result from structural 510 differences, despite both possessing typical catalytic GH33 domains with predicted 6-blade propeller 511 structures the sialidases only share 23 % amino acid sequence identity.

512 Given this, and the ability of SiaPG to cleave sialic acid from complex glycans (FA2G2S2, EPO), we 513 assessed the ability of SiaPG to remove sialoglycans from oral epithelial cells, illustrating removal in 514 both cases and highlighting the broad glycan specificity of this enzyme and the ability of zanamivir to 515 affect this process. As shown previously, NanH was also capable of desialylating oral epithelial cells 516 [22].

517 Considering the ability of zanamivir to inhibit P. gingivalis sialidase activity, in vitro virulence models 518 were carried out. It has been shown previously that NanH is key during biofilm formation in T. 519 forsythia, and that sialidase inhibitors are also detrimental to T. forsythia biofilms [42]. We observed 520 that sialidase inhibitors decreased P. gingivalis biofilm formation on host glycoprotein sources, 521 significantly on saliva and mucin. In addition, biofilm formation of sialidase deficient P. gingivalis on 522 mucin was reduced compared to its parent, and restored upon complementation. Biofilms of 523 sialidase deficient P. gingivalis are also reduced on plastic surfaces [17]. There are various 524 mechanisms by which sialidase activity may contribute to P. gingivalis biofilm formation. Possibilities 525 include decreased proteolysis and subsequent nutrient deficiency or an inability to attach to cleaved 526 proteins, since sialylation may protect against proteolytic activity [43]. Interference with the P. 527 gingivalis capsule is also possible, sialidase expression appears to be important for P. gingivalis 528 capsular synthesis, mutants deficient in SiaPG show reduced capsule formation [17, 18], and yet loss 529 of the capsule has been shown to enhance biofilm formation [44], thus, we might expect zanamivir 530 (or sialidase knockout) to enhance biofilm formation, but here we observed the opposite, possibly 531 because non-capsulated (fimbriated) strains were tested. In any case, we cannot rule out the 532 possibility that sialidase inhibition may interfere with capsular structure or interactions in the case of 533 P. gingivalis strains which are encapsulated. Finally, maturation and processing of gingipains534 proteases central to P. gingivalis virulence and asacharolytic nutrition-is also altered in sialidase 535 deficient mutants [18]. Zanamivir may impact biofilm formation through any, or all, of these 536 mechanisms. It would be particularly interesting to assess the impact of zanamivir on mixed-species 537 biofilms, and in addition to direct impacts on sialic acid-mediated interactions and any indirect 538 effects that may result from sialidase inhibition. For example, it has been previously shown that 539 lysine gingipains (Kgp) of *P. gingivalis* mediate *T. forsythia* levels in mixed species biofilms [45], and . 540 given that sialylation may influence protease access to protein backbones, inhibition of P. gingivalis 541 sialidase may also impact the activity of Kgp, subsequently reproducing the detrimental effect of Kgp 542 deficiency on T. forsythia abundance. Similarly, it has been noted that T. forsythia surface S-layer 543 glycans play a potential role in localisation with *P. gingivalis* within polymicrobial biofilms [46]. 544 These glycans contain terminal nonulosonic analogues of sialic acid (pseudaminic and legianaminic 545 acids) and it is possible that Zanamivir may influence these interactions as a glycan mimic.

546 Given that periodontal pathogen sialidases have been shown to be important during host cell 547 association [17, 18, 22, 42] and zanamivir inhibited host cell surface desialylation by SiaPG, but not T. 548 forsythia NanH, it might be expected that zanamivir would decrease attachment and invasion of 549 host cells by P. gingivalis but not T. forsythia. Surprisingly, host cell association was greatly inhibited 550 in the case of both organisms, and in the sialidase negative F. nucleatum [13]. Of note here is that 551 while our data re-emphasize the role of sialidase in the equivalent *P. gingivalis* capsulated W83 552 strain [18], indicating that this phenotype is widespread in *P. gingivalis*, it would be of interest to 553 establish whether zanamavir has the same effect on capsulated strains. It is also possible that other 554 periodontal pathogen virulence factors are inhibited by zanamivir. For example, the adhesin FadA is 555 highly conserved among oral fusobacteria, and considered important for association with oral 556 epithelial cells [47] and has been shown to bind cell surface E-selectin, whose native ligand is a 557 sialoglycan (sialyl lewis A/X), so perhaps a sialic acid analogue might interfere with FadA-host cell 558 surface interactions.

559 Although the monospecies antibiotic protection assays hinted at the usefulness of zanamivir as an 560 anti-virulence therapeutic, periodontitis is mediated by a dysbiotic polymicrobial community. 561 Therefore, we performed mixed species antibiotic protection assays with combinations of all three 562 of the periodontal pathogens from the monospecies antibiotic protection assays. Importantly, zanamivir influenced invasion in the triple-species infection model with F. nucleatum, P. gingivalis, 563 564 and T. forsythia. Despite evidence of synergy in invasive capability reported in the literature [34, 36, 565 48] we were surprised zanamivir effected all species in these assays, given that F. nucleatum has no 566 sialidase and T. forsythia sialidase is only weakly inhibited by zanamivir. One explanation might be

567 that only a modest reduction in sialidase activity can reduce invasion for T. forsythia. However, 568 zanamivir acts on host sialidases, in particular Neu1 and Neu3, which contribute to the immune 569 response to bacteria via TLRs, where host sialidases are mobilised to the plasma membrane where 570 they activate TLRs 2-, 3- and 4- [49-51], and zanamivir (and other sialidase inhibitors) has been 571 shown to inhibit TLR activation by LPS [52]. Inhibition of host cell sialidases by zanamivir may 572 therefore affect the cell surface charge and glycosylation state of receptors involved in bacterial 573 adhesion and invasion. Indeed our data may suggest that there are host and bacterial sialidase 574 dependent events that not only warrant further study but make a strong case for the potential of 575 these inhibitors to affect periodontal disease pathogenesis, since they may have dual action on host 576 and bacterial processes.

577 Conclusion

In summary, we further characterised the sialidase of P. gingivalis, and provided evidence 578 579 advocating for the development of sialidase inhibitors as therapeutics for periodontitis-and other 580 diseases-where bacterial sialidases play key roles in virulence. Zanamivir was capable of inhibiting 581 sialidase activity and virulence mechanisms of *P. gingivalis*, most prominently host cell association. 582 However, our study raises several questions and directions for further work, most notably that host 583 as well as bacterial sialidase may influence interactions more than currently understood. Ultimately, 584 sialidase inhibition represents a potential novel therapy for periodontal and other diseases, and 585 merits further investigation. Finally, zanamivir is a safe and efficacious drug for treatment of 586 influenza, but clearly only inhibits bacterial sialidases relatively weakly (μ M –mM range). However, 587 its effectiveness in this study suggests that the development of more potent and selective inhibitors 588 of bacterial sialidase should be a focus in upcoming years as they may hold potential as a safe and efficacious treatment in periodontal and other diseases. 589

590 Author statements

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745 Figure legends:

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Figure 1. pH optima of *P. gingivalis* whole cell sialidase activity and purified recombinant SiaPG.
MUNANA was incubated with SiaPG for 1 minute or *P. gingivalis* (ATCC 33277) for 1 hour, in a variety
of buffers with variable pH. Reactions were halted and the pH equalised by addition of an excess of
sodium carbonate-bicarbonate buffer, pH 10.5. Sialidase activity catalysed the production of 4-MU
from MU-NANA, which was quantified by measuring fluorescence of the reactions at excitation and
emission wavelengths of 350 and 450 nm. A) SiaPG (B) *P. gingivalis*. Data shown represent the
mean of one experiment where each condition was repeated three times. Error bars = S.E.M.

754 Figure 2. Reaction kinetics of MUNANA and SiaPG under different conditions. Variable 755 concentrations of MUNANA were exposed to SiaPG, under different pH and salinity conditions. 756 Reactions were quenched by addition of pH 10.5 buffer at 1, 2, and 3 min, and the rate of 4-MU 757 release determined by application of a 4-MU standard curve. A) Michaelis–Menten plot, rate of 4-758 MU release (V0, μ mol MU released min⁻¹ mg⁻¹ SiaPG), plotted against [MUNANA] (μ M) using Prism 7 759 (GraphPad). Error bars = SD (B) Table summarising Michaelis-Menten reaction kinetics and catalytic 760 efficiency of SiaPG and MU-NANA, the table includes the k_{cat} (4-MU release min⁻¹), and k_{cat}/K_M (μ M 761 min^{-1}). Data shown represent the mean (± S.D.) of one experiment, where each condition was 762 repeated three times per experiment.

763 Figure 3. Reaction kinetics of 3- or 6- sialyllactose and SiaPG under different conditions. Variable 764 concentrations of sialyllactose were exposed to SiaPG, under conditions mimicking physiological (pH 765 7.4 200 mM NaCl). Reactions were quenched by commencing the thiobarbituric acid assay at 1, 2, 766 and 3 min, and the rate of Neu5Ac release determined by application of a Neu5Ac standard curve. 767 (A) Michaelis–Menten plot, rate of 4-MU release (V0, μmol MU released min⁻¹ mg⁻¹ SiaPG), plotted 768 against [3- or 6- sialyllactose] (μ M) using Prism 7 (GraphPad). Error bars = SEM. (B) Table 769 summarising Michaelis-Menten reaction kinetics and catalytic efficiency of SiaPG and MU-NANA, the 770 table includes the k_{cat} (4-MU release min⁻¹), and k_{cat}/K_M (μ M min⁻¹). Data shown represent the mean 771 (± S.D.) of one experiment, where each condition was repeated three times per experiment.

772Figure 4. Desialylation of host-relevant glycans by SiaPG. UHPLC chromatogram showing elution of773FA2G2S2 and SLeX with and without digestion by SiaPG. FA2G2S2 contains $\alpha 2$ -6 linked Neu5Ac.774Some contaminating glycans can be observed in the undigested sample, but these also appear to be775desialylated. Gc = N-glycolylneuraminic acid. SLeX contains $\alpha 2$ -3 linked Neu5Ac.

776 Figure 6. Inhibition of periodontal pathogen sialidase activity by zanamivir. MUNANA was exposed 777 to periodontal pathogens or purified sialidases. Sialidase activity was expressed as the difference in 778 4-MU fluorescence relative to conditions with no inhibitor. A) MUNANA was exposed to T. forsythia 779 or P. gingivalis in the presence of zanamivir for 1 hour and 4 hours, respectively. Data represent the 780 mean of two experimental repeats, where each condition was performed three times per 781 experiment. Error bars = SD. B) MUNANA was exposed to SiaPG and NanH in the presence or 782 absence of zanamivir (1-10 mM, 1 mM increments), and the IC50 of zanamivir for both enzymes 783 obtained using the variable slope model in Graphpad Prism 7. Data represent the mean of three 784 experiments, where each condition was repeated three times. Error bars=SEM.

Figure 7. Enzyme synergy in sialic acid release from BSM, and inhibition by zanamivir. BSM was
incubated with either NanS, SiaPG, or NanS + SiaPG, in the presence or absence of 0.5 mM
zanamivir, before undergoing the TBA assay to assess sialic acid release. Data based on the mean of
three experiments, each condition was tested in triplicate during each experiment. Error Bars=SEM.
Significance determined by one-way ANOVA with repeated measures, with Bonferroni correction for
multiple comparisons (*p=<0.05, **p=<0.01).

791 Figure 8. Zanamivir inhibits *P. gingivalis* growth and biofilm formation on sialoglycoproteins.

Bacteria were cultured for 5 days at 37 °C anaerobically, in the presence of Neu5Ac, or on surfaces
coated with the glycoproteins mucin, saliva, or serum. All conditions were performed in the
presence or absence of zanamivir. A) Total growth; OD₆₀₀ of the culture was used to quantify
bacteria in both biofilm and planktonic states. B) Biofilm formation, quantified by resuspension of
biofilms and counting the number of bacteria under microscopy. All data shown represent the mean
of three biological repeats, where conditions were tested three times per experiment. Error bars
=SEM. Significance determined by T-test, **p=>0.01, ***p=>0.005.

Figure 9. Biofilm formation by *P. gingivalis* **381**, sialidase deficient Pg381 Δ siaPG, and SiaPG complemented strain Δ siaPG⁺. *P. gingivalis* strains were resuspended to an OD₆₀₀ 0.5, and seeded into microtitre plates precoated with mucin (BSM), and biofilms were cultured for 24 hours at 37°C anaerobically. Data represent the mean of four replicates per condition. Error bars = SD, statistical significance determined by unpaired T-test compared to the WT strain, **p=<0.01).

Figure 10. Purified sialidases desialylate oral epithelial cell surfaces, and in the case of SiaPG this is inhibited by zanamivir. Cells were stained with lectins for α 2-3 and α 2-6 linked sialic acid in red and green, respectively. Prior to staining, cells were treated with NanH and SiaPG in PBS, in the presence or absence of 10 mM zanamivir (zan), as indicated. All images were visualised using the same

- microscopy and image processing parameters (fluorescence intensity, exposure time and
 background subtraction). Images were captured in three fields of view, and this was repeated in
 three separate experiments. Images shown are representative of each condition. Separate colour
- 811 channels are displayed in the supplementary information (supplemental figure S3).

812 Figure 11. The effect of zanamivir on attachment and invasion of epithelial cells during

813 monospecies infection. Antibiotic protection assays on OKF6 and H357 cell lines were performed in

- the presence or absence of 10 mM zanamivir (thatched and black bars, respectively) with either A) T.
- 815 forsythia, B) P. gingivalis, or C) F. nucleatum. Bacterial attachment, invasion, and total association
- 816 with host cells was normalised to the number of bacteria that were used to infect each condition
- that had survived the duration of the assay (the percentage of viable bacteria). Data represent the
- 818 mean from three independent experimental repeats, and each condition was repeated in triplicate
- 819 during each experiment. Error bars=SEM, Significance determined by paired T-test, *p=<0.05, **

820 Figure 12. The effect of zanamivir on attachment and invasion of epithelial cells during

821 **multispecies infection.** Antibiotic protection assays were performed in the presence or absence of

- 822 10 mM zanamivir (thatched and black bars, respectively) with different combinations of *T. forsythia*,
- 823 *F. nucleatum*, and *P. gingivalis* to establish host cell association in each instance. A) *T. forsythia* + *F.*
- 824 nucleatum B) T. forsythia + P. gingivalis. C) F. nucleatum + P. gingivalis D) T. forsythia, F. nucleatum,
- and *P. gingivalis*. Bacterial attachment, invasion, and total association with host cells was normalised
- to the number of bacteria that were used to infect each condition that had survived the duration of
- the assay (the percentage of viable bacteria). Zanamivir = 10 mM zanamivir present during host cell
- 828 exposure to bacteria. Data represent the mean from three experimental repeats, and each condition
- 829 was repeated in triplicate during each experiment. Error bars=SEM, Significance determined by
- 830 paired T-test, *p=<0.05, **p=<0.01. ***p=<0.001.

831 Table 1. List of primers used in this study

| Primers for sial | PG deletion mutant |
|------------------|--|
| Primer | Sequence (5'>3') |
| #1 | TACGGCATCGCGGTTTTGA |
| #2 | TCGCCATAGAATACAGGATAAGC |
| #3 | gggcaatttcttttttgtcatTGAAAACTATTTTATACCATTTTGGGA |
| #4 | TCCCAAAATGGTATAAAATAGTTTTCAatgacaaaaaaaaaaattgccc |
| #5 | aaaaatttcatccttcgtagGAATAGTGCTTTTTTTATCGAGTTTTTC |
| #6 | GAAAAACTCGATAAAAAAAGCACTATTCctacgaaggatgaaattttt |
| Sequences in u | pper case correspond to <i>siaPG</i> and lower case to <i>ermF</i> . |
| | |
| Primers for sia | PG complementation |
| FWD BamHI | CGCGGATCCG taatacgactcactatagg |
| REV Sall | TTGAC <u>GTCGAC</u> GCGTctagttattgctcagcggtgg |
| Endonuclease | sites are underlined. |





В







| | pH 7.4, 200 mM NaCl | pH 7.4 | pH 5.6, 200 mM NaCl | pH 5.6 |
|--|------------------------|--------------|------------------------|--------------|
| k _{cat} (4-MU release/min) | 789.8 ± 43.6 | 374.8 ± 16.3 | 1916 ± 69.46 | 1408 ± 33.28 |
| KM ([MU-NANA], µM) | 15.8 ± 3.7 | 10.3 ± 2.1 | 11.9 ± 1.9 | 11.3 ± 1.2 |
| k _{cat} /K _M (µM/min) | 49.9 | 36.4 | 161.0 | 124.7 |
| V _{max} (4-MU release, µmol/min/mg NanH) | 14.4 ± 0.8 | 6.8 ± 0.3 | 35.0 ± 1.3 | 25.7 ± 0.6 |



| 356.8 ± 49.64 | 74.42 ± 2.8 |
|-------------------|---|
| 2234 ± 548.8 | 684.9 ± 92.6 |
| 0.16 | 0.11 |
| 124.3 ± 17.3 | 25.9 ± 1.0 |
| | $\begin{array}{r} 356.8 \pm 49.64 \\ 2234 \pm 548.8 \\ \hline 0.16 \\ 124.3 \pm 17.3 \end{array}$ |











Figure 9













С

Native protein

MANNTLLAKTRRYVCLVVFCCLMAMMHLSGQEVTMWGDSHGVAPNQVRRTLVKVALSESLPPGAKQIRIGFSLP KETEEKVTALYLLVSDSLAVRDLPDYKGRVSYDSFPISKEDRTTALSADSVAGRCFFYLAADIGPVASFSRSDTLTARVE ELAVDGRPLPLKELSPASRRLYREYEALFVPGDGGSRNYRIPSILKTANGTLIAMADRRKYNQTDLPEDIDIVMRRST DGGKSWSDPRIIVQGEGRNHGFGDVALVQTQAGKLLMIFVGGVGLWQSTPDRPQRTYISESRDEGLTWSPPRDI THFIFGKDCADPGRSRWLASFCASGQGLVLPSGRVMFVAAIRESGQEYVLNNYVLYSDDEGGTWQLSDCAYHRG DEAKLSLMPDGRVLMSVRNQGRQESRQRFFALSSDDGLTWERAKQFEGIHDPGCNGAMLQVKRNGRNQMLHS LPLGPDGRRDGAVYLFDHVSGRWSAPVVVNSGSSAYSDMTLLADGTIGYFVEEDDEISLVFIRFVLDDLFDARQ

Supplemental figure S1. Purification of SiaPG A) SiaPG purification via affinity chromatography. L= Molecular mass ladder, ins = insoluble fraction of whole bacterial lysate, sol = soluble fraction of whole bacterial lysate, F = flow through, W = wash, elution fractions were 1 ml each. B) Purified SiaPG post-dialysis. C) Native gene encoding SiaPG and resulting encoded protein (PG_0352, obtained from UniProt; Q7MX62), 100 % protein identity between W83 and ATCC 33277 strains). Red = secretion signal site, absent in the codon optimized protein expressed here using *E. coli*.



Supplemental figure S2: Sialidase activity of SiaPG, *P. gingivalis* SiaPG-inactivated isogenic mutant (Pg381ΔsiaPG) and its complemented strain (ΔsiaPG⁺)



Supplemental figure S3. Purified sialidases desialylate oral epithelial cell surfaces.

Cells were stained with lectins for α 2-3 and α 2-6 linked sialic acid in red and green, respectively. Prior to staining, cells were treated with NanH and SiaPG in PBS, in the presence or absence of 10mM zanamivir, as indicated. All images were visualised using the same microscopy and image processing parameters (fluorescence intensity, exposure time and background subtraction). Images were captured in three fields of view, and this was repeated in three separate experiments. Images shown are representative of each condition.



Supplemental figure S4. Viability testing of OKF6 oral epithelial cells exposed to zanamivir.

The oral epithelial cell line OKF6 was exposed to *T. forsythia* in the presence and absence of 10 mM zanamivir for 2.5 hours, followed by an MTT assay on cells, or an LDH assay on culture supernatant, with the absorbance at 490 and 540 nm for LDH and MTT, respectively, used to relatively quantify levels of MTT and LDH. Data shown represent the mean of two experiments, where each condition was repeated three times per experiment. Error bars=SEM. No significant differences were found (where p=<0.05), as determined by one way ANOVA, with repeated measures and Tukey's correction for multiple comparisons.





T. forsythia, P. gingivalis, and F. nucleatum were incubated in the presence and absence of 10mM zanamivir for 2.5 hours, followed by enumeration of viable organisms by agar plate counts, and data were expressed as the change in cell numbers relative to the untreated condition. Data shown represent the mean of three experiments, where each condition was repeated three times per experiment. Error bars=SD. Significance determined by T-test (*p=<0.05).



Supplemental Figure S6. Association of *P. gingivalis* with OBA9 cells. OBA-9 cells were incubated with P. gingivalis 381 or the SiaPG inactivated mutant (Δ PG352), in the presence or absence of 100 and 200 µg of SiaPG. Bacterial association was calculated as percentage of input bacteria. Data represent the mean of two experiments, where each condition was repeated three times per experiment. Error bars = SEM, significant differences between Pg 381 and the other conditions were determined by paired T-test, *p=<0.05, **p=<0.01).



Supplemental figure S7. Colony morphology of *P. gingivalis, F. nucleatum,* and *T. forsythia* during mixed-species enumeration.

Images of mixed-species agar cultures to highlight differences in colony morphology. Left panel = unassisted view, right panel = colony counter microscope view. One colony representative of each species is labelled on the images. *P. gingivalis* (PG) forms opaque black-pigmented colonies, *T. forsythia* (TF) forms translucent grey colonies, and *F. nucleatum* (FN) forms large beige colonies, translucent at the edges with a raised, opaque centre.



Supplemental figure S8. The effect of zanamivir on attachment and invasion of epithelial cells coinfected with *T. forsythia* and *F. nucleatum*.

Antibiotic protection assays were performed in the presence or absence of zanamivir with *T*. forsythia and *F. nucleatum*. A) Level of *T. forsythia*-host cell association, B) Level of *F. nucleatum*-host cell association, or C) Level of both *T. forsythia* and *F. nucleatum*-host cell association. Bacterial attachment, invasion, and total association with host cells was normalised to the number of bacteria that were used to infect each condition that had survived the duration of the assay (the percentage of viable bacteria). Zanamivir= 10mM zanamivir present during host cell exposure to bacteria. Data represent the mean from three experimental repeats, and each condition was repeated in triplicate during each experiment. Error bars=SEM, Significance determined by paired T-test, *p=<0.05, **p=<0.01.



Supplemental figure S9. The effect of zanamivir on attachment and invasion of epithelial cells coinfected with *T. forsythia* and *P. gingivalis*.

Antibiotic protection assays were performed in the presence or absence of zanamivir with *T*. *forsythia* and *P. gingivalis A*) Level of *T. forsythia*-host cell association, B) Level of *P. gingivalis*-host cell association, or C) Level of both *T. forsythia* and *P. gingivalis*-host cell association. Bacterial attachment, invasion, and total association with host cells was normalised to the number of bacteria that were used to infect each condition that had survived the duration of the assay (the percentage of viable bacteria). Zanamivir= 10 mM zanamivir present during host cell exposure to bacteria. Data represent the mean from three experimental repeats, and each condition was repeated in triplicate during each experiment. Error bars=SEM. Significance determined by paired T-test, *p=<0.05.



Supplemental figure S10. The Effect of Zanamivir on Attachment and Invasion of Epithelial Cells Co-infected with *P. gingivalis* and *F. nucleatum*.

Antibiotic protection assays were performed in the presence or absence of zanamivir with *T*. *forsythia* and *F. nucleatum* A) Level of *T. forsythia*-host cell association, B) Level of *F. nucleatum*-host cell association, or C) Level of both *P. gingivalis* and *F. nucleatum*-host cell association. Bacterial attachment, invasion, and total association with host cells was normalised to the number of bacteria that were used to infect each condition that had survived the duration of the assay (the percentage of viable bacteria). Zanamivir=10mM zanamivir present during host cell exposure to bacteria. Data represent the mean from three experimental repeats, and each condition was repeated in triplicate during each experiment. Error bars=SEM, Significance determined by paired T-test, *p=<0.05, **p=<0.01.



Supplemental figure S11. The effect of zanamivir on attachment and invasion of epithelial cells coinfected with *T. forsythia, F. nucleatum*, and *P. gingivalis*.

Antibiotic protection assays were performed in the presence or absence of zanamivir with *T*. *forsythia*, *F. nucleatum*, and *P. gingivalis*. A) *T. forsythia*-host cell association, B) *F. nucleatum*-host cell association, C) *P. gingivalis*-host cell association D) *T. forsythia*, *F. nucleatum*, and *P. gingivalis*-host cell association. Bacterial attachment, invasion, and total association with host cells was normalised to the number of bacteria that were used to infect each condition that had survived the duration of the assay (the percentage of viable bacteria). Zanamivir=10mM zanamivir present during host cell exposure to bacteria. Data represent the mean from three experimental repeats, and each condition was repeated in triplicate during each experiment. Error bars=SEM, Significance determined by paired T-test, *p=<0.05, **p=<0.01. ***p=<0.001.

Supplemental table 1. Composition of EPO glycans.

| | | | | ESI-LC/M | S | | | | | | | | | | | |
|---------|--------|--------------------|-----------------|------------|--------|----------|-------|-----------|-------|--------------|--|--|--|--|--|--|
| UHPLC | | | | Compositio | on | | | | | | | | | | | |
| Peak ID | % Area | Dessible structure | Example Glycan | Hay (H) | HexNAc | Euro (E) | | potential | | | | | | | | |
| | | Possible structure | cartoon | Hex (H) | (N) | FUC (F) | 0 OAc | 1 OAc | 2 OAc | phosphate or | | | | | | |
| 1 | 0.16 | Man4+P | | 4 | 2 | 0 | 0 | 0 | 0 | 1 | | | | | | |
| 2 | 0.45 | FMan4+P | | 4 | 2 | 1 | 0 | 0 | 0 | 1 | | | | | | |
| 3 | 1.23 | Man5+P | | 5 | 2 | 0 | 0 | 0 | 0 | 1 | | | | | | |
| | | Man5+P | ● ●●●●●● | 5 | 2 | 0 | 0 | 0 | 0 | 1 | | | | | | |
| 4 | 0.15 | FA2G2S2(Ac)2 | | 5 | 4 | 1 | 1 | 0 | 1 | 0 | | | | | | |
| 5 | 0.70 | Man5+P | | 5 | 2 | 0 | 0 | 0 | 0 | 1 | | | | | | |
| | | FMan5+P | | 5 | 2 | 1 | 0 | 0 | 0 | 1 | | | | | | |

| 6 | 0.47 | FMan5+P | | 5 | 2 | 1 | 0 | 0 | 0 | 1 |
|-------|------|--------------|---|---|---|---|---|---|---|---|
| | | FA2G2S2(Ac)1 | | 5 | 4 | 1 | 1 | 1 | 0 | 0 |
| 7 | 1.67 | Man6+P | | 6 | 2 | 0 | 0 | 0 | 0 | 1 |
| | 1.16 | FA2G2S1 | | 5 | 4 | 1 | 0 | 0 | 0 | 0 |
| 0,9 | | Man6+P | | 6 | 2 | 0 | 0 | 0 | 0 | 1 |
| 10,11 | 0.62 | FMan6+P | | 6 | 2 | 1 | 0 | 0 | 0 | 1 |
| | | Man6+P | 6 | 6 | 2 | 0 | 0 | 0 | 0 | 1 |

| 12 | 3.05 | FA2G2S2 | + | 5 | 4 | 1 | 2 | 0 | 0 | 0 |
|-------|------|----------------|---|---|---|---|---|---|---|---|
| 42 | | FA2G2S1S1(Ac)2 | | 5 | 4 | 1 | 1 | 0 | 1 | 0 |
| | 0.48 | FA3G3S2(Ac)1 | | 6 | 5 | 1 | 1 | 1 | 0 | 0 |
| 15 | 0.40 | FA2G2S2 | | 5 | 4 | 1 | 2 | 0 | 0 | 0 |
| | | FA3G3S3(Ac)2 | | 6 | 5 | 1 | 3 | 0 | 1 | 0 |
| 14,15 | 0.46 | FA2G2S2 | | 5 | 4 | 1 | 2 | 0 | 0 | 0 |
| 16 | 0.94 | FA3G3S1 | | 6 | 5 | 1 | 1 | 0 | 0 | 0 |

| 17 | 0.54 | FA3G3S1 | 6 | 5 | 1 | 1 | 0 | 0 | 0 |
|----|------|---------------|---|---|---|---|---|---|---|
| | | FA4G4S3(Ac)2. | 7 | 6 | 1 | 1 | 2 | 0 | 0 |
| 40 | 1.96 | FA3G3S2 | 6 | 5 | 1 | 2 | 0 | 0 | 0 |
| 10 | | A3G3S2(Ac)4 | 6 | 5 | 0 | 0 | 0 | 2 | 0 |
| | 1.62 | FA3G3S2 | 6 | 5 | 1 | 2 | 0 | 0 | 0 |
| | | A3G3S2(Ac)4 | 6 | 5 | 0 | 0 | 0 | 2 | 0 |
| 19 | | FA4G4S3(Ac)2 | 7 | 6 | 1 | 1 | 2 | 0 | 0 |
| | | FA4G4S4(Ac)2 | 7 | 6 | 1 | 2 | 2 | 0 | 0 |

| 20 | 5.32 | FA3G3S3 | 6 | 5 | 1 | 3 | 0 | 0 | 0 |
|----|------|--------------------------------|---|---|---|---|---|---|---|
| 21 | 1.76 | FA4G4S1 or FA3G3S1(LacNAc)1 | 7 | 6 | 1 | 1 | 0 | 0 | 0 |
| | | FA3G3S3 | 6 | 5 | 1 | 3 | 0 | 0 | 0 |
| | | FA4G4S4(Ac)2 | 7 | 6 | 1 | 2 | 2 | 0 | 0 |
| | | FA4G4S3Ac | 7 | 6 | 1 | 2 | 1 | 0 | 0 |

| 22 | 6.19 | FA4G4S2 or FA3G3S2(LacNAc)1 | 7 | 6 | 1 | 2 | 0 | 0 | 0 |
|-------|-------|--------------------------------|---|---|---|---|---|---|---|
| | 0.19 | FA4G4S4 | 7 | 6 | 1 | 3 | 1 | 0 | 0 |
| 22 | 11.24 | FA4G4S3 or FA3G3S3(LacNAc)1 | 7 | 6 | 1 | 3 | 0 | 0 | 0 |
| 23 | | FA4G4S4Ac | 7 | 6 | 1 | 3 | 1 | 0 | 0 |
| 24 | 12.96 | FA4G4S4 | 7 | 6 | 1 | 4 | 0 | 0 | 0 |
| 25,26 | 2.78 | FA4G4S2(LacNac)1 | 8 | 7 | 1 | 2 | 0 | 0 | 0 |
| 27 | 9.83 | FA4G4S3(LacNac)1 | 8 | 7 | 1 | 3 | 0 | 0 | 0 |

| 28 | 1 31 | FA4G4S3(LacNac)1 | 8 | 7 | 1 | 3 | 0 | 0 | 0 |
|----|-------|---------------------|----|---|---|---|---|---|---|
| | 1.31 | FA4G4S3Ac2(LacNac)1 | 8 | 7 | 1 | 1 | 2 | 0 | 0 |
| 29 | 13.67 | FA4G4S4(LacNac)1 | 7 | 6 | 1 | 4 | 0 | 0 | 0 |
| 30 | 1.76 | FA4G4S2(LacNac)2 | 7 | 6 | 1 | 2 | 0 | 0 | 0 |
| 31 | 5.50 | FA4G4S3(LacNAc)2 | 9 | 8 | 1 | 3 | 0 | 0 | 0 |
| 32 | 8.26 | FA4G4S4(LacNAc)2 | 9 | 8 | 1 | 4 | 0 | 0 | 0 |
| 33 | 1.33 | FA4G4S3(LacNAc)3 | 10 | 9 | 1 | 3 | 0 | 0 | 0 |
| 34 | 1.72 | FA4G4S4(LacNac)3 | 10 | 9 | 1 | 4 | 0 | 0 | 0 |