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Structure aided design of a Neu5Gc specific lectin

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AB5 toxins exert their effects in a two-step process: (i) binding of the pentameric B subunit to specific glycan receptors on the target cell surface; (ii) internalisation of the AB5 toxin, followed by A subunit-mediated inhibition or corruption of essential host functions. The B subunits of AB5 toxins recognize cell surface glycan receptors, directing internalization and intracellular trafficking of the holotoxin. Specificity of these protein-glycan interactions is critical for pathogenesis, as it determines host susceptibility and tissue tropism. Moreover, the pentavalent interactions between AB5 toxin B subunits and their cognate glycans result in very high affinity binding, making them powerful ligands for glycan detection, a noteworthy example being use of the cholera toxin to demonstrate the binding of cholera toxin B subunit for detection of the ganglioside GM1 in histopathological sections and for labelling of lipid rafts in membranes.

In 2004 Paton et al. described the discovery and initial biological characterization of a new sub-family of bacterial AB5 toxins with the prototype termed subtilase cytotoxin (SubAB). In the case of SubAB, the A subunit (SubA) was found to be a subtilase family serine protease with exquisite specificity for the essential endoplasmic reticulum chaperone BiP/GRP78. Structural studies revealed that unlike most subtilases, SubA possesses an unusually deep active site cleft, explaining its exquisite substrate specificity. Significantly, glycan array analysis has shown that the B subunit of the toxin (SubB) has a high degree of binding specificity for glycans terminating with α2–3 linked Neu5Gc with little discrimination for the penultimate moiety. The crystal structure of the SubB-Neu5Gc complex revealed the basis for this specificity. The additional hydroxyl on the methyl group of the Neu5Gc linkage was changed from α2–3 to α2–6. Using molecular modeling approaches, site directed mutations were made to reduce the α2–3 to α2–6 linkage preference, while maintaining or enhancing the selectivity of SubB for Neu5Gc over Neu5Ac. Surface plasmon resonance and glycan array analysis showed that the SubBΔS106/ΔT107 mutant displayed improved specificity towards Neu5Gc and bound to α2–6 linked Neu5Gc. SubBΔS106/ΔT107 could discriminate Neu5Gc over Neu5Ac-glycoconjugates in ELISA. These data suggest that improved SubB mutants offer a new tool for the testing of biological samples, particularly serum and other fluids from individuals with cancer or suspected of having cancer.

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AB5 toxins with the prototype termed subtilase cytotoxin (SubAB)4. In the case of SubAB, the A subunit for detection of the ganglioside GM1 in histopathological sections and for labelling of lipid rafts in membranes3. Structural studies revealed that unlike most subtilases, SubA possessed an unusually deep active site cleft, explaining its exquisite substrate specificity5. SubA has proven to be a powerful tool for examining the role of BiP in diverse cellular processes and it also has potential as a cancer therapeutic6,7. Significant glycan array analysis has shown that the B subunit of the toxin (SubB) has a high degree of binding specificity for glycans terminating with α2–3 linked N-glycoylneuraminic acid (Neu5Gc), a sialic acid that humans cannot synthesise6. Of all the glycans on the array, the best binding occurred with Neu5Gcα2–3Galβ1–4GlcNAc3–. Binding of labelled toxin to the array was reduced 20-fold if the Neu5Gc was changed to Neu5Ac; over 30-fold if the Neu5Gc linkage was changed from α2–3 to α2–6; and 100-fold if the sialic acid was removed. The overall pattern of binding to structures represented on the array indicated that SubB has a high affinity for terminal α2–3 linked Neu5Gc with little discrimination for the penultimate moiety. The crystal structure of the SubB-Neu5Gc complex revealed the basis for this specificity. The additional hydroxyl on the methyl group of the
Structure-guided mutation of the glycan binding site of SubB. In order to understand the molecular basis for the preference for α2–3-linked structures, we have compared the interaction between SubB and Neu5Gcα2-3Galβ1-3GlcNAc (determined by X-ray crystallography) vs Neu5Gcα2-6Galβ1-3Glc (Fig. 1). Whereas the sub-terminal sugars of the former glycan extend freely out into the solvent, as reported previously\(^8\), the tertiary sugar of the α2–6 structure is folded back onto the SubB surface, making close contact with a loop comprising SubB residues T104-E108. This loop is stabilised by a disulphide bond between C103 and C109. The 2–6 structure is folded back onto the SubB surface, making close contact with a loop comprising SubB residues T104-E108. This loop is stabilised by a disulphide bond between C103 and C109. The resultant steric hindrance distorts the docking of the terminal Neu5Gc into the binding pocket, accounting for the significantly poorer binding of α2–6-linked Neu5Gc structures observed on the original glycan array analysis.

Since α2–6-linked sialic acids are common markers of colon cancer\(^19,20\) and are linked to prognosis in a range of cancers\(^21\), we used molecular engineering to improve binding of α2–6-linked Neu5Gc structures to SubB by designing a series of substitution and/or deletion mutants to reduce the height of the T104-E108 loop. We have modelled the interactions between these SubB mutants and Neu5Gc using molecular dynamics to predict that introduction of charged side chains into the binding pocket would be detrimental to glycan binding by SubB mutants. A Y78F mutation that prevents interactions with the C11 OH group that distinguishes Neu5Gc from Neu5Ac reduced glycan binding by 90% and abolished preference of the mutant SubB protein for Neu5Gc over Neu5Ac\(^8\). An S12A mutation abolished glycan binding completely, while a Y78F mutation that prevents interactions with the C11 OH group that distinguishes Neu5Gc from Neu5Ac reduced glycan binding by 90% and abolished preference of the mutant SubB protein for Neu5Gc over Neu5Ac\(^8\).

Interestingly, the most prominent form of aberrant glycosylation in human cancers is the expression of glycans terminated by Neu5Gc. Neu5Gc is not expressed in significant levels on normal healthy human cells\(^4,5\) as humans cannot synthesise Neu5Gc due to an inactivating mutation in the CMAH gene\(^13\). Nevertheless, research suggests that Neu5Gc presentation in cancer patients can be explained by Neu5Gc absorption through dietary intake of red meat and dairy products, which are the richest sources of Neu5Gc\(^14\). The presence of Neu5Gc is prognostically important, because its expression frequently correlates with invasiveness, metastasis and the tumour grade\(^10\). Preferential display of Neu5Gc glycans on cancer cells may be at least partly explained by the hypoxic tumour environment, which markedly induces expression of the sialic acid transporter sialin, resulting in increased display of Neu5Gc and other sialic acids on the cell surface\(^15\). Due to the fact that sialyl-conjugates regulate adhesion and promote cell mobility, such alterations in surface sialylation may influence the colonisation and metastatic potential of tumour cells\(^16\). Elevated levels of abnormal sialic acids such as Neu5Gc have been observed in breast, ovarian, prostate, colon and lung cancer\(^11,12\). Importantly, incorporation of Neu5Gc in cancer cells is most prominent in soluble glycoproteins found both in the extracellular space and inside the cell, and Neu5Gc is the dominant sialic acid in glycoproteins secreted from cancer cells into the surrounding tissues\(^7\). The expression of Neu5Gc in cancer is also known to drive production of xenautoantibodies against Neu5Gc\(^17,18\). These anti-Neu5Gc antibodies are being investigated to determine their potential for novel diagnostics, prognostics, and therapeutics in human carcinomas\(^17\).

Due to its known involvement in cancer and its normally low level in non-cancerous human tissues, detection of high levels of Neu5Gc in serum and in tissues would be considered abnormal and would be indicative of the presence of a tumour. This raises the possibility of exploiting the specificity of SubB for Neu5Gc to develop a high-throughput diagnostic screening test for a range of cancers. However, the poor affinity for α2–6 linked Neu5Gc might impact on the sensitivity of such a test. In the present study, we have examined the interaction between SubB and glycans terminating in either α2–3-linked, or α2–6 linked, Neu5Gc, with a view to designing a SubB mutant with capacity to recognise both types of structures with high affinity.

Results

Structure-guided mutation of the glycan binding site of SubB.
they would have improved recognition of α2–6-linked Neu5Gc without significantly impacting on α2–3-linked Neu5Gc binding, as shown in Fig. 2. We then constructed recombinant subB genes and expressed and purified the various proteins as C terminal His6-tagged fusion proteins from recombinant E. coli (see Materials and Methods). SubB proteins with single or double amino acid substitutions (T107A and S106A/T107A), a double deletion mutant (ΔS106/ΔT107) and a triple mutant (ΔS106/ΔT107/E108D) were successfully purified.

Surface plasmon resonance of engineered SubB mutants. Purified SubB and the various mutant derivatives were then immobilized on Biacore chips and tested for binding affinities to a range of Neu5Ac- or Neu5Gc-terminating structures (free sialic acid, sialic acid–α2–3-lactose and sialic acid–α2–6-lactose), as well as to human and bovine α1-acid glycoprotein (AGP), by surface plasmon resonance (SPR) (Table 1). The human AGP glycans contain Neu5Ac22, 23 and the bovine AGP glycans contain both Neu5Ac and Neu5Gc23. The MS glycoproteomic analysis (Fig. S1) was performed to confirm the Neu5Ac and Neu5Gc distribution in the human and bovine AGP used in the SPR study. Wild-type SubB was found to have high affinity for α2–3-linked Neu5Gc-lactose and free Neu5Gc, as predicted from the glycan array result, with nanomolar binding affinities observed. No binding was observed for the α2–6-linked Neu5Gc-lactose (tested to a maximum concentration of 25 µM) and 2.2 µM affinity was observed for α2–3-linked Neu5Ac—a more than 300-fold decrease in binding compared to the equivalent Neu5Gc structure. The wild-type SubB also had a 13-fold reduced binding affinity for human AGP compared to bovine AGP. The wild-type SubB had no binding to any non-sialylated glycans tested (Table 1). The mutation in SubB T107A had no significant effect on binding to any of the tested structures compared to the wild-type protein. SubB S106A/T107A had improved binding to α2–6-linked structures, but this improvement was seen for both Neu5Ac and Neu5Gc. The nanomolar range affinities observed for all linked sugars tested including Neu5Acα2–8 (GT2; Table 1) and binding to sulfated Chondroitin (Chondroitin-6-sulfate; Table 1) reveals that SubB S106A/T107A has a relaxed specificity range that included non-sialic acid structures. The SubB ΔS106/ΔT107/E108D mutant had improved recognition of α2–6-linked Neu5Gc without changing the binding to the α2–3-linked Neu5Ac structures. However, the difference in affinity between α2–3-linked Neu5Ac and α2–3-linked Neu5Gc was reduced to 50-fold compared to the 300-fold observed for the wild-type. The SubB ΔS106/ΔT107/E108D mutant was significantly improved for Neu5Gc vs Neu5Ac discrimination compared to the wild-type protein, and had the ability to bind α2–3-linked Neu5Gc and α2–6-linked Neu5Gc with binding affinities that were not significantly different between the two structures (15.3 nM vs 8.5 nM, respectively; P = 0.12). Thus, SubB ΔS106/ΔT107/E108D exhibited the optimum combination of enhanced Neu5Gc vs Neu5Ac discrimination and the capacity to recognise both α2–3- and α2–6-linked Neu5Gc structures. SubB ΔS106/ΔT107 also demonstrated no binding to any of the non-sialylated glycans tested (Table 1). The anti-Neu5Gc antibody produced in chicken was used as a

Figure 2. Surface representation of the wild-type and SubB mutants modeled with Neu5GCα2–6Galβ1–3Glc (shown as a cyan stick). The mutated SubB residues are shown as grey sticks and red and blue residues represent oxygen and nitrogen, respectively.
Table 1. Surface Plasmon Resonance analysis of Neu5Gc binding proteins. Binding affinities of wild type SubB, various mutant derivatives and an anti Neu5Gc IgY antibody, to purified tri- and monosaccharides and/or human or bovine α1-acid glycoprotein (AGP) was determined by SPR, as described in the Materials and Methods. NCDI indicates that no concentration- dependent interaction was observed with concentrations ranging up to 100 μM; n.t.: Not tested; Rmax: the total amount of response units (RU) of the analyte bound to the protein (the higher the number the more the glycan/glycoprotein was bound by the immobilised SubB).

| SubB variant/antibody | Human α1-AGP | Bovine α1-AGP | Neu5Ac-α1-2-3-lac | Neu5Ac-α1-2-6-lac | Neu5Gc-α1-2-3-lac | Neu5Gc-α1-2-6-lac | Free Neu5Ac | Free Neu5Gc | Man5 maltose | Lactose | GT2 | Chondroitin 6-sulfate |
|-----------------------|--------------|--------------|-------------------|-------------------|-------------------|-------------------|-------------|-------------|--------------|----------|-----|---------------------|
| ΔS106/ΔT107          | n.t.         | n.t.         | 249 ± 46 μM       | 2.34 ± 0.85 μM    | n.t.              | n.t.              | NCDI        | 35.7 ± 4.2 μM | n.t.         | n.t.     | n.t. | n.t.                |
| ΔS106/ΔT107/F108D     | 2.82 ± 0.15 μM | (Rmax = 165) | 371 ± 64 nM       | 7.39 ± 0.72 nM    | NC1D              | 3.45 ± 0.87 nM    | NCDI        | 45.1 ± 1.2 nM  | n.t.         | n.t.     | n.t. | n.t.                |
| Wild type SubB        | 2.12 ± 0.56 μM | (Rmax = 125) | 155.8 ± 22 nM     | 6.62 ± 2.17 nM    | NCDI              | NCDI              | NCDI        | 18.1 ± 5.9 nM | NCDI         | NCDI     | NCDI | NCDI                |
| ΔS106/ΔT107          | 723 ± 129 nM  | (Rmax = 142) | 164 ± 10 nM       | 348 ± 52 nM       | 8.05 ± 0.14 nM    | 3.27 ± 0.29 nM    | NCDI        | 6.61 ± 1.6 nM  | NCDI         | NCDI     | NCDI | NCDI                |
| ΔS106/ΔT107/F108D     | 2.25 ± 0.56 μM | (Rmax = 276) | 371 ± 64 nM       | 7.39 ± 0.72 nM    | NC1D              | 3.45 ± 0.87 nM    | NCDI        | 45.1 ± 1.2 nM  | n.t.         | n.t.     | n.t. | n.t.                |

Discussion

Neu5Gc is an important diagnostic and prognostic marker in human carcinomas, with elevated Neu5Gc expression detected in breast, ovarian, prostate, colon and lung cancer. 11, 12 Wild-type SubB had unprecedented control and showed less selectivity and lower affinity for Neu5Gc containing glycans than any of the SubB proteins tested.

Glycan array analysis of wild-type SubB, SubB ΔS106/ΔT107 and SubB ΔS106/ΔT107. To assess whether the preferred, Neu5Gc-specific SubB ΔS106/ΔT107, mutation introduced specificity for non-sialylated structures, not covered by the SPR analysis, glycan array analysis was performed on the SubB wild-type, SubB ΔS106/ΔT107 and SubB ΔS106/ΔT107 mutants (Table S1). Wild-type SubB displayed significant binding to only four of 402 structures on the array and SubB ΔS106/ΔT107 only had displayed significant binding to four structures on the array. These were limited to structures containing Neu5Gc and Neu5Ac. It also recognised sulfated structures including glycosaminoglycans (heparin and chondroitin-6-sulfate) and sulfated lactosamine structures (Table S1). SubB ΔS106/ΔT107 also recognised a range negatively charged of monosaccharides (Neu5Ac, Neu5Gc, 9-NAc-Neu5Ac, 3-O-Su-GlcNAc) on the array.

ELISA of engineered SubB against human and bovine proteins/serum. To assess the ability of the engineered mutants to detect the presence of Neu5Gc in biological samples ELISA assays were performed. Using dishes coated with a dilution series of SubB, labelled serum proteins from human and bovine sources were tested. A two-fold improvement in differential recognition of the Neu5Gc containing serum proteins from bovine was identified with SubB ΔS106/ΔT107 (Fig. 3).

Detection of human vs bovine AGP. To independently verify the capacity to discriminate between human and bovine AGP (only bovine AGP displays significant levels of Neu5Gc-terminating glycans), serially diluted glycoproteins were spotted onto nitrocellulose filters and after washing and blocking, filters were overlayed with a dilution series of SubB, labelled serum proteins from human and bovine sources were tested. The ability to detect Neu5Gc-terminating glycans in human AGP was not detectable even at the maximum amount tested (12.5 µg/spot). This discriminatory power is consistent with the SPR data above.
specificity for glycans terminating in Neu5Gc, but bound poorly to α2–6-linked Neu5Gc and still recognised α2–3-linked Neu5Ac structures albeit weakly 8. To improve the recognition of SubB for α2–6-linked Neu5Gc and make it more specific for Neu5Gc, we engineered SubB using structure-aided modifications, with specific focus on the T104-E108 loop.

Manipulation of this loop had two specific outcomes through the modification of the same two amino acids. Firstly, alanine substitution of S106 and T107 (S106A/T107A) led to a loss of specificity for Neu5Gc, producing a lectin capable of binding to all tested terminally sialylated glycans regardless of linkage (α2–3, α2–6 and α2–8) or sialic acid type (Neu5Ac or Neu5Gc) in SPR studies. Glycan array analysis confirms the relaxed specificity and revealed binding to additional, sulfated glycans. The second was that deletion of the same two amino acids (ΔS106/ΔT107) produced a lectin with exquisite specificity for Neu5Gc regardless of linkage (α2–3 and α2–6). The SubBΔS106/ΔT107 mutant was significantly improved for the recognition Neu5Gc containing structures compared to the wild-type SubB. SubBΔS106/ΔT107 also had no difference in its ability to bind α2–3-linked Neu5Gc or α2–6 linked Neu5Gc structures, making it a significant improvement over the wild-type protein. Further modifications of the SubB protein outside of the S106 and T107 amino acids produced no significant improvement in specificity. The SubBΔS106/ΔT107/E108D mutant protein, which is the SubBΔS106/ΔT107 protein with a E108D mutation also added, was less able to distinguish α2–3-linked Neu5Gc from α2–3-linked Neu5Ac than SubBΔS106/ΔT107 and had stronger binding to the human α1-Acid glycoprotein than the SubBΔS106/ΔT107 mutant (24 fold more protein bound by SubBΔS106/ΔT107/E108D than SubBΔS106/ΔT107).

These improved SubB mutants offer a new tool for the testing of biological samples, particularly serum and other fluids from individuals with cancer or suspected of having cancer.

**Figure 3.** ELISA of engineered SubB against FITC-labelled human and bovine serum. SubB (A) and SubBΔS106/ΔT107 (B) coated onto ELISA plates was able to capture FITC-labelled human and bovine serum proteins. Error bars show ±1 SD from the mean of duplicate assays.

**Figure 4.** Lectin overlay assay. Binding of SubBΔS106/ΔT107 to serial dilutions of human or bovine AGP spotted onto nitrocellulose (total amounts of protein per spot indicated), was determined as described in the Materials and Methods.
added to wells coated with SubB or SubBΔM115.056366].

Instrument (SCIEX) as previously described [Zacchi Schulz MCP 2016 PMID: 27094473 doi:10.1074/mcp.

Wells were washed 3 times with PBS-T. 100µg of protein overnight at 4 °C. Wells were washed 3 times with phosphate-buffered saline, 0.05% Tween-20 (PBS-T) before blocking solution (3% BSA) was added for 1 hour at room temperature. Proteins in normal human serum and bovine serum were fluorescently labelled by

Surface Plasmon Resonance of SubB and engineered SubB mutants. Surface Plasmon resonance (SPR) was run using the Biacore T100 system (GE) as described previously 27. Briefly, SubB, SubB mutants and anti-Neu5Gc IgY (SiaMab; formerly Sialix/GC-Free Inc., San Diego, CA, USA) were immobilized onto flow cell 2–4 of a series S sensor chip CM5 (GE) using the NHS capture kit and flow cell 1 was run as a blank immobilization. Monosaccharides, disaccharides, oligosaccharides and α1-Acid glycoprotein from human and bovine sources (Sigma-Aldrich; See Table 1) were flowed over at 0.01–100 µM on initial range finding experiments. Concentrations were adjusted and all data were analysed using single cycle kinetics using the Biacore T100 Evaluation software.

Mass spectrometric analysis of α1-Acid glycoprotein. AGP from human plasma (Sigma-Aldrich G9885) and bovine plasma (Sigma-Aldrich G3643) (1 mg in 6 M guanidinium chloride, 50 mM Tris-HCl pH 8) was reduced and alkylated with 10 mM dithiothreitol and 25 mM acrylamide, respectively. Protein was then precipitated by adding 4 volumes of 1:1 methanol:acetone, incubating in −20 °C for 16 h and then centrifuged (18,000 rcf, 10 min) to collect the pellet. The precipitated protein was resuspended in 50 µL of 50 mM Tris-HCl pH8 and digested (37 °C, 16 h) with 1 µg trypsin (Trypsin Gold, Promega). Digested peptides were then desalted with C18 ZipTips (Millipore). Mass spectrometry of desalted peptides was performed using a TripleTOF of 5600 instrument (SCIEX) as previously described [Zacchi Schulz MCP 2016 PMID: 27094473 doi:10.1074/mcp.M115.056366].

Glycan array analysis of SubB and engineered SubB mutants. Glycan arrays were printed on SuperEpoxy 3 (ArrayIt) activated substrates using an ArrayIt Spotbot Extreme contact printer as previously described28. For each subarray 2µg of SubB proteins were pre-complexed with anti-His tag antibody (Cell signaling) and Alexa555 secondary and tertiary antibodies (rabbit anti-mouse; goat anti-rabbit) at a ratio of 2:1:0.25:0.25 in a final volume of 500 µL. This 500 µL antibody protein complex was added to a 65 µL gene frame (Thermo Scientific) without a coverslip. Washing and analysis was performed as previously described27.

ELISA analysis of SubB and the engineered SubBΔS106ΔT107 mutant. Wells of black 96-well NUNC Maxisorp plates were coated with SubB or SubBΔS106ΔT107 protein two fold serially diluted in 100 mM bicarbonate/carbonate coating buffer (pH 9.6) starting at 1.25 µg of protein overnight at 4 °C. Wells were washed 3 times with phosphate-buffered saline, 0.05% Tween-20 (PBS-T) before blocking solution (3% BSA) was added for 1 hour at room temperature. Proteins in normal human serum and bovine serum were fluorescently labelled by combining neat serum with 100 µM FITC dye (Peirce) and incubating on ice for 1 hour. Excess dye was removed using a 1 Kda size exclusion spin column. 100 µl of FITC-labelled normal human serum or bovine serum was added to wells coated with SubB or SubBΔS106ΔT107 and wells were incubated for 1 hour at room temperature. Wells were washed 3 times with PBS-T. 100 µL of PBS was added to each well before the fluorescence was measured at 485/535 nm. Fluorescence unit values are shown as the mean of duplicates +/− SD, with the mean fluorescence units obtained for wells containing all reagents except for the SubB proteins subtracted. Any negative value was considered as 0.

SubB overlay experiments. Purified SubBΔS106ΔT107 was labelled with biotin using the EZ-Link® Sulfo-NHS-Biotinylation Kit (Thermo Scientific) according to the manufacturer's instructions. Purified human and bovine α−1 acid glycoprotein (Sigma cat. nos G9885 and G3643) were dissolved in water at 5 mg/ml and 5µl volumes of serial two-fold dilutions were spotted onto nitrocellulose filters and air dried at 37 °C overnight. Filters were then blocked with 5% skim milk in Tris-buffered saline with 0.05% Tween 20 (TTBS) for 2h. After washing three times in TTBS, filters were overlaid with 1 µg/ml biotin-SubBΔS106ΔT107 in TTBS and incubated overnight.
at 4 °C. Filters were then washed three times in TTBS and bound biotin-SubBΔΔΔΔΔΔDeltaT10 was detected using streptavidin-alkaline phosphatase conjugate (Roche). Filters were developed using a chromogenic nitro-blue tetrazolium/X-phosphate substrate system (Roche).

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Author Contributions

A.W.P., J.C.P., C.J.D. and M.P.J. designed the experiments. A.W.P. and B.P.H. made and produced the SubB and SubB mutants for this publication. M.A.H produced the models in Figures 1 and 2. C.J.D., A.W.P., L.K.S., B.P.H. performed the experiments from Table 1 and Figures 3 and 4. F.E.J. and B.L.S. performed and analysed the experiments in Figure S1. C.J.D., M.P.J., A.W.P., J.C.P., F.E.J., B.L.S. and L.K.S. wrote the manuscript.

Additional Information

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Competing Interests: The following authors (A.W.P., J.C.P., C.J.D. and M.P.J.) declare that they inventors on a provisional patent application on matter contained in this manuscript that will be filed soon.

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