Role of the membrane cholesterol-glycosphingolipid complex as a ‘transistor’ to regulate GSL receptor function and signaling of both lipids

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Running title- GSL-cholesterol complex regulates ligand binding
Abstract

Cholesterol and glycosphingolipids (GSL) are the major species that accumulate in plasma membrane lipid rafts. These complexes imbue the membrane with increased order, which in turn, plays a central role in the transmembrane signaling foci lipid rafts provide. In addition, both GSL and cholesterol binding can mediate (separate) signal pathways. We have shown that cholesterol and GSLs however, form a complex in which the GSL sugar is reoriented from a membrane perpendicular to parallel format, becoming largely unavailable for exogenous ligand binding. Similarly, the steroid hydroxyl is masked, restricting access of cholesterol ligands. This was observed in model and cell membranes and in human tumour frozen tissue sections. We now show the order of exogenous ligand binding plays a significant role to determine the extent of GSL or cholesterol receptor activity. Ligand binding to cholesterol enhances subsequent GSL recognition and vice versa, suggesting that ligand binding to “free” receptor (membrane perpendicular GSL carbohydrate, nonmasked cholesterol) can result in partial dissociation of the GSL/cholesterol complex to allow additional GSL ligand and cholesterol ligand binding. Since many GSLs can complex with membrane cholesterol, the binding of a single cholesterol ligand may unmask cholesterol-complexed GSL for increased binding of both a single or multiple GSL-specific ligands. We show that multiple cholesterol-masked GSLs can be coincident in tissues. This provides a mechanism for GSL-dependent signal amplification and diversification, representing a biological ‘transistor’, regulating amplitude and potentially, diversity of GSL signaling. The process represents a new mechanism of ‘cross-talk’ between GSL and cholesterol signaling. This is of clinical importance since we have found cholesterol/GSL masking applies to monoclonal anti GSL antibodies in development and in current use as antineoplastic therapeutics.
Introduction

GSLs are involved in many signal transduction pathways[1], both directly[2] and as modulators of the transduction of other signals[3,4]. Their increased concentration, together with cholesterol[5], in membrane lipids rafts[6] provide the appropriate location for modulating [7] these foci of transmembrane signaling[8].

While aberrant GSL metabolism is the direct cause of the lysosomal GSL storage diseases[9], GSLs play a major role in other human diseases, such that inhibition of their synthesis results in the amelioration of clinical and animal disease model symptoms[10-13]. Errors in cholesterol homeostasis are widespread causes of disease [14-20]. Membrane cholesterol can transduce signals[21] and, via lipid rafts, modulate the activity of many trafficking[22,23]and signaling processes[24-26].

The sugar-lipid conjugate nature of GSL structure results in complex modulation of membrane GSL receptor function [27-30]. One of the best defined mechanisms for this modulation is the masking of membrane GSL expression within the cholesterol/GSL complex [31-34]. This cholesterol -induced conformational change in the GSL carbohydrate, from ligand available, membrane perpendicular (uncomplexed GSL), to ligand unavailable membrane parallel (cholesterol complexed) GSL conformers is dependent on an H-bond network between the steroid hydroxyl and the anomeric oxygen and amino function of the GSL [32]. Cholesterol masking is maximized under conditions of minimal membrane acyl chain mismatch[33]. Increased acyl chain mismatch promotes lipid rafts [35] so cholesterol masking of GSLs may be also modulated at the raft-membrane interface [7]. Due to the increased levels of cholesterol in tumours [36], GSL masking is a common feature observed in human tumour biopsies[34] and may be a means to escape immune surveillance, providing a rationale for cholesterol depletion in tumour immunotherapy.

Significantly, several Mabs approved and in development for human cancer treatment, turn out to target tumour GSLs [37-39]. In our studies, we found that several GSL antigens were cholesterol masked in the same location in tumour frozen sections (e.g. Gb3 and SSEA1 in colon cancer[34]). This led to our consideration of the effect of cholesterol masking of one GSL on the cholesterol masking of another.

From our previous study on the effect of sequential ligand binding to Gb3 and to cholesterol in prostate cancer sections[34], we now propose a model in which cholesterol masking may provide a switch (transistor) for GSL antigen exposure and hence, a network of ligand activated, GSL dependent signal transduction.

Materials and Methods
Tissue staining
Frozen tumor tissues were obtained from the Department of Pediatric Laboratory Medicine at this hospital and the Ontario Institute for Cancer Research Tumor Bank (Toronto, Ont, Canada). Cryosections (6 μm) were dried and fixed with 4% paraformaldehyde in phosphate-buffered saline (PBS) for 10 min then treated ±10 mM MCD in PBS for 30 min at room temperature (RT). Under these conditions, MCD preferentially extracts cholesterol [40] and we routinely observed negative section staining with filipin.

For brightfield microscopy of sections stained with horseradish peroxidase (HRP) endogenous peroxidase blocker (Universal Block, KPL, Inc., Gaithersburg, MD) was added for 20 min, then samples were blocked with 1% normal goat serum (NGS) in PBS (1% NGS) for 20 min. Sections were treated with primary antibodies or VT1-B subunit. All reagents were diluted in 1% NGS, and all toxin/antibody reactions were for 30 min. VT1-B subunit (1 μg/mL) was purified in our laboratory [41]. VTB incubation was followed by rabbit polyclonal anti-VT1-B-6869 [42] diluted 1:1000. Unituxin (dinutuximab anti-GD2 mAb, humanized chimeric mAb 14.18) was generously provided by United Therapeutics Inc NC, USA and Mab F77 was a gift from Dr M Greene, University of Pennsylvania, Pa, USA. Primary antibodies were used at 2 μg/mL. Secondary antibodies conjugated to HRP were applied (1:500): HRP-goat anti-rabbit IgG, HRP-goat anti-mouse IgG, or HRP-goat anti-human IgG (Bio-Rad, 172-1050). HRP-goat anti-mouse IgG (Bio-RAD, Hercules, CA, 170-6516).

Sections were incubated for 5 min with peroxidase substrate (ImmPACT dianaminobenzidine, Vector Laboratories, Burlingame, CA). Nuclei were weakly counterstained with aqueous Mayer’s hematoxylin (Vector Laboratories, Burlingame, CA). Slides were dehydrated in ethanol, cleared with xylene and mounted in Permount (Fisher, Ottawa, ON Canada). Images were recorded with a Nikon microscope (Eclipse E4000) using the Nikon ACT-1 software, version 2.70 and a Sony camera using Infinity Lumenara software, version 6.5.4. Pixel quantification was performed on digital images using Image J software.

For fluorescence microscopy of filipin-stained sections, filipin III (Sigma, St Louis, MO, F4767) was dissolved in dimethyl sulfoxide/PBS (1:4) (50 μg/mL) and added to sections for 30 min. After rinsing, slides were mounted in aqueous fluorescent-mounting medium (Dako Cytomation, Glostrup, Denmark). Images were recorded with a Leica microscope (DMI 6000 B) using the LA SAF software, version 2.2.1. For double staining, sections were stained with filipin and VTB/antibodies. When filipin was used first, it was followed by VTB, primary antibody, then HRP conjugated secondary antibodies. When VTB, antibody was reacted first, filipin was added after the HRP-conjugated secondary antibody, followed by incubation with the peroxidase substrate. All incubations were at RT. Slides were mounted in the aqueous fluorescent-mounting medium and recorded with the Leica microscope.

Results and Discussion
Cholesterol/GSL masking provides a basis for amplification of GSL signaling
Our results on the effect of sequential binding of Verotoxin B subunit (to Gb3 GSL) and filipin (to cholesterol) in prostate cancer biopsy frozen sections[34], have been reconstructed in figure 1A, and provide the basis of our ‘transistor’ model (figure 1B) for the regulation of GSL receptor function in membrane GSL signaling.

An electrical transistor is a semiconductor which provides a switch mechanism for regulating current flow in an external circuit. A small current pulse through a transistor can induce the flow of a much larger pulse in the receiver (amplifier). The transistor can also allow a single electric input pulse to be split to many downstream receivers.

Using serial prostate cancer frozen sections, a binding comparison was made when VTB or filipin was added first or second, and the detection of each ligand separately assessed and compared (fig 1A). The binding of filipin prior to VTB (right panels), resulted in increased detection of Gb3 by VTB compared to the same tubular structures in the tumour stained first with VTB (fig 1A top right, arrowed tubules). Not all VTB stained tubules show this amplification. Similarly, cholesterol detection with filipin was greater following initial VTB binding (left lower panel). Filipin staining was particularly elevated for tubules (fig 1A lower panel left, arrows) not susceptible to filipin-increased VTB binding. This could be due to differential membrane cholesterol vs Gb3 concentrations.

Based on this data, we propose (figure1B) that ligand –receptor binding can partially unmask the cholesterol/GSL complex to amplify GSL or cholesterol mediated signaling pathways. In the GSL/cholesterol complex (represented by the red rhomboid, and blue triangle fig 1B), the availability of each lipid for ligand binding is restricted. This complex will, however, be in equilibrium with the free fraction of the membrane lipids. Ligand binding to either free lipid will drive the equilibrium a little to the right. This complex dissociation will provide additional unmasking of the other lipid to amplify binding of the second ligand. Thus, GSL binding increases the cholesterol available to bind an appropriate ligand (in this case, filipin). Such a ligand could contain a cholesterol binding motif[43,44], for example. Theoretically, due to the widespread nature of GSL masking by cholesterol[31,34], any GSL specific ligand[45] may unmask otherwise unavailable cholesterol. In the reverse case, a cholesterol binding ligand will similarly partially dissociate the GSL/cholesterol complex to partially unmask membrane GSL for increased ligand binding.

Our finding that the filipin cholesterol binding was increased more than VTB Gb3 binding is consistent with the proposal that more than one molecule of cholesterol is involved in the GSL/cholesterol complex[33] i.e. more cholesterol than Gb3 would be unmasked. This increased lipid receptor availability could take the form of allowing a lower concentration of the second ligand to bind the membrane lipid and would thus function as a ‘switch’, activated by the binding of the first ligand.
The signal amplification can be quantitated by $\Delta_{\text{freeGSL}} = k(nL2)$ or $\Delta_{\text{freechol}} = k(nL1)$ where $k$ is the coefficient of complex dissociation and L1 and L2 are the two ligands.

Since many GSLs are cholesterol masked[31], it is likely that any cholesterol binding species[44] could unmask a variety different GSLs if present, allowing diversification and/or amplification of a signaling route.

If two or more masked GSLs are masked within the same cholesterol domain, it is possible that, subsequent to GSL 1 ligand binding, a cholesterol binding ligand may then unmask both GSLs, resulting in signal diversification and the binding of one GSL ligand would result in the unmasking of other GSLs. This signal diversification would permit location dependent signal cross talk.

Cholesterol is also involved in many signal transduction pathways [21,46-48] and this model provides an interface between between GSL recognition and these pathways.

From analysis of figure 1A, we propose by analogy, that a biological ligand binding to a membrane GSL, which itself is either complexed, or not complexed to membrane cholesterol, can both amplify the signal transduced and split the signal into multiple downstream, GSL/cholesterol dependent signaling pathways. Thus, although by entirely different mechanisms, GSL/cholesterol signaling can therefore behave as a biological equivalent of a transistor.

**Mab anti tumour GSLs are subject to cholesterol masking**

To emphasize the clinical relevance, particularly in cancer, we have investigated the effect of cholesterol masking on the tumour binding of two antiGSL Mabs developed for the treatment of human cancers.

**A) Mab F77.** This Mab was raised by immunization with a human prostate cancer cell line[37]. Prostate cancer is the cancer in which cholesterol plays a major role[49,50]. This antibody binds a H-blood group related carbohydrate expressed on GSLs and mucin[51]. Figure 2 shows the binding of Mab F77 to serial frozen tissue sections from a primary human prostate cancer biopsy, before and after beta-methyl cyclodextrin mediated cholesterol extraction. Mab F77 staining to the same tissue components was increased by 350% following MCD cholesterol extraction (primarily tubular structures [52]).

**B) Mab anti GD2 (Unituxin)** is the first FDA approved Mab for pediatric cancer (neuroblastoma [39,53]). Binding of this humanized mouse anti GD2 ganglioside antibody was increased to 156% following cholesterol depletion, (figure 3). This significant increase in tumour binding offers the opportunity to increase the clinical efficacy of this antineoplastic approach. In this regard, it is of interest to note that a cyclodextrin based method to deplete membrane cholesterol in pediatric patients has been developed for Niemann-Pick disease [54], which is without apparent long term side effects.
Coincident unmasking of different GSL antigens. Our transistor model implies that ligand binding to one masked GSL could result in unmasking of a second masked GSL if it were in the same location. We therefore investigated whether two GSL could be masked in the same location. Figure 4 shows the localization of unmasked GD2 and Gb3 in colon cancer serial sections. MCD untreated sections show weak GD2 staining within the tumour. Cholesterol depletion markedly un_masks GD2 which is largely uniformly distributed within the tumour, but is not found in the columnar epithelial cells (fig 4B*). Gb3 is not detectable in untreated sections, except in these epithelial cells (fig 4C*). Significant Gb3 is unmasked (fig 4D) which is more widely distributed and overlaps unmasked GD2 (fig 4B,D arrows). Non-coincident GSL unmasking demonstrates the selectivity of cholesterol-based GSL masking (i.e. it is not a non-selective general enhancement, but rather depends on the presence of specific GSLs). Co- incidental unmasking of different GSLs indicates locations in which our transistor model of signal diversification could operate.

In breast tumour (figure 5), these antigens are, interestingly, partially colocalized within the tubular ducts of the tumour. GD2 is expressed within the tumour stroma and also within the outer layers of ducts (figure 5A). Increased GD2 expression is observed in both locations after cholesterol depletion (figure 5B). In contrast, Gb3 is expressed in a few blood vessels in the untreated section, but following cholesterol extraction, blood vessel staining is strongly increased, together with staining of the middle and inner layers of the duct structures. Thus, these unmasked GSL antigens overlap within the middle layer of the ducts but are distinct in the outer (GD2) and inner (Gb3) tumour duct layers.

Thus, a transistor based signal might be selectively propagated to the middle layer interface cells of the duct structures from either ligation of the outer GD2 masked layer or the inner Gb3 masked layer.

Significance
Our studies demonstrate a new mechanism for the physicochemical regulation of cellular GSL and cholesterol receptor function, whereby molecules within a membrane lipid complex can ‘talk’ to each other. The transistor-like mechanism for signaling we have demonstrated in (what was) living tissue, can likely be mimicked in model membranes, since cholesterol masking of GSLs is readily apparent in such vesicles [31], opening the possibility that such a biological transistor might be useful in bridging biological and semiconductor based computation [55]. Biological computer prototypes have been made based on synthetic DNA[56] or RNA[57] but the present model would be the first ‘transistor’ operating in normal unmodified cells.

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Disclosure statement
The authors declare no conflict of interest in these studies.
Legends

Figure 1. Double labeling of Gb3 and filipin to illustrate function of membrane GSL transistor. (A). Serial sections of a prostate cancer biopsy were co stained either with VTB/peroxidase (to detect Gb3, brown) or filipin-fluorescence (to detect cholesterol, green) or filipin first followed by VTB) [34]. Bar=200µm. Adding VTB first (top left) promoted subsequent filipin staining (bottom left). Filipin staining was most increased for vesicle showed the greatest initial VTB (Gb3) staining (lower left panel, arrows). In contrast, adding filipin first (top right) promoted VTB binding (not to all vesicles but see vesicles arrowed). (B) The explanation for this effect provides the basis of the GSL/cholesterol transistor. Binding of either ligand (VTB or filipin) to the small fraction of non-complexed lipid will result in a shift in the complexed vs. free equilibrium to the right, and thus dissociate a fraction of the complex, to promote the binding of the second ligand. This can be a switch or amplification for the second ligand binding.

Figure 2. F77 Mab binding to prostate tumour is markedly increased by MCD cholesterol depletion. F77 binding to serial prostate cancer biopsy before and after cholesterol depletion. A few tubular structures are labeled before but many more after MCD extraction.

Figure 3 AntiGD2 staining of pediatric neuroblastoma
Anti GD2 (Unituxin) showed extensive staining of neuroblastoma sections (left panels) but cholesterol extraction resulted in a significant increase in tumour reactivity (right panels). Bar=250µm.

Figure 4 Colabelling of unmasked GD2 ganglioside and Gb3 in colon carcinoma biopsy sections. Serial human colon carcinoma frozen sections were stained for GD2 (A) or Gb3 (B) using Unituxin or VTB, prior to (A,B) or after (C,D), MCD cholesterol extraction. A marked increase is staining or these glycolipids within the bulk of the tumour was observed (arrows).* marks columnar epithelial cells. Inverted images are shown. Bar=50µm.

Figure 5 Colabelling of unmasked GD2 ganglioside and Gb3 in breast carcinoma biopsy sections. Serial human breast carcinoma frozen sections were stained for GD2 (A,B) or Gb3 (C,D) using Unituxin or VTB, prior to (A,C) or after (B,D), MCD cholesterol extraction. Little initial VTB blood vessel staining was markedly increased after cholesterol depletion and the central and luminal cells of tumour ducts were labeled. More of the tumour stromal tissue was initially antiGD2 labelled (A*) as were outer layer cells of tumour ducts(A arrow). Both were significantly increased by MCD cholesterol extraction. Bar=100µm.
References

1. Lingwood CA (2011) Glycosphingolipid Functions. Cold Spring Harb Perspect Biol Biology of Lipids: 149-174.
2. Chatterjee S, Pandey A (2008) The Yin and Yang of lactosylceramide metabolism: implications in cell function. Biochim Biophys Acta 1780: 370-382.
3. Shu L, Shayman JA (2003) Src kinase mediates the regulation of phospholipase C-gamma activity by glycosphingolipids. J Biol Chem 278: 31419-31425.
4. Klokk TI, Kavaliauskiene S, Sandvig K (2016) Cross-linking of glycosphingolipids at the plasma membrane: consequences for intracellular signaling and traffic. Cell Mol Life Sci 73: 1301-1316.
5. Macdonald JL, Pike LJ (2005) A simplified method for the preparation of detergent-free lipid rafts. J Lipid Res 46: 1061-1067.
6. Pike LJ (2004) Lipid rafts: heterogeneity on the high seas. Biochem J 378: 281-292.
7. Lingwood D, Simons K (2010) Lipid rafts as a membrane-organizing principle. Science 327: 46-50.
8. Staubach S, Hanisch FG (2011) Lipid rafts: signaling and sorting platforms of cells and their roles in cancer. Expert Rev Proteomics 8: 263-277.
9. Ballabio A, Gieselmann V (2009) Lysosomal disorders: from storage to cellular damage. Biochim Biophys Acta 1793: 684-696.
10. Karman J, Tedstone JL, Gumlaw NK, Zhu Y, Yew N, et al. (2010) Reducing glycosphingolipid biosynthesis in airway cells partially ameliorates disease manifestations in a mouse model of asthma. Int Immunol 22: 593-603.
11. Zhao H, Przybylska M, Wu IH, Zhang J, Maniatis P, et al. (2009) Inhibiting glycosphingolipid synthesis ameliorates hepatic steatosis in obese mice. Hepatology 50: 85-93.
12. Langeveld M, Aerts JM (2009) Glycosphingolipids and insulin resistance. Prog Lipid Res 48: 196-205.
13. Tsukuda Y, Iwasaki N, Seito N, Kanayama M, Fujitani N, et al. (2012) Ganglioside GM3 has an essential role in the pathogenesis and progression of rheumatoid arthritis. PloS one 7: e40136.
14. Magura L, Blanchard R, Hope B, Beal JR, Schwartz GG, et al. (2008) Hypercholesterolemia and prostate cancer: a hospital-based case-control study. Cancer Causes Control 19: 1259-1266.
15. Tierney E, Bukelis I, Thompson RE, Ahmed K, Aneja A, et al. (2006) Abnormalities of cholesterol metabolism in autism spectrum disorders. American journal of medical genetics Part B, Neuropsychiatric genetics: the official publication of the International Society of Psychiatric Genetics 141B: 666-668.
16. del Toro D, Xifro X, Pol A, Humbert S, Saudou F, et al. (2010) Altered cholesterol homeostasis contributes to enhanced excitotoxicity in Huntington’s disease. J Neurochem 115: 153-167.
17. Cai SY, Gautam S, Nguyen T, Soroka CJ, Rahner C, et al. (2009) ATP8B1 deficiency disrupts the bile canalicular membrane bilayer structure in hepatocytes, but...
FXR expression and activity are maintained. Gastroenterology 136: 1060-1069.
18. Sun H, Yuan Y, Sun ZL (2013) Cholesterol Contributes to Diabetic Nephropathy through SCAP-SREBP-2 Pathway. International journal of endocrinology 2013: 592576.
19. Llaverias G, Danilo C, Mercier I, Daumer K, Capozza F, et al. (2011) Role of cholesterol in the development and progression of breast cancer. The American journal of pathology 178: 402-412.
20. Nishimura S, Ehara S, Hasegawa T, Matsumoto K, Yoshikawa J, et al. (2017) Cholesterol crystal as a new feature of coronary vulnerable plaques: An optical coherence tomography study. J Cardiol 69: 253-259.
21. Luchetti G, Sircar R, Kong JH, Nachtergaele S, Sagner A, et al. (2016) Cholesterol activates the G-protein coupled receptor Smoothened to promote Hedgehog signaling. Elife 5.
22. Hoekstra D, van Ijzendoorn SC (2000) Lipid trafficking and sorting: how cholesterol is filling gaps. Curr Opin Cell Biol 12: 496-502.
23. Lippincott-Schwartz J, Phair RD (2010) Lipids and Cholesterol as Regulators of Traffic in the Endomembrane System. Annu Rev Biophys 39: 559-578.
24. Pike L, Casey L (2002) Cholesterol levels modulate EGF receptor-mediated signaling by altering receptor function and trafficking. Biochemistry 41: 10315-10322.
25. Simons K, Ehehalt R (2002) Cholesterol, lipid rafts, and disease. J Clin Invest 110: 597-603.
26. Epand RM (2006) Cholesterol and the interaction of proteins with membrane domains. Progress in lipid research 45: 279-294.
27. Kannagi R, Nudelman E, Hakomori S (1982) Possible role of ceramide in defining structure and function of membrane glycolipids. Proc Natl Acad Sci USA 79: 3470-3474.
28. Kannagi R, Stroup R, Cochran NA, Urdal DL, Young Jr. WW, et al. (1983) Factors affecting expression of glycolipid tumor antigens: Influence of ceramide composition and coexisting glycolipid on the antigenicity of gangliotriaosylceramide in murine lymphoma cells. Cancer Res 43: 4997-5005.
29. Crook SJ, Boggs JM, Vistnes AI, Koshy KM (1986) Factors affecting surface expression of glycolipids influence of lipid environment and ceramide composition on antibody recognition of cerebroside sulfate in liposomes. Biochemistry 25: 7488-7494.
30. Nakakuma H, Ari M, Kawaguchi T, Horikawa K, Hidaka M, et al. (1989) Monoclonal antibody to galactosylceramide: discrimination of structural difference in the ceramide moiety. FEBS Lett 258: 230-232.
31. Mahfoud R, Manis A, Binnington B, Ackerley C, Lingwood CA (2010) A major fraction of glycosphingolipids in model and cellular cholesterol containing membranes are undetectable by their binding proteins. J Biol Chem 285: 36049-36059.
32. Yahi N, Aulas A, Fantini J (2010) How cholesterol constrains glycolipid conformation for optimal recognition of Alzheimer's beta amyloid peptide (Abeta1-40). PLoS One 5: e9079.
33. Lingwood D, Binnington B, Róg T, Vattulainen I, Grzybek M, et al. (2011) Cholesterol modulates glycolipid conformation and receptor activity. Nature Chem Biol 7: 260-262.
34. Novak A, Binnington B, Ngan B, Chadwick K, Fleshner N, et al. (2013) Cholesterol masking membrane glycosphingolipid tumor-associated antigens reduces their immunodetection in human cancer biopsies. Glycobiology 23: 1230-1239.
35. Heberle FA, Petruzielo RS, Pan J, Drazba P, Kucerka N, et al. (2013) Bilayer thickness mismatch controls domain size in model membranes. J Am Chem Soc 135: 6853-6859.
36. Beloribi-Djefaflia S, Vasseur S., F. G (2016) Lipid metabolic reprogramming in cancer cells. Oncogenesis 5.
37. Carroll AM, Zalutsky M, Schatten S, Bhan A, Perry LL, et al. (1984) Monoclonal antibodies to tissue-specific cell surface antigens. I. Characterization of an antibody to a prostate tissue antigen. Clin Immunol Immunopathol 33: 268-281.
38. Zhang G, Zhang H, Wang Q, Lal P, Carroll AM, et al. (2010) Suppression of human prostate tumor growth by a unique prostate-specific monoclonal antibody F77 targeting a glycolipid marker. Proc Natl Acad Sci U S A 107: 732-737.
39. Jones PC, Irie RF (2016) Therapeutic Strategies for Human IgM Antibodies Directed at Tumor-Associated Ganglioside Antigens: Discoveries Made During the Morton Era and Future Directions. Crit Rev Oncog 21: 75-81.
40. Ottico E, Prinetti A, Prioni S, Giannotta C, Basso L, et al. (2003) Dynamics of membrane lipid domains in neuronal cells differentiated in culture. J Lipid Res 44: 2142-2151.
41. Nutikka A, Binnington-Boyd B, Lingwood CA (2003) Methods for the purification of Shiga toxin 1. In: Philpot D, Ebel F, editors. Methods Mol Med. Totowa, NY: Humana Press. pp. 187-195.
42. Boyd B, Richardson S, Gariepy J (1991) Serological responses to the B subunit of Shiga-like toxin 1 and its peptide fragments indicate that the B subunit is a vaccine candidate to counter the action of the toxin. Infect Immun 59: 750-757.
43. Li H, Papadopoulos V (1998) Peripheral-type benzodiazepine receptor function in cholesterol transport. Identification of a putative cholesterol recognition/interaction amino acid sequence and consensus pattern. Endocrinology 139: 4991-4997.
44. Fantini J, Barrantes FJ (2013) How cholesterol interacts with membrane proteins: an exploration of cholesterol-binding sites including CRAC, CARC, and tilted domains. Frontiers in physiology 4: 31.
45. Mahfoud R, Garmy N, Maresca M, Yahi N, Puigserver A, et al. (2002) Identification of a common sphingolipid-binding domain in Alzheimer, prion and HIV-1 proteins. J Biol Chem 277: 11292-11296.
46. Incardona JP, Eaton S (2000) Cholesterol in signal transduction. Curr Opin Cell Biol 12: 193-203.
47. Paila YD, Chattopadhyay A (2010) Membrane cholesterol in the function and organization of G-protein coupled receptors. Subcell Biochem 51: 439-466.
48. Blassberg R, Macrae JI, Briscoe J, Jacob J (2016) Reduced cholesterol levels impair Smoothed activation in Smith-Lemli-Opitz syndrome. Human molecular genetics 25: 693-705.
49. Di Vizio D, Solomon KR, Freeman MR (2008) Cholesterol and cholesterol-rich membranes in prostate cancer: an update. Tumori 94: 633-639.
50. Armandari I, Hamid AR, Verhaegh G, Schalken J (2014) Intratumoral steroidogenesis in castration-resistant prostate cancer: a target for therapy. Prostate International 2: 105-113.
51. Gao C, Liu Y, Zhang H, Zhang Y, Fukuda MN, et al. (2014) Carbohydrate sequence of the prostate cancer-associated antigen F77 assigned by a mucin O-glycome designer array. The Journal of biological chemistry 289: 16462-16477.
52. Nagle RB, Cress AE (2011) Metastasis Update: Human Prostate Carcinoma Invasion via Tubulogenesis. Prostate Cancer 2011: 249290.
53. Mora J (2016) Dinutuximab for the treatment of pediatric patients with high-risk neuroblastoma. Expert Rev Clin Pharmacol 9: 647-653.
54. Vecserynes M, Fenyvesi F, Backsay I, Deli MA, Szente L, et al. (2014) Cyclodextrins, Blood-Brain Barrier, and Treatment of Neurological Diseases. Archives of medical research 45: 711-729.
55. Simon DT, Gabrielsson EO, Tybrandt K, Berggren M (2016) Organic Bioelectronics: Bridging the Signaling Gap between Biology and Technology. Chem Rev 116: 13009-13041.
56. Roquet N, Soleimany AP, Ferris AC, Aaronson S, Lu TK (2016) Synthetic recombinase-based state machines in living cells. Science 353: aad8559.
57. Bonnet J, Yin P, Ortiz ME, Subsoontorn P, Endy D (2013) Amplifying genetic logic gates. Science 340: 599-603.
Fig 1
fig 2

F77 staining of frozen prostate tumour serial sections

cholesterol extracted
Fig 4
Fig 5