Virus recognition by specific natural antibodies and complement results in MHC I cross-presentation

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Natural antibodies (NAb) and complement (C’) are important regulators of immune system activation. We have shown previously that the galactosyl-α1,3-galactosyl (Galα1,3Gal) xenoantigen and the similar ABO histo-blood group antigens are transferred onto virus from the producer cell, resulting in sensitisation of the virus to the respective NAb in a C’-dependent manner. Here we show that measles virus (Mv) that expresses Galα1,3Gal termini can drive the proliferation of human T cells in the presence of serum and autologous DC, whereas without such targets, measles, as expected, suppress T cell reactivity. The use of affinity-purified NAb to Galα1,3Gal and rabbit C’ demonstrated the components in human serum responsible for this effect. Proteasome inhibition and blocking of antigen presentation showed that the increased T cell proliferation was mediated by MHC class I cross-presentation of immune complexes. These results lend further support to the idea that polymorphic carbohydrates of the Galα1,3Gal/ABO type serve as important targets for NAb and C’ and that their expression on virus has influenced their evolution by contributing to protection against viral transmission within as well as between species. The adjuvance effect of this recognition, acting as a bridge between the natural innate and adaptive immune systems, also has important implications for vaccine development.

Introduction

Natural antibodies (NAb) and complement (C’) contribute to an important but not well-defined block of both bacterial and viral spread to vital organs preceding the development of an adaptive and specific immune response [1, 2]. NAb, assisted by C’, serve as endogenous adjuvants for the generation of protective CD8+ T cells after vaccination against leishmaniasis [3]. Studies in mice that lack expression of secreted IgM have shown that IgM NAb are necessary components of an anti-viral response [4]. Furthermore, such NAb recognition contributes not only to direct clearance of virus but also to the stimulation of immune responses requiring macrophages and/or follicular dendritic cells (FDC) [5]. DC play a major role in the initiation of a specific immune response by presenting antigens to and activating naive T cells, and many viruses have, as a consequence, evolved strategies for interfering with their functions [6]. MHC class I cross-presentation of viral antigens by DC, either by uptake of virus-infected cells or viral immune complexes, can serve as a means by which the host cell can circumvent the viral interference [7]. In addition, some viruses do not infect DC and have to be presented by this pathway for efficient T cell activation [8]. In the case of measles virus (Mv), a
The nature of the target for NAb on viruses is much less clear. Since a virus must inherit its glycosylation pattern from the producer cell, questions arise as to how there could possibly be targets for NAb on viral envelopes. The production of specific NAb in only some individuals of a species (e.g. anti-ABO) or in only some species (e.g. anti-Gal1,3Gal) correlates with the absence of the respective carbohydrate epitope. Antibodies against the Gal1,3Gal epitope are produced by splenic B cells in both primates and mice that have been engineered to lack expression of the Gal1,3Gal structure [12, 13]. It is not clear why such polymorphic carbohydrate/NAb systems have evolved. Classically, bacteria have been thought to be chiefly responsible [14]. We, as well as others, have shown previously that human cells can be a target for neutralisation mediated by NAb and C by transfecting pig cDNA [15] encoding a1,3galactosyltransferase (α1,3GT), which synthesizes the Gal1,3Gal epitope [16, 17]. This mimics the situation in xenotransplantation in which recognition of the Gal1,3Gal epitope leads to hyperacute rejection of pig tissue in the presence of human serum [18, 19], and it has subsequently been suggested as a means to induce rejection of tumour cells following de novo expression of the Gal1,3Gal epitope [20, 21]. More recently, this phenomenon was shown to be due to the induction of specific immunity against tumour antigens [22, 23]. In addition, C-type retrovirus produced from α1,3GT-transfected cells became sensitised to human serum containing specific NAb and C [16, 17]. We have also shown that in addition to the Gal1,3Gal epitope, the similar ABO epitopes can appear on and sensitise measles to human non-immune serum containing the relevant NAb and C [24]. More recently we found that this is also true for HIV virus [25]. These findings suggest that polymorphic carbohydrates and the NAb against them may have evolved partly in response to the appearance of such carbohydrates on enveloped virus being transmitted from one individual to another or between species. Further support for this contention was provided in a mathematical modeling study, which showed that the frequencies of ABO histo-blood groups in human populations can be explained by their direct interactions with pathogens (e.g. bacteria) in combination with their appearance on viruses serving as targets for ABO NAb [26].

In this study we examine whether C-assisted specific NAb recognition of measles virus leads to increased T cell reactivity in the presence of DC. If such an effect on T cells is seen, this would point out a potentially important function for polymorphic terminal carbohydrates on viruses beyond the neutralising effect previously observed [16, 17, 24, 25, 27]. Our data show that this is the case and indicate that this mechanism could potentially be utilised to increase cytotoxic T cell responses in viral vaccinations without the need for other adjuvants.

Results

Expression of Galu1,3Gal terminal glycosylation on measles

Carbohydrate termini on HT1080 cells were modified by stable transfection with cDNA encoding α1,3GT, denoted HT-1,3GT, or with the empty plasmid as a control (HT1080–0) [16]. Different measles preparations with (Mv-G1–3G) or without (Mv) Gal1,3Gal terminal glycosylation were produced by infecting HT-1,3GT or HT1080–0 cells. The amount of Gal1,3Gal terminal glycosylation was assessed by a capture-ELISA assay (Fig. 1) [24]. A terminal galactose-specific lectin, BSI-B4, was used to show that measles produced from the Galu1,3Gal-expressing HT-1,3GT cells acquired the terminal glycosylation, whereas those expressed in the HT1080–0 did not. These preparations were then used to determine the effect of terminal Galu1,3Gal glycosylation on human T cell responses in the presence of autologous DC and either autologous serum or affinity-purified NAb against Galu1,3Gal terminal carbohydrates and rabbit C'.

Suppression of T cell proliferation by measles virus is reversed by serum and C'

In a series of experiments, 2.5 × 10^4 DC were incubated together with 2.5 × 10^5 live viral particles as determined by a plaque assay on Vero cells, as previously described [24]. These amounts of DC and viral particles were found to be optimal from several experiments using different amounts and conditions (data not shown). A representative experiment performed in triplicate using these conditions is shown (Fig. 2a). T cell and DC exposure to measles in the absence of human serum resulted in complete obliteration of T cell proliferation, indicating viral suppression of the T cell immunity to infection 1255
response, as previously documented [28–30]. Both viruses, i.e., displaying Galα1,3Gal or not, exhibited a similar effect on T cell proliferation (p<0.001). In the presence of autologous serum deprived of C', exposure to either virus (Mv-G1–3G or Mv) resulted in low proliferation (p<0.006 and p<0.01, respectively). When rabbit C' was added to heat-inactivated serum, more pronounced T cell proliferation was seen; moreover, proliferation was significantly higher when Mv-G1–3G virus (with the NAb epitope) was used as compared to Mv (without the epitope) (p<0.03).

**T cell proliferation in the presence of measles together with NAb and C'**

To ascertain whether NAb and C' alone could result in the triggering of T cell proliferation, we used T cells and DC from the same donor and affinity-purified Ab against Galα1,3Gal together with rabbit C' instead of serum. As shown in Fig. 2b, measles without Ab and C' again resulted in a drastic suppression as compared to the background proliferation without virus. This level of suppression was also seen when Mv (without NAb epitopes) was used in the presence of Galα1,3Gal-specific NAb and C'. However, when Mv-G1–3G (with the NAb epitope) was used, proliferation was restored to background levels (p<0.001). The use of UV-irradiated measles, however, yielded normal or even above background levels of proliferation even in the absence of NAb and C', indicating that the virus needed to replicate in the DC in order to suppress T cell proliferation. We hypothesized that the high background proliferation was due to the presence of 'xenoantigens' in the fetal calf serum (FCS) included in the media. We next assessed whether Galα1,3Gal-specific NAb and C' and their target on measles could increase T cell proliferation on their

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**Figure 1.** Capture ELISA showing expression of the Galα1,3Gal antigen on measles virus. Microwell plates were coated with an anti-measles hemagglutinin mAb. Virus-containing supernatants from cells transfected with an α1,3GT cDNA clone were diluted with respective supernatant from uninfected cells and added at an identical 5 × 10^5 PFU/mL. Subsequently, an HRP-labelled Galα1,3Gal-specific lectin from *B. simplicifolia* (BSI-B4) was added at different concentrations (50, 5 and 0.5 μg/mL), after which the wells were incubated with peroxidase substrate and the absorbance analyzed on a plate reader. The results, expressed as arbitrary units, are normalised to the background value of the virus-free supernatant incubated with 0.5 μg/mL BSI-B4 and are shown as the mean ± SEM of duplicate samples. Significant differences were found between Mv-G1–3G and Mv at both 50 and 5 μg/mL BSI-B4 staining (*p<0.02 and **p<0.05, respectively).

**Figure 2.** Galα1,3Gal on the virus reverses measles-induced suppression of T cell proliferation. Measles virus produced from HT1080–0 (Mv) or HT-1,3GT (Mv-G1–3G) cells were exposed to a source of NAb and C' in triplicate series, and DC-mediated T cell proliferation was measured using [3H]-thymidine incorporation (cpm). (a) Autologous heat-inactivated (HI) human serum with or without added C' was used. Significant differences between T cell proliferation are indicated by the respective p values. The results are representative of several individual experiments. (b) Measles virus, UV-irradiated or not, was exposed to Galα1,3Gal-specific NAb with or without rabbit C' (*p<0.001, T cell proliferation Mv versus Mv-G1–3G, in the presence of purified NAb and C'). The results are representative of two individual experiments.

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own, in the absence of background proliferation caused by FCS-associated antigens.

**T cell proliferation in the presence of measles, serum and reduced FCS concentration**

When a similar T cell proliferation experiment was performed in only 3% FCS, as a means of reducing the amount of xenogeneic background antigens in the cultures, and using autologous serum as the source of NAb and C', 'no measles' virus incubation as well as UV-irradiated Mv (without Gal\(_1,3\)Gal) resulted, as expected, in reduced autologous T cell proliferation (Fig. 3a), as compared to the previous experiment using 10% FCS in combination with purified NAb and C' (Fig. 2b). However, when UV-irradiated Mv-G1–3G virus was used with intact human serum, drastically increased C'-dependent proliferation resulted (\(p<0.001\)).

**T cell proliferation when using NAb, C' and reduced amounts of FCS**

In order to determine that it is the Gal\(_1,3\)Gal moieties in combination with specific NAb and C' that result in increased T cell proliferation in this autologous setting with low background proliferation, we next repeated the experiments using affinity-purified Ab against Gal\(_1,3\)Gal together with rabbit C' instead of serum in the presence of only 3% FCS. It is well known that measles readily infects DC, upon which it efficiently inhibits T cell activation [28–30]; after UV irradiation, however, measles virus does not induce inhibition [28]. We thus conducted these experiments with both live infectious measles and UV-irradiated measles shown to lack any infectious particles as detected by a highly sensitive Vero cell plaque assay (data not shown) [24] to investigate whether infection of the DC (or not) has an effect on the degree of NAb- and C'-dependent T cell activation. As shown in Fig. 3b and 3c, this resulted in significantly higher T cell proliferation only in the presence of NAb against Gal\(_1,3\)Gal and C' and only when using the Mv-G1–3G virus with NAb targets (\(p<0.001\) and \(p<0.05\)). This effect appeared to be largely independent of whether the measles was live and thus infectious or UV-irradiated and therefore non-infectious, thus indicating that opsonisation and uptake of virus-NAb-C' complexes rather than infecting particles resulted in T cell activation. Furthermore, the results when using UV-irradiated non-infectious measles with both serum and affinity-purified NAb and C' in the low background setting indicated that it is very unlikely that the T cell proliferation was simply due to a neutralising effect of the NAb and C'.

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Figure 3. T cell proliferation following exposure of autologous DC differentiated in 3% FCS to differentially glycosylated measles viruses. (a) UV-irradiated non-infectious measles virus produced from HT1080 (Mv) or HT-1,3GT (Mv-G1–3G) cells were exposed to fresh human serum with or without (heat-inactivated, HI) intact C' in triplicate series. (b) Infectious measles viruses (Mv or Mv-G1–3G) were exposed to Gal\(_1,3\)Gal-specific NAb, with or without C', in triplicate series. (c) UV-irradiated non-infectious measles (Mv or Mv-G1–3G) were exposed to Gal\(_1,3\)Gal-specific NAb, with or without C', in triplicate series. In addition, conditions with medium alone were used in all three experiments. DC were subsequently added, and autologous T cell proliferation was measured using \(^{[3]H}\)-thymidine incorporation. *Significant differences in T cell proliferation were observed between Mv and Mv-G1–3G with intact fresh human serum (\(p<0.001\)), with affinity-purified NAb and C' using infectious virus (\(p<0.001\)) and with affinity-purified NAb and C' using UV-irradiated non-infectious virus (\(p<0.05\)).
Cells with Galα1,3Gal can substitute for virus in inducing T cell proliferation

In order to further address the formal possibility that the T cell activation could be due mainly to neutralisation by NAb and C and to compare the measles results with another source of antigens, we next investigated whether UV-irradiated producer cells of the measles virus are able to substitute for the virus in driving T cell proliferation. The results show that the allogeneic HT-1,3GT producer cells of Mv-G1–3G virus were able to substitute for the virus and appeared to provide antigenic stimulation driving T cell proliferation via the DC cultures (Fig. 4a). However, while Mv-G1–3G virus-stimulated proliferation was again strictly C'-dependent, the HT-1,3GT-driven proliferation appeared to be less so. When the Mv producer cells HT1080–0 were used, no T cell proliferation resulted, again mirroring the results using Mv produced from HT1080–0 (Fig. 4b). Proliferation using DC and medium only or no T cells resulted in levels at or below the levels seen in Fig. 3b (data not shown).

DC in the co-culture do not display increased general endocytosis or activation

The increase in T cell proliferation may be related to an altered ability of DC to endocytose in general and/or to become activated. We therefore tested whether or not DC incubated with different measles viruses and/or mAb and/or C' were able to maintain similar levels of endocytosis. We used the fluid phase markers FITC-dextran and rhodamine-BSA in combination with flow cytometric analysis. The FITC-dextran was internalised and accumulated in a time-dependent manner in DC (Fig. 5). Similar results were observed for rhodamine-BSA uptake (data not shown). The mean value of fluorescence was then compared for the different conditions and the different viruses after 20 min of endocytosis; no differences in uptake were observed for the FITC-dextran (Fig. 5). Very similar results were

![Figure 4](image1.png)

**Figure 4.** T cell proliferation following exposure of differing amounts of autologous DC to differentially glycosylated cell lines or measles with autologous human serum. (a) Irradiated HT-1,3GT cells or Mv-G1–3G produced from these cells were exposed to human serum with (solid symbols) or without (heat-inactivated, open symbols) intact C'. (b) Irradiated HT1080–0 cells or Mv produced from these cells were exposed to human serum with (solid symbols) or without (heat-inactivated, open symbols) intact C'. In all experiments DC and T cells were subsequently added, and autologous T cell proliferation was measured using [3H]-thymidine incorporation.

![Figure 5](image2.png)

**Figure 5.** Fluid-phase endocytosis by DC was assessed by FITC-dextran internalisation as quantified by FACS analysis. During FITC-dextran internalisation, DC were incubated with or without control measles virus (Mv) or measles virus expressing Galα1,3Gal epitopes (Mv-G1–3G) in the presence or absence of the purified NAb against Galα1,3Gal epitopes and/or C'. The relative uptake was determined after 20 min incubation. Values on the y-axis indicate the mean fluorescence intensity (MFI).
obtained when using Rhodamine-BSA to study uptake under the different conditions (data not shown).

We then tested whether or not the incubation of DC with measles viruses under the different conditions modulated the membrane expression of markers potentially involved in the activation of T cells. We observed that incubation of DC with the different viruses and under the different conditions was associated with strong activation of DC in all cases. However, no differences in the levels of CD83 and CD86 were observed between Mv and Mv-G1–3G under the different conditions (Fig. 6). Moreover, the expression of CD46, the measles virus receptor at the cell surface of DC, was similar following incubation of DC with measles viruses having the Galα1,3Gal epitope or not and in the presence or absence of NAb and/or C' (data not shown). In addition, no differences were found when comparing the expression levels of MHC class I or class II on the DC (data not shown).

T cell proliferation is the result of MHC class I cross-presentation

In order to examine whether the NAb/C'-mediated T cell proliferation was due to antigen presentation and to assess to what extent such antigen presentation is mediated through the MHC class I pathway, we used lactacystin, a specific inhibitor for this pathway (e.g. after receptor-mediated uptake of virus) [31]. Lactacystin treatment drastically reduced the purified NAb- and C'-mediated proliferation down to background levels, whereas anti-Galα1,3Gal and C' again resulted in significantly higher T cell proliferation with Mv-G1–3G virus as compared to with Mv (Fig. 7a). Since these data strongly indicate that mainly MHC class I presentation of viral antigens is responsible for the T cell proliferation, we next confirmed this finding by showing that a functionally blocking MHC class I mAb (W6/32) [32] was able to also reduce the T cell proliferation to background levels (Fig. 7a). In addition, to exclude the possibility that lactacystin was having a general toxic effect on the cells, thereby causing reduced proliferation of T cells, we confirmed that the same concentration of lactacystin did not significantly inhibit proliferating T cells in an allogeneic mixed lymphocyte reaction (Fig. 7a) or in the presence of anti-CD3 and IL-2 (Fig. 7b). Since it has been shown that lactacystin may inhibit NF-kappa B activity as well as cause apoptosis of some cells [33], we also confirmed that the degree of MHC class I expression on the DC was not significantly affected by lactacystin incubation (Fig. 7c) and that lactacystin did not induce a significant degree of apoptosis in either proliferating T cells or activated mature DC (Fig. 7d).

Mainly CD8+ T cells expand as a result of the Galα1,3Gal recognition

We next determined the phenotype of the expanded population of T cells following incubation with Mv-G1–3G. In the presence of Mv-G1–3G plus NAb and C', the percentage of CD8+ T cells in a co-culture with DC and T cells was increased by 54% as compared to a co-culture with Mv (without the Galα1,3Gal epitope),
representing an absolute increase in CD8$^+$ T cells. No increase in the proportion of CD4$^+$ T cells was seen under the same conditions (Fig. 8a), and no significant T cell death was observed. We also confirmed that a vast majority of the CD8$^+$ T cells following proliferation were TCR$\alpha\beta^+$, thus representing classical T cells; no change was seen after addition of lactacystin (Fig. 8b).

Figure 7. T cell proliferation is dependent on MHC class I presentation. (a) Measles without (Mv) or with (Mv-G1–3G) Gal$\alpha$1,3Gal were exposed to purified NAb with added C' in triplicate series. DC were subsequently added, and autologous T cell proliferation was measured using [$^3$H]-thymidine incorporation in the presence or absence of lactacystin (Lc) or a blocking anti-MHC class I mAb. When present, the lactacystin and mAb were able to significantly inhibit proliferation (both $^*p<0.004$). (b) Proliferating T cell cultures were treated with the same concentration of lactacystin as in (a) and did not significantly differ from the control culture. (c) Non-activated as well as activated DC were shown to have similar MHC class I expression levels with and without lactacystin. (d) No significant increase in apoptotic activity could be seen in either two types of proliferating T cells or in activated DC following lactacystin treatment. CICCP represents control treatment to induce maximal apoptosis.
Discussion

Cells as well as viruses display specific terminal carbohydrate epitopes that serve as targets for NAb. In the presence of C', NAb are able to destroy cells expressing their carbohydrate epitopes and neutralise viruses carrying such epitopes [16]. In vivo, this kind of virus neutralisation can be envisaged to occur in serum or certain other biological fluids (e.g. mucosa) associated with epithelia directly or indirectly exposed to the environment. We addressed the possibility that recognition of specific polymorphic carbohydrates (e.g. Galα1,3Gal) on viruses by NAb, with or without the assistance of C', might lead not only to neutralisation of the virus but also to increased activation of T cells. Our results indicate that expression of Galα1,3Gal histo-blood group-like epitopes for NAb on viruses (i.e. measles) leads to increased T cell proliferation in the presence of human serum and autologous DC. This proliferation appears to be entirely dependent on the presence of intact C'. We then proceeded to show that affinity-purified NAb against Galα1,3Gal together with C' results in the same effect, indicating that these are the components in human serum responsible for driving the T cell proliferation. Since the donors were immune against measles, we conclude that the NAb-mediated proliferation must exceed any specific anti-measles Ab-mediated effects. The fact that UV-irradiated non-infectious measles virus was able to stimulate the same degree of NAb-assisted proliferation when using both human serum and affinity-purified anti-Galα1,3Gal and C' showed that neither the DC nor the T cells needed to be infected by measles virus for the proliferation to occur. One alternative explanation for our measles data would be that the normal suppressive effect of measles is overcome simply by a reduction in the number of measles virus particles available for infection of the DC. We feel that this alternative explanation can be ruled out as the main contributing mechanism, however, since: (i) The measles-driven Galα1,3Gal-dependent proliferation remains significantly above background levels when the FCS concentration is lowered to 3%, in combination with decreased T cell proliferation when DC numbers were titrated down stepwise to zero (Fig. 4 and data not shown); (ii) The α1,3GT-transfected cells alone are able to support similar proliferation, whereas the same cell line without Galα1,3Gal epitopes on the cell surface is not (Fig. 4), a mechanism which has been shown to be the explanation for why the Galα1,3Gal epitope has been successfully used to initiate an anti-tumour cell response in vitro as well as in vivo [20–23]. Since this represents an allogeneic situation, the antigens that are presented to the responding T cells directly or indirectly could be either major (i.e. MHC class I or II molecules) or other minor antigens. The fact that they appeared to be efficiently presented to the T cells only in the presence of NAb and their carbohydrate targets on the cells would indicate that the cells needed to be lysed and/or cell fragments containing antigens needed to form immune complexes, thus essentially mimicking the situation with the measles virus; (iii) We showed that lactacystin, an inhibitor of proteasome-dependent MHC class I antigen presentation, inhibits the proliferation whilst not causing inhibition of an allogeneic mixed lymphocyte reaction, NF-kappa B inhibition or apoptosis. This observation warrants further detailed investigation to elucidate the exact function of the pathway in this system, e.g. by the use of measles-specific T cells. However, the fact that mainly classical CD8+ TCRβ+ T cells proliferated after NAb, C' and Galα1,3Gal target interaction, together with to the fact that both lactacystin treatment and anti-MHC mAb blocking could efficiently inhibit the proliferation and the finding that non-infectious virus could be used, clearly showed that the T cell proliferation is the result of MHC class I antigen presentation of measles antigens by way of the cross-presentation pathway. This pathway has previously been
shown to be responsible for virus-specific antibody-mediated enhancement of viral antigen presentation [31].

Morbilliviruses (i.e. viruses related to measles and distemper) are well known to cross species barriers, often with devastating consequences. NAb recognition of such viruses (as well as potentially other emerging viral types, e.g. SARS and avian influenza) may therefore be an important contributor to protecting against cross-species transmission. It is also very likely that the same effect on T cells would result if the Galα1,3Gal were replaced by the very similar ABO histo-blood group antigens and the NAb directed against them, even if this remains to be demonstrated.

It has been suggested that recognition by NAb and C' can lead to rapid T-independent increases in B cell responses. Such B cell triggering, the mechanisms of which are far from clear, is believed to be partly mediated by macrophages and follicular dendritic cells (FDC) in the marginal zone of lymph nodes [5] following the binding of immune complexes to C' receptors on their respective cell surfaces. It is thus possible that NAb of the anti-Galα1,3Gal histo-blood group-like type recognizing virus could lead to increased T-independent B cell responses using the same mechanism. However, it is even less clear what happens to Ab-group-like type recognizing virus could lead to increased NAb recognition of leishmaniasis [3]. In our case, we also describe natural adjuvance effect of NAb and C' and can be independent of infection of the DC. Such recognition could result in an important ‘head start’ for the adaptive immune response, as assisted by an NAb repertoire determined by factors other than previous viral exposure. In addition, it would ensure that virus types that do not normally infect DC are quickly handled by DC, for example through the recently described Toll-like receptor (TLR) 3 and 7 pathways for recognition of viral RNA [42–44], and would enhance e.g. IL-4, IL-6 and/or IFN-α/β production, in particular in a newly infected naive individual. Measles recognition via this pathway could include the wild-type measles-specific activation of TLR2 signalling [45]. This type of ‘natural recognition’ of viruses constitutes a parallel example to the recently described natural adjuvant effect of NAb and C' recognition of leishmaniasis [3]. In our case, we also define a target for the NAb and C' as a terminal carbohydrate of the histo-blood group-like type.

Viral vaccine formulations are often far from optimal and in some cases simply fail (e.g. HIV). One possible avenue for improvement has involved work with viral subunit vaccines. For such uses, as well as in other vaccine situations, an improved CD8+ T cell-mediated cytotoxic response would be crucial. It has been shown...
that cytotoxic responses can be drastically enhanced by the cross-presentation pathway for MHC class I/CD8 [46], including responses against viruses in vivo [47]. In an influenza virus model, it was shown that intact C′ is of crucial importance for CD4+ as well as CD8+ T cell activation [48]. The recent in vivo demonstration that Galu1,3Gal epitopes conjugated to BSA or gp120 from HIV can result in a very impressive adjuvant effect, with resulting levels of antibody and cytokine as well as cytotoxic responses as good as or even better than in the presence of Freund’s complete adjuvant alone [35, 36], in combination with the involvement of NAb, C′ and MHC class I presentation in the effect on T cell proliferation seen here with live or UV-inactivated measles carrying antigenic carbohydrates, shows that further studies into the effect such recognition may have on vaccination are warranted. It would be particularly interesting to see what effect this glycosylation would have in the context of different subunit or split viral vaccines, where a great need for safe but efficient adjuvant alternatives exists.

Materials and methods

Cells and virus production

Human HT1080 cells were previously transfected with porcine cDNA for α1,3GT, HT-1,3GT; or a mock construct, HT1080–0 [16]. The Loss strain of measles [49] was kindly provided by Dr. D. Brown at the Health Protection Agency, Colindale, UK. This measles strain was passaged through the human HT1080 cells and subsequently used to infect either α1,3GT cDNA-transfected HT-1,3GT or mock-transfected HT1080–0 cells, from which supernatants containing virus (referred to as Mv-G1–3G or Mv, respectively) were harvested upon maximal virus production for 1 h at 25°C containing 0.1% Tween 20 and 1% bovine serum albumin (DBS). Nunc Maxisorp 96-well plates were coated overnight at 4°C with 1 μg anti-measles hemagglutinin mAb (clone ZD6, Biodesign Int., Kennebunk, ME). The wells were washed using 0.1% Tween 20 in PBS, blocked with a PBS solution containing 0.1% TWEEN 20 and 1% bovine serum albumin (DB) for 1 h at 25°C and again washed. Viral supernatants (Mv-G1–3G or Mv), diluted with respective uninfected cell supernatants, were added at identical 5 × 10^5 PFU/mL and aliquots stored at –80°C. The wells were washed; after immunodepletion with anti-HLA-DR, anti-CD19 immunobeads (Dynal, AS, Norway) and used for further differentiation into DC, as previously described [50]. Briefly, the resulting cells were resuspended in RPMI medium, plated in endotoxin-free culture plates and incubated for 2 h to allow monocyte adhesion. Subsequently, adherent cells were incubated for 5 days at 37°C/5% CO2 in RPMI medium containing 10^5 U/mL GM-CSF, 10^4 U/mL IL-4 and 10% FCS, allowing differentiation into immature DC. After 5 days non-adherent cells were removed, washed and immunodepleted using anti-CD3 and anti-CD19 immunobeads (Dynacl AS, Norway). Antigenic T cells were isolated simultaneously from the non-adherent fraction of peripheral blood cells following centrifugation and washes; after immunodepletion with anti-HLA-DR, anti-CD19 and anti-CD14 immunobeads (Dynacl AS, Norway), the cells were stored at –80°C. Local Ethical Committee permission was received for these studies, and informed consent from the participating laboratory worker volunteers was obtained.

T cell proliferation assay

Immature DC (2.5 × 10^5/well unless otherwise stated) were γ-irradiated and incubated for 2 h with 2.5 × 10^5 Mv-G1–3G or Mv particles, UV-irradiated or not, in the presence of 20% human autologous immune serum with or without heat-inactivation of C′ for 30 min at 56°C. Alternatively, 10 μg/mL affinity-purified baboon anti-Galu1,3Gal, a gift from BioTransplant Inc. (Charlestown, MA) and 1% rabbit C serum (Sigma) was used instead of human serum. In one experiment virus was substituted for by 10^5 irradiated cells of the different virus-producing cell lines HT1080–0 or HG-1,3GT, and in another experiment 2.5 × 10^4 allogeneic DC were used as stimulators. Subsequently, 2.5 × 10^4 T cells were added in an equal volume, resulting in a 1:2 dilution of the previously added components; in some experiments lactacyclin (10 μM; Sigma) or 50 μg/mL of the blocking anti-MHC class I mAb W6/32 (a kind gift from P. Kourilsky, Paris) (32) was added to selected cultures. Cells were incubated at 37°C for 4 days in 3% FCS. [3H]-thymidine (1 μCi) was added to cultures, which were then left for an additional 24 h. On day 5, cultures were harvested and assayed using a Wallac Trilux Microbeta scintillation counter. In another proliferation protocol, T cells were obtained from non-adherent PBMC by negative selection using anti-CD20, anti-CD56 and anti-CD14 mAb conjugated to magnetic beads (Miltenyi Biotec, Auburn, CA). The cells were then stimulated for 4 days using 0.25 μg anti-CD3 mAb (OKT3; Ortho Biotec, Bridgewater, NJ) and 100 IU/mL IL-2 (Chiron, Suresnes, France) in RPMI/10% FCS, with or without the addition of 10 μM lactacyclin.

Fluid phase marker endocytosis

DC were incubated for 2 days with or without control measles virus (Mv) or measles virus expressing the Galu1,3Gal epitope (Mv-G1–3G), in the presence or absence of purified baboon antibodies against the Galu1,3Gal epitope and rabbit C′ serum.
Expression of membrane markers on DC and T cells

DC were incubated for 2 days with or without control measles viruses (Mv) or measles viruses expressing the Galα1,3Gal epitope (Mv-G1–3G), in the presence or absence of purified baboon antibodies against the Galα1,3Gal epitope and rabbit C' serum (Sigma). In one experiment DC were incubated with 100 ng/mL LPS (Sigma) and/or 10 μg lactacytisin. Cells were then washed twice with PBS at 4°C and incubated with mAb against human MHC class I (Diaclone, France), CD46, CD80, CD86 (all Santa Cruz Biotechnology, CA), CD4 (Diaclone) or CD8 (Diaclone) directly conjugated with FITC or PE for 45 min at 4°C. Cells were then washed three times with PBS at 4°C and directly analyzed using a Beckman-Coulter Epics XL flow cytometer.

Apoptosis assay

Purified T cells were stimulated with anti-CD3 mAb (0.25 μg/mL OKT3; Ortho Biotech) and IL-2 (100 IU/mL) or with OKT3 and anti-CD28 (1 μg/mL; eBioscience, San Diego, CA) for 4 days and incubated in the presence or absence of 10 μM lactacytisin for 2 days. Living cells were determined by FACS analysis using propidium iodine (PI) added 10 min before analysis at 4°C to evaluate the integrity of the cellular membrane and DiOC6 (a marker of the integrity of the mitochondrial transmembrane potential, ΔΨm). Dying cells were PI+ and/or DiOC6−/−. As an apoptosis control for DiOC6 staining, cells were incubated for 10 min with 5 μM carbonyl cyanide m-chlorophenyl hydracide (CICCP; Sigma), which induced a complete ΔΨm loss. Similarly, mature DC were activated with LPS for 2 days and were treated or not with lactacytisin. The viability of the cells was determined as for lymphocytes, and the percentage of dying cells (PI+ and/or DiOC6−/−) was assessed.

Statistical analysis

Statistical analyses were performed using a paired Student’s t-test for in vitro experiments. A difference was considered as significant when p<0.05.

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